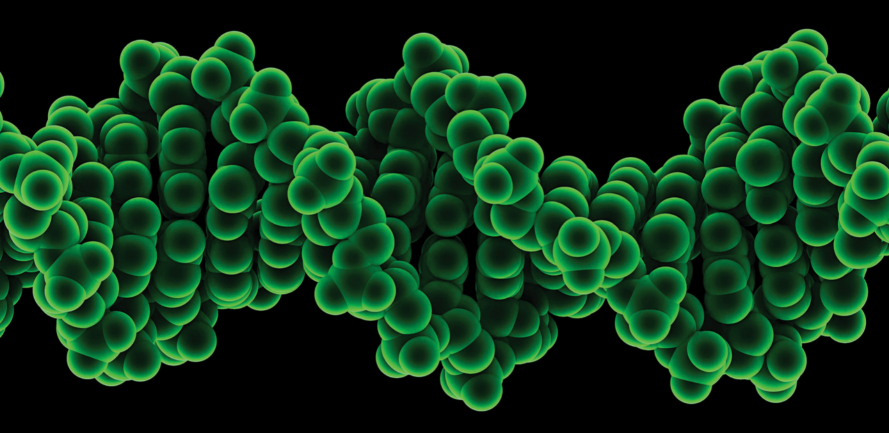
Principles of GENETICS

SIXTH EDITION



SNUSTAD • SIMMONS

Principles of GENETICS

SIXTH EDITION

D. Peter Snustad

University of Minnesota

Michael J. Simmons

University of Minnesota



John Wiley & Sons, Inc.

About the Cover: The cover shows a three-dimensional model of a DNA molecule. The two strands in the molecule are wound around each other to form a double helix.

SENIOR EDITOR Kevin Witt ASSISTANT EDITOR Lauren Morris SENIOR PRODUCTION/ILLUSTRATION EDITOR Elizabeth Swain SENIOR MEDIA EDITOR Linda Muriello MEDIA SPECIALIST Daniela DiMaggio EXECUTIVE MARKETING MANAGER Clay Stone EDITORIAL ASSISTANT Jennifer Dearden SENIOR DESIGNER Maureen Eide INTERIOR AND COVER DESIGNER: *John Michael* GRAPHICS SENIOR PHOTO EDITOR Jennifer MacMillan PHOTO RESEARCHER Lisa Passmore COVER PHOTO Laguna Design/Peter Arnold, Inc./Photolibrary This book was set in 10/12 Janson Text by Aptara Inc. and printed and bound by R.R. Donnelley/Jefferson City. The cover was printed by R.R. Donnelley/Jefferson City.

This book is printed on acid free paper.

Founded in 1807, John Wiley & Sons, Inc. has been a valued source of knowledge and understanding for more than 200 years, helping people around the world meet their needs and fulfill their aspirations. Our company is built on a foundation of principles that include responsibility to the communities we serve and where we live and work. In 2008, we launched a Corporate Citizenship Initiative, a global effort to address the environmental, social, economic, and ethical challenges we face in our business. Among the issues we are addressing are carbon impact, paper specifications and procurement, ethical conduct within our business and among our vendors, and community and charitable support. For more information, please visit our website: http://www.wiley.com/go/citizenship.

Copyright © 2012, 2009, 2006, 2003, 2000, and 1997 John Wiley & Sons, Inc. All rights reserved. No part of this publication may be reproduced, stored in a retrieval system or transmitted in any form or by any means, electronic, mechanical, photocopying, recording, scanning or otherwise, except as permitted under Sections 107 or 108 of the 1976 United States Copyright Act, without either the prior written permission of the Publisher, or authorization through payment of the appropriate per-copy fee to the Copyright Clearance Center, Inc. 222 Rosewood Drive, Danvers, MA 01923, website www.copyright.com. Requests to the Publisher for permission should be addressed to the Permissions Department, John Wiley & Sons, Inc., 111 River Street, Hoboken, NJ 07030-5774, (201)748-6011, fax (201)748-6008, website http://www.wiley.com/go/permissions.

Evaluation copies are provided to qualified academics and professionals for review purposes only, for use in their courses during the next academic year. These copies are licensed and may not be sold or transferred to a third party. Upon completion of the review period, please return the evaluation copy to Wiley. Return instructions and a free of charge return mailing label are available at http://www.wiley.com/go/returnlabel. If you have chosen to adopt this textbook for use in your course, please accept this book as your complimentary desk copy. Outside of the United States, please contact your local sales representative.

Library of Congress Cataloging-in-Publication Data

Snustad, D. Peter.
Principles of genetics / D. Peter Snustad, Michael J. Simmons. — 6th ed. p. cm.
Includes index.
ISBN 978-0-470-90359-9 (cloth)
Binder-ready version ISBN 978-1-11812921-0
1. Genetics. I. Simmons, Michael J. II. Title.
QH430.S68 2012
576.5—dc23

Printed in the United States of America

10987654321

Dedications

To Judy, my wife and best friend. **D.P.S.**

To my family, especially to Benjamin. **M.J.S.**

About the Authors

D. Peter Snustad is a Professor Emeritus at the University of Minnesota, Twin Cities. He received his B.S. degree from the University of Minnesota and his M.S. and Ph.D. degrees from the University of California, Davis. He began his faculty career in the Department of Agronomy and Plant Genetics at Minnesota in 1965, became a charter member of the new Department of Genetics in 1966, and moved to the Department of Plant Biology in 2000. During his 43 years at Minnesota, he taught courses ranging from general biology to biochemical genetics. His initial research focused on the interactions between bacteriophage T4 and its host, *E. coli*. In the 1980s, his research switched to the cytoskeleton of *Arabidopsis* and the glutamine synthetase genes of corn. His honors include the Morse-Amoco and Dagley Memorial teaching awards and election to Fellow of the American Association for the Advancement of Science. A lifelong love of the Canadian wilderness has kept him in nearby Minnesota.

Michael J. Simmons is a Professor in the Department of Genetics, Cell Biology and Development at the University of Minnesota, Twin Cities. He received his B.A. degree in biology from St. Vincent College in Latrobe, Pennsylvania, and his M.S. and Ph.D. degrees in genetics from the University of Wisconsin, Madison. Dr. Simmons has taught a variety of courses, including genetics and population genetics. He has also mentored many students on research projects in his laboratory. Early in his career he received the Morse-Amoco teaching award from the University of Minnesota in recognition of his contributions to undergraduate education. Dr. Simmons's research focuses on the genetic significance of transposable elements in the genome of *Drosophila melanogaster*. He has served on advisory committees at the National Institutes of Health and was a member of the Editorial Board of the journal *Genetics* for 21 years. One of his favorite activities, figure skating, is especially compatible with the Minnesota climate. The science of genetics has been evolving rapidly. The DNA of genomes, even large ones, can now be analyzed in great detail; the functions of individual genes can be studied with an impressive array of techniques; and organisms can be changed genetically by introducing alien or altered genes into their genomes. The ways of teaching and learning genetics have also been changing. Electronic devices to access and transmit information are ubiquitous; engaging new media are being developed; and in many colleges and universities, classrooms are being redesigned to incorporate "active learning" strategies. This edition of *Principles of Genetics* has been created to recognize these scientific and educational advances.

Goals

Principles of Genetics balances new information with foundational material. In preparing this edition, we have been guided by four main goals:

- To focus on the basic principles of genetics by presenting the important concepts of classical, molecular, and population genetics carefully and thoroughly. We believe that an understanding of current advances in genetics and an appreciation for their practical significance must be based on a strong foundation. Furthermore, we believe that the breadth and depth of coverage in the different areas of genetics—classical, molecular, and population—must be balanced, and that the ever-growing mass of information in genetics must be organized by a sturdy—but flexible—framework of key concepts.
- To focus on the scientific process by showing how scientific concepts develop from observation and experimentation. Our book provides numerous examples to show how genetic principles have emerged from the work of different scientists. We emphasize that science is an ongoing process of observation, experimentation, and discovery.
- To focus on human genetics by incorporating human examples and showing the relevance of genetics to societal issues. Experience has shown us that students are keenly interested in the genetics of their own species. Because of this interest, they find it easier to comprehend complex concepts when these concepts are illustrated with human examples. Consequently, we have used human examples to illustrate genetic principles wherever possible. We have also included discussions of the Human Genome Project, human gene mapping, genetic disorders, gene therapy, and genetic counseling throughout the text. Issues such as genetic screening, DNA profiling, genetic engineering, cloning, stem cell research, and gene therapy have sparked vigorous debates about the social, legal, and ethical ramifications of genetics. We believe that it is important to involve students in discussions about these issues, and we hope that this textbook will provide students with the background to engage in such discussions thoughtfully.
- To focus on developing critical thinking skills by emphasizing the analysis of experimental data and problems. Genetics has always been a bit different from other disciplines in biology because of its heavy emphasis on problem solving. In this text, we have fleshed out the analytical nature of genetics in many ways—in the development of principles in classical genetics, in the discussion of experiments in molecular genetics, and in the presentation of calculations in population genetics. Throughout the book we have emphasized the integration of observational and experimental evidence with logical analysis in the development of key concepts. Each chapter has two sets of worked-out problems—the *Basic Exercises* section,

which contains simple problems that illustrate basic genetic analysis, and the *Testing Your Knowledge* section, which contains more complex problems that integrate different concepts and techniques. A set of *Questions and Problems* follows the worked-out problems so that students can enhance their understanding of the concepts in the chapter and develop their analytical skills. Another section, *Genomics on the Web*, poses issues that can be investigated by going to the National Center for Biotechnology Information web site. In this section, students can learn how to use the vast repository of genetic information that is accessible via that web site, and they can apply that information to specific problems. Each chapter also has a *Problem-Solving Skills* feature, which poses a problem, lists the pertinent facts and concepts, and then analyzes the problem and presents a solution. Finally, we have added a new feature, *Solve It*, to provide students with opportunities to test their understanding of concepts as they encounter them in the text. Each chapter poses two Solve It problems; step-by-step explanations of the answers are presented on the book's web site, some in video format.

Content and Organization of the Sixth Edition

The organization of this edition of *Principles of Genetics* is similar to that of the previous edition. However, the content has been sifted and winnowed to allow thoughtful updating. In selecting material to be included in this edition of *Principles of Genetics*, we have tried to be comprehensive but not encyclopedic.

The text comprises 24 chapters—one less than the previous edition. Chapters 1–2 introduce the science of genetics, basic features of cellular reproduction, and some of the model genetic organisms; Chapters 3–8 present the concepts of classical genetics and the basic procedures for the genetic analysis of microorganisms; Chapters 9–13 present the topics of molecular genetics, including DNA replication, transcription, translation, and mutation; Chapters 14–17 cover more advanced topics in molecular genetics and genomics; Chapters 18–21 deal with the regulation of gene expression and the genetic basis of development, immunity, and cancer; Chapters 22–24 present the concepts of quantitative, population, and evolutionary genetics.

As in previous editions, we have tried to create a text that can be adapted to different course formats. Many instructors prefer to present the topics in much the same way as we have, starting with classical genetics, progressing into molecular genetics, and finishing with quantitative, population, and evolutionary genetics. However this text is constructed so that teachers can present topics in different orders. They may, for example, begin with basic molecular genetics (Chapters 9–13), then present classical genetics (Chapters 3–8), progress to more advanced topics in molecular genetics (Chapters 14–21), and finish the course with quantitative, population, and evolutionary genetics (Chapters 22–24). Alternatively, they may wish to insert quantitative and population genetics between classical and molecular genetics.

Pedagogy of the Sixth Edition

The text includes special features designed to emphasize the relevance of the topics discussed, to facilitate the comprehension of important concepts, and to assist students in evaluating their grasp of these concepts.

- *Chapter-Opening Vignette.* Each chapter opens with a brief story that highlights the significance of the topics discussed in the chapter.
- *Chapter Outline.* The main sections of each chapter are conveniently listed on the chapter's first page.
- Section Summary. The content of each major section of text is briefly summarized at the beginning of that section. These opening summaries focus attention on the main ideas developed in a chapter.

- *Key Points.* These learning aids appear at the end of each major section in a chapter. They are designed to help students review for exams and to recapitulate the main ideas of the chapter.
- Focus On Boxes. Throughout the text, special topics are presented in separate *Focus On* boxes. The material in these boxes supports or develops concepts, techniques, or skills that have been introduced in the text of the chapter.
- **On the Cutting Edge Boxes.** The content of these boxes highlights exciting new developments in genetics—often the subject of ongoing research.
- **Problem-Solving Skills Boxes.** Each chapter contains a box that guides the student through the analysis and solution of a representative problem. We have chosen a problem that involves important material in the chapter. The box lists the facts and concepts that are relevant to the problem, and then explains how to obtain the solution. Ramifications of the problem and its analysis are discussed in the Student Companion site.
- **Solve It Boxes.** Each of these boxes poses a problem related to concepts students encounter as they read the text. The step-by-step solution to each of the problems is presented in the Student Companion site, and for selected problems, it is presented in video format. The two Solve It boxes in each chapter allow students to test their understanding of key concepts.
- **Basic Exercises.** At the end of each chapter we present several worked-out problems to reinforce each of the fundamental concepts developed in the chapter. These simple, one-step exercises are designed to illustrate basic genetic analysis or to emphasize important information.
- **Testing Your Knowledge.** Each chapter also has more complicated worked-out problems to help students hone their analytical and problem-solving skills. The problems in this section are designed to integrate different concepts and techniques. In the analysis of each problem, we walk the students through the solution step by step.
- **Questions and Problems.** Each chapter ends with a set of questions and problems of varying difficulty organized according to the sequence of topics in the chapter. The more difficult questions and problems have been designated with colored numbers. These sets of questions and problems provide students with the opportunity to enhance their understanding of the concepts covered in the chapter and to develop their analytical skills. Also, some of the questions and problems—called GO problems—have been selected for interactive solutions on the Student Companion site. The GO problems are designated with a special icon.
- *Genomics on the Web.* Information about genomes, genes, DNA sequences, mutant organisms, polypeptide sequences, biochemical pathways and evolutionary relationships is now freely available on an assortment of web sites. Researchers routinely access this information, and we believe that students should become familiar with it. To this end, we have incorporated a set of questions at the end of each chapter that can be answered by using the National Center for Biotechnology Information (NCBI) web site, which is sponsored by the U. S. National Institutes of Health.
- *Appendices.* Each Appendix presents technical material that is useful in genetic analysis.
- *Glossary.* This section of the book defines important terms. Students find it useful in clarifying topics and in preparing for exams.
- *Answers.* Answers to the odd-numbered Questions and Problems are given at the end of the text.

ONLINE RESOURCES

TEST BANK

The test bank is available on the Instructor Companion site and contains approximately 50 test questions per chapter. It is available online as MS Word files and as a computerized test bank. This easy-to-use test-generation program fully supports graphics, print tests, student answer sheets, and answer keys. The software's advanced features allow you to produce an exam to your exact specifications.

LECTURE POWERPOINT PRESENTATIONS

Highly visual lecture PowerPoint presentations are available for each chapter and help convey key concepts illustrated by imbedded text art. The presentations may be accessed on the Instructor Companion site.

PRE AND POST LECTURE ASSESSMENT

This assessment tool allows instructors to assign a quiz prior to lecture to assess student understanding and encourage reading, and following lecture to gauge improvement and weak areas. Two quizzes are provided for every chapter.

PERSONAL RESPONSE SYSTEM QUESTIONS

These questions are designed to provide readymade pop quizzes and to foster student discussion and debate in class. Available on the Instructor Companion site.

PRACTICE QUIZZES

Available on the Student Companion site, these quizzes contain 20 questions per chapter for students to quiz themselves and receive instant feedback.

MILESTONES IN GENETICS

The *Milestones* are available on the Student Companion site. Each of them explores a key development in genetics usually an experiment or a discovery. We cite the original papers that pertain to the subject of the *Milestone*, and we include two *Questions for Discussion* to provide students with an opportunity to investigate the current significance of the subject. These questions are suitable for cooperative learning activities in the classroom, or for reflective writing exercises that go beyond the technical aspects of genetic analysis.

SOLVE IT

Solve It boxes provide students with opportunities to test their understanding of concepts as they encounter them in the text. Each chapter poses two Solve It problems; step-by-step explanations of the answers are presented on the book's web site, some in video format. Students can view Camtasia videos, prepared by Dubear Kroening at the University of Wisconsin-Fox Valley. These tutorials enhance interactivity and hone problemsolving skills to give students the confidence they need to tackle complex problems in genetics.

ANIMATIONS

These animations illustrate key concepts from the text and aid students in grasping some of the most difficult concepts in genetics. Also included are animations that will give students a refresher in basic biology.

ANSWERS TO QUESTIONS AND PROBLEMS

Answers to odd-numbered Questions and Problems are located at the end of the text for easy access for students. Answers to all Questions and Problems in the text are available only to instructors on the Instructor Companion site.

ILLUSTRATIONS AND PHOTOS

All line illustrations and photos from *Principles of Genetics*, 6^{tb} *Edition*, are available on the Instructor Companion site in both jpeg files and PowerPoint format. Line illustrations are enhanced to provide the best presentation experience.

BOOK COMPANION WEB SITE

(www.wiley.com/college/snustad)

This text-specific web site provides students with additional resources and extends the chapters of the text to the resources of the World Wide Web. Resources include:

- For Students: practice quizzes covering key concepts for each chapter of the text, flashcards, and the Biology NewsFinder.
- For Instructors: Test Bank, PowerPoint Presentations, line art and photos in jpeg and PowerPoint formats, personal response system questions, and all answers to end-of-chapter Questions and Problems.

WILEY RESOURCE KIT

The Wiley Resource Kit fully integrates all content into easyto-navigate and customized modules that promote student engagement, learning, and success. All online resources are housed on this easy-to-navigate website, including:

Animated Solutions to the Solve It prompts in the text utilize Camtasia Studio software, a registered trademark of TechSmith Corporation, and they provide step-by-step solutions that appear as if they are written out by hand as an instructor voiceover explains each step.

GO Problem Tutorials give students the opportunity to observe a problem being worked out and then attempt to solve a similar problem. Working with GO problems will instill the confidence students need to succeed in the Genetics course.

Acknowledgments

As with previous editions, this edition of *Principles of Genetics* has been influenced by the genetics courses we teach. We thank our students for their constructive feedback on both content and pedagogy, and we thank our colleagues at the University of Minnesota for sharing their knowledge and expertise. Genetics professors at other institutions also provided many helpful suggestions. In particular, we acknowledge the help of the following reviewers:

6TH EDITION REVIEWERS

Ann Aguano, Manhattan Marymount College; Mary A. Bedell, University of Georgia; Jonathan Clark, Weber State University; Robert Fowler, San Jose State University; Cheryl Hertz, Loyola Marymount University; Shawn Kaeppler, University of Wisconsin; Todd Kelson, Brigham Young University – Idaho; Richard D. Noyes, University of Central Arkansas; Maria E. Orive, University of Kansas; Rongsun Pu, Kean University

REVIEWERS OF PREVIOUS EDITIONS

Michelle Boissere, Xavier University of Louisiana; Stephen P. Bush, Coastal Carolina University; Sarah Crawford, Southern Connecticut State University; Xiongbin Lu, University of South Carolina – Columbia; Valery N. Soyfer, George Mason University; David Starkey, University of Central Arkansas; Frans Tax, University of Arizona; Tzvi Tzfira, University of Michigan; Harald Vaessin, The Ohio State University – Columbus; Sarah VanVickle-Chavez, Washington University in St. Louis; Willem Vermerris, University of Florida; Alan S. Waldman, University of South Carolina – Columbia Many people contributed to the development and production of this edition. Kevin Witt, Senior Editor, and Michael Palumbo, Assistant Editor, initiated the project and provided ideas about some of the text's features. Dr. Pamela Marshall of Arizona State University suggested many ways in which the previous edition could be improved, and a panel of genetics teachers thoughtfully commented on her suggestions. The panel's members were: Anna Aguano, Manhattan Marymount College; Robert Fowler, San Jose State University; Jane Glazebrook, University of Minnesota; Shawn Kaeppler, University of Wisconsin; Todd Kelson, Brigham Young University – Idaho; and Dwayne A. Wise, Mississippi State University. We are grateful for all the input from these experienced teachers of genetics.

Jennifer Dearden and Lauren Morris helped with many of the logistical details in preparing this edition, and Lisa Passmore researched and obtained many new photographs. Jennifer MacMillan, Senior Photo Editor, skillfully coordinated the entire photo program. We are grateful for all their contributions. We thank Maureen Eide, Senior Designer, for creating a fresh text layout, and we thank Precision Graphics and Aptara for executing the illustrations. Elizabeth Swain, Senior Production Editor, superbly coordinated the production of this edition, Betty Pessagno faithfully copyedited the manuscript, Lilian Brady did the final proofreading, and Stephen Ingle prepared the index. We deeply appreciate the excellent work of all these people. We also thank Clay Stone, Executive Marketing Manager, for helping to get this edition into the hands of prospective users. With an eye toward the next edition, we encourage students, teaching assistants, instructors, and other readers to send us comments on this edition in care of Jennifer Dearden at John Wiley & Sons, Inc., 111 River Street, Hoboken, NJ, 07030.

Contents

CHAPTER 1

The Science of Genetics 1

The Personal Genome 1

An Invitation 2

Three Great Milestones in Genetics 2

MENDEL: GENES AND THE RULES OF INHERITANCE 2 WATSON AND CRICK: THE STRUCTURE OF DNA 3 THE HUMAN GENOME PROJECT: SEQUENCING DNA AND CATALOGUING GENES 4

DNA as the Genetic Material 6

DNA REPLICATION: PROPAGATING GENETIC INFORMATION 6 GENE EXPRESSION: USING GENETIC INFORMATION 7 MUTATION: CHANGING GENETIC INFORMATION 9

Genetics and Evolution 10

Levels of Genetic Analysis 11

CLASSICAL GENETICS 11 MOLECULAR GENETICS 11 POPULATION GENETICS 12

Genetics in the World: Applications of Genetics to Human Endeavors *12*

GENETICS IN AGRICULTURE 12 GENETICS IN MEDICINE 14 GENETICS IN SOCIETY 15

CHAPTER 2

Cellular Reproduction 18

Dolly 18

Cells and Chromosomes 19

THE CELLULAR ENVIRONMENT 19 PROKARYOTIC AND EUKARYOTIC CELLS 20 CHROMOSOMES: WHERE GENES ARE LOCATED 20 CELL DIVISION 23

Mitosis 24

Meiosis 27

SOLVE IT How Much DNA in Human Meiotic Cells 27

MEIOSIS I 27 MEIOSIS II AND THE OUTCOMES OF MEIOSIS 31

SOLVE IT How Many Chromosome Combinations in Sperm *31*

Life Cycles of Some Model Genetic Organisms *32*

SACCHAROMYCES CEREVISIAE, BAKER'S YEAST 32 ARABIDOPSIS THALIANA, A FAST-GROWING PLANT 33 MUS MUSCULUS, THE MOUSE 34

PROBLEM-SOLVING SKILLS Counting Chromosomes and Chromatids *36*

CHAPTER 3

Mendelism: The Basic Principles of Inheritance 40

The Birth of Genetics: A Scientific Revolution 40

Mendel's Study of Heredity 41

MENDEL'S EXPERIMENTAL ORGANISM, THE GARDEN PEA 41 MONOHYBRID CROSSES: THE PRINCIPLES OF DOMINANCE AND SEGREGATION 42

DIHYBRID CROSSES: THE PRINCIPLE OF INDEPENDENT ASSORTMENT 44

Applications of Mendel's Principles 46

THE PUNNETT SQUARE METHOD 46 THE FORKED-LINE METHOD 46 THE PROBABILITY METHOD 47

SOLVE IT Using Probabilities in a Genetic Problem *48*

Testing Genetic Hypotheses 48

THE CHI-SQUARE TEST 50

SOLVE IT Using the Chi-Square Test 52

Mendelian Principles in Human Genetics 52

PEDIGREES 53 MENDELIAN SEGREGATION IN HUMAN FAMILIES 54 GENETIC COUNSELING 54

PROBLEM-SOLVING SKILLS Making Predictions from Pedigrees 56

CHAPTER 4

Extensions of Mendelism 62

Genetics Grows Beyond Mendel's Monastery Garden 62

Allelic Variation and Gene Function 63

INCOMPLETE DOMINANCE AND CODOMINANCE 63 MULTIPLE ALLELES 64 ALLELIC SERIES 65 TESTING GENE MUTATIONS FOR ALLELISM 65

SOLVE IT The Test for Allelism 66

VARIATION AMONG THE EFFECTS OF MUTATIONS 66 GENES FUNCTION TO PRODUCE POLYPEPTIDES 67

FOCUS ON Genetic Symbols 68

WHY ARE SOME MUTATIONS DOMINANT AND OTHERS RECESSIVE? 68

Gene Action: From Genotype to Phenotype 70

INFLUENCE OF THE ENVIRONMENT 70 ENVIRONMENTAL EFFECTS ON THE EXPRESSION OF HUMAN GENES 70 PENETRANCE AND EXPRESSIVITY 71 GENE INTERACTIONS 72 EPISTASIS 72 PLEIOTROPY 75

PROBLEM-SOLVING SKILLS Going from Pathways to Phenotypic Ratios *76*

Inbreeding: Another Look at Pedigrees 77

THE EFFECTS OF INBREEDING 77 GENETIC ANALYSIS OF INBREEDING 78

SOLVE IT Compound Inbreeding 81

MEASURING GENETIC RELATIONSHIPS 82

CHAPTER 5

The Chromosomal Basis of Mendelism 89

Sex, Chromosomes, and Genes 89

Chromosomes 90

CHROMOSOME NUMBER 90 SEX CHROMOSOMES 90

The Chromosome Theory of Heredity 92

EXPERIMENTAL EVIDENCE LINKING THE INHERITANCE OF GENES TO CHROMOSOMES 92

NONDISJUNCTION AS PROOF OF THE CHROMOSOME THEORY 93 THE CHROMOSOMAL BASIS OF MENDEL'S PRINCIPLES OF SEGREGATION AND INDEPENDENT ASSORTMENT 95

SOLVE IT Sex Chromosome Nondisjunction 95

PROBLEM-SOLVING SKILLS Tracking X-Linked and Autosomal Inheritance 97

Sex-Linked Genes in Humans 98

HEMOPHILIA, AN X-LINKED BLOOD-CLOTTING DISORDER 98 COLOR BLINDNESS, AN X-LINKED VISION DISORDER 98 GENES ON THE HUMAN Y CHROMOSOME 100

SOLVE IT Calculating the Risk for Hemophilia *100* GENES ON BOTH THE X AND Y CHROMOSOMES *100*

Sex Chromosomes and Sex Determination *100*

SEX DETERMINATION IN HUMANS 101 SEX DETERMINATION IN DROSOPHILA 102 SEX DETERMINATION IN OTHER ANIMALS 102

Dosage Compensation of X-Linked Genes *104*

HYPERACTIVATION OF X-LINKED GENES IN MALE DROSOPHILA 104 INACTIVATION OF X-LINKED GENES IN FEMALE MAMMALS 104

CHAPTER **6**

Variation in Chromosome Number and Structure 110

Chromosomes, Agriculture, and Civilization 110 Cytological Techniques 111 ANALYSIS OF MITOTIC CHROMOSOMES 111 THE HUMAN KARYOTYPE 113 CYTOGENETIC VARIATION: AN OVERVIEW 114

Polyploidy 115

STERILE POLYPLOIDS 115 FERTILE POLYPLOIDS 116

SOLVE IT Chromosome Pairing in Polyploids *117* TISSUE-SPECIFIC POLYPLOIDY AND POLYTENY *117*

Aneuploidy 119

TRISOMY IN HUMANS 120 MONOSOMY 121

FOCUS ON Amniocentesis and Chorionic Biopsy 123

PROBLEM-SOLVING SKILLS Tracing Sex Chromosome Nondisjunction 124

DELETIONS AND DUPLICATIONS OF CHROMOSOME SEGMENTS 124

Rearrangements of Chromosome Structure *126*

INVERSIONS 126 TRANSLOCATIONS 127 COMPOUND CHROMOSOMES AND ROBERTSONIAN TRANSLOCATIONS 128

SOLVE IT Pollen Abortion in Translocation Heterozygotes *129*

CHAPTER 7

Linkage, Crossing Over, and Chromosome Mapping in Eukaryotes 135

The World's First Chromosome Map 135

Linkage, Recombination, and Crossing Over 136

EARLY EVIDENCE FOR LINKAGE AND RECOMBINATION 136 CROSSING OVER AS THE PHYSICAL BASIS OF

RECOMBINATION 138 EVIDENCE THAT CROSSING OVER CAUSES

RECOMBINATION 139

CHIASMATA AND THE TIME OF CROSSING OVER 140

Chromosome Mapping 141

CROSSING OVER AS A MEASURE OF GENETIC DISTANCE 141 RECOMBINATION MAPPING WITH A TWO-POINT TESTCROSS 141 RECOMBINATION MAPPING WITH A THREE-POINT TESTCROSS 142

SOLVE IT Mapping Two Genes with Testcross Data 143

PROBLEM-SOLVING SKILLS Using a Genetic Map to Predict the Outcome of a Cross 146

RECOMBINATION FREQUENCY AND GENETIC MAP DISTANCE 146

Cytogenetic Mapping 148

LOCALIZING GENES USING DELETIONS AND DUPLICATIONS 148 GENETIC DISTANCE AND PHYSICAL DISTANCE 149

SOLVE IT Cytological Mapping of a *Drosophila* Gene 150

Linkage Analysis in Humans 150

Recombination and Evolution 153

EVOLUTIONARY SIGNIFICANCE OF RECOMBINATION 153 SUPPRESSION OF RECOMBINATION BY INVERSIONS 153 GENETIC CONTROL OF RECOMBINATION 155

CHAPTER 8

The Genetics of Bacteria and Their Viruses 163

Multi-Drug-Resistant Bacteria: A Ticking Timebomb? 163

Viruses and Bacteria in Genetics 164

The Genetics of Viruses 165

BACTERIOPHAGE T4 165 BACTERIOPHAGE LAMBDA 166

The Genetics of Bacteria 169

MUTANT GENES IN BACTERIA 170 UNIDIRECTIONAL GENE TRANSFER IN BACTERIA 171

Mechanisms of Genetic Exchange in Bacteria *172*

TRANSFORMATION 173 CONJUGATION 175 PLASMIDS AND EPISOMES 179

PROBLEM-SOLVING SKILLS Mapping Genes Using Conjugation Data 180

F' FACTORS AND SEXDUCTION 181 TRANSDUCTION 182

SOLVE IT How Can You Map Closely Linked Genes Using Partial Diploids *183*

The Evolutionary Significance of Genetic Exchange in Bacteria *186*

SOLVE IT How Do Bacterial Genomes Evolve? 186

ON THE CUTTING EDGE Antibiotic-Resistant Bacteria *186*

CHAPTER 9

DNA and the Molecular Structure of Chromosomes 192

Discovery of Nuclein 192

Functions of the Genetic Material 193

Proof That Genetic Information Is Stored in DNA 193

PROOF THAT DNA MEDIATES TRANSFORMATION 194 PROOF THAT DNA CARRIES THE GENETIC INFORMATION IN BACTERIOPHAGE T2 195

PROOF THAT RNA STORES THE GENETIC INFORMATION IN SOME VIRUSES 197

The Structures of DNA and RNA 197

NATURE OF THE CHEMICAL SUBUNITS IN DNA AND RNA 198 DNA STRUCTURE: THE DOUBLE HELIX 199

PROBLEM-SOLVING SKILLS Calculating Base

Content in DNA 202

DNA STRUCTURE: ALTERNATE FORMS OF THE DOUBLE HELIX 203

SOLVE IT What Are Some Important Features of Double-Stranded DNA? 203

DNA STRUCTURE: NEGATIVE SUPERCOILS IN VIVO 204

Chromosome Structure in Prokaryotes and Viruses 205

Chromosome Structure in Eukaryotes 207

CHEMICAL COMPOSITION OF EUKARYOTIC CHROMOSOMES 207 ONE LARGE DNA MOLECULE PER CHROMOSOME 208 THREE LEVELS OF DNA PACKAGING IN EUKARYOTIC CHROMOSOMES 208

SOLVE IT How Many Nucleosomes in One Human X Chromosome *210*

CENTROMERES AND TELOMERES 211

REPEATED DNA SEQUENCES 214

ON THE CUTTING EDGE The 1000 Genomes Project 216

CHAPTER 10

Replication of DNA and Chromosomes 220

Monozygotic Twins: Are They Identical? 220

Basic Features of DNA Replication In Vivo 221

SEMICONSERVATIVE REPLICATION 221

SOLVE IT Understanding the Semiconservative Replication of DNA 224

UNIQUE ORIGINS OF REPLICATION 224

PROBLEM-SOLVING SKILLS Predicting Patterns of ³H Labeling in Chromosomes 226

VISUALIZATION OF REPLICATION FORKS BY AUTORADIOGRAPHY 227 BIDIRECTIONAL REPLICATION 228

DNA Replication in Prokaryotes 231

FOCUS ON DNA Synthesis In Vitro 231

CONTINUOUS SYNTHESIS OF ONE STRAND; DISCONTINUOUS SYNTHESIS OF THE OTHER STRAND 232

COVALENT CLOSURE OF NICKS IN DNA BY DNA LIGASE 232

INITIATION OF DNA REPLICATION 234

INITIATION OF DNA CHAINS WITH RNA PRIMERS 234

UNWINDING DNA WITH HELICASES, DNA-BINDING PROTEINS, AND TOPOISOMERASES 236 MULTIPLE DNA POLYMERASES AND PROOFREADING 238 THE PRIMOSOME AND THE REPLISOME 242 ROLLING-CIRCLE REPLICATION 243

Unique Aspects of Eukaryotic Chromosome Replication *244*

THE CELL CYCLE 245 MULTIPLE REPLICONS PER CHROMOSOME 245

SOLVE IT Understanding Replication of the Human X Chromosome 246

TWO OR MORE DNA POLYMERASES AT A SINGLE REPLICATION FORK 246

DUPLICATION OF NUCLEOSOMES AT REPLICATION FORKS 247 TELOMERASE: REPLICATION OF CHROMOSOME TERMINI 248 TELOMERE LENGTH AND AGING IN HUMANS 249

CHAPTER 11

Transcription and RNA Processing 256

Storage and Transmission of Information with Simple Codes 256

Transfer of Genetic Information: The Central Dogma *257*

TRANSCRIPTION AND TRANSLATION 257 FIVE TYPES OF RNA MOLECULES 258

The Process of Gene Expression 259

AN mRNA INTERMEDIARY 259 GENERAL FEATURES OF RNA SYNTHESIS 261

PROBLEM-SOLVING SKILLS Distinguishing RNAs Transcribed from Viral and Host DNAs 262

Transcription in Prokaryotes 263

RNA POLYMERASES: COMPLEX ENZYMES 263 INITIATION OF RNA CHAINS 264 ELONGATION OF RNA CHAINS 264 TERMINATION OF RNA CHAINS 265 CONCURRENT TRANSCRIPTION, TRANSLATION, AND mRNA DEGRADATION 266

Transcription and RNA Processing in Eukaryotes *267*

FIVE RNA POLYMERASES/FIVE SETS OF GENES 267

ON THE CUTTING EDGE Chromatin Remodeling and Gene Expression *269*

INITIATION OF RNA CHAINS 270

SOLVE IT Initiation of Transcription by RNA Polymerase II in Eukaryotes *270* RNA CHAIN ELONGATION AND THE ADDITION OF 5' METHYL GUANOSINE CAPS 271

TERMINATION BY CHAIN CLEAVAGE AND THE ADDITION OF 3' POLY(A) TAILS 272

SOLVE IT Formation of the 3'-Terminus of an RNA Polymerase II Transcript *273*

RNA EDITING: ALTERING THE INFORMATION CONTENT OF mRNA MOLECULES 273

Interrupted Genes in Eukaryotes: Exons and Introns 274

SOME VERY LARGE EUKARYOTIC GENES 276 INTRONS: BIOLOGICAL SIGNIFICANCE? 276

Removal of Intron Sequences by RNA Splicing 277

tRNA PRECURSOR SPLICING: UNIQUE NUCLEASE AND LIGASE ACTIVITIES 278

AUTOCATALYTIC SPLICING 278

PRE-mRNA SPLICING: snRNAs, snRNPs, AND THE SPLICEOSOME 279

CHAPTER 12

Translation and the Genetic Code 285

Sickle-Cell Anemia: Devastating Effects of a Single Base-Pair Change 285

Protein Structure 286

POLYPEPTIDES: TWENTY DIFFERENT AMINO ACID SUBUNITS 286

PROTEINS: COMPLEX THREE-DIMENSIONAL STRUCTURES 287

One Gene—One Colinear Polypeptide 289

BEADLE AND TATUM: ONE GENE—ONE ENZYME 289 COLINEARITY BEWEEN THE CODING SEQUENCE OF A GENE AND ITS POLYPEPTIDE PRODUCT 291

Protein Synthesis: Translation 293

OVERVIEW OF PROTEIN SYNTHESIS 293 COMPONENTS REQUIRED FOR PROTEIN SYNTHESIS: RIBOSOMES 294

COMPONENTS REQUIRED FOR PROTEIN SYNTHESIS: TRANSFER RNAs 296

TRANSLATION: THE SYNTHESIS OF POLYPEPTIDES USING mRNA TEMPLATES 298

SOLVE IT Control of Translation in Eukaryotes 304

The Genetic Code 306

PROPERTIES OF THE GENETIC CODE: AN OVERVIEW 306 THREE NUCLEOTIDES PER CODON 306 DECIPHERING THE CODE 307 INITIATION AND TERMINATION CODONS 309 A DEGENERATE AND ORDERED CODE 310

PROBLEM-SOLVING SKILLS Predicting Amino Acid

Substitutions Induced by Mutagens 311

A NEARLY UNIVERSAL CODE 312

Codon-tRNA Interactions 312

RECOGNITION OF CODONS BY tRNAs: THE WOBBLE HYPOTHESIS 312

SUPPRESSOR MUTATIONS THAT PRODUCE tRNAs WITH ALTERED CODON RECOGNITION 313

SOLVE IT Effects of Base-Pair Substitutions in the Coding Region of the *HBB* Gene *314*

ON THE CUTTING EDGE Selenocysteine, the 21st Amino Acid *315*

CHAPTER 13

Mutation, DNA Repair, and Recombination 320

Xeroderma Pigmentosum: Defective Repair of Damaged DNA in Humans 320

Mutation: Source of the Genetic Variability Required for Evolution *321*

The Molecular Basis of Mutation 321

SOLVE IT Nucleotide-Pair Substitutions in the Human *HBB* Gene *323*

INDUCED MUTATIONS 324 MUTATIONS INDUCED BY CHEMICALS 326 MUTATIONS INDUCED BY RADIATION 328

PROBLEM-SOLVING SKILLS Predicting Amino Acid Changes Induced by Chemical Mutagens 329

MUTATIONS INDUCED BY TRANSPOSABLE GENETIC ELEMENTS 331

EXPANDING TRINUCLEOTIDE REPEATS AND INHERITED HUMAN DISEASES 331

Mutation: Basic Features of the Process 332

MUTATION: SOMATIC OR GERMINAL 332 MUTATION: SPONTANEOUS OR INDUCED 333 MUTATION: USUALLY A RANDOM, NONADAPTIVE PROCESS 333 MUTATION: A REVERSIBLE PROCESS 335

Mutation: Phenotypic Effects 337

MUTATIONS WITH PHENOTYPIC EFECTS: USUALLY DELETERIOUS AND RECESSIVE 337 EFFECTS OF MUTATIONS IN HUMAN GLOBIN GENES 338 MUTATION IN HUMANS: BLOCKS IN METABOLIC PATHWAYS 339

ON THE CUTTING EDGE Screening Eight-cell

Pre-embryos for Tay-Sachs Muitations 340

CONDITIONAL LETHAL MUTATIONS: POWERFUL TOOLS FOR GENETIC STUDIES 340

Assigning Mutations to Genes by the Complementation Test *342*

SOLVE IT How Can You Assign Mutations to Genes? 344

Screening Chemicals for Mutagenicity: The Ames Test *346*

DNA Repair Mechanisms 348

LIGHT-DEPENDENT REPAIR 348 EXCISION REPAIR 348 OTHER DNA REPAIR MECHANISMS 349

Inherited Human Diseases with Defects in DNA Repair *351*

DNA Recombination Mechanisms 354

RECOMBINATION: CLEAVAGE AND REJOINING OF DNA MOLECULES 354

GENE CONVERSION: DNA REPAIR SYNTHESIS ASSOCIATED WITH RECOMBINATION 356

CHAPTER 14

The Techniques of Molecular Genetics 366

Treatment of Pituitary Dwarfism with Human Growth Hormone 366

Basic Techniques Used to Identify, Amplify, and Clone Genes *367*

SOLVE IT How Many *Not*l Restriction Fragments in Chimpanzee DNA? *368*

THE DISCOVERY OF RESTRICTION ENDONUCLEASES 368

- THE PRODUCTION OF RECOMBINANT DNA MOLECULES IN VITRO 371
- AMPLIFICATION OF RECOMBINANT DNA MOLECULES IN CLONING VECTORS 372
- CLONING LARGE GENES AND SEGMENTS OF GENOMES IN BACs, PACs, AND YACs 374
- AMPLIFICATION OF DNA SEQUENCES BY THE POLYMERASE CHAIN REACTION (PCR) 374

Construction and Screening of DNA Libraries 377

CONSTRUCTION OF GENOMIC LIBRARIES 377 CONSTRUCTION OF cDNA LIBRARIES 378 SCREENING DNA LIBRARIES FOR GENES OF INTEREST 378 **SOLVE IT** How Can You Clone a Specific *Not* Restriction Fragment from the Orangutan Genome? *380*

The Molecular Analysis of DNA, RNA, and Protein *380*

ANALYSIS OF DNAs BY SOUTHERN BLOT HYBRIDIZATIONS 381 ANALYSIS OF RNAs BY NORTHERN BLOT HYBRIDIZATIONS 382

FOCUS ON Detection of a Mutant Gene Causing Cystic Fibrosis 383

ANALYSIS OF RNAs BY REVERSE TRANSRIPTASE-PCR (RT-PCR) 384

ANALYSIS OF PROTEINS BY WESTERN BLOT TECHNIQUES 384

The Molecular Analysis of Genes and Chromosomes *386*

PHYSICAL MAPS OF DNA MOLECULES BASED ON RESTRICTION ENZYME CLEAVAGE SITES 386

NUCLEOTIDE SEQUENCES OF GENES AND CHROMOSOMES 387

PROBLEM-SOLVING SKILLS Determining the Nucleotide Sequences of Genetic Elements 390

CHAPTER 15

Genomics 397

The Neanderthal Genome: What It Reveals about Our Ancestors 397

FOCUS ON GenBank 400

Genomics: An Overview 402

Correlated Genetic, Cytological, and Physical Maps of Chromosomes *402*

RESTRICTION FRAGMENT-LENGTH POLYMORPHISM (RFLP) AND SHORT TANDEM REPEAT (STR) MAPS 403 CYTOGENETIC MAPS 405 PHYSICAL MAPS AND CLONE BANKS 405

Map Position-Based Cloning of Genes 407

CHROMOSOME WALKS AND JUMPS 408

The Human Genome Project 409

MAPPING THE HUMAN GENOME 409 SEQUENCING THE HUMAN GENOME 410 THE HUMAN HAPMAP PROJECT 414

RNA and Protein Assays of Genome Function *415*

EXPRESSED SEQUENCES 416 MICROARRAYS AND GENE CHIPS 416

THE GREEN FLUORESCENT PROTEIN AS A REPORTER OF PROTEIN SYNTHESIS 419

Comparative Genomics 420

BIOINFORMATICS 421

PROBLEM-SOLVING SKILLS Using Bioinformatics to Investigate DNA Sequences 422

PROKARYOTIC GENOMES 424

A LIVING BACTERIUM WITH A CHEMICALLY SYNTHESIZED GENOME 425

THE GENOMES OF CHLOROPLASTS AND MITOCHONDRIA 426

SOLVE IT What Do We Know about the Mitochondrial Genome of the Extinct Woolly Mammoth? *429*

EUKARYOTIC GENOMES 429

SOLVE IT What Can You Learn about DNA Sequences using Bioinformatics? *431*

GENOME EVOLUTION IN THE CEREAL GRASSES 431 GENOME EVOLUTION IN MAMMALS 432

CHAPTER 16

Applications of Molecular Genetics 439

Gene Therapy Improves Sight in Child with Congenital Blindness 439

Use of Recombinant DNA Technology to Identify Human Genes and Diagnose Human Diseases 440

HUNTINGTON'S DISEASE 440

FOCUS ON Fragile X Syndrome and Expanded Trinucleotide Repeats 443

PROBLEM-SOLVING SKILLS Testing for Mutant Alleles that Cause Fragile X Mental

Retardation 445

CYSTIC FIBROSIS 445

Molecular Diagnosis of Human Diseases 448

Human Gene Therapy 450

DNA Profiling 455

PATERNITY TESTS 459

SOLVE IT How Can DNA Profiles Be Used to Establish Identity? *459*

FORENSIC APPLICATIONS 459

Production of Eukaryotic Proteins in Bacteria *461*

HUMAN GROWTH HORMONE 461 PROTEINS WITH INDUSTRIAL APPLICATIONS 462

Transgenic Plants and Animals 463

TRANSGENIC ANIMALS: MICROINJECTION OF DNA INTO FERTILIZED EGGS AND TRANSFECTION OF EMBRYONIC STEM CELLS 463

TRANSGENIC PLANTS: THE TI PLASMID OF AGROBACTERIUM TUMEFACIENS 464

Reverse Genetics: Dissecting Biological Processes by Inhibiting Gene Expression 467

KNOCKOUT MUTATIONS IN THE MOUSE 467 T-DNA AND TRANSPOSON INSERTIONS 469 RNA INTERFERENCE 471

SOLVE IT How Might RNA Interference Be Used to Treat Burkitt's Lymphoma? 471

CHAPTER 17

Transposable Genetic Elements 477

Maize: A Staple Crop with a Cultural Heritage 477

Transposable Elements: An Overview 478

Transposable Elements in Bacteria 479

IS ELEMENTS 479

SOLVE IT Accumulating Drug-Resistance Genes 481 COMPOSITE TRANSPOSONS 481 THE Tn3 ELEMENT 481

Cut-and-Paste Transposons in Eukaryotes *483*

Ac AND Ds ELEMENTS IN MAIZE 483 P ELEMENTS AND HYBRID DYSGENESIS IN DROSOPHILA 485

PROBLEM-SOLVING SKILLS Analyzing Transposon Activity in Maize 486

ON THE CUTTING EDGE Small RNAs Repress *P* Element Activity 487

Retroviruses and Retrotransposons 488

RETROVIRUSES 488 RETROVIRUSLIKE ELEMENTS 492 RETROPOSONS 493

Transposable Elements in Humans 494

The Genetic and Evolutionary Significance of Transposable Elements 496

TRANSPOSONS AS MUTAGENS 496 GENETIC TRANSFORMATION WITH TRANSPOSONS 496

SOLVE IT Transposon-Mediated Chromosome Rearrangements *498*

TRANSPOSONS AND GENOME ORGANIZATION 498

CHAPTER 18

Regulation of Gene Expression in Prokaryotes 504

D'Hérelle's Dream of Treating Dysentery in Humans by Phage Therapy 504

Constitutive, Inducible, and Repressible Gene Expression *506*

Positive and Negative Control of Gene Expression 507

Operons: Coordinately Regulated Units of Gene Expression *509*

The Lactose Operon in *E. coli*: Induction and Catabolite Repression *511*

INDUCTION 513

SOLVE IT Constitutive Mutations in the *E. coli lac* Operon *513*

CATABOLITE REPRESSION 514

PROBLEM SOLVING SKILLS Testing Your

Understanding of the *lac* Operon *516* PROTEIN-DNA INTERACTIONS THAT CONTROL TRANSCRIPTION OF THE *LAC* OPERON *517*

The Tryptophan Operon in *E. coli*: Repression and Attenuation *519*

REPRESSION 519 **ATTENUATION** 520

SOLVE IT Regulation of the Histidine Operon of Salmonella typhimurium 522

ON THE CUTTING EDGE The Lysine Riboswitch 524

Translational Control of Gene Expression 525

Posttranslational Regulatory Mechanisms *526*

CHAPTER 19

Regulation of Gene Expression in Eukaryotes 531

African Trypanosomes: A Wardrobe of Molecular Disguises 531

Ways of Regulating Eukaryotic Gene Expression: An Overview *532*

DIMENSIONS OF EUKARYOTIC GENE REGULATION 532 CONTROLLED TRANSCRIPTION OF DNA 532 ALTERNATE SPLICING OF RNA 533

SOLVE IT Counting mRNAs 533

CYTOPLASMIC CONTROL OF MESSENGER RNA STABILITY 533

Induction of Transcriptional Activity by Environmental and Biological Factors 534

TEMPERATURE: THE HEAT-SHOCK GENES 535 SIGNAL MOLECULES: GENES THAT RESPOND TO HORMONES 535

Molecular Control of Transcription in Eukaryotes *537*

DNA SEQUENCES INVOLVED IN THE CONTROL OF TRANSCRIPTION 537

PROTEINS INVOLVED IN THE CONTROL OF TRANSCRIPTION: TRANSCRIPTION FACTORS 538

PROBLEM-SOLVING SKILLS Defining the

Sequences Required for a Gene's Expression 539

Posttranscriptional Regulation of Gene Expression by RNA Interference *541*

RNAi PATHWAYS 541 SOURCES OF SHORT INTERFERING RNAs AND MicroRNAs 543

Gene Expression and Chromatin Organization *544*

SOLVE IT Using RNAi in Cell Research 545

EUCHROMATIN AND HETEROCHROMATIN 545 MOLECULAR ORGANIZATION OF TRANSCRIPTIONALLY ACTIVE DNA 545 CHROMATIN REMODELING 546 DNA METHYLATION 547 IMPRINTING 548

ON THE CUTTING EDGE The Epigenetics of Twins 549

Activation and Inactivation of Whole Chromosomes 550

INACTIVATION OF X CHROMOSOMES IN MAMMALS 551 HYPERACTIVATION OF X CHROMOSOMES IN DROSOPHILA 552 HYPOACTIVATION OF X CHROMOSOMES IN CAENORHABDITIS 553

CHAPTER 20

The Genetic Control of Animal Development 558

Stem-Cell Therapy 558

A Genetic Perspective on Development 559

Maternal Gene Activity in Development 561

MATERNAL-EFFECT GENES 561

SOLVE IT A Maternal-Effect Mutation in the *cinnamon* Gene 562

DETERMINATION OF THE DORSAL-VENTRAL AND ANTERIOR-POSTERIOR AXES 562

Zygotic Gene Activity in Development 565

BODY SEGMENTATION 565 ORGAN FORMATION 567 SPECIFICATION OF CELL TYPES 569

SOLVE IT Cave Blindness 569

PROBLEM-SOLVING SKILLS The Effects of Mutations during Eye Development *571*

Genetic Analysis of Development in Vertebrates *571*

VERTEBRATE HOMOLOGUES OF INVERTEBRATE GENES 571 THE MOUSE: RANDOM INSERTION MUTATIONS AND GENE-SPECIFIC KNOCKOUT MUTATIONS 572 STUDIES WITH MAMMALIAN STEM CELLS 573 REPRODUCTIVE CLONING 574

GENETIC CHANGES IN THE DIFFERENTIATION OF VERTEBRATE IMMUNE CELLS 575

CHAPTER 21

The Genetic Basis of Cancer 581

A Molecular Family Connection 581

Cancer: A Genetic Disease 582

THE MANY FORMS OF CANCER 582 CANCER AND THE CELL CYCLE 583 CANCER AND PROGRAMMED CELL DEATH 584 A GENETIC BASIS FOR CANCER 584

Oncogenes 585

TUMOR-INDUCING RETROVIRUSES AND VIRAL ONCOGENES 585

SOLVE IT The *v*-*erbB* and *v*-*fms* Viral Oncogenes 586

CELLULAR HOMOLOGUES OF VIRAL ONCOGENES: THE PROTO-ONCOGENES 586 MUTANT CELLULAR ONCOGENES AND CANCER 587 CHROMOSOME REARRANGEMENTS AND CANCER 589

Tumor Suppressor Genes 590

INHERITED CANCERS AND KNUDSON'S TWO-HIT HYPOTHESIS 590

PROBLEM-SOLVING SKILLS Estimating Mutation Rates in Retinoblastoma *593*

CELLULAR ROLES OF TUMOR SUPPRESSOR PROTEINS 593 pRB 593 p53 595

SOLVE IT Downstream of p53 595

pAPC 597

phMSH2 598 pBRCA1 and pBRCA2 599 FOCUS ON Cancer and Genetic Counseling 600

Genetic Pathways to Cancer 600

CHAPTER 22

Inheritance of Complex Traits 607

Cardiovascular Disease: A Combination of Genetic and Environmental Factors 607

Complex Traits 608

QUANTIFYING COMPLEX TRAITS 608 GENETIC AND ENVIRONMENTAL FACTORS INFLUENCE QUANTITATIVE TRAITS 608

MULTIPLE GENES INFLUENCE QUANTITATIVE TRAITS 608

THRESHOLD TRAITS 610

Statistics of Quantitative Genetics 611

FREQUENCY DISTRIBUTIONS 611 THE MEAN AND THE MODAL CLASS 612 THE VARIANCE AND THE STANDARD DEVIATION 612

Analysis of Quantitative Traits 613

THE MULTIPLE FACTOR HYPOTHESIS 614 PARTITIONING THE PHENOTYPIC VARIANCE 614

SOLVE IT Estimating Genetic and Environmental Variance Components 615

BROAD-SENSE HERITABILITY 615 NARROW-SENSE HERITABILITY 616 PREDICTING PHENOTYPES 617

SOLVE IT Using the Narrow-Sense Heritability *618*

ARTIFICIAL SELECTION 618

FOCUS ON Artificial Selection 619 QUANTITATIVE TRAIT LOCI 620

PROBLEM-SOLVING SKILLS Detecting Dominance at a QTL 623

Correlations Between Relatives 624

CORRELATING QUANTITATIVE PHENOTYPES BETWEEN RELATIVES 625

INTERPRETING CORRELATIONS BETWEEN RELATIVES 626

Quantitative Genetics of Human Behavioral Traits *628*

INTELLIGENCE 628 PERSONALITY 629

CHAPTER 23

Population Genetics 634

A Remote Colony 634

The Theory of Allele Frequencies 635

ESTIMATING ALLELE FREQUENCIES 635 RELATING GENOTYPE FREQUENCIES TO ALLELE FREQUENCIES: THE HARDY-WEINBERG PRINCIPLE 636 APPLICATIONS OF THE HARDY-WEINBERG PRINCIPLE 638 EXCEPTIONS TO THE HARDY-WEINBERG PRINCIPLE 638

SOLVE IT The Effects of Inbreeding on Hardy-Weinberg Frequencies 639

USING ALLELE FREQUENCIES IN GENETIC COUNSELING 640

Natural Selection 641

THE CONCEPT OF FITNESS 641 NATURAL SELECTION AT THE LEVEL OF THE GENE 642

SOLVE IT Selection Against a Harmful Recessive Allele *643*

Random Genetic Drift 645

RANDOM CHANGES IN ALLELE FREQUENCIES 645 THE EFFECTS OF POPULATION SIZE 646

PROBLEM-SOLVING SKILLS Applying Genetic Drift to Pitcairn Island 647

Populations in Genetic Equilibrium 647

BALANCING SELECTION 648 MUTATION-SELECTION BALANCE 649 MUTATION-DRIFT BALANCE 650

CHAPTER 24

Evolutionary Genetics 656

D'ou venons nous? Que sommes nous? Ou allons nous? 656

The Emergence of Evolutionary Theory 657

DARWIN'S THEORY OF EVOLUTION 657 EVOLUTIONARY GENETICS 658

Genetic Variation in Natural Populations 659

VARIATION IN PHENOTYPES 659 VARIATION IN CHROMOSOME STRUCTURE 660 VARIATION IN PROTEIN STRUCTURE 661 VARIATION IN NUCLEOTIDE SEQUENCES 661

Molecular Evolution 662

MOLECULES AS "DOCUMENTS OF EVOLUTIONARY HISTORY" 663 MOLECULAR PHYLOGENIES 664

RATES OF MOLECULAR EVOLUTION 664

PROBLEM-SOLVING SKILLS Using Mitochondrial DNA to Establish a Phylogeny 665 THE MOLECULAR CLOCK 667

SOLVE IT Calculating Divergence Times 667 VARIATION IN THE EVOLUTION OF PROTEIN SEQUENCES 667 VARIATION IN THE EVOLUTION OF DNA SEQUENCES 668 THE NEUTRAL THEORY OF MOLECULAR EVOLUTION 669

SOLVE IT Evolution by Mutation and Genetic Drift 670

MOLECULAR EVOLUTION AND PHENOTYPIC EVOLUTION 670

Speciation 672

WHAT IS A SPECIES? 672 MODES OF SPECIATION 674

Human Evolution 676

HUMANS AND THE GREAT APES 676 HUMAN EVOLUTION IN THE FOSSIL RECORD 676 DNA SEQUENCE VARIATION AND HUMAN ORIGINS 677

Appendices

Appendix A: The Rules of Probability 685 Appendix B: Binomial Probabilities 687 Appendix C: *In Situ* Hybridization 689 Appendix D: Evidence for an Unstable Messenger RNA 691 Appendix E: Evolutionary Rates 693

Answers to Odd-Numbered Questions and Problems 697

Glossary 720

Photo Credits 743

Illustration Credits 745

Index 746

The Science of Genetics



CHAPTER OUTLINE

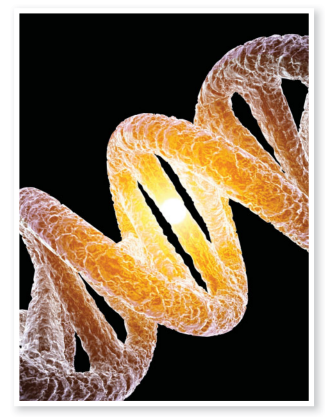
- An Invitation
- Three Great Milestones in Genetics
- DNA as the Genetic Material
- Genetics and Evolution
- Levels of Genetic Analysis
- Genetics in the World: Applications of Genetics to Human Endeavors

time a cell divides, its DNA is replicated and apportioned equally to two daughter cells. The DNA content of these cells—what we call the genome—is thereby conserved. This genome is a master set of instructions, in fact a whole library of information, that cells use to maintain the living state. Ultimately, all the activities of a cell depend on it. To know the DNA is therefore to know the cell, and, in a larger sense, to know the organism to which that cell belongs.

Given the importance of the DNA, it should come as no surprise that great efforts have been expended to study it, down to the finest details. In fact, in the last decade of the twentieth century a worldwide campaign, the Human Genome Project, took shape, and in 2001 it produced a comprehensive analysis of human DNA samples that had been collected from a small number of anonymous donors. This work—stunning in scope and significance—laid the foundation for all future research on the human genome. Then, in 2007, the analysis of human DNA took a new turn. Two of the architects of the Human Genome Project had their own DNA decoded. The technology for analyzing complete genomes has advanced significantly, and the cost for this analysis is no longer exorbitant. In fact, it may soon be possible for each of us to have our own genome analyzed—a prospect that is sure to influence our lives and change how we think about ourselves.

The Personal Genome

Each of us is composed of trillions of cells, and each of those cells contains very thin fibers a few centimeters long that play a major role in who we are, as human beings and as persons. These all-important intracellular fibers are made of DNA. Every



Computer artwork of deoxyribonucleic acid (DNA).

An Invitation

This book is about genetics, the science that deals with DNA. Genetics is also one of the sciences that has a profound impact on us. Through applications in agriculture and medicine, it helps to feed us and keep us healthy. It also provides insight into what makes us human and into what distinguishes each of us as individuals. Genetics is a relatively young science—it emerged only at the beginning of the twentieth century, but it has grown in scope and significance, so much so that it now has a prominent, and some would say commanding, position in all of biology.

Genetics began with the study of how the characteristics of organisms are passed from parents to offspring-that is, how they are inherited. Until the middle of the twentieth century, no one knew for sure what the hereditary material was. However, geneticists recognized that this material had to fulfill three requirements. First, it had to replicate so that copies could be transmitted from parents to offspring. Second, it had to encode information to guide the development, functioning, and behavior of cells and the organisms to which they belong. Third, it had to change, even if only once in a great while, to account for the differences that exist among individuals. For several decades, geneticists wondered what the hereditary material could be. Then in 1953 the structure of DNA was elucidated and genetics had its great clarifying moment. In a relatively short time, researchers discovered how DNA functions as the hereditary material-that is, how it replicates, how it encodes and expresses information, and how it changes. These discoveries ushered in a new phase of genetics in which phenomena could be explained at the molecular level. In time, geneticists learned how to analyze the DNA of whole genomes, including our own. This progress-from studies of heredity to studies of whole genomes-has been amazing.

As practicing geneticists and as teachers, we have written this book to explain the science of genetics to you. As its title indicates, this book is designed to convey the principles of genetics, and to do so in sufficient detail for you to understand them clearly. We invite you to read each chapter, to study its illustrations, and to wrestle with the questions and problems at the chapter's end. We all know that learning and research, teaching, and writing too—takes effort. As authors, we hope your effort studying this book will be rewarded with a good understanding of genetics.

This introductory chapter provides an overview of what we will explain in more detail in the chapters to come. For some of you, it will be a review of knowledge gained from studying basic biology and chemistry. For others, it will be new fare. Our advice is to read the chapter without dwelling on the details. The emphasis here is on the grand themes that run through genetics. The many details of genetics theory and practice will come later.

Three Great Milestones in Genetics

Genetics is rooted in the research of Gregor Mendel, a monk who discovered how traits are inherited. The molecular basis of heredity was revealed when James Watson and Francis Crick elucidated the structure of DNA. The Human Genome Project is currently engaged in the detailed analysis of human DNA. Scientific knowledge and understanding usually advance incrementally. In this book we will examine the advances that have occurred in genetics during its short history—barely a hundred years. Three great milestones stand out in this history: (1) the discovery of rules governing the inheritance of traits in organisms; (2) the identification of the material responsible for this inheritance and the elucidation of its structure; and (3) the comprehensive analysis of the hereditary material in human beings and other organisms.

MENDEL: GENES AND THE RULES OF INHERITANCE

Although genetics developed during the twentieth century, its origin is rooted in the work of *Gregor Mendel* (**Figure 1.1**), a Moravian monk who lived in the nineteenth

century. Mendel carried out his path-breaking research in relative obscurity. He studied the inheritance of different traits in peas, which he grew in the monastery garden. His method involved interbreeding plants that showed different traits—for example, short plants were bred with tall plants—to see how the traits were inherited by the offspring. Mendel's careful analysis enabled him to discern patterns, which led him to postulate the existence of hereditary factors responsible for the traits he studied. We now call these factors genes.

Mendel studied several genes in the garden pea. Each of the genes was associated with a different trait—for example, plant height, or flower color, or seed texture. He discovered that these genes exist in different forms, which we now call **alleles**. One form of the gene for height, for example, allows pea plants to grow more than 2 meters tall; another form of this gene limits their growth to about half a meter.

Mendel proposed that pea plants carry two copies of each gene. These copies may be the same or different. During reproduction, one of the copies is randomly incorporated into each sex cell or gamete. The female gametes (eggs) unite with the male gametes (sperm) at fertilization to produce single cells, called zygotes, which then develop into new plants. The reduction in gene copies from two to one during gamete formation and the subsequent restoration of two copies during fertilization underlie the rules of inheritance that Mendel discovered.

Mendel emphasized that the hereditary factors—that is, the genes—are discrete entities. Different alleles of a gene can be brought together in the same plant through hybridization and can then be separated from each other during the production of gametes. The

coexistence of alleles in a plant therefore does not compromise their integrity. Mendel also found that alleles of different genes are inherited independently of each other.

These discoveries were published in 1866 in the proceedings of the Natural History Society of Brünn, the journal of the scientific society in the city where Mendel lived and worked. The article was not much noticed, and Mendel went on to do other things. In 1900, sixteen years after he died, the paper finally came to light, and the science of genetics was born. In short order, the type of analysis that Mendel pioneered was applied to many kinds of organisms, and with notable success. Of course, not every result fit exactly with Mendel's principles. Exceptions were encountered, and when they were investigated more fully, new insights into the behavior and properties of genes emerged. We will delve into Mendel's research and its applications to the study of inheritance, including heredity in humans, in Chapter 3, and we will explore some ramifications of Mendel's ideas in Chapter 4. In Chapters 5, 6, and 7 we will see how Mendel's principles of inheritance are related to the behavior of chromosomes—the cellular structures where genes reside.

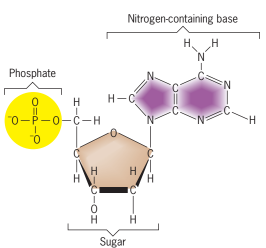
WATSON AND CRICK: THE STRUCTURE OF DNA

The rediscovery of Mendel's paper launched a plethora of studies on inheritance in plants, animals, and microorganisms. The big question on everyone's mind was "What is a gene?" In the middle of the twentieth century, this question was finally answered. Genes were shown to consist of complex molecules called **nucleic acids**.

Nucleic acids are made of elementary building blocks called **nucleotides** (**Figure 1.2**). Each nucleotide has three components: (1) a sugar molecule; (2) a phosphate molecule, which has acidic chemical properties; and (3) a nitrogen-containing molecule, which has slightly basic chemical properties. In **ribonucleic acid**, or RNA, the constituent sugar is ribose; in **deoxyribonucleic acid**, or DNA, it is deoxyribose. Within RNA or DNA, one nucleotide is distinguished from another by its nitrogen-containing base. In RNA,



FIGURE 1.1 Gregor Mendel.



■ **FIGURE 1.2** Structure of a nucleotide. The molecule has three components: a phosphate group, a sugar (in this case deoxyribose), and a nitrogen-containing base (in this case adenine).

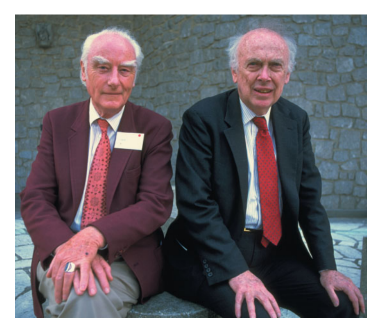


FIGURE 1.3 Francis Crick and James Watson.

the four kinds of bases are adenine (A), guanine (G), cytosine (C), and uracil (U); in DNA, they are A, G, C, and thymine (T). Thus, in both DNA and RNA there are four kinds of nucleotides, and three of them are shared by both types of nucleic acid molecules.

The big breakthrough in the study of nucleic acids came in 1953 when *James Watson* and *Francis Crick* (**Figure 1.3**) deduced how nucleotides are organized within DNA. Watson and Crick knew that the nucleotides are linked, one to another, in a chain. The linkages are formed by chemical interactions between the phosphate of one nucleotide and the sugar of another nucleotide. The nitrogencontaining bases are not involved in these interactions. Thus, a chain of nucleotides consists of a phosphate-sugar backbone to which bases are attached, one base to each sugar in the backbone. From one end of the chain to the other, the bases form a linear sequence characteristic of that particular chain. This sequence of bases is what distinguishes one gene from another. Watson and Crick proposed that DNA molecules consist of two chains of nucleotides (**Figure 1.4a**). These chains are held together by weak chemical attractions-called hydrogen bonds-between particular pairs of bases; A pairs with T, and G pairs with C. Because of these base-pairing rules, the sequence of one nucleotide chain in a double-stranded DNA molecule

can be predicted from that of the other. In this sense, then, the two chains of a DNA molecule are complementary.

A double-stranded DNA molecule is often called a duplex. Watson and Crick discovered that the two strands of a DNA duplex are wound around each other in a helical configuration (**Figure 1.4b**). These helical molecules can be extraordinarily large. Some contain hundreds of millions of nucleotide pairs, and their end-to-end length exceeds 10 centimeters. Were it not for their extraordinary thinness (about a hundredmillionth of a centimeter), we would be able to see them with the unaided eye.

RNA, like DNA, consists of nucleotides linked one to another in a chain. However, unlike DNA, RNA molecules are usually single-stranded. The genes of most organisms are composed of DNA, although in some viruses they are made of RNA. We will examine the structures of DNA and RNA in detail in Chapter 9, and we will investigate the genetic significance of these macromolecules in Chapters 10, 11, and 12.

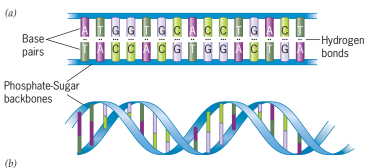
THE HUMAN GENOME PROJECT: SEQUENCING DNA AND CATALOGUING GENES

If geneticists in the first half of the twentieth century dreamed about identifying the stuff that genes are made of, geneticists in the second half of that century dreamed

about ways of determining the sequence of bases in DNA molecules. Near the end of the century, their dreams became reality as projects to determine DNA base sequences in several organisms, including humans, took shape. Obtaining the sequence of bases in an organism's DNA—that is, *sequencing the DNA*—should, in principle, provide the information needed to analyze all that organism's genes. We refer to the collection of DNA molecules that is characteristic of an organism as its genome. Sequencing the genome is therefore tantamount to sequencing all the organism's genes—and more, for we now know that some of the DNA does not comprise genes. The function of this nongenic DNA is not always clear; however, it is present in many genomes, and sometimes it is abundant. A Milestone in Genetics: ΦX174, the First

DNA Genome Sequenced describes how genome sequencing got started. You can find this account in the Student Companion site.

The paragon of all the sequencing programs is the **Human Genome Project**, a worldwide effort to determine the sequence of approximately 3 billion nucleotide pairs in



■ FIGURE 1.4 DNA, a double-stranded molecule held together by hydrogen bonding between paired bases. (a) Two-dimensional representation of the structure of a DNA molecule composed of complementary nucleotide chains. (b) A DNA molecule shown as a double helix. human DNA. As initially conceived, the Human Genome Project was to involve collaborations among researchers in many different countries, and much of the work was to be funded by their governments. However, a privately funded project initiated by Craig Venter, a scientist and entrepreneur, soon developed alongside the publicly funded project. In 2001 all these efforts culminated in the publication of two lengthy articles about the human genome. The articles reported that 2.7 billion nucleotide pairs of human DNA had been sequenced. Computer analysis of this DNA suggested that the human genome contained between 30,000 and 40,000 genes. More recent analyses have revised the human gene number downward, to around 20,500. These genes have been catalogued by location, structure, and potential function. Efforts are now focused on studying how they influence the myriad characteristics of humans.

The genomes of many other organisms—bacteria, fungi, plants, protists, and animals—have also been sequenced. Much of this work has been done under the auspices of the Human Genome Project, or under projects closely allied to it. Initially the sequencing efforts were focused on organisms that are especially favorable for genetic

research. In many places in this book, we explore ways in which researchers have used these model organisms to advance genetic knowledge. Current sequencing projects have moved beyond the model organisms to diverse plants, animals, and microbes. For example, the genomes of the mosquito and the malaria parasite that it carries have both been sequenced, as have the genomes of the honeybee, the poplar tree, and the sea squirt. Some of the targets of these sequencing projects have a medical, agricultural, or commercial significance; others simply help us to understand how genomes are organized and how they have diversified during the history of life on Earth.

All the DNA sequencing projects have transformed genetics in a fundamental way. Genes can now be studied at the molecular level with relative ease, and vast numbers of genes can be studied simultaneously. This approach to genetics, rooted in the analysis of the DNA sequences that make up a genome, is called genomics. It has been made possible by advances in DNA sequencing technology, robotics, and computer science (**Figure 1.5**). Researchers are now able to construct and scan enormous databases containing DNA sequences to address questions about genetics. Although there are a large number of useful databases currently available, we will focus on the databases assembled by the National Center for Biotechnology Information (NCBI), maintained by the U.S. National Institutes of Health. The NCBI databases—available free on the web at http:// www.ncbi.nih.gov-are invaluable repositories of information about genes, proteins, genomes, publications, and other important data in the fields of genetics, biochemistry, and molecular biology. They contain the complete nucleotide sequences of all genomes that have been sequenced to date, and they are continually updated. In addition, the NCBI web site contains tools that can be used to search for specific items of interest-gene and protein sequences, research articles, and so on. In Chapter 15, we will introduce you to some of these tools, and throughout this book, we will encourage you to visit the NCBI web site at the end of each chapter to answer specific questions.

- Gregor Mendel postulated the existence of particulate factors—now called genes—to explain how traits are inherited.
- Alleles, the alternate forms of genes, account for heritable differences among individuals.
- James Watson and Francis Crick elucidated the structure of DNA, a macromolecule composed of two complementary chains of nucleotides.
- DNA is the hereditary material in all life forms except some types of viruses, in which RNA is the hereditary material.
- The Human Genome Project determined the sequence of nucleotides in the DNA of the human genome.
- Sequencing the DNA of a genome provides the data to identify and catalogue all the genes of an organism.



FIGURE 1.5 A researcher loading samples into an automated DNA sequencer.

KEY POINTS

DNA as the Genetic Material

In biology information flows from DNA to RNA to protein.

In all cellular organisms, the genetic material is DNA. This material must be able to *replicate* so that copies can be transmitted from cell to cell and from parents to offspring; it must contain *information* to direct cellular activities and to guide the development, functioning,

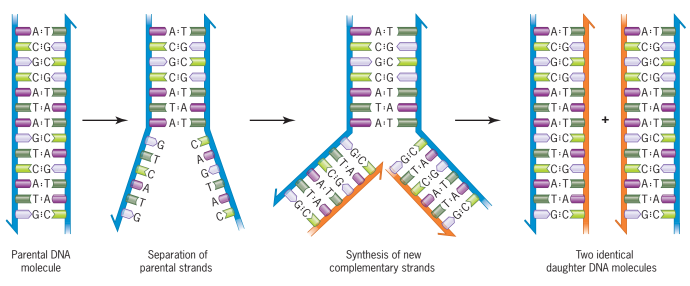
and behavior of organisms; and it must be able to *change* so that over time, groups of organisms can adapt to different circumstances.

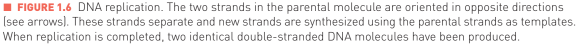
DNA REPLICATION: PROPAGATING GENETIC INFORMATION

The genetic material of an organism is transmitted from a mother cell to its daughters during cell division. It is also transmitted from parents to their offspring during reproduction. The faithful transmission of genetic material from one cell or organism to another is based on the ability of double-stranded DNA molecules to be replicated. DNA replication is extraordinarily exact. Molecules consisting of hundreds of millions of nucleotide pairs are duplicated with few, if any, mistakes.

The process of DNA replication is based on the complementary nature of the strands that make up duplex DNA molecules (**Figure 1.6**). These strands are held together by relatively weak hydrogen bonds between specific base pairs—A paired with T, and G paired with C. When these bonds are broken, the separated strands can serve as templates for the synthesis of new partner strands. The new strands are assembled by the stepwise incorporation of nucleotides opposite to nucleotides in the template strands. This incorporation conforms to the base-pairing rules. Thus, the sequence of nucleotides in a strand being synthesized is dictated by the sequence of nucleotides in the template strand. At the end of the replication process, each template strand is paired with a newly synthesized partner strand. Thus, two identical DNA duplexes are created from one original duplex.

The process of DNA replication does not occur spontaneously. Like most biochemical processes, it is catalyzed by enzymes. We will explore the details of DNA replication, including the roles played by different enzymes, in Chapter 10.





GENE EXPRESSION: USING GENETIC INFORMATION

DNA molecules contain information to direct the activities of cells and to guide the development, functioning, and behavior of the organisms that comprise these cells. This information is encoded in sequences of nucleotides within the DNA molecules of the genome. Among cellular organisms, the smallest known genome is that of Mycoplasma genitalium: 580,070 nucleotide pairs. By contrast, the human genome consists of 3.2 billion nucleotide pairs. In these and all other genomes, the information contained within the DNA is organized into the units we call genes. An M. genitalium has 482 genes, whereas a human sperm cell has around 20,500. Each gene is a stretch of nucleotide pairs along the length of a DNA molecule. A particular DNA molecule may contain thousands of different genes. In an M. genitalium cell, all the genes are situated on one DNA molecule-the single chromosome of this organism. In a human sperm cell, the genes are situated on 23 different DNA molecules corresponding to the 23 chromosomes in the cell. Most of the DNA in *M. genitalium* comprises genes, whereas most of the DNA in humans does not-that is, most of the human DNA is nongenic. We will investigate the genic and nongenic composition of genomes in many places in this book, especially in Chapter 15.

How is the information within individual genes organized and expressed? This question is central in genetics, and we will turn our attention to it in Chapters 11 and 12. Here, suffice it to say that most genes contain the instructions for the synthesis of proteins. Each protein consists of one or more chains of amino acids. These chains are called **polypeptides**. The 20 different kinds of amino acids that occur naturally can be combined in myriad ways to form polypeptides. Each polypeptide has a characteristic sequence of amino acids. Some polypeptides are short—just a few amino acids long whereas others are enormous—thousands of amino acids long.

The sequence of amino acids in a polypeptide is specified by a sequence of elementary coding units within a gene. These elementary coding units, called **codons**, are triplets of adjacent nucleotides. A typical gene may contain hundreds or even thousands of codons. Each codon specifies the incorporation of an amino acid into a polypeptide. Thus, the information encoded within a gene is used to direct the synthesis of a polypeptide, which is often referred to as the gene's product. Sometimes, depending on how the coding information is utilized, a gene may encode several polypeptides; however, these polypeptides are usually all related by sharing some common sequence of amino acids.

The expression of genetic information to form a polypeptide is a two-stage process (**Figure 1.7**). First, the information contained in a gene's DNA is copied into a molecule of RNA. The RNA is assembled in stepwise fashion along one of the strands of the DNA duplex. During this assembly process, A in the RNA pairs with T in the DNA, G in the RNA pairs with C in the DNA, C in the RNA pairs with G in the DNA, and U in the RNA pairs with A in the DNA. Thus, the nucleotide sequence of the RNA is determined by the nucleotide sequence of a strand of DNA in the gene. The process that produces this RNA molecule is called **transcription**, and the RNA itself is called a **transcript**. The RNA transcript eventually separates from its DNA template and, in some organisms, is altered by the addition, deletion, or modification of nucleotides. The finished molecule, called the **messenger RNA** or simply **mRNA**, contains all the information needed for the synthesis of a polypeptide.

The second stage in the expression of a gene's information is called **translation**. At this stage, the gene's mRNA acts as a template for the synthesis of a polypeptide. Each of the gene's codons, now present within the sequence of the mRNA, specifies the incorporation of a particular amino acid into the polypeptide chain. One amino acid is added at a time. Thus, the polypeptide is synthesized stepwise by reading the codons in order. When the polypeptide is finished, it dissociates from the mRNA, folds into a precise three-dimensional shape, and then carries out its role in the cell. Some polypeptides are altered by the removal of the first amino acid, which is usually methionine, in the sequence.

We refer to the collection of all the different proteins in an organism as its **proteome**. Humans, with around 20,500 genes, may have hundreds of thousands of different proteins

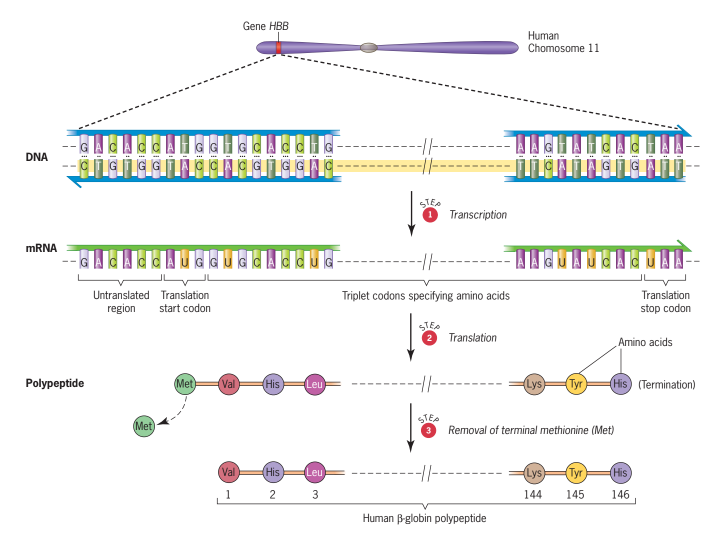


FIGURE 1.7 Expression of the human gene *HBB* coding for the β -globin polypeptide of hemoglobin. During transcription (step 1), one strand of the *HBB* DNA (here the bottom strand shown highlighted) serves as a template for the synthesis of a complementary strand of RNA. After undergoing modifications, the resulting mRNA (messenger RNA) is used as a template to synthesize the β -globin polypeptide. This process is called translation (step 2). During translation each triplet codon in the mRNA specifies the incorporation of an amino acid in the polypeptide chain. Translation is initiated by a start codon, which specifies the incorporation of the amino acid methionine (met), and it is terminated by a stop codon, which does not specify the incorporation of any amino acid. After translation is completed, the initial methionine is removed (step 3) to produce the mature β -globin polypeptide.

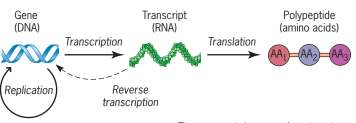
in their proteome. One reason for the large size of the human proteome is that a particular gene may encode several different, but related, polypeptides, and these polypeptides may combine in complex ways to produce different proteins. Another reason is that proteins may be produced by combining polypeptides encoded by different genes. If the number of genes in the human genome is large, the number of proteins in the human proteome is truly enormous.

The study of all the proteins in cells—their composition, the sequences of amino acids in their constituent polypeptides, the interactions among these polypeptides and among different proteins, and, of course, the functions of these complex molecules—is called **proteomics**. Like genomics, proteomics has been made possible by advances in the technologies used to study genes and gene products, and by the development of computer programs to search databases and analyze amino acid sequences.

From all these considerations, it is clear that information flows from genes, which are composed of DNA, to polypeptides, which are composed of amino acids, through an intermediate, which is composed of RNA (**Figure 1.8**). Thus, in the broad sense, the flow of information is DNA \rightarrow RNA \rightarrow polypeptide, a progression often spoken of as the *central dogma of molecular biology*. In several chapters we will see circumstances in which the first part of this progression is reversed—that is, RNA is used as a template for the synthesis of DNA. This process, called *reverse transcription*, plays an important role in the activities of certain types of viruses, including the virus that causes acquired immune deficiency

syndrome, or AIDS; it also profoundly affects the content and structure of the genomes of many organisms, including the human genome. We will examine the impact of reverse transcription on genomes in Chapter 17.

It was once thought that all or nearly all genes encode polypeptides. However, recent research has shown this idea to be incorrect. Many genes do not encode polypeptides; instead, their end products are RNA molecules that play important roles within cells. We will explore these RNAs and the genes that produce them in Chapters 11 and 19.



■ FIGURE 1.8 The central dogma of molecular biology showing how genetic information is propagated (through DNA replication) and expressed (through transcription and translation). In reverse transcription, RNA is used as a template for the synthesis of DNA.

MUTATION: CHANGING GENETIC INFORMATION

DNA replication is an extraordinarily accurate process, but it is not perfect. At a low but measurable frequency, nucleotides are incorporated incorrectly into growing DNA chains. Such changes have the potential to alter or disrupt the information encoded in genes. DNA molecules are also sometimes damaged by electromagnetic radiation or by chemicals. Although the damage induced by these agents may be repaired, the repair processes often leave scars. Stretches of nucleotides may be deleted or duplicated, or they may be rearranged within the overall structure of the DNA molecule. We call all these types of changes **mutations**. Genes that are altered by the occurrence of mutations are called mutant genes.

Often mutant genes cause different traits in organisms (**Figure 1.9**). For example, one of the genes in the human genome encodes the polypeptide known as β -globin. This polypeptide, 146 amino acids long, is a constituent of hemoglobin, the protein that transports oxygen in the blood. The 146 amino acids in β -globin correspond to 146 codons in the β -globin gene. The sixth of these codons specifies the incorporation of glutamic acid into the polypeptide. Countless generations ago, in the germ line of some nameless individual, the middle nucleotide pair in this codon was changed from A:T to T:A, and the resulting mutation was passed on to the individual's descendants. This mutation, now widespread in some human populations, altered the sixth codon so that it specifies the incorporation of value into the β -globin polypeptide. This seemingly insignificant change has a deleterious effect on the structure of the cells that make and store hemoglobin-the red blood cells. People who carry two copies of the mutant version of the β -globin gene have sickle-shaped red blood cells, whereas people who carry two copies of the nonmutant version of this gene have disc-shaped red blood cells. The sickle-shaped cells do not transport oxygen efficiently through the body. Consequently, people with sickle-shaped red blood cells develop a serious disease, so serious in fact that they may eventually die from it. This sickle-cell disease is therefore traceable to a mutation in the β -globin gene. We will investigate the nature and causes of mutations like this one in Chapter 13.

The process of mutation has another aspect—it introduces variability into the genetic material of organisms. Over time, the mutant genes created by mutation may spread through a population. For example, you might wonder why the mutant β -globin

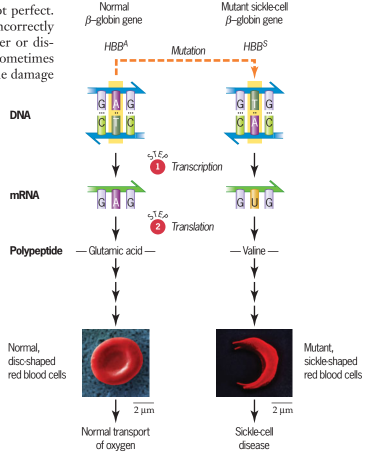


FIGURE 1.9 The nature and consequence of a mutation in the gene for human β -globin. The mutant gene (HBB^{s} top right) responsible for sickle-cell disease resulted from a single base-pair substitution in the β -globin gene (HBB^{A} top left). Transcription and translation of the mutant gene produce a β -globin polypeptide containing the amino acid valine (center right) at the position where normal β -globin contains glutamic acid (center left). This single amino acid change results in the formation of sickle-shaped red blood cells (bottom right) rather than the normal disc-shaped cells (bottom left). The sickle-shaped cells cause a severe form of anemia.

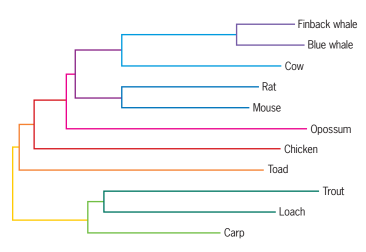
gene is relatively common in some human populations. It turns out that people who carry both a mutant and a nonmutant allele of this gene are less susceptible to infection by the blood parasite that causes malaria. These people therefore have a better chance of surviving in environments where malaria is a threat. Because of this enhanced survival, they produce more children than other people, and the mutant allele that they carry can spread. This example shows how the genetic makeup of a population—in this case, the human population—can evolve over time.

KEY POINTS • When DNA replicates, each strand of a duplex molecule serves as the template for the synthesis of a complementary strand.

- When genetic information is expressed, one strand of a gene's DNA duplex is used as a template for the synthesis of a complementary strand of RNA.
- For most genes, RNA synthesis (transcription) generates a molecule (the RNA transcript) that becomes a messenger RNA (mRNA).
- Coded information in an mRNA is translated into a sequence of amino acids in a polypeptide.
- Mutations can alter the DNA sequence of a gene.
- The genetic variability created by mutation is the basis for biological evolution.

Genetics and Evolution

Genetics has much to contribute to the scientific study of evolution.



■ FIGURE 1.10 Phylogenetic tree showing the evolutionary relationships among 11 different vertebrates. This tree was constructed by comparing the sequences of the gene for cytochrome b, a protein involved in energy metabolism. The 11 different animals have been positioned in the tree according to the similarity of their cytochrome b gene sequences. This tree is consistent with other information (e.g., data obtained from the study of fossils), except for the positions of the three fish species. The loach is actually more closely related to the carp than it is to the trout. This discrepancy points out the need to interpret the results of DNA sequence comparisons carefully.

As mutations accumulate in the DNA over many generations, we see their effects as differences among organisms. Mendel's strains of peas carried different mutant genes, and so do people from different ancestral groups. In almost any species, at least some of the observable variation has an underlying genetic basis. In the middle of the nineteenth century, *Charles Darwin* and *Alfred Wallace*, both contemporaries of Mendel, proposed that this variation makes it possible for species to change—that is, to evolve—over time.

The ideas of Darwin and Wallace revolutionized scientific thought. They introduced an historical perspective into biology and gave credence to the concept that all living things are related by virtue of descent from a common ancestor. However, when these ideas were proposed, Mendel's work on heredity was still in progress and the science of genetics had not yet been born. Research on biological evolution was stimulated when Mendel's discoveries came to light at the beginning of the twentieth century, and it took a new turn when DNA sequencing techniques emerged at the century's end. With DNA sequencing we can see similarities and differences in the genetic material of diverse organisms. On the assumption that sequences of nucleotides in the DNA are the result of historical processes, it is possible to interpret these similarities and differences in a temporal framework. Organisms with very similar DNA sequences are descended from a recent common ancestor, whereas organisms with less similar DNA sequences are descended from a more remote common ancestor. Using this logic, researchers can establish the historical relationships among organisms (**Figure 1.10**). We call these relationships a phylogenetic tree, or more simply, a **phylogeny**, from Greek words meaning "the origin of tribes."

Today the construction of phylogenetic trees is an important part of the study of evolution. Biologists use the burgeoning DNA sequence data from the genome projects and other research ventures, such as the United States National Science Foundation's "Tree of Life" program, in combination with anatomical data collected from living and fossilized organisms to discern the evolutionary relationships among species. We will explore the genetic basis of evolution in Chapters 23 and 24.

- Evolution depends on the occurrence, transmission, and spread of mutant genes in groups of organisms.
- DNA sequence data provide a way of studying the historical process of evolution.

Levels of Genetic Analysis

Genetic analysis is practiced at different levels. The oldest type of genetic analysis follows in Mendel's footsteps by focusing on how traits are inherited when different strains of organisms are hybridized. Another type of genetic analysis follows in the footsteps of WatGeneticists approach their science from different points of view—from that of a gene, a DNA molecule, or a population of organisms.

KEY POINTS

son and Crick and the army of people who have worked on the various genome projects by focusing on the molecular makeup of the genetic material. Still another type of genetic analysis imitates Darwin and Wallace by focusing on entire populations of organisms. All these levels of genetic analysis are routinely used in research today. Although we will encounter them in many different places in this book, we provide brief descriptions of them here.

CLASSICAL GENETICS

The period prior to the discovery of the structure of DNA is often spoken of as the era of *classical genetics*. During this time, geneticists pursued their science by analyzing the outcomes of crosses between different strains of organisms, much as Mendel had done in his work with peas. In this type of analysis, genes are identified by studying the inheritance of trait differences—tall pea plants versus short pea plants, for example—in the offspring of crosses. The trait differences are due to the alternate forms of genes. Sometimes more than one gene influences a trait, and sometimes environmental conditions—for example, temperature and nutrition—exert an effect. These complications can make the analysis of inheritance difficult.

The classical approach to the study of genes can also be coordinated with studies on the structure and behavior of chromosomes, which are the cellular entities that contain the genes. By analyzing patterns of inheritance, geneticists can localize genes to specific chromosomes. More detailed analyses allow them to localize genes to specific positions within chromosomes—a practice called chromosome mapping. Because these studies emphasize the transmission of genes and chromosomes from one generation to the next, they are often referred to as exercises in *transmission genetics*. However, classical genetics is not limited to the analysis of gene and chromosome transmission. It also studies the nature of the genetic material—how it controls traits and how it mutates. We present the essential features of classical genetics in Chapters 3–8.

MOLECULAR GENETICS

With the discovery of the structure of DNA, genetics entered a new phase. The replication, expression, and mutation of genes could now be studied at the molecular level. This approach to genetic analysis was raised to a new level when it became possible to sequence DNA molecules easily. Molecular genetic analysis is rooted in the study of DNA sequences. Knowledge of a DNA sequence and comparisons to other DNA sequences allow a geneticist to define a gene chemically. The gene's internal components coding sequences, regulatory sequences, and noncoding sequences—can be identified, and the nature of the polypeptide encoded by the gene can be predicted.

But the molecular approach to genetic analysis is much more than the study of DNA sequences. Geneticists have learned to cut DNA molecules at specific sites. Whole genes, or pieces of genes, can be excised from one DNA molecule and inserted into another DNA molecule. These "recombinant" DNA molecules can be replicated in bacterial cells or even in test tubes that have been supplied with appropriate enzymes. Milligram quantities of a particular gene can be generated in the laboratory in an afternoon. In short, geneticists have learned how to manipulate genes more or less at will. This artful manipulation has allowed researchers to study genetic phenomena in great detail. They have even learned how to transfer genes from one organism to another. We present examples of molecular genetic analysis in many chapters in this book.

POPULATION GENETICS

Genetics can also be studied at the level of an entire population of organisms. Individuals within a population may carry different alleles of a gene; perhaps they carry different alleles of many genes. These differences make individuals genetically distinct, possibly even unique. In other words, the members of a population vary in their genetic makeup. Geneticists seek to document this variability and to understand its significance. Their most basic approach is to determine the frequencies of specific alleles in a population and then to ascertain if these frequencies change over time. If they do, the population is evolving. The assessment of genetic variability in a population is therefore a foundation for the study of biological evolution. It is also useful in the effort to understand the inheritance of complex traits, such as body size or disease susceptibility. Often complex traits are of considerable interest because they have an agricultural or a medical significance. We discuss genetic analysis at the population level in Chapters 22, 23, and 24.

KEY POINTS • In classical genetic analysis, genes are studied by following the inheritance of traits in crosses between different strains of an organism.

- In molecular genetic analysis, genes are studied by isolating, sequencing, and manipulating DNA and by examining the products of gene expression.
- In population genetic analysis, genes are studied by assessing the variability among individuals in a group of organisms.

Genetics in the World: Applications of Genetics to Human Endeavors

Genetics is relevant in many venues outside the research laboratory.

Modern genetic analysis began in a European monastic enclosure; today, it is a worldwide enterprise. The significance and international scope of genetics are evident in today's scientific journals, which showcase the work of geneticists from many dif-

ferent countries. They are also evident in the myriad ways in which genetics is applied in agriculture, medicine, and many other human endeavors all over the world. We will consider some of these applications in Chapters 14, 15, 16, 23, and 24. Some of the highlights are introduced in this section.

GENETICS IN AGRICULTURE

By the time the first civilizations appeared, humans had already learned to cultivate crop plants and to rear livestock. They had also learned to improve their crops and livestock by selective breeding. This pre-Mendelian application of genetic principles had telling effects. Over thousands of generations, domesticated plant and animal species



Angus





Simmental



Charolais

FIGURE 1.11 Breeds of beef cattle.

came to be quite different from their wild ancestors. For example, cattle were changed in appearance and behavior (**■** Figure 1.11), and corn, which is descended from a wild grass called teosinte (**■** Figure 1.12), was changed so much that it could no longer grow without human cultivation.

Selective breeding programs—now informed by genetic theory—continue to play important roles in agriculture. High-yielding varieties of wheat, corn, rice, and many other plants have been developed by breeders to feed a growing human population. Selective breeding techniques have also been applied to animals such as beef and dairy cattle, swine, and sheep, and to horticultural plants such as shade trees, turf grass, and garden flowers.

Beginning in the 1980s, classical approaches to crop and livestock improvement were supplemented—and in some cases, supplanted—by approaches from molecular genetics. Detailed genetic maps of the chromosomes of several species were constructed to pinpoint genes of agricultural significance. By locating genes for traits such as grain yield or disease resistance, breeders could now design schemes to incorporate particular alleles into agricultural varieties. These mapping projects have been carried on relentlessly and for a few species have culminated in the complete sequencing of the genome. Other crop and livestock genome sequencing projects are still in progress. All sorts of potentially useful genes are being identified and studied in these projects.

Plant and animal breeders are also employing the techniques of molecular genetics to introduce genes from other species into crop plants and livestock. This process of changing the genetic makeup of an organism was initially developed using test species such as fruit flies. Today it is widely used to augment the genetic material of many kinds of creatures. Plants and animals that have been altered by the introduction of foreign genes are called *GMOs—genetically modified organisms*. BT corn is an example. Many corn varieties now grown in the United States carry a gene from the bacterium





■ **FIGURE 1.12** Ears of corn (right) and its ancestor, teosinte (left).





(0)

FIGURE 1.13 Use of a genetically modified plant in agriculture. (*a*) European corn borer eating away the stalk of a corn plant. (*b*) Side-by-side comparison of corn stalks from plants that are resistant (top) and susceptible (bottom) to the corn borer. The resistant plant is expressing a gene for an insecticidal protein derived from *Bacillus thuringiensis*.

Bacillus thuringiensis. This gene encodes a protein that is toxic to many insects. Corn strains that carry the gene for BT toxin are resistant to attacks by the European corn borer, an insect that has caused enormous damage in the past (**Figure 1.13**). Thus, BT corn plants produce their own insecticide.

The development and use of GMOs has stirred up controversy worldwide. For example, African and European countries have been reluctant to grow BT corn or to purchase BT corn grown in the United States. Their reluctance is due to several factors, including the conflicting interests of small farmers and large agricultural corporations, and concerns about the safety of consuming genetically modified food. There is also a concern that BT corn might kill nonpest

species of insects such as butterflies and honeybees. Advances in molecular genetics have provided the tools and the materials to change agriculture profoundly. Today, policy makers are wrestling with the implications of these new technologies.

GENETICS IN MEDICINE

Classical genetics has provided physicians with a long list of diseases that are caused by mutant genes. The study of these diseases began shortly after Mendel's work was rediscovered. In 1909 Sir Archibald Garrod, a British physician and biochemist, published a book entitled *Inborn Errors of Metabolism*. In this book Garrod documented how metabolic abnormalities can be traced to mutant alleles. His research was seminal, and in the next several decades, a large number of inherited human disorders were identified and catalogued. From this work, physicians have learned to diagnose genetic diseases, to trace them through families, and to predict the chances that particular individuals might inherit them. Today some hospitals have professionals known as *genetic counselors* who are trained to advise people about the risks of inheriting or transmitting genetic diseases. We will discuss some aspects of genetic counseling in Chapter 3.

Genetic diseases like the ones that Garrod studied are individually rather rare in most human populations. For example, among newborns, the incidence of phenylketonuria, a disorder of amino acid metabolism, is only one in 10,000. However, mutant genes also contribute to more prevalent human maladies—heart disease and cancer, for example. In Chapter 22 we will explore ways of assessing genetic risks for complex traits such as the susceptibility to heart disease, and in Chapter 21 we will investigate the genetic basis of cancer.

Advances in molecular genetics are providing new ways of detecting mutant genes in individuals. Diagnostic tests based on analysis of DNA are now readily available. For example, a hospital lab can test a blood sample or a cheek swab for the presence of a mutant allele of the *BRCA1* gene, which strongly predisposes its carriers to develop breast cancer. If a woman carries the mutant allele, she may be advised to undergo a mastectomy to prevent breast cancer from occurring. The application of these new molecular genetic technologies therefore often raises difficult issues for the people involved.

Molecular genetics is also providing new ways to treat diseases. For decades diabetics had to be given insulin obtained from animals—usually pigs. Today, perfect human insulin is manufactured in bacterial cells that carry the human insulin gene. Vats of these cells are grown to produce the insulin polypeptide on an industrial scale. Human growth hormone, previously isolated from cadavers, is also manufactured in bacterial cells. This hormone is used to treat children who cannot make sufficient amounts of the hormone themselves because they carry a mutant allele of the growth hormone gene. Without the added hormone, these children would be affected with dwarfism. Many other medically important proteins are now routinely produced in bacterial cells that have been supplied with the appropriate human gene. The large-scale production of such proteins is one facet of the burgeoning biotechnology industry. We will explore ways of producing human proteins in bacterial cells in Chapter 16.

Human gene therapy is another way in which molecular genetic technologies are used to treat diseases. The strategy in this type of therapy is to insert a healthy, functional copy of a particular gene into the cells of an individual who carries only mutant copies of that gene. The inserted gene can then compensate for the faulty genes that the individual inherited. To date, human gene therapy has had mixed results. Efforts to cure individuals with cystic fibrosis (CF), a serious respiratory disorder, by introducing copies of the normal CF gene into lung cells have not been successful. However, medical geneticists have had some success in treating immune system and blood cell disorders by introducing the appropriate normal genes into bone marrow cells, which later differentiate into immune cells and blood cells. We will discuss the emerging technologies for human gene therapy and some of the risks involved in Chapter 16.

GENETICS IN SOCIETY

Modern societies depend heavily on the technology that emerges from research in the basic sciences. Our manufacturing and service industries are built on technologies for mass production, instantaneous communication, and prodigious information processing. Our lifestyles also depend on these technologies. At a more fundamental level, modern societies rely on technology to provide food and health care. We have already seen how genetics is contributing to these important needs. However, genetics impacts society in other ways too.

One way is economic. Discoveries from genetic research have initiated countless business ventures in the biotechnology industry. Companies that market pharmaceuticals and diagnostic tests, or that provide services such as DNA profiling, have contributed to worldwide economic growth. Another way is legal. DNA sequences differ among individuals, and by analyzing these differences, people can be identified uniquely. Such analyses are now routinely used in many situations—to test for paternity, to convict the guilty and to exonerate the innocent of crimes for which they are accused, to authenticate claims to inheritances, and to identify the dead. Evidence based on analysis of DNA is now commonplace in courtrooms all over the world.

But the impact of genetics goes beyond the material, commercial, and legal aspects of our societies. It strikes the very core of our existence because, after all, DNA—the subject of genetics—is a crucial part of us. Discoveries from genetics raise deep, difficult, and sometimes disturbing existential questions. Who are we? Where do we come from? Does our genetic makeup determine our nature? our talents? our ability to learn? our behavior? Does it play a role in setting our customs? Does it affect the ways we organize our societies? Does it influence our attitudes toward other people? Will knowledge about our genes and how they influence us affect our ideas about morality and justice, innocence and guilt, freedom and responsibility? Will this knowledge change how we think about what it means to be human? Whether we like it or not, these and other probing questions await us in the not-so-distant future.

- Discoveries in genetics are changing procedures and practices in agriculture and medicine.
- Advances in genetics are raising ethical, legal, political, social, and philosophical questions.

KEY POINTS

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. How is genetic information expressed in cells?
- Answer: The genetic information is encoded in sequences in the DNA. Initially, these sequences are used to synthesize RNA complementary to them—a process called transcription— and then the RNA is used as a template to specify the incorporation of amino acids in the sequence of a polypeptide— a process called translation. Each amino acid in the polypeptide corresponds to a sequence of three nucleotides in the DNA. The triplets of nucleotides that encode the different amino acids are called codons.
- 2. What is the evolutionary significance of mutation?
- **Answer:** Mutation creates variation in the DNA sequences of genes (and in the nongenic components of genomes as well). This variation accumulates in populations of organisms over time and may eventually produce observable differences among the organisms. One population may come to differ from another according to the kinds of mutations that have accumulated over time. Thus, mutation provides the input for different evolutionary outcomes at the population level.

Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. Suppose a gene contains 10 codons. How many coding nucleotides does the gene contain? How many amino acids are expected to be present in its polypeptide product? Among all possible genes composed of 10 codons, how many different polypeptides could be produced?
- **Answer:** The gene possesses 30 coding nucleotides. Its polypeptide product is expected to contain 10 amino acids, each corresponding to one of the codons in the gene. If each codon can specify one of 20 naturally occurring amino acids, among all possible gene sequences 10 codons long, we can imagine a total of 20¹⁰ polypeptide products—a truly enormous number!

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **1.1** In a few sentences, what were Mendel's key ideas about inheritance?
- **1.2** Both DNA and RNA are composed of nucleotides. What molecules combine to form a nucleotide?
- **1.3** Which bases are present in DNA? Which bases are present in RNA? Which sugars are present in each of these nucleic acids?
- 1.4 What is a genome?
- **1.5** The sequence of a strand of DNA is ATTGCCGTC. If this strand serves as the template for DNA synthesis, what will be the sequence of the newly synthesized strand?
- **1.6** A gene contains 141 codons. How many nucleotides are present in the gene's coding sequence? How many amino acids are expected to be present in the polypeptide encoded by this gene?
- **1.7** The template strand of a gene being transcribed is CTTGCCAGT. What will be the sequence of the RNA made from this template?
- **1.8** What is the difference between transcription and translation?

- **1.9** RNA is synthesized using DNA as a template. Is DNA ever synthesized using RNA as a template? Explain.
- **1.10** The gene for α -globin is present in all vertebrate species. Over millions of years, the DNA sequence of this gene has changed in the lineage of each species. Consequently, the amino acid sequence of α -globin has also changed in these lineages. Among the 141 amino acid positions in this polypeptide, human α -globin differs from shark α -globin in 79 positions; it differs from carp α -globin in 68 and from cow α -globin in 17. Do these data suggest an evolutionary phylogeny for these vertebrate species?
- 1.11 Sickle-cell disease is caused by a mutation in one of the codons in the gene for β -globin; because of this mutation the sixth amino acid in the β -globin polypeptide is a valine instead of a glutamic acid. A less severe disease is caused by a mutation that changes this same codon to one specifying lysine as the sixth amino acid in the β -globin polypeptide. What word is used to describe the two mutant forms of this gene? Do you think that an individual carrying these two mutant forms of the β -globin gene would suffer from anemia? Explain.

1.12 Hemophilia is an inherited disorder in which the bloodclotting mechanism is defective. Because of this defect, people with hemophilia may die from cuts or bruises, especially if internal organs such as the liver, lungs, or kidneys have been damaged. One method of treatment involves injecting a blood-clotting factor that has been purified from blood donations. This factor is a protein encoded by a human gene. Suggest a way in which modern genetic technology could be used to produce this factor on an industrial scale. Is there a way in which the inborn error of hemophilia could be corrected by human gene therapy?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

You might enjoy using the NCBI web site to explore the Human Genome Project. Click on More about NCBI and then on Outreach and Education. From there click on Recommended Links to get to the National Human Genome Research Institute's page. Once there, click on Education to bring up material on the Human Genome Project.



Cellular Reproduction

CHAPTER OUTLINE

- Cells and Chromosomes
- Mitosis
- Meiosis
- Life Cycles of Some Model Genetic Organisms

Dolly

Sheep have grazed on the hard-scrabble landscape of Scotland for centuries. Finn Dorsets and Scottish Blackfaces are some of the breeds raised by shepherds there. Every spring, the lambs that were conceived during the fall are born. They grow quickly and take their places in flocks—or in butcher shops. Early in 1997, a lamb unlike any other came into the world. This lamb, named Dolly, did not have a father, but she did have three mothers; furthermore, her genes were identical to those of one of her mothers. In a word, Dolly was a clone.

Scientists at the Roslin Institute near Edinburgh, Scotland produced Dolly by fusing an egg from a Blackface ewe (the egg cell

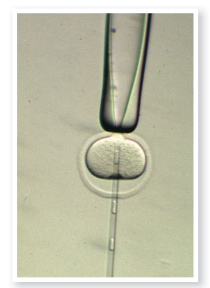


Dolly, the first cloned mammal. The photo on the right shows the cloning process.

mother) with a cell from the udder of a Finn Dorset ewe (the genetic mother). The genetic material in the Blackface ewe's egg had been removed prior to fusing the egg with the udder cell. Subsequently, the newly endowed egg was stimulated to divide. It produced an embryo,

which was implanted in the uterus of another Blackface ewe (the gestational or surrogate mother). This embryo grew and developed, and when the surrogate mother's pregnancy came to term, Dolly was born.

The technology that produced Dolly emerged from a century of basic research on the cellular basis of reproduction. In the ordinary course of events, an egg cell from a female is fertilized by a sperm cell from a male, and the resulting zygote divides to produce genetically identical cells. These cells then divide many times to produce a multicellular organism. Within that organism, a particular group of cells embarks on a different mode of division to produce specialized reproductive



The nuclei of three cells are inside a long, thin micropipette. The topmost nucleus with its genetic material is being injected into an enucleated egg that is being held in place by a wider pipette.

cells—either eggs or sperm. An egg from one such organism then unites with a sperm from another such organism to produce a new offspring. The offspring grows up and the cycle continues, generation after generation. But Dolly, the first cloned mammal, was created by sidestepping this entire process.

Cells and Chromosomes

In the early part of the nineteenth century, a few decades before Gregor Mendel carried out his experiments with peas, biologists established the principle that living things are composed of cells. Some organIn both prokaryotic and eukaryotic cells, the genetic material is organized into chromosomes.

isms consist of just a single cell. Others consist of trillions of cells. Each cell is a complicated assemblage of molecules that can acquire materials, recruit and store energy, and carry out diverse activities, including reproduction. The simplest life forms, viruses, are not composed of cells. However, viruses must enter cells in order to function. Thus, all life has a cellular basis. As preparation for our journey through the science of genetics, we now review the biology of cells. We also discuss chromosomes—the cellular structures in which genes reside.

THE CELLULAR ENVIRONMENT

Living cells are made of many different kinds of molecules. The most abundant is water. Small molecules—for example, salts, sugars, amino acids, and certain vitamins—readily dissolve in water, and some larger molecules interact favorably with it. All these sorts of substances are said to be hydrophilic. Other kinds of molecules do not interact well with water. They are said to be hydrophobic. The inside of a cell, called the **cytoplasm**, contains both hydrophilic and hydrophobic substances.

The molecules that make up cells are diverse in structure and function. Carbohydrates such as starch and glycogen store chemical energy for work within cells. These molecules are composed of glucose, a simple sugar. The glucose subunits are attached one to another to form long chains, or polymers. Cells obtain energy when glucose molecules released from these chains are chemically degraded into simpler compounds-ultimately, to carbon dioxide and water. Cells also possess an assortment of lipids. These molecules are formed by chemical interactions between glycerol, a small organic compound, and larger organic compounds called fatty acids. Lipids are important constituents of many structures within cells. They also serve as energy sources. Proteins are the most diverse molecules within cells. Each protein consists of one or more polypeptides, which are chains of amino acids. Often a protein consists of two polypeptides—that is, it is a dimer; sometimes a protein consists of many different structures. They also catalyze chemical reactions. We call these catalytic proteins enzymes. Cells also contain nucleic acids-DNA and RNA, which, as already described in Chapter 1, are central to life.

Cells are surrounded by a thin layer called a **membrane**. Many different types of molecules make up cell membranes; however, the primary constituents are lipids and proteins. Membranes are also present inside cells. These internal membranes may divide a cell into compartments, or they may help to form specialized structures called **organelles**. Membranes are fluid and flexible. Many of the molecules within a membrane are not rigidly held in place by strong chemical forces. Consequently, they are able to slip by one another in what amounts to an ever-changing molecular sea. Some kinds of cells are surrounded by tough, rigid walls, which are external to the membrane. Plant cell walls are composed of cellulose, a complex carbohydrate. Bacterial cell walls are composed of a different kind of material called murein.

Walls and membranes separate the contents of a cell from the outside world. However, they do not seal it off. These structures are porous to some materials, and they selectively allow other materials to pass through them via channels and gates. The transport of materials in and through walls and membranes is an important activity of cells. Cell membranes also contain molecules that interact with materials in a cell's external environment. Such molecules provide a cell with vital information about conditions in the environment, and they also mediate important cellular activities.

PROKARYOTIC AND EUKARYOTIC CELLS

When we survey the living world, we find two basic kinds of cells: prokaryotic and eukaryotic (**Figure 2.1**). **Prokaryotic cells** are usually less than a thousandth of a millimeter long, and they typically lack a complicated system of internal membranes and membranous organelles. Their hereditary material—that is, the DNA—is not isolated in a special subcellular compartment. Organisms with this kind of cellular organization are called prokaryotes. Examples include the bacteria, which are the most abundant life forms on Earth, and the archaea, which are found in extreme environments such as salt lakes, hot springs, and deep-sea volcanic vents. All other organisms—plants, animals, protists, and fungi—are eukaryotes.

Eukaryotic cells are larger than prokaryotic cells, usually at least 10 times bigger, and they possess complicated systems of internal membranes, some of which are associated with conspicuous, well-organized organelles. For example, eukaryotic cells typically contain one or more **mitochondria** (singular, mitochondrion), which are ellipsoidal organelles dedicated to the recruitment of energy from foodstuffs. Algal and plant cells contain another kind of energy-recruiting organelle called the **chloroplast**, which captures solar energy and converts it into chemical energy. Both mitochondria and chloroplasts are surrounded by membranes.

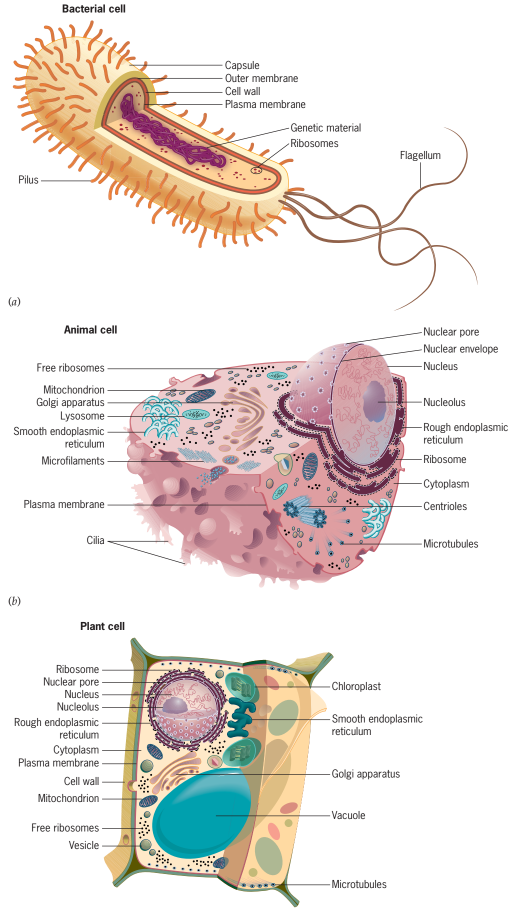
The hallmark of all eukaryotic cells is that their hereditary material is contained within a large, membrane-bounded structure called the **nucleus**. The nuclei of eukaryotic cells provide a safe haven for the DNA, which is organized into discrete structures called **chromosomes**. Individual chromosomes become visible during cell division, when they condense and thicken. In prokaryotic cells, the DNA is usually not housed within a well-defined nucleus. We will investigate the ways in which chromosomal DNA is organized in prokaryotic and eukaryotic cells in Chapter 9. Some of the DNA within a eukaryotic cell is not situated within the nucleus. This extranuclear DNA is located in the mitochondria and chloroplasts. We will examine its structure and function in Chapter 15.

Both prokaryotic and eukaryotic cells possess numerous ribosomes, which are small organelles involved in the synthesis of proteins, a process that we will investigate in Chapter 12. Ribosomes are found throughout the cytoplasm. Although ribosomes are not composed of membranes, in eukaryotic cells they are often associated with a system of membranes called the endoplasmic reticulum. The reticulum may be connected to the Golgi complex, a set of membranous sacs and vesicles that are involved in the chemical modification and transport of substances within cells. Other small, membrane-bound organelles may also be found in eukaryotic cells. In animal cells, lysosomes are produced by the Golgi complex. These organelles contain different kinds of digestive enzymes that would harm the cell if they were released into the cytoplasm. Both plant and animal cells contain perioxisomes, which are small organelles dedicated to the metabolism of substances such as fats and amino acids. The internal membranes and oganelles of eukaryotic cells create a system of subcellular compartments that vary in chemical conditions such as pH and salt content. This variation provides cells with different internal environments that are adapted to the many processes that cells carry out.

The shapes and activities of eukaryotic cells are influenced by a system of filaments, fibers, and associated molecules that collectively form the **cytoskeleton**. These materials give form to cells and enable some types of cells to move through their environment—a phenomenon referred to as cell **motility**. The cytoskeleton holds organelles in place, and it plays a major role in moving materials to specific locations within cells—a phenomenon called **trafficking**.

CHROMOSOMES: WHERE GENES ARE LOCATED

Each chromosome consists of one double-stranded DNA molecule plus an assortment of proteins; RNA may also be associated with chromosomes. Prokaryotic cells typically contain only one chromosome, although sometimes they also possess many smaller DNA molecules called **plasmids**. Most eukaryotic cells contain several



different chromosomes—for example, human sperm cells have 23. The chromosomes of eukaryotic cells are also typically larger and more complex than those of prokaryotic cells. The DNA molecules in prokaryotic chromosomes and plasmids are circular, as are most of the DNA molecules found in the mitochondria and chloroplasts of eukaryotic cells. By contrast, the DNA molecules found in the chromosomes in the nuclei of eukaryotic cells are linear.

Many eukaryotic cells possess two copies of each chromosome. This condition, referred to as the **diploid** state, is characteristic of the cells in the body of a eukaryote— that is, the **somatic cells**. By contrast, the sex cells or **gametes** usually possess only one copy of each chromosome, a condition referred to as the **haploid** state. Gametes are produced from diploid cells located in the **germ line**, which is the reproductive tissue of an organism. In some creatures, such as plants, the germ line produces both sperm and eggs. In other creatures, such as humans, it produces one kind of gamete or the other. When a male and a female gamete unite during fertilization, the diploid state is reestablished, and the resulting zygote develops into a new organism. During animal development, a small number of cells are set aside to form the germ line. All the gametes that will ever be produced are derived from these few cells. The remaining cells form the somatic tissues of the animal. In plants, development is less determinate. Tissues taken from part of a plant—for example, a stem or a leaf—can be used to produce a whole plant, including the reproductive organs. Thus, in plants the distinction between somatic tissues and germ tissues is not as clear-cut as it is in animals.

Chromosomes can be examined by using a microscope. Prokaryotic chromosomes can only be seen with the techniques of electron microscopy, whereas eukaryotic chromosomes can be seen with a light microscope (\blacksquare Figure 2.2). Some eukaryotic chromosomes are large enough to be viewed with low magnification (20×); others require considerably more power (>500×).

Eukaryotic chromosomes are most clearly seen during cell division when each chromosome condenses into a smaller volume. At this time the greater density of the chromosomes makes it possible to discern certain structural features. For example, each chromosome may appear to consist of two parallel rods held together at a common point (**■** Figure 2.2*b*). Each of the rods is an identical copy of the chromosome created during a duplication process that precedes condensation, and the common point,

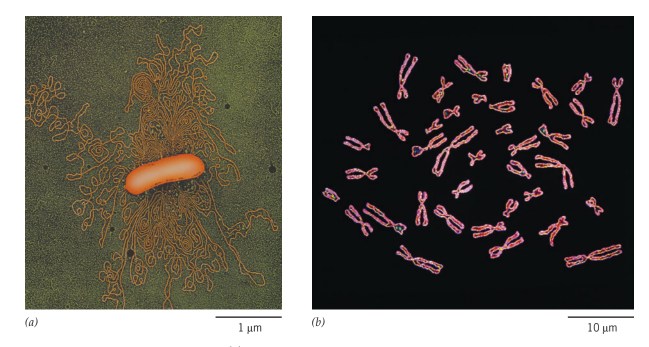


FIGURE 2.2 (a) Electron micrograph showing a bacterial chromosome extruded from a cell. (b) Light micrograph of human chromosomes during cell division. The constriction in each of the duplicated chromosomes is the centromere, the point at which spindle fibers attach to move the chromosome during cell division.

called the **centromere**, becomes associated with an apparatus that moves chromosomes during cell division. We will explore the structures of eukaryotic chromosomes as revealed by light microscopy in Chapter 6.

The discovery that genes are located in chromosomes was made in the first decade of the twentieth century. In Chapter 5 we will examine the experimental evidence for this discovery, and in Chapters 7 and 8 we will study some of the techniques for locating genes within chromosomes.

CELL DIVISION

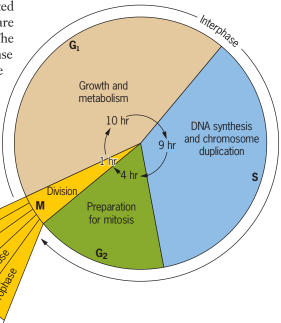
Among the many activities carried out by living cells, division is the most astonishing. A cell can divide into two cells, each of which can also divide into two, and so on through time, to create a population of cells called a **clone**. Barring errors, all the cells within a clone are genetically identical. Cell division is an integral part of the growth of multicellular organisms, and it is also the basis of reproduction.

A cell that is about to divide is called a **mother cell**, and the products of division are called **daughter cells**. When prokaryotic cells divide, the contents of the mother cell are more or less equally apportioned between the two daughter cells. This process is called **fission**. The mother cell's chromosome is duplicated prior to fission, and copies of it are bequeathed to each of the daughter cells. Under optimal conditions, a prokaryote such as the intestinal bacterium *Escherichia coli* divides every 20 to 30 minutes. At this rate, a single *E. coli* cell could form a clone of approximately 2^{50} cells—more than a quadrillion—in just one day. In reality, of course, *E. coli* cells do not sustain this high rate of division. As cells accumulate, the rate of division declines because nutrients are exhausted and waste products pile up. Nevertheless, a single *E. coli* cell call call call to form a mass visible to the unaided eye. We call such a mass of cells a **colony**.

The division of eukaryotic cells is a more elaborate process than the division of prokaryotic cells. Typically many chromosomes must be duplicated, and the duplicates must be distributed equally and exactly to the daughter cells. Organelles—mitochondria, chloroplasts, endoplasmic reticulum, Golgi complex, and so on—must also be distributed to the daughter cells. However, for these entities the distribution process is not equal and exact. Mitochondria and chloroplasts are randomly apportioned to the daughter cells. The endoplasmic reticulum and the Golgi complex are fragmented at the time of division and later are re-formed in the daughter cells.

Each time a eukaryotic cell divides, it goes through a series of phases that collectively form the **cell cycle** (**Figure 2.3**). The progression of phases is denoted $G_1 \rightarrow S \rightarrow G_2 \rightarrow M$. In this progression, S is the period in which the chromosomes are duplicated—an event that requires DNA *synthesis*, to which the label "S" refers. The M phase in the cell cycle is the time when the mother cell actually divides. This phase usually has two components: (1) **mitosis**, which is the process that distributes the duplicated chromosomes equally and exactly to the daughter cells, and (2) **cytokinesis**, which is the process that physically separates the two daughter cells from each other. The label "M" refers to the term *mitosis*, which is derived from a Greek word for thread; during mitosis, the chromosomes appear as threadlike bodies inside cells. The G_1 and G_2 phases are "gaps" between the S and M phases.

The length of the cell cycle varies among different types of cells. In embryos, where growth is rapid, the cycle may be as short as 30 minutes. In slow-growing adult tissues, it may last several months. Some cells, such as those in nerve and muscle tissues, cease to divide once they have acquired their specialized functions. The progression of eukaryotic cells through their cycle is tightly controlled by different types of proteins. When the activities of these proteins are disrupted, cells divide in an unregulated fashion. This deregulation of cell division may lead to cancer, which is a major cause of death among people today. In Chapter 21 we will investigate the genetic basis of cancer. ■ **FIGURE 2.3** The cycle of an animal cell. This cycle is 24 hours long. The duration of the cycle varies among different types of eukaryotic cells.



KEY POINTS

- Cells, the basic units of all living things, are enclosed by membranes.
 - Chromosomes, the cellular structures that carry the genes, are composed of DNA, RNA, and protein.
 - In eukaryotes, chromosomes are contained within a membrane-bounded nucleus; in prokaryotes they are not.
 - Eukaryotic cells possess complex systems of internal membranes as well as membranous organelles such as mitochondria, chloroplasts, and the endoplasmic reticulum.
 - Haploid eukaryotic cells possess one copy of each chromosome; diploid cells possess two copies.
 - Prokaryotic cells divide by fission; eukaryotic cells divide by mitosis and cytokinesis.
 - Eukaryotic chromosomes duplicate when a cell's DNA is synthesized; this event, which precedes mitosis, is characteristic of the S phase of the cell cycle.

Mitosis

When eukaryotic cells divide, they distribute their genetic material equally and exactly to their offspring.

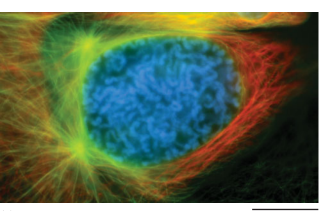
The orderly distribution of duplicated chromosomes in a mother cell to its daughter cells is the essence of mitosis. Each chromosome in a mother cell is duplicated prior to the onset of mitosis, specifically during the S phase. At this time individual chromosomes cannot be identified because they are too extended and too thin. The network of thin strands formed

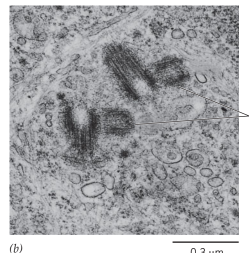
by all the chromosomes within the nucleus is referred to as chromatin. During mitosis, the chromosomes shorten and thicken-that is, they "condense" out of the chromatin network-and individual chromosomes become recognizable. After mitosis, the chromosomes "decondense" and the chromatin network is re-formed. Biologists often refer to the period when individual chromosomes cannot be seen as interphase. This period, which may be quite lengthy, is the time between successive mitotic events.

When mitosis begins, each chromosome has already been duplicated. The duplicates, called sister chromatids, remain intimately associated with each other and are joined at the chromosome's centromere. The term *sister* is something of a misnomer because these chromatids are copies of the original chromosome-therefore, they are more closely related than sisters. Perhaps the word "twin" would describe the situation better. However, "sister" is commonly used, and we will use it here.

The distribution of duplicated chromosomes to the daughter cells is organized and executed by **microtubules**, which are components of the cytoskeleton. These fibers, composed of proteins called tubulins, attach to the chromosomes and move them about within the dividing mother cell. During mitosis the microtubules assemble into a complex array called the **spindle** (**IFigure 2.4***a*). The formation of the spindle is associated with microtubule organizing centers (MTOCs), which are found in the cytoplasm of

FIGURE 2.4 (a) The mitotic spindle in a cultured animal cell, which has been stained to show the microtubules (green) emanating from the two asters. (b) Electron micrograph showing two pairs of centrioles.





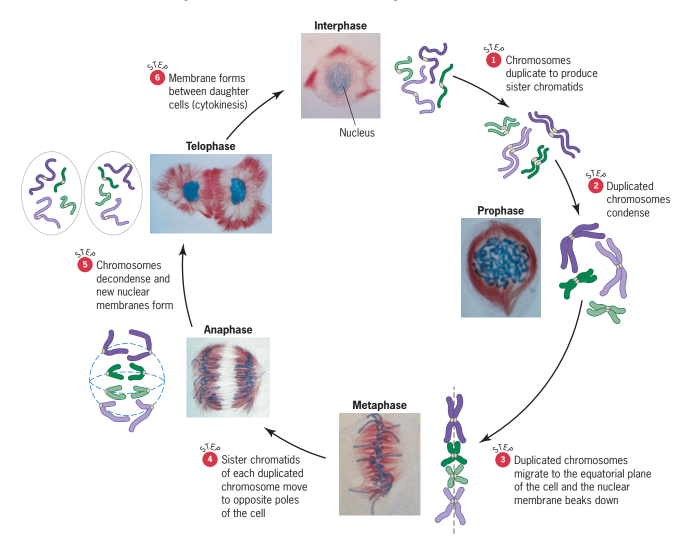
Two pairs of centrioles

20 µm

0.3 µm

eukaryotic cells, usually near the nucleus. In animal cells, the MTOCs are differentiated into small organelles called **centrosomes**; these organelles are not present in plant cells. Each centrosome contains two barrel-shaped **centrioles**, which are aligned at right angles to each other (**Figure 2.4b**). The centrioles are surrounded by a diffuse matrix called the pericentriolar material, which initiates the formation of the microtubules that will make up the mitotic spindle. The single centrosome that exists in an animal cell is duplicated during interphase. As the cell enters mitosis, microtubules develop around each of the daughter centrosomes to form a sunburst pattern called an **aster**. These centrosomes then move around the nucleus to opposite positions in the cell, where they establish the axis of the upcoming mitotic division. The final positions of the centrosomes define the poles of the dividing mother cell. In plant cells, MTOCs that do not have distinct centrosomes define these poles and establish the mitotic spindle.

The initiation of spindle formation and the condensation of duplicated chromosomes from the diffuse network of chromatin are hallmarks of the first stage of mitosis, called **prophase** (**Trigure 2.5**). Formation of the spindle is accompanied by fragmentation of many intracellular organelles—for instance, the endoplasmic reticulum and the Golgi complex. The **nucleolus**, a dense body involved in RNA synthesis within the nucleus, also disappears; however, other types of organelles such as mitochondria and chloroplasts remain intact. Concomitant with the fragmentation of the endoplasmic reticulum, the nuclear membrane (also known as the nuclear envelope) breaks up into many small vesicles, and microtubules formed within the cytoplasm invade the nuclear space. Some of these microtubules attach to the **kinetochores**, which are protein structures associated with the centromeres of the duplicated chromosomes. Attachment of spindle microtubules

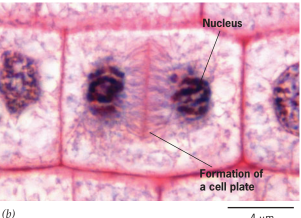






(mag \times 30)

FIGURE 2.6 Cytokinesis in animal (a) and plant (b) cells. The animal cell is a fertilized egg, which is dividing for the first time. Cytokinesis is accomplished by constricting the dividing cell around its middle. This constriction creates a cleavage furrow, which is seen here on one side of the dividing cell. In plant cells, cytokinesis is accomplished by the formation of a membranous cell plate between the daughter cells; eventually, walls composed of cellulose are built on either side of the cell plate.



to the kinetochores indicates that the cell is entering the **metaphase** of mitosis.

During metaphase the duplicated chromosomes move to positions midway between the spindle poles. This movement is leveraged by changes in the length of the spindle microtubules and by the action of force-generating motor proteins that work near the kinetochores. The spindle apparatus also contains microtubules that are not attached to kinetochores. These additional

microtubules appear to stabilize the spindle apparatus. Through the operation of the spindle apparatus, the duplicated chromosomes come to lie in a single plane in the middle of the cell. This equatorial plane is called the metaphase plate. At this stage, each sister chromatid of a duplicated chromosome is connected to a different pole via microtubules attached to its kinetochore. This polar alignment of the sister chromatids is crucial for the equal and exact distribution of genetic material to the daughter cells.

4 um

The sister chromatids of duplicated chromosomes are separated from each other during the **anaphase** of mitosis. This separation is accomplished by shortening the microtubules attached to the kinetochores and by degrading materials that hold the sister chromatids together. As the microtubules shorten, the sister chromatids are pulled to opposite poles of the cell. The separated sister chromatids are now referred to as chromosomes. While the chromosomes are moving toward the poles, the poles themselves also begin to move apart. This double movement cleanly separates the two sets of chromosomes into distinct spaces within the dividing cell. Once this separation has been achieved, the chromosomes decondense into a network of chromatin fibers, and the organelles that were lost at the onset of mitosis re-form. Each set of chromosomes becomes enclosed by a nuclear membrane. The decondensation of the chromosomes and the restoration of the internal organelles are characteristic of the telophase of mitosis. When mitosis is complete, the two daughter cells are separated by the formation of membranes between them. In plants, a wall is also laid down between the daughter cells. This physical separation of the daughter cells is called cytokinesis (**Figure 2.6**).

The daughter cells that are produced by the division of a mother cell are genetically identical. Each daughter has a complete set of chromosomes that were derived by duplicating the chromosomes originally present in the mother cell. The genetic material is therefore transmitted fully and faithfully to the daughter cells from the mother cell. Occasionally, however, mistakes are made during mitosis. A chromatid may become detached from the mitotic spindle and may not be incorporated into one of the daughter cells, or chromatids may become entangled, leading to breakage and the subsequent loss of chromatid parts. These types of events cause genetic differences between the daughter cells. We will consider some of their consequences in Chapter 6 and again in Chapter 21.

KEY POINTS

- As a cell enters mitosis, its duplicated chromosomes condense into rod-shaped bodies (prophase).
- As mitosis progresses, the chromosomes migrate to the equatorial plane of the cell (metaphase).
- Later in mitosis, the centromere that holds the sister chromatids of a duplicated chromosome together splits, and the sister chromatids separate (or disjoin) from each other (anaphase).
- As mitosis comes to an end, the chromosomes decondense and a nuclear membrane re-forms around them (telophase).
- Each daughter cell produced by mitosis and cytokinesis has the same set of chromosomes; thus, daughter cells are genetically identical.

Meiosis

If we denote the number of chromosomes in a gamete by the letter n, then the zygote produced by the union of two gametes has 2n chromosomes. We refer to the n chromosomes of a gamete as the haploid state, and the

2n chromosomes of the zygote as the diploid state. Meiosis—from a Greek word meaning "diminution"—is the process that reduces the diploid state to the haploid state—that is, it reduces the number of chromosomes in a cell by half. The resulting haploid cells either directly become gametes or divide to produce cells that later become gametes. Meiosis therefore plays a key role in reproduction among eukaryotes. Without it, organisms would double their chromosome number every generation—a situation that would quickly become unsupportable given the obvious limitations on the size and metabolic capacity of cells.

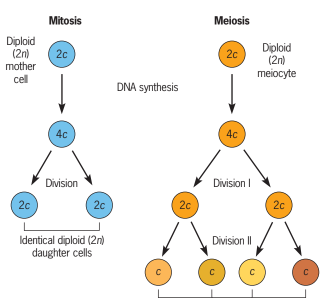
If we look at the chromosomes in a diploid cell, we find that they come in pairs (**Figure 2.7**). For example, somatic human cells have 23 pairs of chromosomes. Each pair is distinct. Different pairs of chromosomes carry different sets of genes. The members of a pair are called homologous chromosomes, or simply **homologues**, from a Greek word meaning "in agreement with." Homologues carry the same set of genes, although as we will see in Chapter 5, they may carry different alleles of these genes. Chromosomes from different pairs are called **heterologues**. During meiosis, homologues associate intimately with each other. This association is the basis of an orderly process that ultimately reduces the chromosome number to the haploid state. The reduction in chromosome number occurs in such a way that each of the resulting haploid cells receives exactly one member of each chromosome pair.

The process of meiosis involves two cell divisions (**■** Figure 2.8). Chromosome duplication, which is associated with DNA synthesis, occurs prior to the first of these divisions. It does not occur between the two divisions. Thus, the progression of events is: chromosome duplication \rightarrow meiotic division I \rightarrow meiotic division II. If we represent the haploid amount of DNA by the letter *c*, then in sequence, these events double the amount of DNA (from 2*c* to 4*c*), cut it in half (from 4*c* to 2*c*), and finally cut it in half again (from 2*c* to *c*). The overall effect is to reduce the diploid chromosome number (2*x*)

chromosome number (2*n*) to the haploid chromosome number (*n*). You can test your understanding of this overall process by working through Solve It: How Much DNA in Human Meiotic Cells?

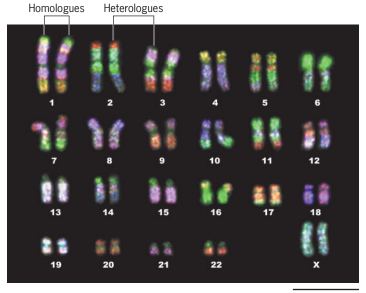
MEIOSIS I

The events in the two meiotic divisions are illustrated in Figure 2.9. The first meiotic division is complicated and protracted. When it begins, the chromosomes have already been duplicated; consequently, each of them consists of two



Nonidentical haploid (n) cells

Sexual reproduction involves a mechanism that reduces the number of chromosomes by half.



10 µm

FIGURE 2.7 The 23 pairs of homologous chromosomes found in human cells.

Solve It!

How Much DNA in Human Meiotic Cells?

If a human sperm cell contains 3.2 billion base pairs of DNA, how many base pairs are present in (a) a diploid cell that has duplicated its DNA in preparation to enter meiosis, (b) a cell emerging from the first meiotic division, and (c) a cell emerging from the second meiotic division?

To see the solution to this problem, visit the Student Companion site.

FIGURE 2.8 Comparison between mitosis and meiosis; *c* is the haploid amount of DNA in the genome.

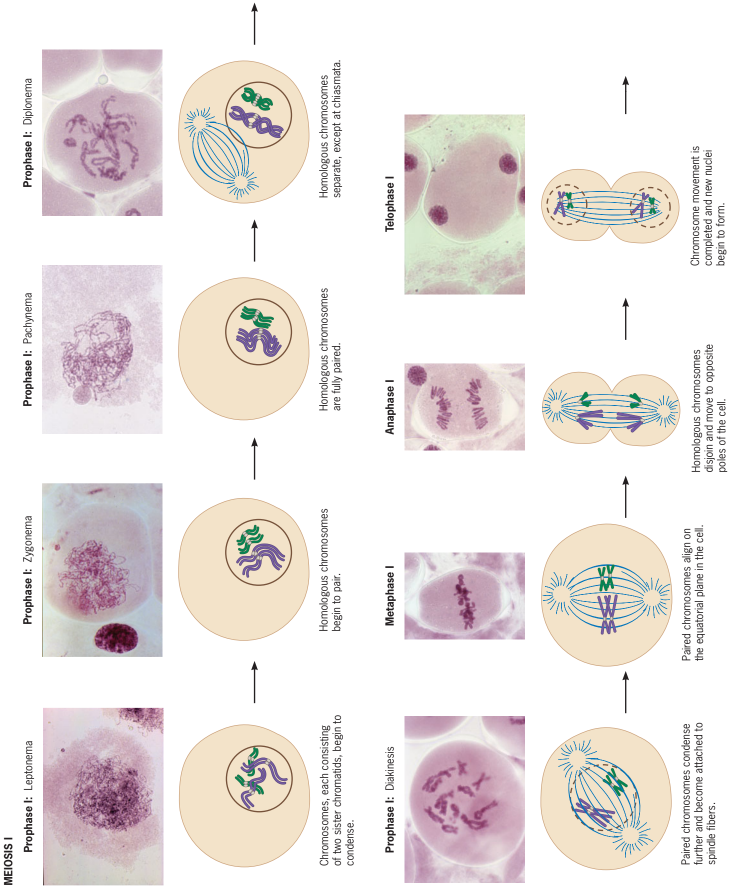


FIGURE 2.9 Meiosis in the plant Lilium longiflorum.

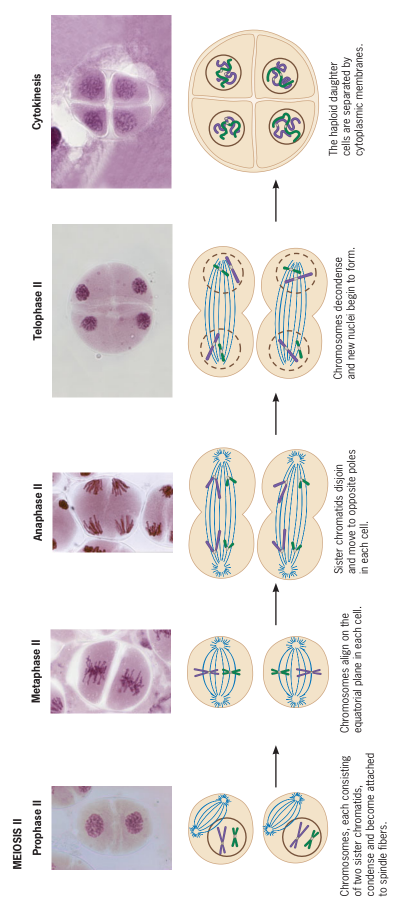


FIGURE 2.9 [continued]

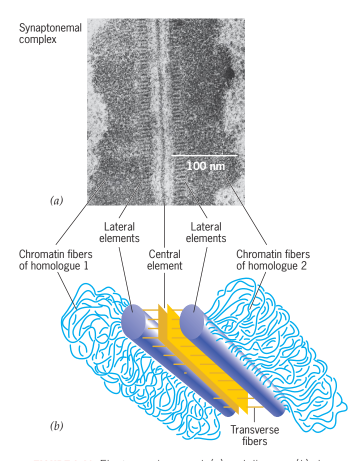


FIGURE 2.10 Electron micrograph (*a*) and diagram (*b*) showing the structure of the synaptonemal complex that forms between homologous chromosomes during prophase I of meiosis.

sister chromatids. The prophase of meiosis I—or simply, **prophase I** is divided into five stages, each denoted by a Greek term. These terms convey key features about the appearance or behavior of the chromosomes.

Leptonema, from Greek words meaning "thin threads," is the earliest stage of prophase I. During leptonema (also referred to as the leptotene stage) the duplicated chromosomes condense out of the diffuse chromatin network. With a light microscope, individual chromosomes can barely be seen, but with an electron microscope, each of the chromosomes appears to consist of two sister chromatids. As chromosome condensation continues, the cell progresses into zygonema (from Greek words meaning "paired threads"). During zygonema (also the zygotene stage), homologous chromosomes come together intimately. This process of pairing between homologues is called synapsis. In some species, synapsis begins at the ends of chromosomes and then spreads toward their middle regions. Synapsis is usually accompanied by the formation of a protein structure between the pairing chromosomes (■ Figure 2.10). This structure, called the synaptonemal complex, consists of three parallel rods—one associated with each of the chromosomes (called the lateral elements) and one located midway between them (called the central element)-and a large number of ladderlike rungs connecting the lateral elements with the central element. The role of the synaptonemal complex in chromosome pairing and in subsequent meiotic events is not fully understood. In some types of meiotic cells it does not even appear. Thus, it may not be absolutely essential for pairing during prophase I. The process by which homologues find each other in prophase I also is not well understood. Recent studies suggest that homologues may actually begin to pair early in meiosis I, during leptonema. This pairing may be facilitated by a tendency for homologous chromosomes to remain in the same region of the nucleus during interphase. Thus, homologues may not have far to go to find each other.

As synapsis progresses, the duplicated chromosomes continue to condense into smaller volumes. The thickened chromosomes that result from this process are characteristic of pachynema (from Greek words for "thick threads"). At pachynema (also the pachytene stage), paired chromosomes can easily be seen with a light microscope. Each pair consists of two duplicated homologues, which themselves consist of two sister chromatids. If we count homologues, the pair is referred to as a **bivalent** of chromosomes, whereas if we count strands, it is referred to as a tetrad of chromatids. During pachynema—or perhaps a bit before or after, the paired chromosomes may exchange material (**Figure 2.11**). We will explore this phenomenon, called **cross**ing over, and its consequences in Chapter 7. Here, suffice it to say that individual sister chromatids may be broken during pachynema, and the broken pieces may be swapped between chromatids within a tetrad. The breakage and reunion that occur during crossing over may therefore lead to recombination of genetic material between the paired chromosomes. The fact that these types of exchanges have occurred can be seen as the cell progress to the next stage of meiosis I, diplonema (from Greek words for "two threads"). During diplonema (also the diplotene stage), the paired chromosomes separate slightly. However, they remain in close contact where they have crossed over. These contact points are called chiasmata (singular, chiasma, from a Greek word meaning "cross"). Close examination of the chiasmata indicates that each of them involves only two of the four chromatids in the tetrad. The diplotene stage may last a very long time. In human females, for example, it may persist for more than 40 years.

Near the end of prophase I, the chromosomes condense further, the nuclear membrane fragments, and a spindle apparatus forms. Spindle microtubules penetrate into the nuclear space and attach to the kinetochores of the chromosomes.

The chromosomes, still held together by the chiasmata, then move to a central plane of the cell that is perpendicular to the axis of the spindle apparatus. This movement is characteristic of the last stage of prophase I, called **diakinesis** (from Greek words meaning "movement through").

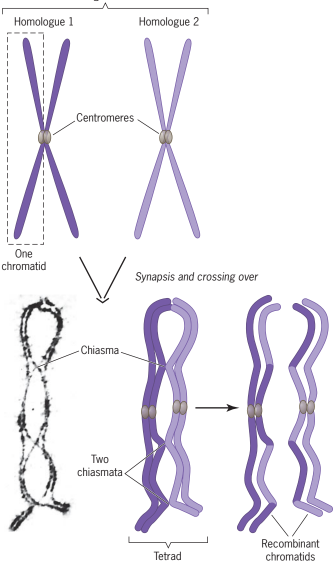
During metaphase I, the paired chromosomes orient toward opposite poles of the spindle. This orientation ensures that when the cell divides, one member of each pair will go to each pole. At the end of prophase I and during metaphase I, the chiasmata that hold the bivalents together slip away from the centromeres toward the ends of the chromosomes. This phenomenon, called terminalization, reflects the growing repulsion between the members of each chromosome pair. During anaphase I, the paired chromosomes separate from each other definitively. This separation, called *chromosome disjunction*, is mediated by the spindle apparatus acting on each of the bivalents in the cell. As the separating chromosomes gather at opposite poles, the first meiotic division comes to an end. During the next stage, called **telophase I**, the spindle apparatus is disassembled, the daughter cells are separated from each other by membranes, the chromosomes are decondensed, and a nucleus is formed around the chromosomes in each daughter cell. In some species, chromosome decondensation is incomplete, the daughter nuclei do not form, and the daughter cells proceed immediately into the second meiotic division. The cells produced by meiosis I contain the haploid number of chromosomes; however, each chromosome still consists of two sister chromatids, which may not be genetically identical because they might have exchanged material with their pairing partners during prophase I.

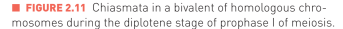
MEIOSIS II AND THE OUTCOMES OF MEIOSIS

During meiosis II, the chromosomes condense and become attached to a new spindle apparatus (prophase II). They then move to positions in the equatorial plane of the cell (metaphase II), and their centromeres split to allow the constituent sister chromatids to move to opposite poles (anaphase II), a phenomenon called *chromatid disjunction*. During telophase II, the separated chromatids—now called chromosomes—gather at the poles and daughter nuclei form around them. Each daughter nucleus contains a haploid set of chromosomes. Mechanistically, meiosis II is therefore much like mitosis. However, its products are haploid, and unlike the products of mitosis, the cells that emerge from meiosis II are not genetically identical.

One reason these cells differ is that homologous chromosomes pair and disjoin from each other during meiosis I. Within each pair of chromosomes, one homologue was inherited from the organism's mother, and the other was inherited from its father. During meiosis I, the maternally and paternally inherited homologues come together and synapse. They are positioned on the meiotic spindle and become oriented randomly with respect to the spindle's poles. Then they disjoin. For each pair of chromosomes, half the daughter cells produced by the first meiotic division receive the maternally inherited homologue, and the other half receive the paternally inherited homologue. Thus, from the end of the first meiotic division, the products of meiosis are destined to be different. These differences are compounded by the number of chromosome pairs that disjoin during meiosis I. Each of the pairs disjoins independently. Thus, if there are 23 pairs of chromosomes, as there are in humans, meiosis I can produce 2²³ chromosomally different daughter cells—that is, more than 8 million possibilities. To test your understanding of this concept go to Solve It: How Many Chromosome Combinations in Sperm?







Solve It!

How Many Chromosome Combinations in Sperm?

The fruit fly *Drosophila melanogaster* has four pairs of chromosomes in its somatic cells. In the female fly, crossing over occurs between maternally and paternally inherited homologues during prophase I of meiosis. In the male fly, crossing over does not occur. Given this fact, how many chromosomally distinct types of sperm can be produced by a male fruit fly?

To see the solution to this problem, visit the Student Companion site. Another reason the cells that emerge from meiosis differ is that during meiosis I, homologous chromosomes exchange material by crossing over. This process can create countless different combinations of genes. When we superimpose the variability created by crossing over on the variability created by the random disjunction of homologues, it is easy to see that no two products of meiosis are likely to be the same.

KEY POINTS •

- Diploid eukaryotic cells form haploid cells by meiosis, a process involving one round of chromosome duplication followed by two cell divisions (meiosis I and meiosis II).
 - During meiosis I, homologous chromosomes pair (synapse), exchange material (cross over), and separate (disjoin) from each other.
 - During meiosis II, chromatids disjoin from each other.

Life Cycles of Some Model Genetic Organisms

Geneticists focus their research on microorganisms, plants, and animals well suited to experimentation.

When genetics began, the organisms that were used for research were the ones that came to hand from the garden or the barnyard. Some early geneticists branched out to study inheritance in other types of creatures—moths and canaries, for example—and as genetics progressed, research became focused on organisms that were well suited

for controlled experimentation in laboratories or field plots. Today a select group of microorganisms, plants, and animals are favored in genetic research. These creatures, often called **model organisms**, lend themselves well to genetic analysis. For the most part, they are easily cultured in the laboratory, their life cycles are relatively short, and they are genetically variable. In addition, through work over many years, geneticists have established large collections of mutant strains for these organisms. We will encounter the model genetic organisms many times in this book. **Table 2.1** summarizes information about several of them, and in the sections that follow, we discuss the life cycles of three of these genetically important species.

SACCHAROMYCES CEREVISIAE, BAKER'S YEAST

Baker's yeast came into genetics research in the first half of the twentieth century. However, long before it was commonplace in genetics laboratories, this organism was used in kitchens as a leavening agent for making bread. Yeast is a unicellular fungus, although under some conditions, its cells divide to form long filaments. Yeast cells can

TABLE 2.1

Some Important Model Genetic Organisms

Organism	Haploid Chromosome Number	Genome Size (in millions of base pairs)	Gene Number
Saccharomyces cerevisiae (yeast)	16	12	6,268
Arabidopsis thaliana (flowering plant)	5	157	27,706
Caenorhabitis elegans (worm)	5	100	21,733
Drosophila melanogaster (fly)	4	170	17,000
Danio rerio (zebra fish) Mus musculus (mouse)	25 20	1,600 2,900	23,524 25,396

be cultured on simple media in the laboratory, and large numbers of cells can be obtained from a single mother cell in just a few days. In addition, mutant strains with different growth characteristics can be readily isolated.

Saccharomyces cerevisiae reproduces both sexually and asexually (**Figure 2.12**). Asexual reproduction occurs by a process called budding, which involves a mitotic division of the haploid nucleus. After this division, one daughter nucleus moves into a small "bud" or progeny cell. Eventually, the bud is separated from the mother cell by cytokinesis. Sexual reproduction in *S. cerevisiae* occurs when haploid cells of opposite mating types (denoted *a* and *alpha*) come together—an event referred to as mating—to fuse and form a diploid cell, which then undergoes meiosis. The four haploid products of meiosis are created in a sac called the ascus (plural, asci), and each of the products is called an ascospore. By dissecting this sac, a researcher can isolate each meiotic product and place it in a culture dish to start a new yeast colony.

Asexual "budding" Asexual "budding" Asexual "budding" Asexual "budding" Asexual "budding" Asexual "Mating" Asexual Ase

ARABIDOPSIS THALIANA, A FAST-GROWING PLANT

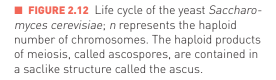
Garden plants were the first organisms to be studied genetically. Today geneticists focus their attention on *Arabidopsis thaliana*, a weed sometimes called the mouse ear cress. This fast-growing species is related to food plants such as radish, cabbage, and canola; however, it has no agronomic or horticultural value.

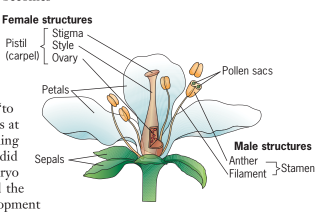
The reproductive organs of *Arabidopsis* are located in its flowers (**■** Figure 2.13). The male gametes are produced by meiosis in **anthers**, which are atop the **stamens**. The female gametes are produced by meiosis in the **ovary**, which is located within the **pistil** at the center of the flower. In plants such as *Arabidopsis*, these meiotic products are usually referred to as **microspores** (from male meiosis) or as **megaspores** (from female meiosis).

Compared to yeast, *Arabidopsis* reproduction is complex (**Figure 2.14**). The mature plant is called a **sporophyte** because it produces microspores and megaspores; the suffix "phyte" in this term is derived from the Greek word for plant. On the male side of *Arabidopsis* reproduction, each diploid microspore mother cell—alas, this type of cell is not called a microspore father cell as you might expect—undergoes meiosis to produce four haploid microspores. Each microspore then undergoes mitosis to produce a **pollen grain**, which contains two generative or sperm cells located within a vegetative cell; the nuclei in the sperm cells and the vegetative cell are all haploid and identical to each other. This trio of nuclei within the pollen grain constitutes the **male gametophyte** of *Arabidopsis*. The botanical term "gametophyte" derives from the fact that the pollen is, in effect, a very tiny plant that holds the male gametes.

On the female side of *Arabidopsis* reproduction, each diploid megaspore mother cell undergoes meiosis to produce four haploid cells; however, three of these cells subsequently degenerate, leaving only one functional meiotic product, which becomes

a megaspore. The haploid nucleus in the megaspore then undergoes three mitotic divisions to produce a total of eight identical haploid nuclei within a structure called the embryo sac. When cytokinesis occurs, six of Pistil these eight nuclei become separated from each other by cell membranes. Three of the resulting cells move to the top of the embryo sac and three move to the bottom. One of the cells at the bottom becomes the egg and the other two become synergid cells, named from Greek words meaning "to work together" because these cells remain alongside the egg. The three cells at the top of the embryo sac are called antipodal cells, from Greek words meaning "on the opposite side of." They will soon degenerate. The two nuclei that did not become enclosed by cell membranes remain in the center of the embryo sac. These polar nuclei subsequently fuse to form a diploid nucleus, called the secondary endosperm nucleus, which will later play a key role in the development of nutritive tissue in the seed. The cells and nuclei within the embryo sac make up the female gametophyte of Arabidopsis.







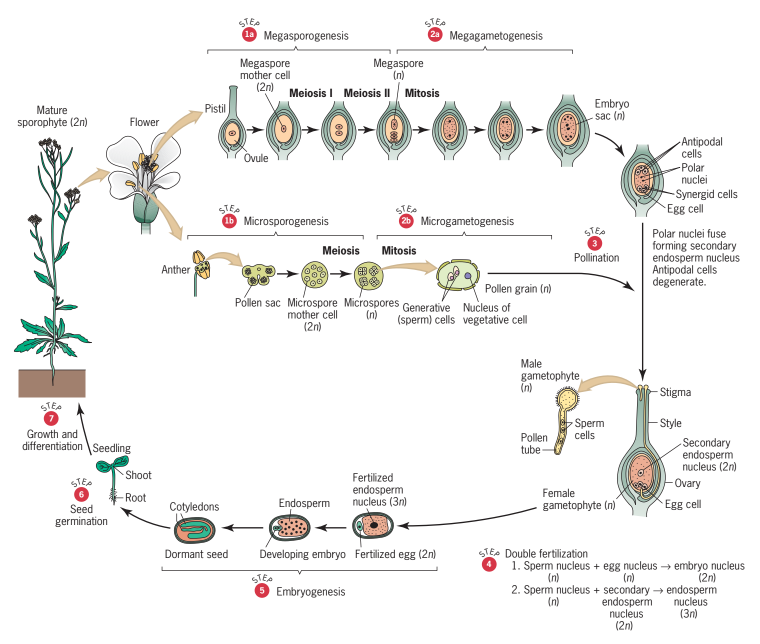


FIGURE 2.14 The life cycle of the model plant, Arabidopsis thaliana.

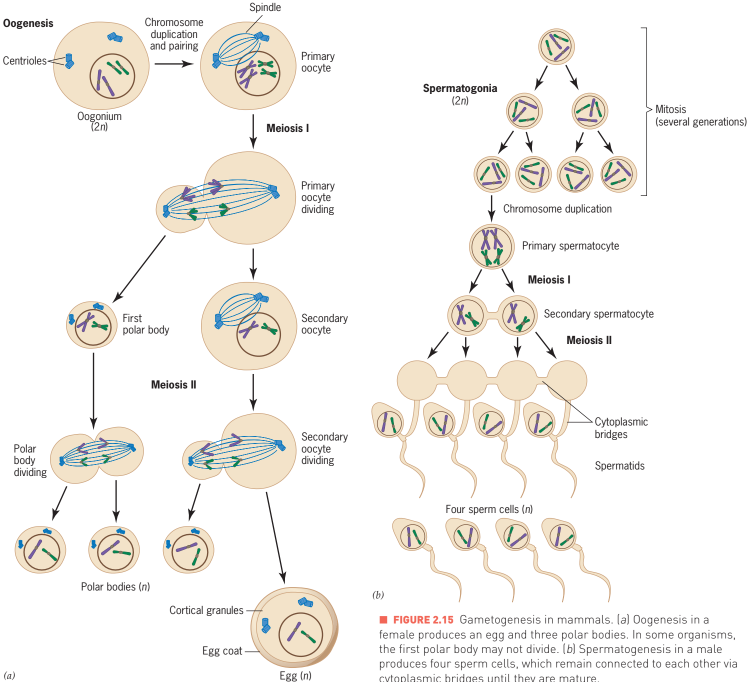
When a mature pollen grain lands on the stigma atop the pistil, a pollen tube grows down through the style to an egg cell within the ovary. In plants such as *Arabidopsis*, fertilization involves two events. (1) One sperm cell within the pollen tube fuses with the egg cell in the female gametophyte to form the *diploid zygote*, which will subsequently grow into an embryo. (2) The other sperm cell nucleus combines with the diploid secondary endosperm nucleus in the female gametophyte to form the *triploid endosperm nucleus*, which will subsequently direct the development of nutritive tissue (the endosperm) to feed the embryo when the seed that surrounds it germinates. It takes about five weeks for an *Arabidopsis* plant to reach maturity—short compared to other flowering plants. Scientists who work with *Arabidopsis* can therefore make fairly rapid progress in their research projects.

MUS MUSCULUS, THE MOUSE

The mouse has been especially important in biomedical research. Mice have been the subjects of innumerable projects to ascertain the effects of drugs, chemicals, foods,

and other materials relevant to human health. Mouse genetics began early in the twentieth century with studies on the inheritance of coat color, and since those days, it has developed into an impressive enterprise.

Mice, like humans, have separate sexes. The formation of gametes—a process called gametogenesis—occurs in the gonads of each sex. Oogenesis, the formation of eggs, occurs in the ovaries, which are the female gonads, and spermatogenesis, the formation of sperm, occurs in the testes, which are the male gonads. These processes begin when undifferentiated diploid cells, called oogonia or spermatogonia, undergo meiosis to produce haploid cells. The haploid cells then differentiate into mature gametes (**Figure 2.15**). Usually, only one of the four haploid cells from female meiosis becomes an egg, or ovum; the other three cells, called polar bodies, degenerate. By contrast, all four of the haploid cells from male meiosis



cytoplasmic bridges until they are mature.

PROBLEM-SOLVING SKILLS

Counting Chromosomes and Chromatids

THE PROBLEM

The cat (*Felis domesticus*) has 36 pairs of chromosomes in its somatic cells. (a) How many chromosomes are present in a cat's mature sperm cells? (b) How many sister chromatids are present in a cell that is entering the first meiotic division? (c) In a cell that is entering the second meiotic divison?

FACTS AND CONCEPTS

- Chromosomes come in pairs—that is, there are two homologous chromosomes in each pair.
- **2.** Chromosome duplication creates *two* sister chromatids for each chromosome in the cell.
- **3.** The first meiotic division reduces the number of duplicated chromosomes (and the number of sister chromatids present) by a factor of *two*.
- **4.** The second meiotic division reduces the number of sister chromatids by another factor of *two*.

ANALYSIS AND SOLUTION

- a. If the cat has 36 pairs of chromosomes in its diploid somatic cells—that is, 2 × 36 = 72 chromosomes altogether—a haploid sperm cell, which is an end product of meiosis, should have half as many chromosomes—that is, 72/2 = 36, or one chromosome from each homologous pair.
- **b.** A cell that is entering the first meiotic division has just duplicated its 72 chromosomes. Because each chromosome now consists of two sister chromatids, altogether $72 \times 2 = 144$ sister chromatids are present in this cell.
- **c.** A cell that is entering the second meiotic division has one homologue from each of the 36 homologous chromosome pairs, and each of these homologues consists of two sister chromatids. Consequently, such a cell has $36 \times 2 = 72$ sister chromatids.

For further discussion visit the Student Companion site.

develop into sperm. The process of gametogenesis is similar in other mammals. To assess your understanding of how the number of chromosomes is reduced during this process, work through Problem-Solving Skills: Counting Chromosomes and Chromatids.

Mice are sexually mature by about 7–8 weeks of age. Some research institutions maintain large breeding colonies to provide animals for various projects. As you might imagine, research that involves mice is significantly more time-consuming and expensive than research with other model organisms. However, because the mouse is the model most closely related to humans, research with it can provide important insights into issues of human health and disease.

Unlike yeast, *Arabidopsis*, or mice, our own species cannot be subjected to genetic experimentation. In the strictest sense, *Homo sapiens* is therefore not a model organism. However, we have learned to grow human cells in culture, and this advance has made it possible to study human genetic material in the laboratory. A Milestone in Genetics: Culturing Human Cells, which you can find in the Student Companion site, provides some details.

KEY POINTS

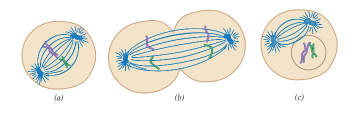
 In yeast, haploid cells with opposite mating types fuse to form a diploid zygote, which then undergoes meiosis to produce four haploid cells.

- Meiosis in the reproductive organs of Arabidopsis produces microspores and megaspores, which subsequently develop into male and female gametophytes.
- The double fertilization that occurs during Arabidopsis reproduction creates a diploid zygote, which develops into an embryo, and a triploid endosperm, which develops into nutritive tissue in the seed.
- In mice and other mammals, one cell from female meiosis becomes the egg, whereas all four cells from male meiosis become sperm.

Basic Exercises

Illustrate Basic Genetic Analysis

1. Identify the stages of mitosis in the following drawings.

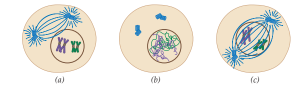


Answer: (*a*) metaphase; (*b*) anaphase; (*c*) prophase

- **2.** Why does a diploid mother cell that undergoes meiosis produce four haploid cells?
- Answer: During meiosis, chromosome duplication precedes two division events. If the number of chromosomes in the diploid mother cell is 2n, then after duplication, the cell contains 4n chromatids. During the first meiotic division, homologous chromosomes pair and then are separated into different daughter cells, each of which receives 2n chromatids. During the second meiotic division, the centromere that holds the two chromatids of each chromosome together splits and the chromatids are separated into different daughter cells. Each of the four cells resulting from these successive meiotic divi-

sions therefore contains n chromatids (now called chromosomes). Thus, the diploid state of the mother cell is reduced to the haploid state in the four cells that emerge from meiosis.

3. Identify the stages of prophase I of meiosis in the following drawings.



Answer: (*a*) diplonema; (*b*) leptonema; (*c*) diakinesis

- 4. Twenty pairs of chromosomes are present in a somatic cell of the mouse. How many sister chromatids are present in (a) a primary oocyte, (b) a secondary spermatocyte, (c) a mature sperm cell?
- Answer: (a) 80, because each of the 40 chromosomes (20 pairs × 2 chromosomes/pair) had been duplicated prior to the cell's entry into meiosis I; (b) 40, because homologous chromosomes (each still consisting of two sister chromatids) were apportioned to different cells during the first meiotic division; (c) 20, the haploid chromosome number.

Testing Your Knowledge

Integrate Different Concepts and Techniques

- **1.** What are the principal differences between mitosis and meiosis?
- Answer: In mitosis, one division event follows one round of chromosome duplication. In meiosis, two division events follow one round of chromosome duplication. Furthermore, during the first meiotic division, homologous chromosomes pair with each other. This homology-based pairing does not normally occur during mitosis. The two cells produced by a mitotic division are identical to each other and to the mother cell from which they were derived. The four cells produced by the two successive meiotic divisions are not identical to each other or to the mother cell from which they were derived. When a diploid cell undergoes mitosis, the two cells derived from it will also be diploid. When a diploid cell undergoes meiosis, the four cells derived from it will be haploid.
- 2. Caenorbabditis elegans, a small nonparastic worm, is used in genetics research. Some of these worms are hermaphrodites capable of producing both eggs and sperm. C. elegans hermaphrodites have 5 pairs of chromosomes. How many chromosomes are present (a) in a sperm cell from a hermaphrodite? (b) in a fertilized egg from a hermaphrodite? How many sister chromatids are present in a hermaphrodite's cell that (c) is entering the first meiotic division? (d) is entering the second meiotic division? (e) has completed the second meiotic division?
- **Answer:** (a) 5, because sperm are haploid. (b) 10, because a fertilized egg contains chromosomes from the egg and the sperm that fertilized it. (c) 20, because each of the 10 chromosomes in a cell entering meiosis I has been duplicated to produce two sister chromatids. (d) 10, because homologous chromosomes have been apportioned to different cells during the

first meiotic division; however, the sister chromatids of each homologue are still held together by a common centromere. (e) 5, because the end products of meiosis are haploid.

- 3. A human sperm cell contains about $3.2 \times 10^{\circ}$ nucleotide pairs of DNA. How much DNA is present in each of the following: (a) a primary human spermatocyte; (b) a secondary human spermatocyte; (c) the first polar body produced by division of a primary oocyte?
- **Answer:** (a) $4 \times 3.2 \times 10^9 = 12.8 \times 10^9$ nucleotide pairs because a primary spermatocyte contains the 4*c* amount of DNA; (b) $2 \times 3.2 \times 10^9 = 6.4 \times 10^9$ nucleotide pairs because a secondary spermatocyte contains the 2*c* amount of DNA; (c) $2 \times 3.2 \times 10^9 = 6.4 \times 10^9$ nucleotide pairs because a first polar body contains the 2*c* amount of DNA.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **2.1** Carbohydrates and proteins are linear polymers. What types of molecules combine to form these polymers?
- **2.2** All cells are surrounded by a membrane; some cells are surrounded by a wall. What are the differences between cell membranes and cell walls?
- **2.3** What are the principal differences between prokaryotic and eukaryotic cells?
- **2.4** Distinguish between the haploid and diploid states. What types of cells are haploid? What types of cells are diploid?
- **2.5** Compare the sizes and structures of prokaryotic and eukaryotic chromosomes.
- **2.6** With a focus on the chromosomes, what are the key events during interphase and M phase in the eukaryotic cell cycle?
- **2.7** Which typically lasts longer, interphase or M phase? Can you explain why one of these phases lasts longer than the other?
- **2.8** In what way do the microtubule organizing centers of plant and animal cells differ?
- 2.9 Match the stages of mitosis with the events they encompass: Stages: (1) anaphase, (2) metaphase, (3) prophase, (4) telophase. Events: (a) re-formation of the nucleolus, (b) disappearance of the nuclear membrane, (c) condensation of the chromosomes, (d) formation of the mitotic spindle, (e) movement of chromosomes to the equatorial plane, (f) movement of chromosomes to the poles, (g) decondensation of the chromosomes, (h) splitting of the centromere, (i) attachment of microtubules to the kinetochore.
- 2.10 Arrange the following events in the correct temporal sequence during eukaryotic cell division, starting with the earliest: (a) condensation of the chromosomes, (b) movement of chromosomes to the poles, (c) duplication of the chromosomes, (d) formation of the nuclear membrane, (e) attachment of microtubules to the kinetochores, (f) migration of centrosomes to positions on opposite sides of the nucleus.
- 2.11 In humans, the gene for β -globin is located on chromosome 11, and the gene for α -globin, which is another component

of the hemoglobin protein, is located on chromosome 16. Would these two chromosomes be expected to pair with each other during meiosis? Explain your answer.

- 2.12 The A sperm cell from the fruit fly *Drosophila melanogaster* contains four chromosomes. How many *chromosomes* would be present in a spermatogonial cell about to enter meiosis? How many *chromatids* would be present in a spermatogonial cell at metaphase I of meiosis? How many would be present at metaphase II?
- **2.13** Does crossing over occur before or after chromosome duplication in cells going through meiosis?
- **2.14** What visible characteristics of chromosomes indicate that they have undergone crossing over during meiosis?
- **2.15** During meiosis, when does *chromosome* disjunction occur? When does *chromatid* disjunction occur?
- **2.16** In *Arabidopsis*, is leaf tissue haploid or diploid? How many nuclei are present in the female gametophyte? How many are present in the male gametophyte? Are these nuclei haploid or diploid?
- **2.17** From the information given in Table 1 in this chapter, is there a relationship between genome size (measured in base pairs of DNA) and gene number? Explain.
- **2.18** Are the synergid cells in an *Arabidopsis* female gametophyte genetically identical to the egg cell nestled between them?
- **2.19** A cell of the bacterium *Escherichia coli*, a prokaryote, contains one chromosome with about 4.6 million base pairs of DNA comprising 4288 protein-encoding genes. A cell of the yeast *Saccharomyces cerevisiae*, a eukaryote, contains about 12 million base pairs of DNA comprising 6268 genes, and this DNA is distributed over 16 distinct chromosomes. Are you surprised that the chromosome of a prokaryote is larger than some of the chromosomes of a eukaryote? Explain your answer.
- **2.20** Given the way that chromosomes behave during meiosis, is there any advantage for an organism to have an even number of chromosome pairs (such as the fruit fly *Drosophila* does), as opposed to an odd number of chromosome pairs (such as humans do)?

- **2.21** In flowering plants, two nuclei from the pollen grain participate in the events of fertilization. With which nuclei from the female gametophyte do these nuclei combine? What tissues are formed from the fertilization events?
- **2.22** The mouse haploid genome contains about 2.9×10^9 nucleotide pairs of DNA. How many nucleotide pairs of DNA are present in each of the following mouse cells: (a) somatic cell, (b) sperm cell, (c) fertilized egg,

(d) primary oocyte, (e) first polar body, (f) secondary spermatocyte?

2.23 *Arabidopsis* plants have 10 chromosomes (5 pairs) in their somatic cells. How many chromosomes are present in each of the following: (a) egg cell nucleus in the female gametophyte, (b) generative cell nucleus in a pollen grain, (c) fertilized endosperm nucleus, (d) fertilized egg nucleus?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

- 1. Find out more about the model organisms mentioned in this chapter by clicking on More about NCBI and then on the Model Organisms Guide. From there, investigate mammalian, nonmammalian, and other model organisms.
- **2.** Use the links on the NCBI web site to locate web sites dedicated to each of the eukaryotic model organisms mentioned

in this chapter: SGD (*Saccharomyces* Genome Database), Flybase, WormBase, ZIRC (Zebrafish International Resource Center), MGI (Mouse Genomic Informatics), and TAIR (The *Arabidopsis* Information Resource).



Mendelism

The Basic Principles of Inheritance

CHAPTER OUTLINE

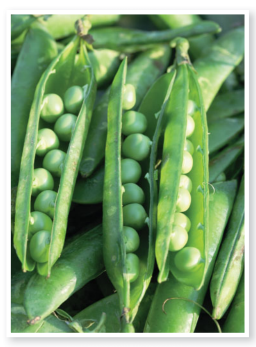
- Mendel's Study of Heredity
- Applications of Mendel's Principles
- Testing Genetic Hypotheses
- Mendelian Principles in Human Genetics

The Birth of Genetics: A Scientific Revolution

Science is a complex endeavor involving careful observation of natural phenomena, reflective thinking about these phenomena, and formula-

tion of testable ideas about their causes and effects. Progress in science often depends on the work of a single insightful individual. Consider, for example, the effect that Nicolaus Copernicus had on astronomy, that Isaac Newton had on physics, or that Charles Darwin had on biology. Each of these individuals altered the course of his scientific discipline by introducing radically new ideas. In effect, they began scientific revolutions.

In the middle of the nineteenth century, the Austrian monk Gregor Mendel, a contemporary of Darwin, laid the foundation for another revolution in biology, one that eventually produced an entirely new science—genetics. Mendel's ideas, published in 1866 under the title "Experiments in Plant Hybridization," endeavored to explain how the characteristics of organisms are inherited. Many people had attempted such an explanation previously but without much success. Indeed, Mendel commented on their failures in the opening paragraphs of his article:



Pisum sativum, the subject of Gregor Mendel's experiments.

[However,] Those who survey the work in this department will arrive at the conviction that among all the numerous experiments made, not one has been carried out to such an extent and in such a way as to make it possible to determine the number of different forms under which the offspring of the hybrids appear, or to arrange

> these forms with certainty according to their separate generations, or definitely to ascertain their statistical relations.¹

He then described his own efforts to elucidate the mechanism of heredity:

It requires indeed some courage to undertake a labor of such far-reaching extent; this appears, however, to be the only right way by which we can finally reach the solution of a question the importance of which cannot be overestimated in connection with the history of the evolution of organic forms.

The paper now presented records the results of such a detailed experiment. This experiment was practically confined to a small plant group, and is now, after eight years' pursuit, concluded in all essentials. Whether the plan upon which the separate experiments were conducted and carried out was the best suited to attain the desired end is left to the friendly decision of the reader.²

To this object, numerous careful observers, such as Kölreuter, Gärtner, Herbert, Lecoq, Wichura and others, have devoted a part of their lives with inexhaustible perseverance....

^{1,2}Peters, J. A., ed. 1959. *Classic Papers in Genetics*. Prentice-Hall, Englewood Cliffs, NJ.

Mendel's Study of Heredity

The life of Gregor Johann Mendel (1822–1884) spanned the middle of the nineteenth century. His parents were farmers in Moravia, then a part of the Hapsburg Empire in Central Europe. A rural upbring-

Gregor Mendel's experiments with peas elucidated how traits are inherited.

ing taught him plant and animal husbandry and inspired an interest in nature. At the age of 21, Mendel left the farm and entered a Catholic monastery in the city of Brünn (today, Brno in the Czech Republic). In 1847 he was ordained a priest, adopting the clerical name Gregor. He subsequently taught at the local high school, taking time out between 1851 and 1853 to study at the University of Vienna. After returning to Brünn, he resumed his life as a teaching monk and began the genetic experiments that eventually made him famous.

Mendel performed experiments with several species of garden plants, and he even tried some experiments with honeybees. His greatest success, however, was with peas. He completed his experiments with peas in 1864. In 1865, Mendel presented the results before the local Natural History Society, and the following year, he published a detailed report in the society's proceedings. Unfortunately, this paper languished in obscurity until 1900, when it was rediscovered by three botanists—Hugo de Vries in Holland, Carl Correns in Germany, and Eric von Tschermak-Seysenegg in Austria. As these men searched the scientific literature for data supporting their own theories of heredity, each found that Mendel had performed a detailed and careful analysis 35 years earlier. Mendel's ideas quickly gained acceptance, especially through the promotional efforts of a British biologist, William Bateson. This champion of Mendel's discoveries coined a new term to describe the study of heredity: genetics, from the Greek word meaning "to generate."

MENDEL'S EXPERIMENTAL ORGANISM, THE GARDEN PEA

One reason for Mendel's success is that he chose his experimental material astutely. The garden pea, *Pisum sativum*, is easily grown in experimental gardens or in pots in a greenhouse. Pea flowers contain both male and female organs. The male organs, called anthers, produce sperm-containing pollen, and the female organ, called the ovary, produces eggs.

One peculiarity of pea reproduction is that the petals of the flower close down tightly, preventing pollen grains from entering or leaving. This enforces a system of self-fertilization, in which male and female gametes from the same flower unite with each other to produce seeds. As a result, individual pea strains are highly inbred, displaying little if any genetic variation from one generation to the next. Because of this uniformity, we say that such strains are *true-breeding*.

At the outset, Mendel obtained many different true-breeding varieties of peas, each distinguished by a particular characteristic. In one strain, the plants were 2 meters high, whereas in another they measured only a half meter. Another variety produced green seeds, and still another produced yellow seeds. Mendel took advantage of these contrasting traits to determine how the characteristics of pea plants are inherited. His focus on these singular differences between pea strains allowed him to study the inheritance of one trait at a time—for example, plant height. Other biologists had attempted to follow the inheritance of many traits simultaneously, but because the results of such experiments were complex, they were unable to discover any fundamental principles about heredity. Mendel succeeded where these biologists had failed because he focused his attention on contrasting differences between plants that were otherwise the same—tall versus short, green seeds versus yellow seeds, and so forth. In addition, he kept careful records of the experiments that he performed.

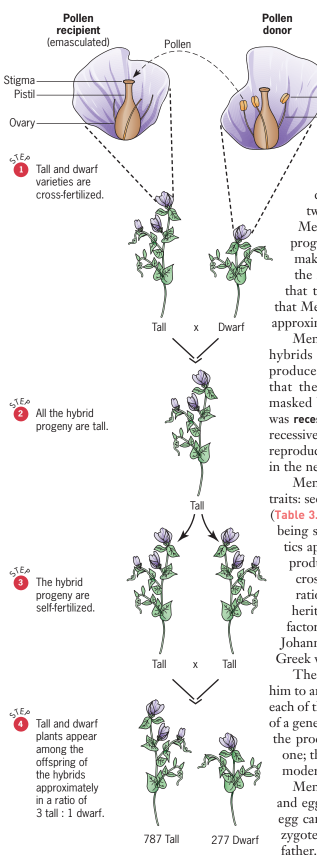


FIGURE 3.1 Mendel's crosses involving tall and dwarf varieties of peas.

MONOHYBRID CROSSES: THE PRINCIPLES **OF DOMINANCE AND SEGREGATION**

In one experiment, Mendel cross-fertilized—or, simply, crossed tall and dwarf pea plants to investigate how height was inherited (**Figure 3.1**). He carefully removed the anthers from one variety before its pollen had matured and then applied pollen from Stamen the other variety to the stigma, a sticky organ on top of the pistil that leads to the ovary. The seeds that resulted from these crossfertilizations were sown the next year, yielding hybrids that were uniformly

tall. Mendel obtained tall plants regardless of the way he performed the cross (tall male with dwarf female or dwarf male with tall female); thus, the two reciprocal crosses gave the same results. Even more significantly, however, Mendel noted that the dwarf characteristic seemed to have disappeared in the progeny of the cross, for all the hybrid plants were tall. To explore the hereditary makeup of these tall hybrids, Mendel allowed them to undergo self-fertilizationthe natural course of events in peas. When he examined the progeny, he found that they consisted of both tall and dwarf plants. In fact, among 1064 progeny that Mendel cultivated in his garden, 787 were tall and 277 were dwarf-a ratio of approximately 3:1.

Anther

Mendel was struck by the reappearance of the dwarf characteristic. Clearly, the hybrids that he had made by crossing tall and dwarf varieties had the ability to produce dwarf progeny even though they themselves were tall. Mendel inferred that these hybrids carried a latent genetic factor for dwarfness, one that was masked by the expression of another factor for tallness. He said that the latent factor was recessive and that the expressed factor was dominant. He also inferred that these recessive and dominant factors separated from each other when the hybrid plants reproduced. This enabled him to explain the reappearance of the dwarf characteristic in the next generation.

Mendel performed similar experiments to study the inheritance of six other traits: seed texture, seed color, pod shape, pod color, flower color, and flower position (Table 3.1). In each experiment—called a monohybrid cross because a single trait was being studied-Mendel observed that only one of the two contrasting characteristics appeared in the hybrids and that when these hybrids were self-fertilized, they produced two types of progeny, each resembling one of the plants in the original crosses. Furthermore, he found that these progeny consistently appeared in a ratio of 3:1. Thus, each trait that Mendel studied seemed to be controlled by a heritable factor that existed in two forms, one dominant, the other recessive. These factors are now called genes, a word coined by the Danish plant breeder Wilhelm Johannsen in 1909; their dominant and recessive forms are called **alleles**—from the Greek word meaning "of one another." Alleles are alternate forms of a gene.

The regular numerical relationships that Mendel observed in these crosses led him to another important conclusion: that genes come in pairs. Mendel proposed that each of the parental strains that he used in his experiments carried two identical copies of a gene—in modern terminology, they are diploid and homozygous. However, during the production of gametes, Mendel proposed that these two copies are reduced to one; that is, the gametes that emerge from meiosis carry a single copy of a gene—in modern terminology, they are haploid.

Mendel recognized that the diploid gene number would be restored when sperm and egg unite to form a zygote. Furthermore, he understood that if the sperm and egg came from genetically different plants-as they did in his crosses-the hybrid zygote would inherit two different alleles, one from the mother and one from the father. Such an offspring is said to be heterozygous. Mendel realized that the different alleles that are present in a heterozygote must coexist even though one is dominant and the other recessive, and that each of these alleles would have an equal chance of entering a gamete when the heterozygote reproduces. Furthermore, he realized that random fertilizations with a mixed population of gametes-half carrying the dominant allele and half carrying the recessive allele-would produce some

TABLE 3.1		
Results of Mendel's Monohybrid Cross	25	
Parental Strains	F ₂ Progeny	Ratio
Tall plants × dwarf plants Round seeds × wrinkled seeds Yellow seeds × green seeds Violet flowers × white flowers Inflated pods × constricted pods Green pods × yellow pods Axial flowers × terminal flowers	787 tall, 277 dwarf 5474 round, 1850 wrinkled 6022 yellow, 2001 green 705 violet, 224 white 882 inflated, 299 constricted 428 green, 152 yellow 651 axial, 207 terminal	2.84:1 2.96:1 3.01:1 3.15:1 2.95:1 2.82:1 3.14:1

zygotes in which both alleles were recessive. Thus, he could explain the reappearance of the recessive characteristic in the progeny of the hybrid plants.

Mendel used symbols to represent the hereditary factors that he postulated—a methodological breakthrough. With symbols, he could describe hereditary phenomena clearly and concisely, and he could analyze the results of crosses mathematically. He could even make predictions about the outcome of future crosses. Although the practice of using symbols to analyze genetic problems has been much refined since Mendel's time, the basic principles remain the same. The symbols stand for genes (or, more precisely, for their alleles), and they are manipulated according to the rules of inheritance that Mendel discovered. These manipulations are the essence of formal genetic analysis. As an introduction to this subject, let's consider the symbolic representation of the cross between tall and dwarf peas (**■** Figure 3.2).

The two true-breeding varieties, tall and dwarf, are homozygous for different alleles of a gene controlling plant height. The allele for dwarfness, being recessive, is symbolized by a lowercase letter d; the allele for tallness, being dominant, is symbolized by the corresponding uppercase letter D. In genetics, the letter that is chosen

to denote the alleles of a gene is usually taken from the word that describes the recessive trait (*d*, for *d*warfness). Thus, the tall and dwarf pea strains are symbolized by *DD* and *dd*, respectively. The allelic constitution of each strain is said to be its **genotype**. By contrast, the physical appearance of each strain—the tall or dwarf characteristic—is said to be its **phenotype**.

As the **parental** strains, the tall and dwarf pea plants form the **P** generation of the experiment. Their hybrid progeny are referred to as the first filial generation, or F_1 , from a Latin word meaning "son" or "daughter." Because each parent contributes equally to its offspring, the genotype of the F_1 plants must be Dd; that is, they are heterozygous for the alleles of the gene that controls plant height. Their phenotype, however, is the same as that of the DD parental strain because D is dominant over d. During meiosis, these F_1 plants produce two kinds of gametes, D and d, in equal proportions. Neither allele is changed by having coexisted with the other in a heterozygous genotype; rather, they separate, or **segregate**, from each other during gamete formation. This process of allele segregation is perhaps the most important discovery that Mendel made.

Upon self-fertilization, the two kinds of gametes produced by heterozygotes can unite in all possible ways. Thus, they produce four kinds of zygotes (we write the contribution of the egg first): *DD*, *Dd*, *dD*, and *dd*. However, because of dominance, three of these genotypes have the same phenotype. Thus, in the next generation, called the F_{2} the plants are either tall or dwarf, in a ratio of 3:1.

Mendel took this analysis one step further. The F_2 plants were self-fertilized to produce an F_3 . All the dwarf F_2 plants produced only dwarf offspring, demonstrating that they were homozygous for the *d* allele, but the tall F_2 plants

demonstrating that they were homozygous for the d allele, but the tall F_2 plants comprised two categories. Approximately one-third of them produced only tall offspring,

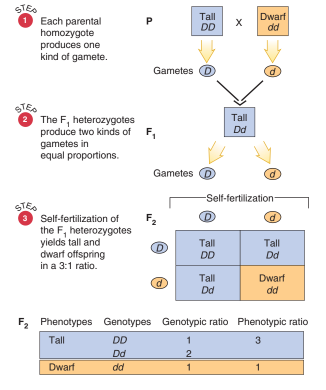


FIGURE 3.2 Symbolic representation of the cross between tall and dwarf peas.

whereas the other two-thirds produced a mixture of tall and dwarf offspring. Mendel concluded that the third that were true-breeding were DD homozygotes and that the two-thirds that were segregating were Dd heterozygotes. These proportions, 1/3 and 2/3, were exactly what his analysis predicted because, among the tall F_2 plants, the DD and Dd genotypes occur in a ratio of 1:2.

We summarize Mendel's analysis of this and other monohybrid crosses by stating two key principles that he discovered:

- **1.** The Principle of Dominance: *In a heterozygote, one allele may conceal the presence of another*. This principle is a statement about genetic function. Some alleles evidently control the phenotype even when they are present in a single copy. We consider the physiological explanation for this phenomenon in later chapters.
- **2.** The Principle of Segregation: In a heterozygote, two different alleles segregate from each other during the formation of gametes. This principle is a statement about genetic transmission. An allele is transmitted faithfully to the next generation, even if it was present with a different allele in a heterozygote. The biological basis for this phenomenon is the pairing and subsequent separation of homologous chromosomes during meiosis, a process we discussed in Chapter 2. We will consider the experiments that led to this chromosome theory of heredity in Chapter 5.

DIHYBRID CROSSES: THE PRINCIPLE OF INDEPENDENT ASSORTMENT

Mendel also performed experiments with plants that differed in two traits (**Figure 3.3**). He crossed plants that produced yellow, round seeds with plants that produced green, wrinkled seeds. The purpose of the experiments was to see if the two seed traits, color and texture, were inherited independently. Because the F_1 seeds were all yellow and round, the alleles for these two characteristics were dominant. Mendel grew plants from these seeds and allowed them to self-fertilize. He then classified the F_2 seeds and counted them by phenotype.

The four phenotypic classes in the F_2 represented all possible combinations of the color and texture traits. Two classes—yellow, round seeds and green, wrinkled seeds—resembled the parental strains. The other two—green, round seeds and yellow, wrinkled seeds—showed new combinations of traits. The four classes had an approximate ratio of 9 yellow, round:3 green, round:3 yellow, wrinkled:1 green, wrinkled (Figure 3.3). To Mendel's insightful mind, these numerical relationships suggested a simple explanation: Each trait was controlled by a different gene segregating two alleles, and the two genes were inherited independently.

Let's analyze the results of this two-factor, or **dihybrid cross**, using Mendel's methods. We denote each gene with a letter, using lower case for the recessive allele and uppercase for the dominant (**True Figure 3.4**). For the seed color gene, the two alleles are g (for green) and G (for yellow), and for the seed texture gene, they are w (for wrinkled) and W (for round). The parental strains, which were true-breeding, must have been doubly homozygous; the yellow, round plants were GG WW and the green, wrinkled plants were gg ww. Such two-gene genotypes are customarily written by separating pairs of alleles with a space.

The haploid gametes produced by a diploid plant contain one copy of each gene. Gametes from GG WW plants therefore contain one copy of the seed color gene (the *G* allele) and one copy of the seed texture gene (the *W* allele). Such gametes are symbolized by *G W*. By similar reasoning, the gametes from gg ww plants are written g w. Cross-fertilization of these two types of gametes produces F_1 hybrids that are doubly heterozygous, symbolized by Gg Ww, and their yellow, round phenotype indicates that the *G* and *W* alleles are dominant.

The Principle of Segregation predicts that the F_1 hybrids will produce four different gametic genotypes: (1) *G W*, (2) *G w*, (3) *g W*, and (4) *g w*. If each gene segregates its alleles independently, these four types will be equally

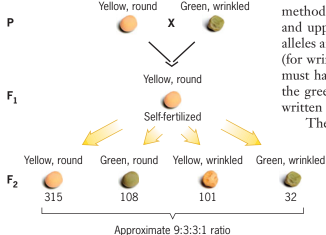


FIGURE 3.3 Mendel's crosses between peas with yellow, round seeds and peas with green, wrinkled seeds.

frequent; that is, each will be 25 percent of the total. On this assumption, self-fertilization in the F_1 will produce an array of 16 equally frequent zygotic genotypes. We obtain the zygotic array by systematically combining the gametes, as shown in Figure 3.4. We then obtain the phenotypes of these F_2 genotypes by noting that *G* and *W* are the dominant alleles. Altogether, there are four distinguishable phenotypes, with relative frequencies indicated by the number of positions occupied in the array. For absolute frequencies, we divide each number by the total, 16: STED

1

STED

2

STED

yellow, round	9/16
yellow, wrinkled	3/16
green, round	3/16
green, wrinkled	1/16

This analysis is predicated on two assumptions: (1) that each gene segregates its alleles, and (2) that these segregations are independent of each other. The

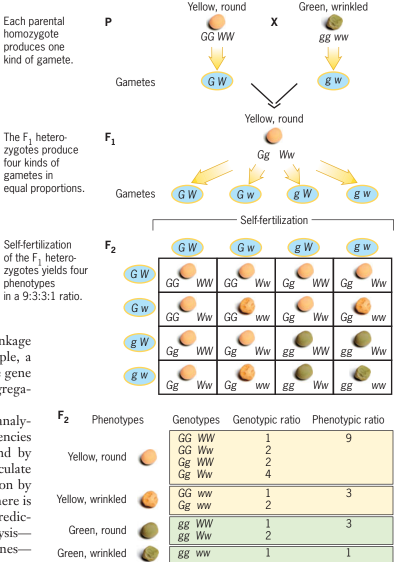
second assumption implies that there is no connection or linkage between the segregation events of the two genes. For example, a gamete that receives W through the segregation of the texture gene is just as likely to receive G as it is to receive g through the segregation of the color gene.

Do the experimental data fit with the predictions of our analysis? **Figure 3.5** compares the predicted and observed frequencies of the four F_2 phenotypes in two ways—by proportions and by numerical frequencies. For the numerical frequencies, we calculate the predicted numbers by multiplying the predicted proportion by the total number of F_2 seeds examined. With either method, there is obviously good agreement between the observations and the predictions. Thus, the assumptions on which we have built our analysis independent segregation of the seed color and seed texture genes are consistent with the observed data.

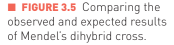
Mendel conducted similar experiments with other combinations of traits and in each case observed that the genes segregated independently. The results of these experiments led him to a third key principle:

3. The Principle of Independent Assortment: *The alleles of different genes segregate, or as we sometimes say, assort, independently of each other.* This principle is another rule of genetic transmission, based, as we will see in Chapter 5, on the behavior of different pairs of chromosomes during meiosis. However, not all genes abide by the Principle of Independent Assortment. In Chapter 7 we consider some important exceptions.

	F ₂ phenotypes	Observed Number Proportion			Expected Number Proportion	
0	Yellow, round	315	0.567	313	0.563	
C	Green, round	108	0.194	104	0.187	
C	Yellow, wrinkled	101	0.182	104	0.187	
C	Green, wrinkled	32	0.057	35	0.063	
	Total	556	1 000	556	1 000	



[■] FIGURE 3.4 Symbolic representation of Mendel's dihybrid cross.



KEY POINTS

- Mendel studied the inheritance of seven different traits in garden peas, each trait being controlled by a different gene.
 - Mendel's research led him to formulate three principles of inheritance: (1) the alleles of a gene are either dominant or recessive, (2) different alleles of a gene segregate from each other during the formation of gametes, and (3) the alleles of different genes assort independently.

Applications of Mendel's Principles

Mendel's principles can be used to predict the outcomes of crosses between different strains of organisms. If the genetic basis of a trait is known, Mendel's principles can be used to predict the outcome of crosses. There are three general procedures, two relying on the systematic enumeration of all the zygotic genotypes or phenotypes and one relying on mathematical insight.

THE PUNNETT SQUARE METHOD

For situations involving one or two genes, it is possible to write down all the gametes and combine them systematically to generate the array of zygotic genotypes. Once these have been obtained, the Principle of Dominance can be used to determine the associated phenotypes. This procedure, called the *Punnett square method* after the British geneticist R. C. Punnett, is a straightforward way of predicting the outcome of crosses. We have used it to analyze the zygotic output of the cross with Mendel's yellow, round F_1 hybrids—a type of mating commonly called an intercross (Figure 3.4). However, in more complicated situations, like those involving more than two genes, the Punnett square method is unwieldy. We will see in Figure 3.8 how the Punnett square method is related to an approach to genetic problems that uses the concept of probability.

THE FORKED-LINE METHOD

Another procedure for predicting the outcome of a cross involving two or more genes is the *forked-line method*. However, instead of enumerating the progeny in a square by combining the gametes systematically, we tally them in a diagram of branching lines. As an example, let us consider an intercross between peas that are heterozygous for three independently assorting genes—one controlling plant height, one controlling seed color, and one controlling seed texture. This is a trihybrid cross— $Dd \ Gg \ Ww \ Dd$ $Gg \ Ww$ —that can be partitioned into three monohybrid crosses— $Dd \ Cog \ Cog \ Cog$, $Gg \ Cog \ Co$

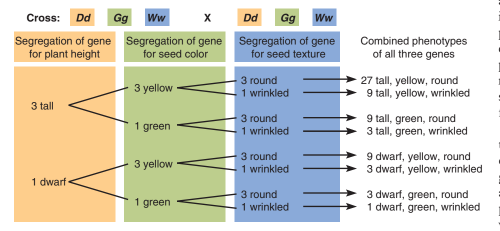


FIGURE 3.6 The forked-line method for predicting the outcome of an intercross involving three independently assorting genes in peas.

and $Ww \times Ww$ —because all the genes assort independently. For each gene, we expect the phenotypes to appear in a 3:1 ratio. Thus, for example, $Dd \times Dd$ will produce a ratio of 3 tall plants:1 dwarf plant. Using the forked-line method (**■** Figure 3.6), we can combine these separate ratios into an overall phenotypic ratio for the offspring of the cross.

We can also use this method to analyze the results of a cross between multiply heterozygous individuals and multiply homozygous individuals. This type of cross is called a **testcross**. For example, if *Dd Gg Ww* pea plants are crossed with *dd gg ww* pea plants, we can predict the phenotypes of the progeny by noting that each of the three genes in the heterozygous parent segregates dominant

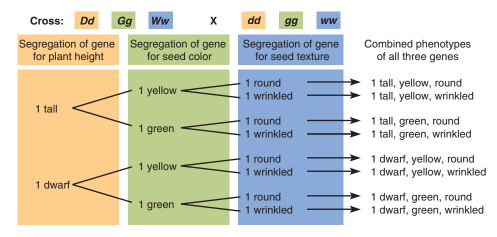


FIGURE 3.7 The forked-line method for predicting the outcome of a testcross involving three independently assorting genes in peas.

and recessive alleles in a 1:1 ratio, and that the homozygous parent transmits only recessive alleles of these genes. Thus, the genotypes—and ultimately the phenotypes— of the offspring of this cross depend on which alleles the heterozygous parent transmits (**Figure 3.7**).

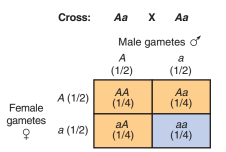
THE PROBABILITY METHOD

An alternative method to the Punnett square and forked-line methods-and a quicker one—is based on the principle of **probability**. Mendelian segregation is like a coin toss; when a heterozygote produces gametes, half contain one allele and half contain the other. The probability that a particular gamete contains the dominant allele is therefore 1/2, and the probability that it contains the recessive allele is also 1/2. These probabilities are the frequencies of the two types of gametes produced by the heterozygote. Can we use these frequencies to predict the outcome of crossing two heterozygotes? In such a cross, the gametes will be combined randomly to produce the next generation. Let's suppose the cross is $Aa \times Aa$ (**Figure 3.8**). The chance that a zygote will be AA is simply the probability that each of the uniting gametes contains A, or $(1/2) \times (1/2) = (1/4)$, since the two gametes are produced independently. The chance for an *aa* homozygote is also 1/4. However, the chance for an Aa heterozygote is 1/2 because there are two ways of creating a heterozygote—A may come from the egg and a from the sperm, or vice versa. Because each of these events has a one-quarter chance of occurring, the total probability that an offspring is heterozygous is (1/4) + (1/4) = (1/2). We therefore obtain the following probability distribution of the genotypes from the mating $Aa \times Aa$:

AA	1/4
Aa	1/2
aa	1/4

By applying the Principle of Dominance, we conclude that (1/4) + (1/2) = (3/4) of the progeny will have the dominant phenotype and 1/4 will have the recessive.

For such a simple situation, using the probability method to predict the outcome of a cross may seem unnecessary. However, in more complicated situations, it is clearly the most practical approach. Consider, for example, a cross between plants heterozygous for four different genes, each assorting independently. What fraction of the progeny will be homozygous for all four recessive alleles? To answer this question, we consider the genes one at a time. For the first gene, the fraction of offspring that will be recessive homozygotes is 1/4, as it will be for the second, third, and fourth genes. Therefore, by the Principle of Independent Assortment, the fraction of offspring that will be quadruple recessive homozygotes is $(1/4) \times (1/4) \times (1/4) = (1/256)$.



Progeny: Genotype Frequency Phenotype Frequency

AA	1/4	3	Dominant	3/4
Aa	1/2		Dominant	0/1
aa	1/4		Recessive	1/4

■ FIGURE 3.8 An intercross showing the probability method in the context of a Punnett square. The frequency of each genotype from the cross is obtained from the frequencies in the Punnett square, which are, in turn, obtained by multiplying the frequencies of the two types of gametes produced by the hetero-zygous parents.

Solve It!

Using Probabilities in a Genetic Problem

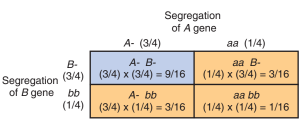
Mendel found that three traits in peasheight, flower color and pod shape-are determined by different genes, and that these genes assort independently. Suppose that tall plants with violet flowers and inflated pods are crossed to dwarf plants with white flowers and constricted pods, and that all the F_1 plants are tall, with violet flowers and inflated pods. If these F_1 plants are self-fertilized, what fraction of their offspring are expected to (a) show all three dominant phenotypes, (b) be tall, with white flowers and constricted pods, (c) be heterozygous for all three genes, (d) have at least one dominant allele of each gene in the genotype?

• To see the solution to this problem, visit the Student Companion site.

Surely, using the probability method is a better approach than diagramming a Punnett square with 256 entries!

Now let's consider an even more difficult question. What fraction of the offspring will be homozygous for all four genes? Before computing any probabilities, we must first decide what genotypes satisfy the question. For each gene there are two types of homozygotes, the dominant and the recessive, and together they constitute half the progeny. The fraction of progeny that will be homozygous for all four genes will therefore be $(1/2) \times (1/2) \times (1/2) \times$ (1/2) = (1/16).To see the full power

of the probability method,



Aa Bb

Х

Aa Bb

Cross:

Progeny:	Genotype	Frequency	Phenotype	Frequency
----------	----------	-----------	-----------	-----------

A- B-	9/16	Dominant for both genes	9/16
aa B-	3/16	Recessive for	7/16
A- bb	3/16	at least one gene	//10
aa bb	1/16		

■ FIGURE 3.9 Application of the probability method to an intercross involving two genes. In this cross, each gene segregates dominant and recessive phenotypes, with probabilities 3/4 and 1/4, respectively. Because the segregations occur independently, the frequencies of the combined phenotypes within the square are obtained by multiplying the marginal probabilities. The frequency of progeny showing the recessive phenotype for at least one of the genes is obtained by adding the frequencies in the relevant cells (tan color).

we need to consider one more question. Suppose the cross is $Aa \ Bb \times Aa \ Bb$ and we want to know what fraction of the progeny will show the recessive phenotype for at least one gene (**True 3.9**). Three kinds of genotypes would satisfy this condition: (1) *A*- *bb* (the dash stands for either *A* or *a*), (2) *aa B*-, and (3) *aa bb*. The answer to the question must therefore be the sum of the probabilities corresponding to each of these genotypes. The probability for *A*- *bb* is $(3/4) \times (1/4) = (3/16)$, that for *aa B*- is $(1/4) \times (3/4) = (3/16)$, and that for *aa bb* is $(1/4) \times (1/4) = (1/16)$. Adding these together, we find that the answer is 7/16. For more insights into this way of analyzing genetic problems, study Appendix A: The Rules of Probability at the back of this book. There you will find two simple rules—the Multiplicative Rule and the Additive Rule—along with some helpful examples. Then try working out the answers to the questions posed in Solve It: Using Probabilities in a Genetic Problem.

KEY POINTS

• The outcome of a cross can be predicted by the systematic enumeration of genotypes using a Punnett square.

• When more than two genes are involved, the forked-line or probability methods are used to predict the outcome of a cross.

Testing Genetic Hypotheses

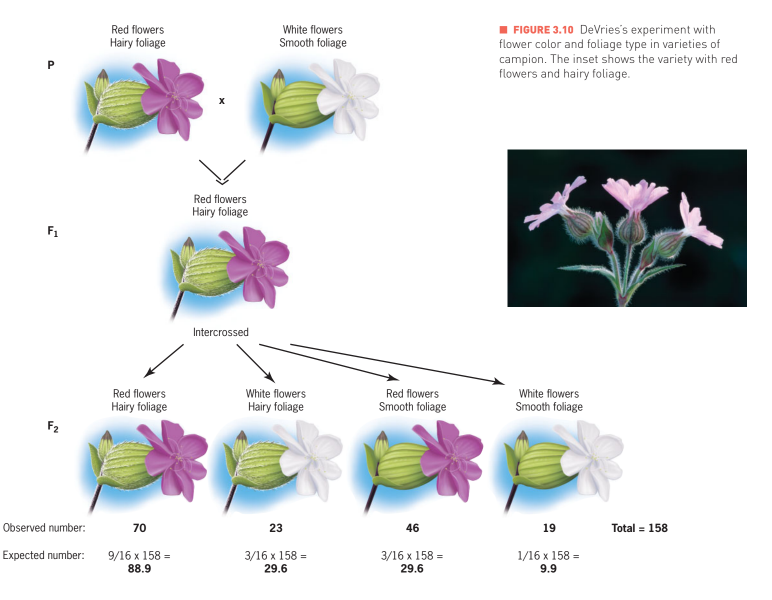
The chi-square test is a simple way of evaluating whether the predictions of a genetic hypothesis agree with data from an experiment. A scientific investigation always begins with observations of a natural phenomenon. The observations lead to ideas or questions about the phenomenon, and these ideas or questions are explored more fully by conducting further observations or by performing experiments. A well-formulated scientific idea is called a **hypothesis**. Data collected

from observations or from experimentation enable scientists to test hypotheses—that is, to determine if a particular hypothesis should be accepted or rejected.

In genetics, we are usually interested in deciding whether or not the results of a cross are consistent with a hypothesis. As an example, let's consider the data that Mendel

obtained from his dihybrid cross involving the color and texture of peas. In the F_2 , 556 peas were examined and sorted into four phenotypic classes (Figure 3.3). From the data, Mendel hypothesized that pea color and texture were controlled by different genes, that each of the genes segregated two alleles—one dominant, the other recessive—and that the two genes assorted independently. Are the data from the experiment actually consistent with this hypothesis? To answer this question, we need to compare the results of the experiment with the predictions of the hypothesis. The comparison laid out in Figure 3.5 suggests that the experimental results are indeed consistent with the hypothesis. Across the four phenotypic classes, the discrepancies between the observed and expected numbers are small, so small in fact that we are comfortable attributing them to chance. The hypothesis that Mendel conceived to explain his data therefore fits well with the results of his dihybrid cross. If it did not, we would have reservations about accepting the hypothesis and the whole theory of Mendelism would be in doubt. We consider another possibility—that Mendel's data fit his hypothesis too well—in A Milestone in Genetics: Mendel's 1866 Paper, which you can find in the Student Companion site.

Unfortunately, the results of a genetic experiment do not always agree with the predictions of a hypothesis as clearly as Mendel's did. Take, for example, data obtained by Hugo DeVries, one of the rediscoverers of Mendel's work. DeVries crossed different varieties of the campion, a plant that grew in his experimental garden. One variety had red flowers and hairy foliage; the other had white flowers and smooth foliage. The F_1 plants all had red flowers and hairy foliage, and when intercrossed, they produced F_2 plants that sorted into four phenotypic classes (**■** Figure 3.10). To explain the results of



these crosses, DeVries proposed that flower color and foliage type were controlled by two different genes, that each gene segregated two alleles—one dominant, the other recessive—and that the two genes assorted independently; that is, he simply applied Mendel's hypothesis to the campion. However, when we compare DeVries's data with the predictions of the Mendelian hypothesis, we find some disturbing discrepancies. Are these discrepancies large enough to raise questions about the experiment or the hypothesis?

THE CHI-SQUARE TEST

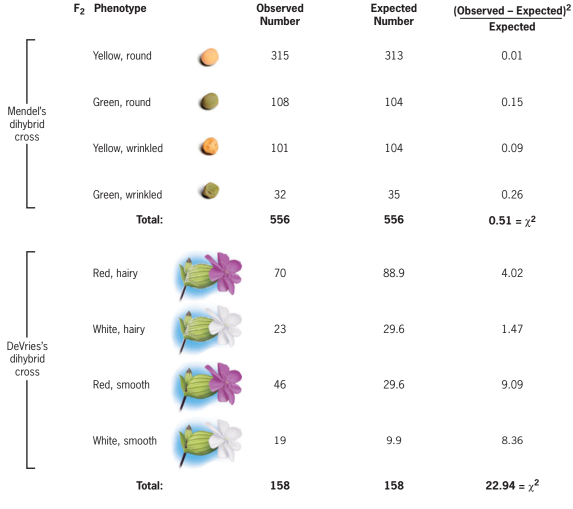
With DeVries's data, and with other genetic data as well, we need an objective procedure to compare the results of the experiment with the predictions of the underlying hypothesis. This procedure has to take into account how chance might affect the outcome of the experiment. Even if the hypothesis is correct, we do not anticipate that the results of the experiment will exactly match the predictions of the hypothesis. If they deviate a bit, as Mendel's data did, we would ascribe the deviations to chance variation in the outcome of the experiment. However, if they deviate grossly, we would suspect that something was amiss. The experiment might have been executed poorly—for example, the crosses might have been improperly carried out, or the data might have been incorrectly recorded—or, perhaps, the hypothesis is simply wrong. The possible discrepancies between observations and expectations obviously lie on a continuum from small to large, and we must decide how large they need to be for us to entertain doubts about the execution of the experiment or the acceptability of the hypothesis.

One procedure for assessing these discrepancies uses a statistic called **chi-square** (χ^2). A *statistic* is a number calculated from data—for example, the mean of a set of examination scores. The χ^2 statistic allows a researcher to compare data, such as the numbers we get from a breeding experiment, with their predicted values. If the data are not in line with the predicted values, the χ^2 statistic will exceed a critical number and we will decide either to reevaluate the experiment—that is, look for a mistake in technique—or reject the underlying hypothesis. If the χ^2 statistic is below this number, we tentatively conclude that the results of the experiment are consistent with the predictions of the hypothesis. The χ^2 statistic therefore reduces hypothesis testing to a simple, objective procedure.

As an example, let's consider the data from the experiments of Mendel and DeVries. Mendel's F_2 data seemed to be consistent with the underlying hypothesis, whereas DeVries's F_2 data showed some troubling discrepancies. **Figure 3.11** outlines the calculations.

For each phenotypic class in the F_2 , we compute the difference between the observed and expected numbers of offspring and square these differences. The squaring operation eliminates the canceling effects of positive and negative values among the four phenotypic classes. Then we divide each squared difference by the corresponding expected number of offspring. This operation scales each squared difference by the size of the expected number. If two classes have the same squared difference, the one with the smaller expected number contributes relatively more in the calculation. Finally, we sum all the terms to obtain the χ^2 statistic. For Mendel's data, the χ^2 statistic is 0.51 and for DeVries's data it is 22.94. These statistics summarize the discrepancies between the observed and expected numbers across the four phenotypic classes in each experiment. If the observed and expected numbers are in basic agreement with each other, the χ^2 statistic will be small, as it happens to be with Mendel's data. If they are in serious disagreement, it will be large, as it happens to be with DeVries's data. Clearly, we must decide what value of χ^2 on the continuum between small and large casts doubt on the experiment or the hypothesis. This critical value is the point where the discrepancies between observed and expected numbers are not likely to be due to chance.

To determine the critical value, we need to know how chance affects the χ^2 statistic. Assume for the moment that the underlying genetic hypothesis is true. Now imagine



Formula for chi-square statistic to test for agreement between observed and expected numbers:

 $\chi^{2} = \sum \frac{(\text{Observed} - \text{Expected})^{2}}{\text{Expected}}$

FIGURE 3.11 Calculating χ^2 for Mendel's and DeVries's F_2 data.

carrying out the experiment-carefully and correctly-many times, and each time, calculating a χ^2 statistic. All these statistics can be compiled into a graph that shows how often each value occurs. We call such a graph a *frequency distribution*. Fortunately, the χ^2 frequency distribution is known from statistical theory (**Figure 3.12**)—so we don't actually need to carry out many replications of the experiment to get it. The critical value is the point that cuts off the upper 5 percent of the distribution. By chance alone, the χ^2 statistic will exceed this value 5 percent of the time. Thus, if we perform an experiment once, compute a χ^2 statistic, and find that the statistic is greater than the critical value, we have either observed a rather unlikely set of results-something that happens less than 5 percent of the time—or there is a problem with the way the experiment was executed or with the appropriateness of the hypothesis. Assuming that the experiment was done properly, we are inclined to reject the hypothesis. Of course we must realize that with this procedure we will reject a true hypothesis 5 percent of the time.

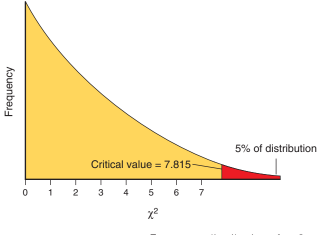


FIGURE 3.12 Frequency distribution of a χ^2 statistic.

Solve It!

Using the Chi-Square Test

When true-breeding tomato plants with spherical fruit were crossed to truebreeding plants with ovoid fruit, all the F₁ plants had spherical fruit. These F₁ plants were then intercrossed to produce an F₂ generation that comprised 73 plants with spherical fruit and 11 with ovoid fruit. Are these results consistent with the hypothesis that fruit shape in tomatoes is controlled by a single gene?

To see the solution to this problem, visit the Student Companion site.

TABLE 3.2 Table of Chi-Square (χ^2) 5% Critical Values^a

Degrees of Freedom	5% Critical Value		
1	3.841		
2	5.991		
3	7.815		
4	9.488		
5	11.070		
6	12.592		
7	14.067		
8	15.507		
9	16.919		
10	18.307		
15	24.996		
20	31.410		
25	37.652		
30	43.773		
^a Selected entries from R. A. Fisher and Yates, 1943, <i>Statistical Tables for Biological, Agricultural, and Medical Research</i> . Oliver and Boyd, London.			

Thus, as long as we know the critical value, the χ^2 testing procedure leads us to a decision about the fate of the hypothesis. However, this critical value-and the shape of the associated frequency distribution—depends on the number of phenotypic classes in the experiment. Statisticians have tabulated critical values according to the **degrees of freedom** associated with the χ^2 statistic (**Table 3.2**). This index to the set of χ^2 distributions is determined by subtracting one from the number of phenotypic classes. In each of our examples, there are 4 - 1 = 3 degrees of freedom. The critical value for the χ^2 distribution with 3 degrees of freedom is 7.815. For Mendel's data, the calculated χ^2 statistic is 0.51, much less than the critical value and therefore no threat to the hypothesis being tested. However, for DeVries's data the calculated χ^2 statistic is 22.94, very much greater than the critical value. Thus, the observed data do not fit with the genetic hypothesis. Ironically, when DeVries presented these data in 1905, he judged them to be consistent with the genetic hypothesis. Unfortunately, he did not perform a χ^2 test. DeVries also argued that his data provided further evidence for the correctness and widespread applicability of Mendel's ideas-not the only time that a scientist has come to the right conclusion for the wrong reason. To solidify your understanding of the χ^2 procedure, answer the question posed in Solve It: Using the Chi-Square Test.

- **KEY POINTS** The chi-square statistic is $\chi^2 = \Sigma$ (observed number expected number)²/ expected number, with the sum computed over all categories comprising the data.
 - Each chi-square statistic is associated with an index, the degrees of freedom, which is equal to the number of data categories minus one.

Mendelian Principles in Human Genetics

Mendel's principles can be applied to study the inheritance of traits in humans.

The application of Mendelian principles to human genetics began soon after the rediscovery of Mendel's paper in 1900. However, because it is not possible to make controlled crosses

with humans, progress was obviously slow. The analysis of human heredity depends on family records, which are often incomplete. In addition, humans-unlike experimental

organisms—do not produce many progeny, making it difficult to discern Mendelian ratios, and humans are not maintained and observed in a controlled environment. For these and other reasons, human genetic analysis has been a difficult endeavor. Nonetheless, the drive to understand human heredity has been very strong, and today, despite all the obstacles, we have learned about thousands of human genes. Table 3.3 lists some of the conditions they control. We discuss many of these conditions in later chapters of this book.

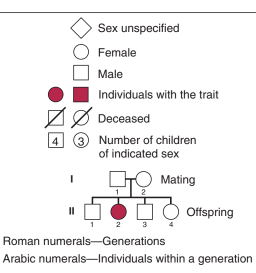
PEDIGREES

Pedigrees are diagrams that show the relationships among the members of a family (**■** Figure 3.13*a*). It is customary to represent males as squares and females as circles. A horizontal line connecting a circle and a square represents a mating. The offspring of the mating are shown beneath the mates, starting with the first born at the left and proceeding through the birth order to the right. Individuals that have a genetic condition are indicated by coloring or shading. The generations in a pedigree are usually denoted by Roman numerals, and particular individuals within a generation are referred to by Arabic numerals following the Roman numeral.

Traits caused by dominant alleles are the easiest to identify. Usually, every individual who carries the dominant allele manifests the trait, making it possible to trace the transmission of the dominant allele through the pedigree (**■** Figure 3.13*b*). Every affected individual is expected to have at least one affected parent, unless, of course, the dominant allele has just appeared in the family as a result of a new mutation—a change in the gene itself. However, the frequency of most new mutations is very low—on the order of one in a million; consequently, the spontaneous appearance of a dominant condition is an extremely rare event. Dominant traits that are associated with reduced viability or fertility never become frequent in a population. Thus, most of the people who show such traits are heterozygous for the dominant allele. If their spouses do not have the trait, half their children should inherit the condition.

Recessive traits are not so easy to identify because they may occur in individuals whose parents are not affected. Sometimes several generations of pedigree data are needed to trace the transmission of a recessive allele (■ Figure 3.13*c*). Nevertheless, a large number of recessive traits have been observed in humans-at last count, over 4000. Rare recessive traits are more likely to appear in a pedigree when spouses are related to each other-for example, when they are first cousins. This increased incidence occurs because relatives share alleles by virtue of their common ancestry. Siblings share one-half their alleles, half siblings one-fourth their alleles, and first

cousins one-eighth their alleles. Thus, when such relatives mate, they have a greater chance of producing a child who is homozygous for a particular recessive allele than do unrelated parents. Many of the classical studies in human genetics have relied on the analysis of matings between relatives, principally first cousins. We will consider this subject in more detail in Chapter 4.



(a) Pedigree conventions

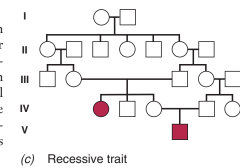


TABLE 3.3

Inherited Conditions in Humans Dominant Traits

Achondroplasia (dwarfism) Brachydactyly (short fingers)

Congenital night blindness

- Ehler-Danlos syndrome (a connective tissue disorder)
- Huntington's disease (a neurological disorder)
- Marfan syndrome (tall, gangly stature) Neurofibromatosis (tumorlike growths on the body)

Phenylthiocarbamide (PTC) tasting Widow's peak Woolly hair

Recessive Traits

- Albinism (lack of pigment)
 Alkaptonuria (a disorder of amino acid metabolism)
 Ataxia telangiectasia (a neurological disorder)
 Cystic fibrosis (a respiratory disorder)
 Duchenne muscular dystrophy
 Galactosemia (a disorder of carbohydrate metabolism)
 Glycogen storage disease
 Phenylketonuria (a disorder of amino acid metabolism)
 Sickle-cell disease (a hemoglobin disorder)
 Tay-Sachs disease (a linid storage
- Tay-Sachs disease (a lipid storage disorder)

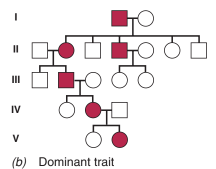


FIGURE 3.13 Mendelian inheritance in human pedigrees. (a) Pedigree conventions. (b) Inheritance of a dominant trait. The trait appears in each generation. (c) Inheritance of a recessive trait. The two affected individuals are the offspring of relatives.

MENDELIAN SEGREGATION IN HUMAN FAMILIES

In humans, the number of children produced by a couple is typically small. Today in the United States, the average is around two. In developing countries, it is six to seven. Such numbers provide nothing close to the statistical power that Mendel had in his experiments with peas. Consequently, phenotypic ratios in human families often deviate significantly from their Mendelian expectations.

As an example, let's consider a couple who are each heterozygous for a recessive allele that, in homozygous condition, causes cystic fibrosis, a serious disease in which breathing is impaired by an accumulation of mucus in the lungs and respiratory tract. If the couple were to have four children, would we expect *exactly* three to be unaffected and one to be affected by cystic fibrosis? The answer is no. Although this is a possible outcome, it is not the only one. There are, in fact, five distinct possibilities:

- 1. Four unaffected, none affected.
- 2. Three unaffected, one affected.
- 3. Two unaffected, two affected.
- 4. One unaffected, three affected.
- 5. None unaffected, four affected.

Intuitively, the second outcome seems to be the most likely, since it conforms to Mendel's 3:1 ratio. We can calculate the probability of this outcome, and of each of the others, by using Mendel's principles and by treating each birth as an independent event (**I Figure 3.14**).

For a particular birth, the chance that the child will be unaffected is 3/4. The probability that all four children will be unaffected is therefore $(3/4) \times (3/4) \times (3/4) \times (3/4) = (3/4)^4 = 81/256$. Similarly, the chance that a particular child will be affected is 1/4; thus, the probability that all four will be affected is $(1/4)^4 = 1/256$. To find the probabilities for the three other outcomes, we need to recognize that each actually represents a collection of distinct events. The outcome of three unaffected children and one affected child, for instance, comprises four distinct events; if we let U symbolize an unaffected child and A an affected child, and if we write the children in their order of birth, we can represent these events as

UUUA, UUAU, UAUU, and AUUU

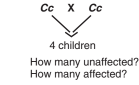
Because each has probability $(3/4)^3 \times (1/4)$, the total probability for three unaffected children and one affected, regardless of birth order, is $4 \times (3/4)^3 \times (1/4)$. The coeffi-

cient 4 is the number of ways in which three children could be unaffected and one could be affected in a family with four children. Similarly, the probability for two unaffected children and two affected is $6 \times (3/4)^2 \times (1/4)^2$, since in this case there are six distinct events. The probability for one unaffected child and three affected is $4 \times (3/4) \times (1/4)^3$, since in this case there are four distinct events. Figure 3.14 summarizes the calculations in the form of a probability distribution. As anticipated, three unaffected children and one affected child is the most probable outcome (probability 108/256).

In this example the children fall into two possible phenotypic classes. Because there are only two classes, the probabilities associated with the various outcomes are called **binomial probabilities**. Appendix B: Binomial Probabilities at the back of the book generalizes the method of analyzing this example so that you can apply it to other situations involving two phenotypic classes.

GENETIC COUNSELING

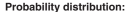
The diagnosis of genetic conditions is often a difficult process. Typically, diagnoses are made by physicians who have been trained in genetics. The study of these conditions requires a great deal of careful research, including examining patients, interviewing relatives, and sifting through vital statistics on births, deaths,



Number of children that are:

Parents

Unaffected	Affected	Probability
4	0	1 x (3/4) x (3/4) x (3/4) x (3/4) = 81/256
3	1	$4 \times (3/4) \times (3/4) \times (3/4) \times (1/4) = 108/256$
2	2	6 x (3/4) x (3/4) x (1/4) x (1/4) = 54/256
1	3	4 x (3/4) x (1/4) x (1/4) x (1/4) = 12/256
0	4	$1 \times (1/4) \times (1/4) \times (1/4) \times (1/4) = 1/256$



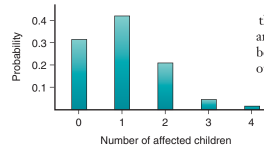


FIGURE 3.14 Probability distribution for families with four children segregating a recessive trait.

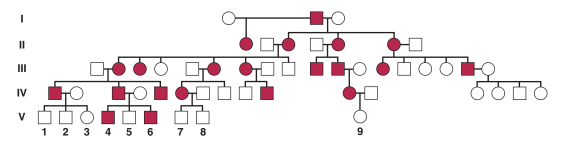


FIGURE 3.15 Pedigree showing the inheritance of hereditary nonpolypoid colorectal cancer.

and marriages. The accumulated data provide the basis for defining the condition clinically and for determining its mode of inheritance.

Parents may want to know whether their children are at risk to inherit a particular condition, especially if other family members have been affected. It is the responsibility of the *genetic counselor* to assess such risks and to explain them to the prospective parents. Risk assessment requires familiarity with probability and statistics, as well as a thorough knowledge of genetics.

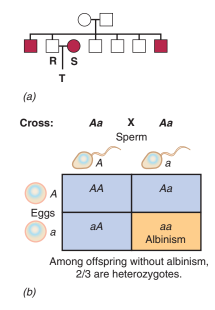
As an example, let's consider a pedigree showing the inheritance of **nonpolypoid colorectal cancer** (**Figure 3.15**). This disease is one of several types of cancer that are inherited. It is due to a dominant mutation that affects about 1 in 500 individuals in the general population. The median age when hereditary nonpolypoid colorectal cancer appears in an individual who carries the mutation is 42. In the pedigree we see that the cancer is manifested in at least one individual in each generation and that every affected individual has an affected parent. These facts are consistent with the dominant mode of inheritance of this disease.

The counseling issue arises in generation V. Among the nine individuals shown, two are affected and seven are not. Yet each of the seven unaffected individuals had one affected parent who must have been heterozygous for the cancer-causing mutation. Some of these seven unaffected individuals may therefore have inherited the mutation and would be at risk to develop nonpolypoid colorectal cancer later in life. Only time will tell. As the unaffected individuals age, those who carry the mutation will be at increased risk to develop the disease. Thus, the longer they remain unaffected, the greater the probability that they are actually not carriers. In this situation, the risk is a function of an individual's age and must be ascertained empirically from data on the age of onset of the disease among individuals from the same population, if possible from the same family. Each of the seven unaffected individuals will, of course, have to live with the anxiety of being a possible carrier of the cancer-causing mutation. Furthermore, at some point they will have to decide if they wish to reproduce and risk transmitting the mutation to their children.

As another example, consider the situation shown in **Figure 3.16.** A couple, denoted R and S in Figure 3.16*a*, is concerned about the possibility that they will have a child (T) with **albinism**, a recessive condition characterized by a complete absence of melanin pigment in the skin, eyes, and hair. S, the prospective mother, has albinism, and R, the prospective father, has two siblings with albinism. It would therefore seem that the child has some risk of being born with albinism.

This risk depends on two factors: (1) the probability that R is a heterozygous carrier of the albinism allele (a), and (2) the probability that he will transmit this allele to T if he actually is a carrier. S, who is obviously homozygous for the albinism allele, must transmit this allele to her offspring.

To determine the first probability, we need to consider the possible genotypes for R. One of these, that he is homozygous for the recessive allele (*aa*), is excluded because we know that he does not have albinism himself. However, the other two genotypes, *AA* and *Aa*, remain distinct possibilities. To calculate the probabilities associated with each of these, we note that both of R's parents must be heterozygotes because they have had two children with albinism. The mating that produced R was therefore $Aa \times Aa$, and from such a mating we would expect 2/3 of the offspring without albinism to be *Aa* and 1/3 to be *AA* (Figure 3.16*b*). Thus, the probability that



■ FIGURE 3.16 Genetic counseling in a family with albinism. (a) Pedigree showing the inheritance of albinism. (b) Punnett square showing that among offspring without albinism, the frequency of heterozygotes is 2/3. R is a heterozygous carrier of the albinism allele is 2/3. To determine the probability that he will transmit this allele to his child, we simply note that *a* will be present in half of his gametes.

In summary, the risk that T will be *aa*

= [Probability that R is Aa] × [Probability that R transmits *a*, assuming that R is Aa] = $(2/3) \times (1/2) = (1/3)$

The example in Figure 3.16 illustrates a simple counseling situation in which the risk can be determined precisely. Often the circumstances are much more complicated, making the task of risk assessment quite difficult. The genetic counselor's responsibility is to analyze the pedigree information and determine the risk as precisely as possible. For practice in calculating genetic risks, work through the example in Problem-Solving Skills: Making Predictions from Pedigrees.

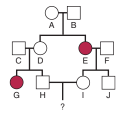
Today, genetic counseling is a well-established profession. Each genetic counselor has a master's degree and has been certified to practice by the American Board of Genetic Counseling, an oversight organization that also accredits genetic counseling training programs. There are roughly 2500 certified genetic counselors in the United States. Genetic counselors are trained to obtain and evaluate family histories to assess the risk for genetic disease. They are also trained to educate people about genetic diseases and to provide advice about how to prevent or cope with these diseases. Genetic counselors practice as part of a health care team, and their expertise is often valued by other health care professionals, who may not be so well informed about the genetic causes of disease. Genetic counselors must know about the ethical and legal ramifications of their work, and they must be sensitive to the psychological, social, cultural, and religious needs of their patients. Genetic counselors must also be good communicators. In the course of their work, they must explain complicated issues to their patients, who may not know much about the principles of inheritance or have the mathematical skills to understand how genetic risks are calculated. In the future, the ever-expanding fund of genetic information, much of it deriving from the ongoing Human Genome Project, will likely make the work of genetic counselors even more challenging.

PROBLEM-SOLVING SKILLS

Making Predictions from Pedigrees

THE PROBLEM

This pedigree shows the inheritance of a recessive trait in humans. Individuals that have the trait are homozygous for a recessive allele *a*. If H and I, who happen to be first cousins, marry and have a child, what is the chance that this child will have the recessive trait?



FACTS AND CONCEPTS

- **1.** The child can show a recessive trait only if both of its parents carry the recessive allele.
- 2. One parent (H) has a sister (G) with the trait.
- 3. The other parent (I) has a mother (E) with the trait.

- **4.** The chance that a heterozygote will transmit a recessive allele to its offspring is 1/2.
- In a mating between two heterozygotes, 2/3 of the offspring that do not show the trait are expected to be heterozygotes (see Figure 3.16b).

ANALYSIS AND SOLUTION

I must be a heterozygous carrier of the recessive allele because her mother E is homozygous for it, but she herself does not show the trait. I therefore has a 1/2 chance of transmitting the recessive allele to her child. Because H's sister has the trait, both of her parents must be heterozygotes. H, who does not show the trait, therefore has a 2/3 chance of being a heterozygote, and if he is, there is a 1/2 chance that he will transmit the recessive allele to his child. Putting all these factors together, we calculate the chance that the child of H and I will show the trait as 1/2 (the chance that I transmits the recessive allele) \times 2/3 (the chance that H is a heterozygote) \times 1/2 (the chance that H transmits the recessive allele assuming that he is a heterozygote) = 1/6, which is a fairly substantial risk.

For further discussion visit the Student Companion site.

- Pedigrees are used to identify dominant and recessive traits in human families.
- The analysis of pedigrees allows genetic counselors to assess the risk that an individual will inherit a particular trait.

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. Two highly inbred strains of mice, one with black fur and the other with gray fur, were crossed, and all of the offspring had black fur. Predict the outcome of intercrossing the offspring.
- **Answer:** The two strains of mice are evidently homozygous for different alleles of a gene that controls fur color: *G* for black fur and *g* for gray fur; the *G* allele is dominant because all the F_1 animals are black. When these mice, genotypically *Gg*, are intercrossed, the *G* and *g* alleles will segregate from each other to produce an F_2 population consisting of three genotypes, *GG*, *Gg*, and *gg*, in the ratio 1:2:1. However, because of the dominance of the *G* allele, the *GG* and *Gg* genotypes will have the same phenotype (black fur); thus, the phenotypic ratio in the F_2 will be 3 black:1 gray.
- 2. A plant heterozygous for three independently assorting genes, *Aa Bb Cc*, is self-fertilized. Among the offspring, predict the frequency of (a) *AA BB CC* individuals, (b) *aa bb cc* individuals, (c) individuals that are either *AA BB CC* or *aa bb cc*, (d) *Aa Bb Cc* individuals, (e) individuals that are not heterozygous for all three genes.
- Answer: Because the genes assort independently, we can analyze them one at a time to obtain the answers to each of the questions. (a) When Aa individuals are selfed, 1/4 of the offspring will be AA; likewise, for the B and C genes, 1/4 of the individuals will be BB and 1/4 will be CC. Thus, we can calculate the frequency (that is, the probability) of AA BB CC offspring as $(1/4) \times (1/4) \times (1/4) = 1/64$. (b) The frequency of *aa bb cc* individuals can be obtained using similar reasoning. For each gene the frequency of recessive homozygotes among the offspring is 1/4. Thus, the frequency of triple recessive homozygotes is $(1/4) \times (1/4) \times (1/4) = 1/64$. (c) To obtain the frequency of offspring that are either triple dominant homozygotes or triple recessive homozygotes-these are mutually exclusive events—we sum the results of (a) and (b): 1/64 + 1/64 = 2/64 = 1/32. (d) To obtain the frequency of offspring that are triple heterozygotes, again we multiply probabilities. For each gene, the frequency of heterozygous offspring is 1/2; thus, the frequency of triple heterozygotes should be $(1/2) \times (1/2) \times (1/2) = 1/8$. (e) Offspring that are not heterozygous for all three genes occur with a frequency that is one minus the frequency calculated in (d). Thus, the answer is 1 - 1/8 = 7/8.
- **3.** Two true-breeding strains of peas, one with tall vines and violet flowers and the other with dwarf vines and white flowers, were crossed. All the F₁ plants were tall and pro-

duced violet flowers. When these plants were backcrossed to the dwarf, white parent strain, the following offspring were obtained: 53 tall, violet; 48 tall, white; 47 dwarf, violet; 52 dwarf, white. Do the genes that control vine length and flower color assort independently?

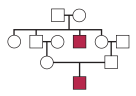
KEY POINTS

Answer: The hypothesis of independent assortment of the vine length and flower color genes must be evaluated by calculating a chi-square test statistic from the experimental results. To obtain this statistic, the results must be compared to the predictions of the genetic hypothesis. Under the assumption that the two genes assort independently, the four phenotypic classes in the F_2 should each be 25 percent of the total (200); that is, each should contain 50 individuals. To compute the chi-square statistic, we must obtain the difference between each observation and its predicted value, square these differences, divide each squared difference by the predicted value, and then sum the results:

$$\chi^2 = (53 - 50)^2 / 50 + (48 - 50)^2 / 50 + (47 - 50)^2 / 50 + (52 - 50)^2 / 50 = 0.52$$

This statistic must then be compared to the critical value of the chi-square frequency distribution for 3 degrees of freedom (calculated as the number of phenotypic classes minus one). Because the computed value of the chi-square statistic (0.52) is much less than the critical value (7.815; see Table 3.2), there is no evidence to reject the hypothesis of independent assortment of the vine length and flower color genes. Thus, we may tentatively accept the idea that these genes assort independently.

4. Is the trait that is segregating in the following pedigree due to a dominant or a recessive allele?



- **Answer:** Both affected individuals have two unaffected parents, which is inconsistent with the hypothesis that the trait is due to a dominant allele. Thus, the trait appears to be due to a recessive allele.
- 5. In a family with three children, what is the probability that two are boys and one is a girl?

Answer: To answer this question, we must apply the theory of binomial probabilities. For any one child, the probability that it is a boy is 1/2 and the probability that it is a girl is 1/2. Each child is produced independently. Thus, the probability of two boys and one girl is $(1/2)^3$ times the

Testing Your Knowledge

Integrate Different Concepts and Techniques

- Phenylketonuria, a metabolic disease in humans, is caused by a recessive allele, k. If two heterozygous carriers of the allele marry and plan a family of five children: (a) What is the chance that all their children will be unaffected? (b) What is the chance that four children will be unaffected and one affected with phenylketonuria? (c) What is the chance that at least three children will be unaffected? (d) What is the chance that the first child will be an unaffected girl?
- **Answer:** Before answering each of the questions, note that from a mating between two heterozygotes, the probability that a particular child will be unaffected is 3/4, and the probability that a particular child will be affected is 1/4. Furthermore, for any one child born, the chance that it will be a boy is 1/2 and the chance that it will be a girl is 1/2.
- (a) To calculate the chance that all five children will be unaffected, use the Multiplicative Rule of Probability (Appendix A). For each child, the chance that it will be unaffected is 3/4, and all five children are independent. Consequently, the probability of five unaffected children is $(3/4)^5 = 0.237$. This is the first term of the binomial probability distribution (see Appendix B) with p = 3/4 and q = 1/4.
- (b) To calculate the chance that four children will be unaffected and one affected, compute the second term of the binomial distribution using the formula in Appendix B:

$$= [5!/(4! 1!)] \times (3/4)^4 \times (1/4)^1 = 5 \times (81/1024) = 0.399$$

(c) To find the probability that at least three children will be unaffected, calculate the third term of the binomial distribution and add it to the first and second terms:

Event	Binomial Formula	Probability
5 unaffected, 0 affected	$\frac{[(5!) / (5! \ 0!) \times}{(3/4)^5 \ (1/4)^0} =$	0.237
4 unaffected, 1 affected	$[(5!)/(4! 1!)] \times (3/4)^4 (1/4)^1 =$	0.399
3 unaffected, 2 affected	$[(5!)/(3! 2!)] \times (3/4)^3 (1/4)^2 =$	0.264
	Total	0.900

(d) To determine the probability that the first child will be an unaffected girl, use the Multiplicative Rule: P(unaffected child and girl) = P(unaffected child) × P(girl) = $(3/4) \times (1/2) = (3/8)$.

number of ways in which two boys and one girl can appear in the birth order. By enumerating all the possible birth orders—BBG, BGB, and GBB—we find that the number of ways is 3. Thus, the final answer is $3 \times (1/2)^3 = 3/8$.

- 2. Mice from wild populations typically have gray-brown (or *agouti*) fur, but in one laboratory strain, some of the mice have yellow fur. A single yellow male is mated to several agouti females. Altogether, the matings produce 40 progeny, 22 with agouti fur and 18 with yellow fur. The agouti F₁ animals are then intercrossed with each other to produce an F₂, all of which are agouti. Similarly, the yellow F₁ animals are intercrossed with each other, but their F₂ progeny segregate into two classes; 30 are agouti and 54 are yellow. Subsequent crosses between yellow F₂ animals also segregate yellow and agouti progeny. What is the genetic basis of these coat color differences?
- **Answer:** We note that the cross agouti \times agouti produces only agouti animals and that the cross yellow \times yellow produces a mixture of yellow and agouti. Thus, a reasonable hypothesis is that yellow fur is caused by a dominant allele, A, and that agouti fur is caused by a recessive allele, a. According to this hypothesis, the agouti females used in the initial cross would be aa and their yellow mate would be Aa. We hypothesize that the male was heterozygous because he produced approximately equal numbers of agouti and yellow F_1 offspring. Among these, the agouti animals should be *aa* and the yellow animals *Aa*. These genotypic assignments are borne out by the F_2 data, which show that the F_1 agouti mice have bred true and the F_1 yellow mice have segregated. However, the segregation ratio of yellow to agouti (54:30) seems to be out of line with the Mendelian expectation of 3:1. Is this lack of fit serious enough to reject the hypothesis?

We can use the χ^2 procedure to test for disagreement between the data and the predictions of the hypothesis. According to the hypothesis, 3/4 of the F₂ progeny from the yellow × yellow intercross should be yellow and 1/4 should be agouti. Using these proportions, we can calculate the expected numbers of progeny in each class and then calculate a χ^2 statistic with 2 - 1 = 1 degree of freedom.

F ₂ Phenotype	Obs	Exp	(Obs – Exp) ² /Exp
yellow (AA and Aa)	54	$(3/4) \times 84 = 63$	1.286
agouti (aa)	30	$(1/4) \times 84 = 21$	3.857
Total	84	84	5.143

The χ^2 statistic (5.143) is much greater than the critical value (3.841) for a χ^2 distribution with 1 degree of freedom.

Consequently, we reject the hypothesis that the coat colors are segregating in a 3:1 Mendelian fashion.

What might account for the failure of the coat colors to segregate as hypothesized? We obtain a clue by noting that subsequent yellow \times yellow crosses failed to establish a true-breeding yellow strain. This suggests that the yellow animals are all Aa heterozygotes and that the AA homozygotes produced by matings between heterozygotes do not survive to the adult stage. Embryonic death is, in fact, why the yellow mice are underrepresented in the F₂ data. Examination of the uteruses of pregnant females reveals that about 1/4 of the embryos are dead. These dead embryos must be genotypically AA. Thus, a single copy of the Aallele produces a visible phenotypic effect (yellow fur), but two copies cause death. Taking this embryonic mortality into account, we can modify the hypothesis and predict that 2/3 of the live-born F_2 progeny should be yellow (*Aa*) and 1/3 should be agouti (*aa*). We can then use the χ^2 procedure to test this modified hypothesis for consistency with the data.

F ₂ Phenotype	Obs	Exp	(Obs – Exp) ² /Exp
yellow (Aa)	54	$(2/3) \times 84 = 56$	0.071
agouti (aa)	30	$(1/3) \times 84 = 28$	0.143
Total	84	84	0.214

This χ^2 statistic is less than the critical value for a χ^2 distribution with 1 degree of freedom. Thus, the data are in agreement with the predictions of the modified hypothesis.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **3.1** On the basis of Mendel's observations, predict the results from the following crosses with peas:
 - (a) A tall (dominant and homozygous) variety crossed with a dwarf variety.
 - (b) The progeny of (a) self-fertilized.
 - (c) The progeny from (a) crossed with the original tall parent.
 - (d) The progeny of (a) crossed with the original dwarf parent.
- **3.2** Mendel crossed pea plants that produced round seeds with those that produced wrinkled seeds and self-fertilized the progeny. In the F_2 , he observed 5474 round seeds and 1850 wrinkled seeds. Using the letters W and w for the seed texture alleles, diagram Mendel's crosses, showing the genotypes of the plants in each generation. Are the results consistent with the Principle of Segregation?
- **3.3** A geneticist crossed wild, gray-colored mice with white (albino) mice. All the progeny were gray. These progeny were intercrossed to produce an F₂, which consisted of 198 gray and 72 white mice. Propose an hypothesis to explain these results, diagram the crosses, and compare the results with the predictions of the hypothesis.
- **3.4** A woman has a rare abnormality of the eyelids called ptosis, which prevents her from opening her eyes completely. This condition is caused by a dominant allele, *P*. The woman's father had ptosis, but her mother had normal eyelids. Her father's mother had normal eyelids.
 - (a) What are the genotypes of the woman, her father, and her mother?
 - (b) What proportion of the woman's children will have ptosis if she marries a man with normal eyelids?
- **3.5** In pigeons, a dominant allele *C* causes a checkered pattern in the feathers; its recessive allele *c* produces a plain

pattern. Feather coloration is controlled by an independently assorting gene; the dominant allele B produces red feathers, and the recessive allele b produces brown feathers. Birds from a true-breeding checkered, red variety are crossed to birds from a true-breeding plain, brown variety.

- (a) Predict the phenotype of their progeny.
- (b) If these progeny are intercrossed, what phenotypes will appear in the F₂, and in what proportions?
- **3.6 (C)** In mice, the allele C for colored fur is dominant over the allele c for white fur, and the allele V for normal behavior is dominant over the allele v for waltzing behavior, a form of discoordination. Give the genotypes of the parents in each of the following crosses:
 - (a) Colored, normal mice mated with white, normal mice produced 29 colored, normal and 10 colored, waltzing progeny.
 - (b) Colored, normal mice mated with colored, normal mice produced 38 colored, normal, 15 colored, waltzing, 11 white, normal, and 4 white, waltzing progeny.
 - (c) Colored, normal mice mated with white, waltzing mice produced 8 colored, normal, 7 colored, waltzing, 9 white, normal, and 6 white, waltzing progeny.
- **3.7** In rabbits, the dominant allele *B* causes black fur and the recessive allele *b* causes brown fur; for an independently assorting gene, the dominant allele *R* causes long fur and the recessive allele *r* (for *rex*) causes short fur. A homozygous rabbit with long, black fur is crossed with a rabbit with short, brown fur, and the offspring are intercrossed. In the F_2 , what proportion of the rabbits with long, black fur will be homozygous for both genes?
- **3.8** In shorthorn cattle, the genotype *RR* causes a red coat, the genotype *rr* causes a white coat, and the genotype *Rr* causes a roan coat. A breeder has red, white, and roan cows

and bulls. What phenotypes might be expected from the following matings, and in what proportions?

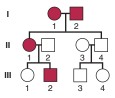
- (a) red \times red;
- (b) red \times roan;
- (c) red \times white;
- (d) roan \times roan.
- **3.9** How many different kinds of F_1 gametes, F_2 genotypes, and F_2 phenotypes would be expected from the following crosses:
 - (a) $AA \times aa;$
 - (b) $AA BB \times aa bb;$
 - (c) $AA BB CC \times aa bb cc?$
 - (d) What general formulas are suggested by these answers?
- **3.10 (a)** A researcher studied six independently assorting genes in a plant. Each gene has a dominant and a recessive allele: *R* black stem, *r* red stem; *D* tall plant, *d* dwarf plant; *C* full pods, *c* constricted pods; *O* round fruit, *o* oval fruit; *H* hairless leaves, *b* hairy leaves; *W* purple flower, *w* white flower. From the cross (P1) *Rr Dd cc Oo Hh Ww* × (P2) *Rr dd Cc oo Hh ww*,
 - (a) How many kinds of gametes can be formed by P1?
 - (b) How many genotypes are possible among the progeny of this cross?
 - (c) How many phenotypes are possible among the progeny?
 - (d) What is the probability of obtaining the *Rr Dd cc Oo hh ww* genotype in the progeny?
 - (e) What is the probability of obtaining a black, dwarf, constricted, oval, hairy, purple phenotype in the progeny?
- **3.11** For each of the following situations, determine the degrees of freedom associated with the χ^2 statistic and decide whether or not the observed χ^2 value warrants acceptance or rejection of the hypothesized genetic ratio.

	Hypothesized Ratio	Observed χ^2
(a)	3:1	7.0
(b)	1:2:1	7.0
(c)	1:1:1:1	7.0
(d)	9:3:3:1	5.0

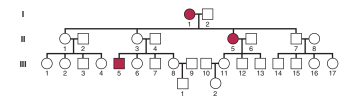
- **3.12 (co)** Mendel testcrossed pea plants grown from yellow, round F₁ seeds to plants grown from green, wrinkled seeds and obtained the following results: 31 yellow, round; 26 green, round; 27 yellow, wrinkled; and 26 green, wrinkled. Are these results consistent with the hypothesis that seed color and seed texture are controlled by independently assorting genes, each segregating two alleles?
- **3.13** Perform a chi-square test to determine if an observed ratio of 30 tall: 20 dwarf pea plants is consistent with an expected ratio of 1:1 from the cross $Dd \times dd$.
- **3.14** Seed capsules of the Shepherd's purse are either triangular or ovoid. A cross between a plant with triangular seed capsules and a plant with ovoid seed capsules yielded F_1 hybrids that all had triangular seed capsules. When these F_1 hybrids were intercrossed, they produced 80 F_2 plants, 72 of which

had triangular seed capsules and 8 of which had ovoid seed capsules. Are these results consistent with the hypothesis that capsule shape is determined by a single gene with two alleles?

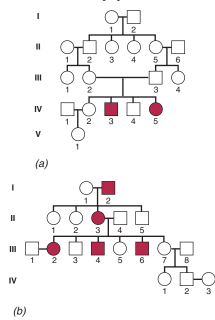
- **3.15** Albinism in humans is caused by a recessive allele *a*. From marriages between people known to be carriers (*Aa*) and people with albinism (*aa*), what proportion of the children would be expected to have albinism? Among three children, what is the chance of one without albinism and two with albinism?
- 3.16 If both husband and wife are known to be carriers of the allele for albinism, what is the chance of the following combinations in a family of four children: (a) all four unaffected; (b) three unaffected and one affected; (c) two unaffected and two affected; (d) one unaffected and three affected?
- 3.17 In humans, cataracts in the eyes and fragility of the bones are caused by dominant alleles that assort independently. A man with cataracts and normal bones marries a woman without cataracts but with fragile bones. The man's father had normal eyes, and the woman's father had normal bones. What is the probability that the first child of this couple will (a) be free from both abnormalities; (b) have cataracts but not have fragile bones; (c) have fragile bones but not have cataracts; (d) have both cataracts and fragile bones?
- **3.18** In generation V in the pedigree in Figure 3.15, what is the probability of observing seven children without the cancer-causing mutation and two children with this mutation among a total of nine children?
- **3.19** If a man and a woman are heterozygous for a gene, and if they have three children, what is the chance that all three will also be heterozygous?
- **3.20** If four babies are born on a given day:
 - (a) What is the chance that two will be boys and two girls?
 - (b) What is the chance that all four will be girls?
 - (c) What combination of boys and girls among four babies is most likely?
 - (d) What is the chance that at least one baby will be a girl?
- **3.21** In a family of six children, what is the chance that at least three are girls?
- 3.22 The following pedigree shows the inheritance of a dominant trait. What is the chance that the offspring of the following matings will show the trait: (a) III-1 × III-3; (b) III-2 × III-4?



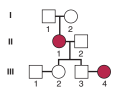
3.23 The following pedigree shows the inheritance of a recessive trait. Unless there is evidence to the contrary, assume that the individuals who have married into the family do not carry the recessive allele. What is the chance that the offspring of the following matings will show the trait: (a) III-1 \times III-12; (b) III-4 \times III-14; (c) III-6 \times III-13; (d) IV-1 \times IV-2?



3.24 In the following pedigrees, determine whether the trait is more likely to be due to a dominant or a recessive allele. Assume the trait is rare in the population.



- **3.25** In pedigree (*b*) of Problem 3.24, what is the chance that the couple III-1 and III-2 will have an affected child? What is the chance that the couple IV-2 and IV-3 will have an affected child?
- **3.26 (a)** Peas heterozygous for three independently assorting genes were intercrossed.
 - (a) What proportion of the offspring will be homozygous for all three recessive alleles?
 - (b) What proportion of the offspring will be homozygous for all three genes?
 - (c) What proportion of the offspring will be homozygous for one gene and heterozygous for the other two?
 - (d) What proportion of the offspring will be homozygous for the recessive allele of at least one gene?
- **3.27** The following pedigree shows the inheritance of a recessive trait. What is the chance that the couple III-3 and III-4 will have an affected child?



- 3.28 a geneticist crosses tall pea plants with short pea plants. All the F_1 plants are tall. The F_1 plants are then allowed to self-fertilize, and the F_2 plants are classified by height: 62 tall and 26 short. From these results, the geneticist concludes that shortness in peas is due to a recessive allele (s) and that tallness is due to a dominant allele (S). On this hypothesis, 2/3 of the tall F_2 plants should be heterozygous Ss. To test this prediction, the geneticist uses pollen from each of the 62 tall plants to fertilize the ovules of emasculated flowers on short pea plants. The next year, three seeds from each of the 62 crosses are sown in the garden and the resulting plants are grown to maturity. If none of the three plants from a cross is short, the male parent is classified as having been homozygous SS; if at least one of the three plants from a cross is short, the male parent is classified as having been heterozygous Ss. Using this system of progeny testing, the geneticist concludes that 29 of the 62 tall F₂ plants were homozygous SS and that 33 of these plants were heterozygous Ss.
 - (a) Using the chi-square procedure, evaluate these results for goodness of fit to the prediction that 2/3 of the tall F_2 plants should be heterozygous.
 - (b) Informed by what you read in A Milestone in Genetics: Mendel's 1866 Paper (which you can find in the Student Companion site), explain why the geneticist's procedure for classifying tall F₂ plants by genotype is not definitive.
 - (c) Adjust for the uncertainty in the geneticist's classification procedure and calculate the expected frequencies of homozygotes and heterozygotes among the tall F₂ plants.
 - (d) Evaluate the predictions obtained in (c) using the chisquare procedure.
- **3.29** A researcher who has been studying albinism has identified a large group of families with four children in which at least one child shows albinism. None of the parents in this group of families shows albinism. Among the children, the ratio of those without albinism to those with albinism is 1.7:1. The researcher is surprised by this result because he thought that a 3:1 ratio would be expected on the basis of Mendel's Principle of Segregation. Can you explain the apparently non-Mendelian segregation ratio in the researcher's data?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

- 1. Gregor Mendel worked out the rules of inheritance by performing experiments with peas (*Pisum sativum*). Has the genome of this organism been sequenced, or is it currently being sequenced?
- 2. Which plant genomes have been sequenced completely?
- **3.** What is the scientific or agricultural significance of the plants whose genomes have been sequenced completely?

Hint: At the web site, click on Genomes and Maps, then on Genome Project, and finally on Plant Genomes.

Extensions of Mendelism

CHAPTER OUTLINE

- Allelic Variation and Gene Function
- **Gene Action: From Genotype to Phenotype**
- Inbreeding: Another Look at Pedigrees

Genetics Grows beyond Mendel's <mark>Mon</mark>astery Garden

In 1902, enthused by what he read in Mendel's paper, the British biologist William Bateson published an English translation of Mendel's German text and appended to it a brief account of what he called "Mendelism—the Principles of Dominance, Segregation, and Independent Assortment." Later, in 1909, he published *Mendel's Principles of Heredity*, in which he summarized all the evidence then available to support Mendel's findings. This book was remarkable for two reasons. First, it examined the results of breeding experiments with many different plants and animals and in each case demonstrated that Mendel's principles applied. Second, it considered the implications of these experiments and raised questions about the fundamental nature of genes, or, as Bateson called them, "unit-characters." At the time Bateson's book was published, the word "gene" had not yet been invented.

Bateson's book played a crucial role in spreading the principles of Mendelism to the scientific world. Botanists, zoologists, naturalists, horticulturalists, and animal breeders got the message in plain and simple language: Mendel's principles—tested by experiments with peas, beans, sunflowers, cotton, wheat, barley, tomatoes, maize, and assorted ornamental plants, as well as with cattle, sheep, cats, mice, rabbits, guinea pigs, chickens, pigeons, canaries, and moths were universal. In the preface to his book, Bateson remarked that "The study of heredity thus becomes an organized branch of physiological science, already abundant in results, and in promise unsurpassed."1

¹Bateson, W. 1909. *Mendel's Principles of Heredity*. University Press, Cambridge, England.



Diverse species of plants growing in a garden. Experiments with many different plants extended Mendel's Principles of Dominance, Segregation, and Independent Assortment.

Allelic Variation and Gene Function

Mendel's experiments established that genes can exist in alternate forms. For each of the seven traits that he studied—seed color, seed texture, plant height, flower color, flower position, pod shape, and pod

color—Mendel identified two alleles, one dominant, the other recessive. This discovery suggested a simple functional dichotomy between alleles, as if one allele did nothing and the other did everything to determine the phenotype. However, research early in the twentieth century demonstrated this to be an oversimplification. Genes can exist in more than two allelic states, and each allele can have a different effect on the phenotype.

INCOMPLETE DOMINANCE AND CODOMINANCE

An allele is dominant if it has the same phenotypic effect in heterozygotes as in homozygotes—that is, the genotypes Aa and AA are phenotypically indistinguishable. Sometimes, however, a heterozygote has a phenotype different from that of either of its associated homozygotes. Flower color in the snapdragon, *Antirrbinum majus*, is an example. White and red varieties are homozygous for different alleles of a color-determining gene; when crossed, they produce heterozygotes that have pink flowers. The allele for red color (W) is therefore said to be **incompletely**, or **partially**, **dominant** over the allele for white color (w). The most likely explanation is that the intensity of pigmentation in this species depends on the amount of a product specified by the color gene (**Figure 4.1**). If the W allele specifies this product as Ww heterozygotes do and will therefore show deeper color. When the heterozygote's phenotype is midway between the phenotypes of the two homozygotes, as it is here, the partially dominant allele is sometimes said to be **semidominant** (from the Latin word for "half"—thus half-dominant).

Another exception to the principle of simple dominance arises when a heterozygote shows characteristics found in each of the associated homozygotes. This occurs with human blood types, which are identified by testing for special cellular products called *antigens*. An antigen is detected by its ability to react with factors obtained from the serum portion of the blood. These factors, which are produced by the immune system, recognize antigens quite specifically. Thus, for example, one serum, called anti-M, recognizes only the M antigen on human blood cells; another serum, called

anti-N, recognizes only the N antigen on these cells (**■** Figure 4.2). When one of these sera detects its specific antigen in a blood-typing test, the cells clump together in a reaction called *agglutination*. Thus, by testing cells for agglutination with different sera, a medical technologist can identify which antigens are present and thereby determine the blood type.

The ability to produce the M and N antigens is determined by a gene with two alleles. One allele allows the M antigen to be produced; the other allows the N antigen to be produced. Homozygotes for the M allele produce only the M antigen, and homozygotes for the N allele produce only the N antigen. However, heterozygotes for these two alleles produce both kinds of antigens. Because the two alleles appear to contribute independently to the phenotype of the heterozygotes, they are said to be **codominant.** Codominance implies that there is an independence of allele function. Neither allele is dominant, or even partially dominant, over the other. It would therefore be inappropriate to distinguish the alleles by

upper- and lowercase letters, as we have in all previous examples. Instead, codominant alleles are represented by superscripts on the symbol for the gene, which in this case is the letter L—a tribute to Karl Landsteiner, the discoverer of blood-typing. Thus, the

PhenotypeGenotypeAmount of
gene productRedWW2xPinkWwxWhiteww0

■ FIGURE 4.1 Genetic basis of flower color in snapdragons. The allele *W* is incompletely dominant over *w*. Differences among the phenotypes could be due to differences in the amount of the product specified by the *W* allele.

Genotype	Blood type (antigen present)	Reactions w	ith anti-sera
		Anti-M serum	Anti-N serum
LM LM	M (M)		•••••
LM LN	M N (M and N)	4. 4. X.	14. 4. 4 17. 17
LN LN	N (N)		

■ FIGURE 4.2 Detection of the M and N antigens on blood cells by agglutination with specific anti-sera. With the anti-M and anti-N sera, three blood types can be identified.

The diverse kinds of alleles of genes affect phenotypes in different ways.

M	Genotype	Phenotype
Carlo Carlo	сс	White hairs over the entire body
Albino		
	c ^h c ^h	Black hairs on the extremities; white hairs everywhere else
Himalayan		
	c ^{ch} c ^{ch}	White hair with black tips on the body
Chinchilla		
	C ⁺ C ⁺	Colored hairs over the entire body
Wild-type		

FIGURE 4.3 Coat colors in rabbits. The different phenotypes are caused by four different alleles of the *c* gene.

M allele is L^M and the N allele is L^N . Figure 4.2 shows the three possible genotypes formed by the L^M and L^N alleles, and their associated phenotypes.

MULTIPLE ALLELES

The Mendelian concept that genes exist in no more than two allelic states had to be modified when genes with three, four, or more alleles were discovered. A classic example of a gene with **multiple alleles** is the one that controls coat color in rabbits (**True Figure 4.3**). The color-determining gene, denoted by the lowercase letter *c*, has four alleles, three of which are distinguished by a superscript: *c (albino)*, *c^b (bimalayan)*, *c^{cb} (cbimchilla)*, and *c⁺* (wild-type). In homozygous condition, each allele has a characteristic effect on the coat color. Because most rabbits in wild populations are homozygous for the *c*⁺ allele, this allele is called the **wild type**. In genetics it is customary to represent wild-type alleles by a superscript plus sign after the letter for the gene. When the context is clear, the letter is sometimes omitted and only the plus sign is used; thus, *c*⁺ may be abbreviated simply as +.

The other alleles of the *c* gene are **mutants**—altered forms of the wildtype allele that must have arisen sometime during the evolution of the rabbit. The *himalayan* and *chinchilla* alleles are denoted by superscripts, but the *albino* allele is denoted simply by the letter *c* (for colorless, another word for the albino condition). This notation reflects another custom in genetics nomenclature: genes are often named for a mutant allele, usually the allele associated with the most abnormal phenotype. The convention of naming a gene for a mutant allele is generally consistent with the convention we discussed in Chapter 3—that of naming genes for a recessive allele—because most mutant alleles are recessive. However, sometimes a mutant allele is dominant, in which case the gene is named after its associated phenotype. For example, a gene in mice controls the length of the tail. The first mutant allele of this gene that was discovered caused a shortening of the tail in heterozygotes. This dominant mutant was therefore symbolized by *T*, for *t*ail-length. All other alleles of this gene—and there are many—have been denoted by an uppercase or lowercase letter, depending on whether they are dominant or recessive; different alleles are distinguished from each other by superscripts.

Another example of multiple alleles comes from the study of human blood types. The A, B, AB, and O blood types, like the M, N, and MN blood types discussed previously, are identified by testing a blood sample with different sera. One serum detects the A antigen, another the B antigen. When only the A antigen is present on the cells, the blood is type A; when only the B antigen is present, the blood is type B. When both antigens are present, the blood is type AB, and when neither antigen is present, it is type O. Blood-typing for the A and B antigens is completely independent of blood-typing for the M and N antigens.

The gene responsible for producing the A and B antigens is denoted by the letter *I*. It has three alleles: I^A , I^B , and *i*. The I^A allele specifies the production of the A antigen, and the I^B allele specifies the production of the B antigen. However, the *i* allele does not specify an antigen. Among the six possible genotypes, there are four distinguishable phenotypes—the A, B, AB, and O blood types (Table 4.1). In this system, the I^A and

TABLE 4.1

Genotypes, Ph	enotypes, and	Frequencies in th	e ABO Blood-Typi	ng System
Genotype	Blood Type	A Antigen Present	B Antigen Present	Frequency in U.S. White Population (%)
l ^a l ^a or l ^a i	А	+	_	41
l ^B l ^B or l ^B i	В	—	+	11
$I^{A}I^{B}$	AB	+	+	4
ii	0	-	-	44

 I^{B} alleles are codominant, since each is expressed equally in the I^{A} I^{B} heterozygotes, and the *i* allele is recessive to both the I^{A} and I^{B} alleles. All three alleles are found at appreciable frequencies in human populations; thus, the *I* gene is said to be **polymorphic**, from the Greek words for "having many forms." We consider the population and evolutionary significance of genetic polymorphisms in Chapter 24.

ALLELIC SERIES

The functional relationships among the members of a series of multiple alleles can be studied by making heterozygous combinations through crosses between homozygotes. For example, the four alleles of the *c* gene in rabbits can be combined with each other to make six different kinds of heterozygotes: $c^b c$, $c^{cb} c$, $c^+ c$, $c^{cb} c^b$, $c^+ c^b$, and $c^+ c^{cb}$. These heterozygotes allow the dominance relations among the alleles to be studied (**■** Figure 4.4). The wild-type allele is completely dominant over all the other alleles in the series; the *chinchilla* allele is partially dominant over the *himalayan* and *albino* alleles, and the *himalayan* allele is completely dominant over the *albino* allele. These dominance relations can be summarized as $c^+ > c^{cb} > c^b > c$.

Notice that the dominance hierarchy parallels the effects that the alleles have on coat color. A plausible explanation is that the *c* gene controls a step in the formation of black pigment in the fur. The wild-type allele is fully functional in this process, producing colored hairs throughout the body. The *chinchilla* and *himalayan* alleles are only partially functional, producing some colored hairs, and the *albino* allele is not functional at all. Nonfunctional alleles are said to be **nutt** or **amorphic** (from the Greek words for "without form"); they are almost always completely recessive. Partially functional alleles are said to be **hypomorphic** (from the Greek words for "beneath form"); they are recessive to alleles that are more functional, including (usually) the wild-type allele. Later in this chapter we consider the biochemical basis for these differences.

TESTING GENE MUTATIONS FOR ALLELISM

A mutant allele is created when an existing allele changes to a new genetic state—a process called **mutation**. This event always involves a change in the physical composition of the gene (see Chapter 13) and sometimes produces an allele that has a detectable phenotypic effect. If, for example, the c^+ allele mutated to a null allele, a rabbit homozygous for this mutation would have the albino phenotype. However, it is not always possible to assign a new mutation to a gene on the basis of its phenotypic effect. In rabbits, for example, several genes determine coat color, and a mutation in any one of them could reduce, alter, or abolish pigmentation in the hairs. Thus, if a new coat color appears in a population of rabbits, it is not immediately clear which gene has been mutated.

A simple test can be used to determine the allelic identity of a new mutation, providing that the new mutation is recessive. The procedure involves crosses to combine the new recessive mutation with recessive mutations of known genes (**Figure 4.5**). If the hybrid progeny show a mutant phenotype, then the new mutation and the tester mutation *are* alleles of the same gene. If the hybrid progeny show a wild phenotype,

then the new mutation and the tester mutation *are not* alleles of the same gene. This test is based on the principle that mutations of the same gene impair the same genetic function. If two such mutations are combined, the organism will be abnormal for this function and will show a mutant phenotype, even if the two mutations had an independent origin.

It is important to remember that this test applies only to recessive mutations. Dominant mutations cannot be tested in this way because they exert their effects even if a wild-type copy of the gene is present.

As an example, let's consider the analysis of two recessive mutations affecting eye color in the fruit fly, *Drosophila melanogaster* (**Figure 4.6**). This organism has been investigated by geneticists for a century, and a great many different mutations have

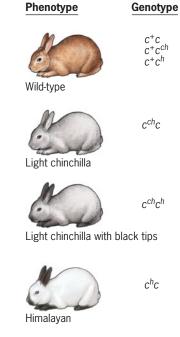


FIGURE 4.4 Phenotypes of different combinations of *c* alleles in rabbits. The alleles form a series, with the wild-type allele, c^+ , dominant over all the other alleles and the null allele, *c* (*albino*), recessive to all the other alleles; one hypomorphic allele, c^{ch} (*chinchilla*), is partially dominant over the other, c^h (*himalayan*).

New recessive mutation	Tester genotype	Hybrid phenotype	Conclusion
c*c* X ≤	aa 🛶 bb →	Wild-typeWild-type	a and c* not alleles b and c* not alleles
		Mutant	c and c* alleles
	dd →	 Wild-type 	d and c [*] not alleles

■ FIGURE 4.5 A general scheme to test recessive mutations for allelism. Two mutations are alleles if a hybrid that contains both of them has the mutant phenotype.

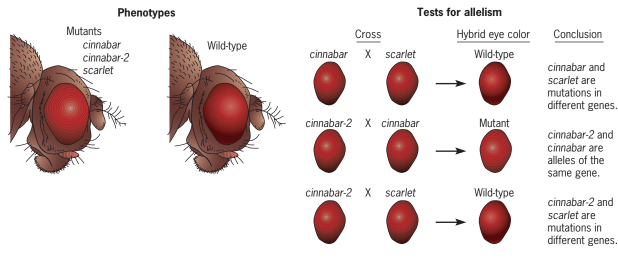


FIGURE 4.6 A test for allelism involving recessive eye color mutations in *Drosophila*. Three phenotypically identical mutations, *cinnabar*, *scarlet*, and *cinnabar*-2, are tested for allelism by making pairwise crosses between flies homozygous for different mutations. The phenotypes of the hybrids show that the *cinnabar* and *cinnabar*-2 mutations are alleles of a single gene and that the *scarlet* mutation is not an allele of this gene.

Solve It!

The Test for Allelism

Two researchers working independently have each discovered an albino mouse in their large breeding colonies of wild-type animals. Genetic testing indicates that each of these mice is homozygous for a recessive mutation that prevents pigment formation. An albino mouse from one colony is crossed to an albino mouse from the other colony, and all the offspring have wild-type body color. Are the two albino mutations allelic?

To see the solution to this problem, visit the Student Companion site. been identified. Two independently isolated recessive mutations, called *cinnabar* and *scarlet*, are phenotypically indistinguishable, each causing the eyes to be bright red. In wild-type flies, the eyes are dark red. We wish to know whether the *cinnabar* and *scarlet* mutations are alleles of a single color-determining gene or if they are mutations in two different genes. To find the answer, we must cross the homozygous mutant strains with each other to produce hybrid progeny. If the hybrids have bright red eyes, we will conclude that *cinnabar* and *scarlet* are alleles of the same gene. If they have dark red eyes, we will conclude that they are mutations in different genes.

The hybrid progeny turn out to have dark red eyes; that is, they are wild type rather than mutant. Thus, *cinnabar* and *scarlet* are not alleles of the same gene but, rather, mutations in two different genes, each apparently involved in the control of eye pigmentation. When we test a third mutation, called *cinnabar*-2, for allelism with the *cinnabar* and *scarlet* mutations, we find that the hybrid combination of *cinnabar*-2 and *scarlet* has the mutant phenotype (bright red eyes) and that the hybrid combination of *cinnabar*-2 and *scarlet* has the wild phenotype (dark red eyes). These results tell us that the mutations *cinnabar* and *cinnabar*-2 are alleles of one color-determining gene and that the *scarlet* mutation is not an allele of this gene. Rather, the *scarlet* mutation defines another color-determining gene.

The test to determine whether mutations are alleles of a particular gene is based on the phenotypic effect of combining the mutations in the same individual. If the hybrid combination is mutant, we conclude that the mutations are alleles; if it is wildtype, we conclude that they are not alleles. Chapter 13 discusses how this test—called the *complementation test* in modern terminology—enables geneticists to define the functions of individual genes. To solidify your understanding of the concepts discussed here, try Solve It: The Test for Allelism.

VARIATION AMONG THE EFFECTS OF MUTATIONS

Genes are identified by mutations that alter the phenotype in some conspicuous way. For instance, a mutation may change the color or shape of the eyes, alter a behavior, or cause sterility or even death. The tremendous variation among the effects of individual mutations suggests that each organism carries many different kinds of genes and that each of these can mutate in different ways. In nature, mutations provide the raw material for evolution (see Chapter 24).

Mutations that alter some aspect of morphology, such as seed texture or color, are called *visible mutations*. Most visible mutations are recessive, but a small number

of them are dominant. Geneticists have learned much about genes by analyzing the properties of these mutations. We will encounter many examples of this analysis throughout this textbook. Mutations that limit reproduction are called *sterile mutations*. Some sterile mutations affect both sexes, but most affect either males *or* females. As with visible mutations, steriles can be either dominant or recessive. Some steriles completely prevent reproduction, whereas others only impair it slightly.

Mutations that interfere with necessary vital functions are called *lethal mutations*. Their phenotypic effect is death. We know that many genes are capable of mutating to the lethal state. Thus, each of these genes is absolutely essential for life. Dominant lethals that act early in life are lost one generation after they occur because the individuals that carry them die; however, dominant lethals that act later in life, after reproduction, can be passed on to the next generation. Recessive lethals may linger a long time in a population because they can be hidden in heterozygous condition by a wild-type allele. Recessive lethal mutations are detected by observing unusual segregation ratios in the progeny of heterozygous carriers. An example is the *yellow-lethal* mutation, A^Y , in the mouse (**Figure 4.7**). This mutation is a dominant visible, causing the fur to be yellow instead of gray-brown (the wild-type color, also known as *agouti*, which is determined by the allele A^+). In addition, the A^Y mutation is a recessive lethal, killing $A^Y A^Y$ homozygotes early in their development. A cross between $A^Y A^+$ heterozygotes produces two kinds of viable progeny, yellow ($A^Y A^+$) and gray-brown ($A^+ A^+$), in a ratio of 2:1. The $A^Y A^Y$ homozygotes die during embryonic development.

Geneticists have used different conventions to symbolize genes and their mutations. Mendel began the practice of using letters to denote genes. However, he simply started with the letter A and proceeded through the alphabet as symbols were needed to represent genes in his crosses. William Bateson was the first person to use letters mnemonically to symbolize genes. For the symbol, Bateson chose the first letter of the word that described the gene's phenotypic effect—thus, B for a gene causing blue flowers, L for a gene causing long pollen grains. As the number of known genes grew, it became necessary to use two or more letters to represent newly discovered genes. Unfortunately, geneticists do not all follow the same conventions when they represent genes and alleles. Some of their practices are discussed in Focus on Genetic Symbols.

GENES FUNCTION TO PRODUCE POLYPEPTIDES

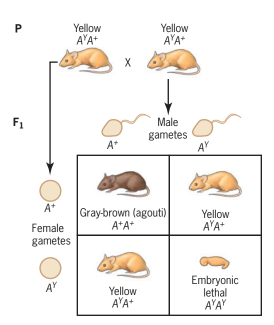
The extensive variation revealed by mutations indicates that organisms contain many different genes and that these genes can exist in multiple allelic states. However, it does not tell us how genes actually affect the phenotype. What is it about a gene that enables it to influence a trait such as eye color, seed texture, or plant height?

The early geneticists had no answer to this question. However, today it is clear that most genes specify a product that subsequently affects the phenotype. This idea, which was discussed in Bateson's book and which was supported by the research of many scientists, including, most notably, the British physician Sir Archibald Garrod (see A Milestone in Genetics: Garrod's Inborn Errors of Metabolism in the Student Companion site), was forcefully brought out in the middle of the twentieth century when George Beadle and Edward Tatum discovered that the prod-

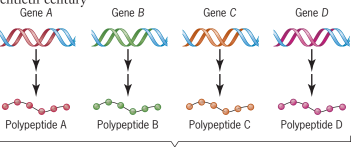
ucts of genes are *polypeptides* (**■** Figure 4.8).

Polypeptides are macromolecules built of a linear chain of *amino acids*. Every organism makes thousands of different polypeptides, each characterized by a specific amino acid sequence. These polypeptides are the fundamental constituents of *proteins*. Two or more polypeptides may combine to form a protein. Some proteins, called *enzymes*, function as catalysts in biochemical reactions; others form the structural components of cells; and still others are responsible for transporting substances within and between

cells. Beadle and Tatum proposed that each gene is responsible for the synthesis of a particular polypeptide. When a gene is mutated, its polypeptide product either is not made or is altered in such a way that its role in the organism is changed. Mutations that eliminate or alter a polypeptide are often associated with a phenotypic effect.



■ **FIGURE 4.7** A^{*Y*}, the *yellow-lethal* mutation in mice: a dominant visible that is also a recessive lethal. A cross between carriers of this mutation produces yellow heterozygotes and gray-brown (agouti) homozygotes in a ratio of 2:1. The yellow homozygotes die as embryos.



Aspects of phenotype

FIGURE 4.8 Relationship between genes and polypeptides. Each gene specifies a different polypeptide. These polypeptides then function to influence the organism's phenotype.

FOCUS ON

GENETIC SYMBOLS

We will a state of the practice of choosing gene symbols mnemonically. In discussing Mendel's work, for example, he symbolized the dominant allele for tall pea plants as *T* and the recessive allele for short plants as *t*. Later, when it became customary to choose allele symbols based on the mutant trait, these symbols were changed to *D* (for tall) and *d* (for *d*warf). This convention provided a simple and consistent notation in which the dominant and recessive alleles of a particular gene were represented by a single letter, and that letter was mnemonic for the trait influenced by the gene. Bateson also coined the words *genetics*, *alleleomorph* (which was later shortened to *allele*), *homozygote*, and *heterozygote*, and he introduced the practice of denoting the generations in a breeding scheme as P, F₁, F₂, and so forth.

The gene-naming system that Bateson developed worked well until the number of genes that had been identified exceeded the capacity of the English alphabet; thereupon it became necessary to use two or more letters to symbolize a gene. For example, a particular mutant allele in *Drosophila* causes the eyes to be carmine instead of red. When this allele was discovered, it was given the symbol *cm* because the single letter *c* had already been used to represent a mutant allele that causes the wings to be curved instead of straight.

The discovery of multiple alleles made genetic notation even more complicated; because upper- and lowercase letters were no longer adequate to distinguish among alleles, geneticists began to combine a basic gene symbol with an identification symbol. *Drosophila* geneticists were the first to apply this procedure. They made the identification symbol a superscript on the basic gene symbol. Usually, both the gene symbol and the superscript had some mnemonic significance. Thus, for example, *cn*² was used to symbolize the *second cinn*abar eye color allele that was discovered in *Drosophila*; and *ey^o* was used to symbolize a *d*ominant allele that causes *Drosophila* to be *eyeless*. Plant geneticists adopted a variation of this practice. They use hyphenated symbols to identify mutant alleles; for example, *sh2–6801* represents a mutant allele for *sh*runken maize kernels that was discovered in 1968.

As genetic nomenclature developed, it became necessary to use a special symbol to represent the wild-type allele. The early *Drosophila* geneticists proposed using a plus sign (+), sometimes written as a superscript on the basic gene symbol (for example, c^+). This simple notation conveys the idea that the wild-type allele is the standard, or normal, allele of the gene, and is widely used today. However other gene-naming practices persist. Plant geneticists tend to use the gene symbol itself to represent the wild-type allele, but to make it stand out, they capitalize the first letter. Thus, *Sh2* is the wild-type allele of the second shrunken kernel gene discovered in maize, whereas *sh2* is a mutant allele.

Genetic nomenclature has been further complicated by the discovery of genes through the polypeptides they specify. These discoveries have introduced gene symbols that are mnemonic for polypeptide gene products. For example, the human gene that specifies the polypeptide *hypoxanthine-guanine phosphoribosyl transferase* is symbolized by *HPRT*, and the plant gene that specifies the polypeptide *a*lcohol *dehy*drogenase is symbolized by *Adh*. Whether uppercase letters are used throughout the gene symbol or only for the first letter depends on the organism.

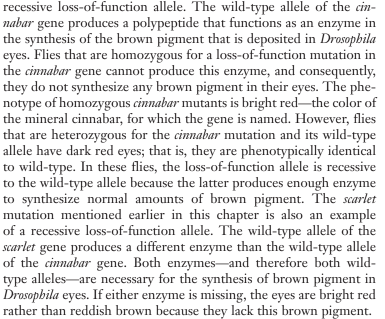
Today there are many specialized systems for symbolizing genes and alleles. Researchers who work with different organisms— *Drosophila*, mice, plants, or humans—speak slightly different languages. Later, we will see that still other genetic dialects have been created to describe the genes of viruses, bacteria, and fungi. These different systems of nomenclature indicate that the symbols in genetics have evolved in response to new discoveries—visible evidence of growth in a dynamic, young science.

Whether this effect is dominant or recessive depends on the nature of the mutation. In Chapter 12 we consider the details of how genes produce polypeptides, and in Chapter 13 we discuss the molecular basis of mutation.

WHY ARE SOME MUTATIONS DOMINANT AND OTHERS RECESSIVE?

The discovery that genes specify polypeptides provides insight into the nature of dominant and recessive mutations. Dominant mutations have phenotypic effects in heterozygotes as well as in homozygotes, whereas recessive mutations have these effects only in homozygotes. What accounts for this striking difference in expression?

Recessive mutations often involve a loss of gene function, that is, when the gene no longer specifies a polypeptide or when it specifies a nonfunctional or underfunctional polypeptide (**Figure 4.9**). Recessive mutations are therefore typically **loss-of-function** alleles. Such alleles have little or no discernible effect in heterozygous condition with a wild-type allele because the wild-type allele specifies a functional polypeptide that will carry out its normal role in the organism. The phenotype of a mutant/wild heterozygote will therefore be the same, or essentially the same, as that of a wild-type homozygote. The *cinnabar* mutation in *Drosophila* is an example of a



Some recessive mutations result in a partial loss of gene function. For example, the *himalayan* allele of the coat color gene in mammals such as rabbits and cats specifies a polypeptide that functions only in the parts of the body where the temperature is reduced. This partial loss of function explains why animals homozygous for the *himalayan* allele have pigmented hair on their extremities—tail, legs, ears, and tip of the nose—but not on the rest of their bodies. In the extremities, the polypeptide specified by this allele is functional, whereas in the rest of the body, it is not. The expression of the *himalayan* allele is therefore temperature-sensitive.

Some dominant mutations may also involve a loss of gene function. If the phenotype controlled by a gene is sensitive to the amount of gene product, a loss-of-function mutation can evoke a mutant phenotype in heterozygous condition with a wild-type

allele. In such cases, the wild-type allele, by itself, is not able to supply enough gene product to provide full, normal function. In effect, the loss-of-function mutation reduces the level of gene product below the level that is needed for the wild phenotype.

Other dominant mutations actually interfere with the function of the wild-type allele by specifying polypeptides that inhibit, antagonize, or limit the activity of the wild-type polypeptide (Figure 4.9). Such mutations are called **dominant-negative mutations**. Some of the mutations of the T gene in the mouse are examples of dominant-negative mutations. We have already seen that in heterozygous condition, these mutations cause a shortening of the tail. In homozygous condition, they are lethal. The wild-type allele of the T gene is therefore essential for life. At the cellular level, the polypeptide product of this allele regulates important events during embryological development. Dominant-negative T alleles produce slightly shorter polypeptides than the wild-type T allele. In heterozygotes, these shorter polypeptides interfere with the function of the wild-type polypeptide. The result is a completely tailless mouse.

Some dominant mutations cause a mutant phenotype in heterozygous condition with a wild-type allele because they enhance the function of the gene product. The enhanced function may arise because the mutation specifies a novel polypeptide or because it causes the wild-type polypeptide to be produced where or when it should not be. Dominant mutations that work in these ways are called **gain-of-function mutations**. In *Drosophila*, the mutation known as *Antennapedia (Antp)* is a dominant gain-of-function mutation. In heterozygous condition with a wild-type allele, *Antp* causes legs to develop in place of the antennae on the head of the fly. The reason for this bizarre anatomical transformation is that the *Antp* mutation causes the polypeptide product of the *Antennapedia* gene to be produced in the head, where, ordinarily,



Allelic Variation and Gene Function

69

Recessive amorphic loss-of-function allele does not produce a functional polypeptide.

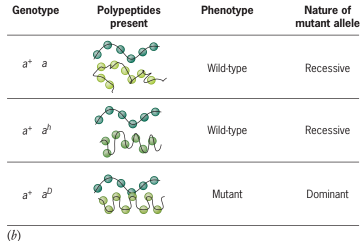


Recessive hypomorphic loss-of-function allele produces a partially functional polypeptide.



Dominant-negative allele produces a polypeptide that interferes with the wild-type polypeptide.





■ FIGURE 4.9 Differences between recessive loss-of-function mutations and dominant gainof-function mutations. (a) Polypeptide products of recessive and dominant mutations. (b) Phenotypes of heterozygotes carrying a wild-type allele and different types of mutant alleles. it is not produced; the *Antennapedia* gene product has therefore expanded the domain of its function.

We should note that not all genes produce polypeptides as the work of Beadle and Tatum implied. Modern research has identified many genes whose end products are RNA molecules rather than polypeptides. We will explore these kinds of genes later in this book.

KEY POINTS • Genes often have multiple alleles.

- Mutant alleles may be dominant, recessive, incompletely dominant, or codominant.
- If a hybrid that inherited a recessive mutation from each of its parents has a mutant phenotype, then the recessive mutations are alleles of the same gene; if the hybrid has a wild phenotype, then the recessive mutations are alleles of different genes.
- Most genes encode polypeptides.
- In homozygous condition, recessive mutations often abolish or diminish polypeptide activity.
- Some dominant mutations produce a polypeptide that interferes with the activity of the polypeptide encoded by the wild-type allele of a gene.

Gene Action: From Genotype to Phenotype

Phenotypes depend on both environmental and genetic factors.

At the beginning of the twentieth century, geneticists had imprecise ideas about how genes evoke particular phenotypes. They knew noth-

ing about the chemistry of gene structure or function, nor had they developed the techniques to study it. Everything that they proposed about the nature of gene action was inferred from the analysis of phenotypes. These analyses showed that genes do not act in isolation. Rather, they act in the context of an environment and in concert with other genes. These analyses also showed that a particular gene can influence many different traits.

INFLUENCE OF THE ENVIRONMENT

A gene must function in the context of both a biological and a physical environment. The factors in the physical environment are easier to study, for particular genotypes can be reared in the laboratory under controlled conditions, allowing an assessment of the effects of temperature, light, nutrition, and humidity. As an example, let's consider the *Drosophila* mutation known as *shibire*. At the normal culturing temperature, 25°C, *shibire* flies are viable and fertile, but are extremely sensitive to a sudden shock. When a *shibire* culture is shaken, the flies—temporarily paralyzed—fall to the bottom of the culture. Indeed, *shibire* is the Japanese word for "paralysis." However, if a culture of *shibire* flies is placed at a slightly higher temperature, 29°C, all the flies fall to the bottom and die, even without a shock. Thus, the phenotype of the *shibire* mutation is temperature-sensitive. At 25°C, the mutation is viable, but at 29°C, it is lethal. A plausible explanation is that at 25°C, the mutant gene makes a partially functional protein, but at 29°C, this protein is totally nonfunctional.

ENVIRONMENTAL EFFECTS ON THE EXPRESSION OF HUMAN GENES

Human genetic research provides an example of how the physical environment can influence a phenotype. **Phenylketonuria** (PKU) is a recessive disorder of amino acid metabolism. Infants homozygous for the mutant allele accumulate toxic substances in their brains that can impair mental ability by affecting the brain's development.

The harmful aspects of PKU are traceable to a particular amino acid, phenylalanine, which is ingested in the diet. Though not toxic itself, phenylalanine is metabolized into other substances that are. Infants with PKU who are fed normal diets ingest enough phenylalanine to bring out the worst manifestations of the disease. However, infants who are fed low-phenylalanine diets usually mature without serious mental impairment. Because PKU can be diagnosed in newborn babies, the clinical impact of this disease can be reduced if infants that are PKU homozygotes are placed on a low-phenylalanine diet shortly after birth. This example illustrates how an environmental factor—diet—can be manipulated to modify a phenotype that would otherwise become a personal tragedy.

The biological environment can also influence the phenotypic expression of genes. **Pattern baldness** in humans is a well-known example. Here the relevant biological factor is gender. Premature pattern baldness is due to an allele that is expressed differently in the two sexes. In males, both homozygotes and heterozygotes for this allele develop bald patches, whereas in females, only the homozygotes show a tendency to become bald, and this is usually limited to general thinning of the hair. The expression of this allele is probably triggered by the male hormone **testosterone**. Females produce much less of this hormone and are therefore seldom at risk to develop bald patches. The sex-influenced nature of pattern baldness shows that biological factors can control the expression of genes.

PENETRANCE AND EXPRESSIVITY

When individuals do not show a trait even though they have the appropriate genotype, the trait is said to exhibit *incomplete penetrance*. An example of incomplete penetrance in humans is **polydactyly**—the presence of extra fingers and toes (**Theorem Figure 4.10a**). This condition is due to a dominant mutation, *P*, that is manifested in some of its carriers. In the pedigree in **Figure 4.10b**, the individual denoted III-2 must be a carrier even though he does not have extra fingers or toes. The reason is that both his mother and three of his children are polydactylous—an indication of the transmission of the mutation through III-2. Incomplete penetrance can be a serious problem in pedigree analysis because it can lead to the incorrect assignment of genotypes.

The term *expressivity* is used if a trait is not manifested uniformly among the individuals that show it. The dominant *Lobe* eye mutation (**Figure 4.11**) in *Drosophila* is an example. The phenotype associated with this mutation is extremely variable. Some heterozygous flies have tiny compound eyes, whereas others have large, lobulated eyes; between these extremes, there is a full range of phenotypes. The *Lobe* mutation is therefore said to have *variable expressivity*.

Incomplete penetrance and variable expressivity indicate that the pathway between a genotype and its phenotypes is subject to considerable modulation. Geneticists know that some of this modulation is due to environmental factors, but

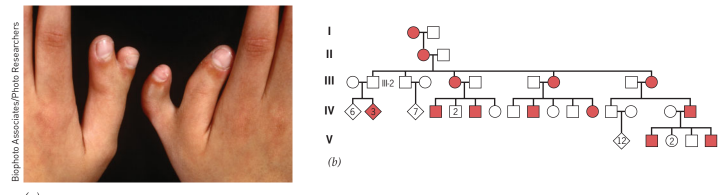
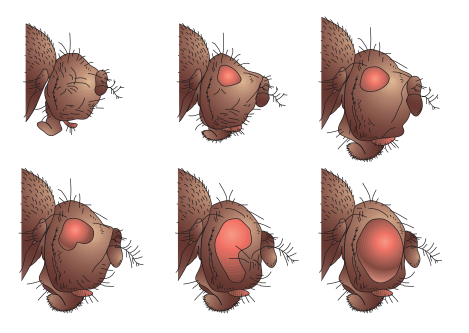




FIGURE 4.10 Polydactyly in humans. (a) Phenotype showing extra fingers. (b) Pedigree showing the inheritance of this incompletely penetrant dominant trait.



■ **FIGURE 4.11** Variable expressivity of the *Lobe* mutation in *Drosophila*. Each fly is heterozygous for this dominant mutation; however, the phenotypes vary from complete absence of the eye to a nearly wild-type eye.





(b)



(c)



■ FIGURE 4.12 Comb shapes in chickens of different breeds. (a) Rose, Wyandottes; (b) pea, Brahmas; (c) walnut, hybrid from cross between chickens with rose and pea combs; (d) single, Leghorns. some is also due to factors in the genetic background. Clearcut evidence for such factors comes from breeding experiments showing that two or more genes can affect a particular trait.

GENE INTERACTIONS

Some of the first evidence that a trait can be influenced by more than one gene was obtained by Bateson and Punnett from breeding experiments with chickens. Their work was carried out shortly after the rediscovery of Mendel's paper. Domestic breeds of chickens have different comb shapes (**Figure 4.12**): Wyandottes have "rose" combs, Brahmas have "pea" combs, and Leghorns have "single" combs. Crosses between Wyandottes and Brahmas produce chickens that have yet another type of comb, called "walnut." Bateson and Punnett discovered that comb type is determined by two independently assorting genes, *R* and *P*, each with two alleles (■ Figure 4.13). Wyandottes (with rose combs) have the genotype RR pp, and Brahmas (with pea combs) have the genotype rr PP. The F_1 hybrids between these two varieties are therefore Rr Pp, and phenotypically they have walnut combs. If these hybrids are intercrossed with each other, all four types of combs appear in the progeny: 9/16 walnut (*R*-*P*-), 3/16 rose (*R*-*pp*), 3/16 pea (rr P-), and 1/16 single (rr pp). The Leghorn breed, which has the single-comb type, must therefore be homozygous for both of the recessive alleles.

The work of Bateson and Punnett demonstrated that two independently assorting genes can affect a trait. Different combinations of alleles from the two genes resulted in different phenotypes, presumably because of interactions between their products at the biochemical or cellular level.

EPISTASIS

When two or more genes influence a trait, an allele of one of them may have an overriding effect on the phenotype. When an allele has such an overriding effect, it is said to be *epistatic* to the other genes that are involved; the term **epistasis** comes from Greek words meaning to "stand above." For example, we know that eye pigmentation in *Drosophila* involves a large number of genes. If a fly is homozygous for a null allele in any one of these genes, the pigment-synthesizing pathway can be blocked, and an abnormal eye color will be produced. This allele essentially nullifies the work of all the other genes, masking their contributions to the phenotype.

A mutant allele of one gene is epistatic to a mutant allele of another gene if it conceals the latter's presence in the genotype. We have already seen that a recessive mutation in the *cinnabar* gene of *Drosophila* causes the eyes of the fly to be bright red. A recessive mutation in a different gene causes the eyes to be white. When both of these mutations are made homozygous in the same fly, the eye color is white. Thus, the *white* mutation is epistatic to the *cinnabar* mutation.

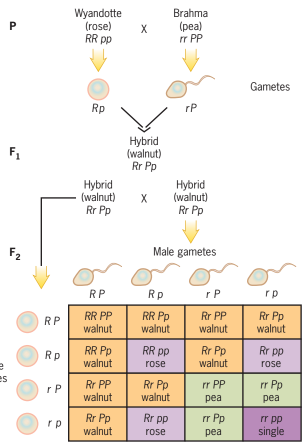
What physiological mechanism makes the *white* mutation epistatic to the *cinnabar* mutation? The polypeptide product of the wild-type allele of the *white* gene transports pigment into the *Drosophila* eye. When this gene is mutated, the transporter polypeptide is not made. Flies that are homozygous for the *cinnabar* mutation cannot synthesize brown pigment, but they can synthesize red pigment. When these flies are also homozygous for the *white* mutation, the red pigment cannot be transported into the eyes. Consequently, flies that are homozygous for both the *cinnabar* and *white* mutations have white eyes.

The analysis of epistatic relationships such as the one between *cinnabar* and *white* can suggest ways in which genes control a phenotype. A classic example of this analysis is again from the work of Bateson and Punnett, who studied the genetic control of flower color in the sweet pea, *Lathyrus odoratus* (**■** Figure 4.14a). The flowers in this plant are either purple or white—purple if they contain anthocyanin pigment and white if they do not. Bateson and Punnett crossed two different varieties with white flowers to obtain F_1 hybrids, which all had purple flowers. When these hybrids were intercrossed, Bateson and Punnett obtained a ratio of 9 purple: 7 white plants in the F_2 . They explained the results by proposing that two independently assorting genes, *C* and *P*, are involved in anthocyanin synthesis and that each gene has a recessive allele that abolishes pigment production (**■** Figure 4.14b).

Given this hypothesis, the parental varieties must have had complementary genotypes: *cc PP* and *CC pp*. When the two varieties were crossed, they produced *Cc Pp* double heterozygotes that had purple flowers. In this system, a dominant allele from each gene is necessary for the synthesis of anthocyanin pigment. In the F_2 , 9/16 of the plants are *C- P-* and have purple flowers; the remaining 7/16 are homozygous for at least one of the recessive alleles and have white flowers. Notice that the double recessive homozygotes, *cc pp*, are not phenotypically different from either of the single recessive homozygotes. Bateson and Punnett's work established that each of the recessive alleles is epistatic over the dominant allele of the other gene. A plausible explanation is that each dominant allele produces an enzyme that controls a step in the synthesis of anthocyanin from a biochemical precursor. If a dominant allele is not present, its step in the biosynthetic pathway is blocked and anthocyanin is not produced:

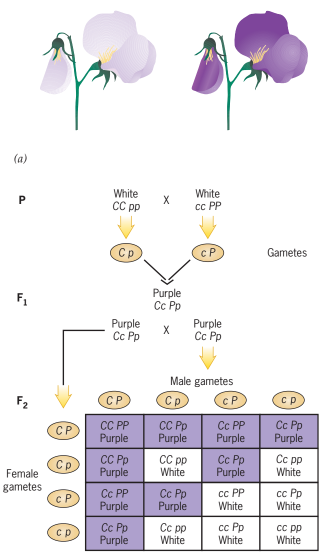
Gene	Procursor	C → Intermediate →	P
Genotype	Flecuisor —		
C-P-	+	+	+
cc P–	+	_	_
С–рр	+	+	_
сс рр	+	-	_

Notice that Bateson's and Punnett's first cross was a test for allelism between two white-flowered strains of the sweet pea. Each strain was homozygous for a recessive mutation in a gene involved in the production of purple pigment. When the two white



Summary: 9/16 walnut, 3/16 rose, 3/16 pea, 1/16 single

■ FIGURE 4.13 Bateson and Punnett's experiment on comb shape in chickens. The intercross in the F₁ produces four phenotypes, each highlighted by a different color in the Punnett square, in a 9:3:3:1 ratio.



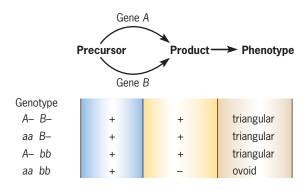
Summary: 9/16 purple, 7/16 white

(b)

FIGURE 4.14 Inheritance of flower color in sweet peas. (a) White and purple flowers of the sweet pea. (b) Bateson and Punnett's experiment on the genetic control of flower color in sweet peas.

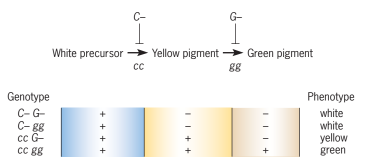
strains were crossed, the F_1 plants had purple flowers. This result tells us that the white strains were homozygous for mutations in different genes involved in the synthesis of purple pigment.

Another classic study of epistasis was performed by George Shull using a weedy plant called the shepherd's purse, Bursa bursa-pastoris (**Figure 4.15a**). The seed capsules of this plant are either triangular or ovoid in shape. Ovoid capsules are produced only if a plant is homozygous for the recessive alleles of two genes-that is, if it has the genotype *aa bb.* If the dominant allele of either gene is present, the plant produces triangular capsules. The evidence for this conclusion comes from crosses between doubly heterozygous plants (**Figure 4.15***b*). Such crosses produce progeny in a ratio of 15 triangular:1 ovoid, indicating that the dominant allele of one gene is epistatic over the recessive allele of the other. The data suggest that capsule shape is determined by duplicate developmental pathways, either of which can produce a triangular capsule. One pathway involves the dominant allele of the A gene, and the other the dominant allele of the *B* gene. A precursor substance can be converted into a product that leads to a triangular seed capsule through either of these pathways. Only when both pathways are blocked by homozygous recessive alleles is the triangular phenotype suppressed and an ovoid capsule produced:



In other cases of epistasis, the product of one gene may inhibit the expression of another gene. Consider, for example, the inheritance of fruit color in summer squash plants. Plants that carry the dominant allele C produce white fruit, whereas plants that are homozygous for the recessive allele c produce colored fruit. If a squash plant is also homozygous for the recessive allele g of an independently assorting gene, the fruit will be green. However, if it carries the dominant allele G of this gene, the fruit will be yellow. These observations suggest that the two genes control steps in the synthesis of green pigment. The first step converts a colorless pre-

cursor into a yellow pigment, and the second step converts this yellow pigment into a green pigment. If the first step is blocked (by the presence of the C allele), neither of the pigments is produced and the fruit will be white. If only the second step is blocked (by the presence of the G allele), the yellow pigment cannot be converted into the green pigment and the fruit will be yellow. We can summarize these ideas with a diagram that shows the genetic control of pigment synthesis in this biochemical pathway:





The arrows in the diagram show the steps in the pathway. The genotype below an arrow allows that step to occur, whereas the genotype above an arrow inhibits that step from occurring. It is customary in genetics to symbolize the inhibitory effect of a genotype by drawing a blunted arrow (\neg) from the genotype to the relevant step in the pathway. In this example, the *C* allele inhibits the first step and the *G* allele inhibits the second step. Because of its role as an inhibitor of the first step, the *C* allele is epistatic to both of the alleles of the other gene. No matter which of the alleles of this other gene is present in a plant, the *C* allele will cause that plant to produce white fruit.

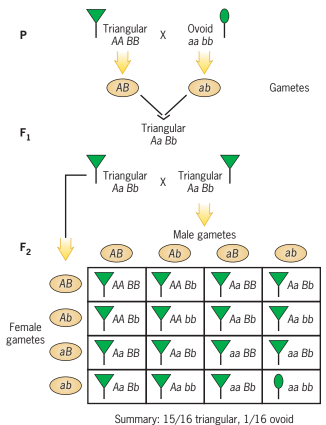
Figure 4.16 shows the outcome of a cross between plants heterozygous for both of the fruit-color-determining genes. When *Cc Gg* plants are intercrossed, they produce progeny that sort into three phenotypic classes: white, yellow, and green. The offspring with green fruit are homozygous for the recessive alleles of both genes; that is, they are α gg, and their frequency is 1/16. The offspring with yellow fruit are homozygous for *c*, and they carry at least one copy of *G*; their frequency is 3/16. The offspring with white fruit carry at least one copy of *C*; the rest of the genotype does not matter. The frequency of the white-fruited plants is 12/16. To test your ability to make genetic predictions from a biochemical pathway, work through the exercise in Problem-Solving Skills: Going from Pathways to Phenotypic Ratios.

These examples indicate that a particular phenotype is often the result of a process controlled by more than one gene. Each gene governs a step in a pathway that is part of the process. When a gene is mutated to a nonfunctional or partially functional state, the process can be disrupted, leading to a mutant phenotype. Much of modern genetic analysis is devoted to the investigation of pathways involved in important biological processes such as metabolism and development. Studying the epistatic relationships among genes can help to sort out the role that each gene plays in these processes.

PLEIOTROPY

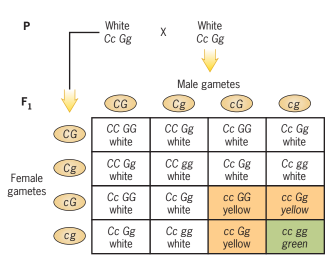
Not only is it true that a phenotype can be influenced by many genes, but it is also true that a gene can influence many phenotypes. When a gene affects many aspects of the phenotype, it is said to be **pleiotropic**, from the Greek words for "to take many turns." The gene for phenylketonuria in humans is an example. The primary effect of recessive mutations in this gene is to cause toxic substances to accumulate in the brain, leading to mental impairment. However, these mutations also interfere with the syn-

thesis of melanin pigment, lightening the color of the hair; therefore, individuals with PKU frequently have light brown or blond hair. Biochemical tests also reveal that the



■ **FIGURE 4.15** Inheritance of seed capsule shape in the shepherd's purse. (a) The shepherd's purse, *Bursa bursa-pastoris*. (b) Crosses showing duplicate gene control of seed capsule shape in the shepherd's purse.

(b)



Summary: 12/16 white, 3/16 yellow, 1/16 green

FIGURE 4.16 Segregation in the offspring of a cross between summer squash plants hetero-zygous for two genes controlling fruit color.

PROBLEM-SOLVING SKILLS

Going from Pathways to Phenotypic Ratios

THE PROBLEM

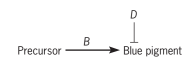
Flower color in a plant is determined by two independently assorting genes, *B* and *D*. The dominant allele *B* allows a pigment precursor to be converted into blue pigment. In homozygous condition, the recessive allele of this gene, *b*, blocks this conversion, and without blue pigment, the flowers are white. The dominant allele of the other gene, *D*, causes the blue pigment to degrade, whereas the recessive allele of this gene, *d*, has no effect. True-breeding blue and white strains of the plant were crossed, and all the F_1 plants had white flowers. (a) What was the genotype of the F_1 plants? (b) What were the genotypes of the plants used in the initial cross? (c) If the F_1 plants are self-fertilized, what phenotypes will appear in the F_2 , and in what proportions?

FACTS AND CONCEPTS

- The dominant allele (D) of one gene is epistatic to both alleles (B and b) of the other gene.
- **2.** Plants with blue flowers must carry at least one *B* allele, but they cannot carry even one *D* allele.
- **3.** Plants with white flowers can be *bb*, or they can be *BB* or *Bb* as long as they also carry at least one *D* allele.
- 4. True-breeding strains are homozygous for their genes.
- **5.** When genes assort independently, we multiply the probabilities associated with the components of the complete genotype.

ANALYSIS AND SOLUTION

A good place to start the analysis is to diagram the biochemical pathway—that is, to transform the "word problem" into a diagram that will guide our search for a solution.



The positive action of the B allele is required for the synthesis of blue pigment. The negative action of the dominant allele D is indicated by a blunted arrow pointed at this pigment. Now we can address the questions in the problem.

- a. The key observation is that the flowers of the F₁ plants are white. Because these plants had a true-breeding blue parent, they must carry the *B* allele, but the blue pigment produced through the action of this allele must be degraded. The F₁ plants must therefore also carry the *D* allele. However, they cannot be homozygous for it because their blue parent could not have carried it. Thus, the F₁ plants must be heterozygous for the *D* allele. Genotypically, they are either *BB Dd* or *Bb Dd*. From the information given in the problem, we cannot distinguish between these two possibilities.
- **b.** The blue plants used in the cross must have been *BB dd*. The white plants could have been either *BB DD* or *bb DD*—we cannot be certain which of these genotypes they were.
- **c.** If the F₁ plants are *BB Dd*, then when they are selfed only the *D* and *d* alleles will segregate, and 1/4 of their offspring will be blue (*BB dd*) and 3/4 will be white (*BB DD* or *BB Dd*). If the F₁ plants are *Bb Dd*, then when they are selfed, both genes will segregate dominant and recessive alleles. Among the offspring, those that are *BB dd* or *Bb dd* will be blue. This phenotypic class will constitute $(3/4) \times (1/4) = 3/16$ of the total. All the other offspring, 1 3/16 = 13/16 of the total, will be white.

For further discussion visit the Student Companion site.

blood and urine of PKU patients contain compounds that are rare or absent in normal individuals. This array of phenotypic effects is typical of most genes and results from interconnections between the biochemical and cellular pathways that the genes control.

Another example of pleiotropy comes from the study of mutations affecting the formation of bristles in *Drosophila*. Wild-type flies have long, smoothly curved bristles on the head and thorax. Flies homozygous for the *singed* bristle mutation have short, twisted bristles on these body parts—as if they had been scorched. Thus, the wild-type *singed* gene product is needed for the proper formation of bristles. It is also needed for the production of healthy, fertile eggs. We know this fact because females that are homozygous for certain *singed* mutations are completely sterile; they lay flimsy, ill-formed eggs that never hatch. However, these mutations have no adverse effect on male fertility. Thus, the *singed* gene pleiotropically controls the formation of both bristles and eggs in females and the formation of bristles in males.

KEY POINTS

- Gene action is affected by biological and physical factors in the environment.
 - Two or more genes may influence a trait.
 - A mutant allele is epistatic to a mutant allele of another gene if it has an overriding effect on the phenotype.
 - A gene is pleiotropic if it influences many different phenotypes.

Inbreeding: Another Look at Pedigrees

Geneticists have always been interested in the phenomenon of inbreeding, whether to make truebreeding strains or to reveal the homozygous effects of recessive alleles. In addition, when inbreeding Geneticists use a simple statistic, the inbreeding coefficient, to analyze the effects of matings between relatives.

occurs in nature, it can affect the character of plant and animal populations. In this section we consider ways to analyze the effects of inbreeding. We also introduce the techniques needed to study common ancestry in pedigrees.

THE EFFECTS OF INBREEDING

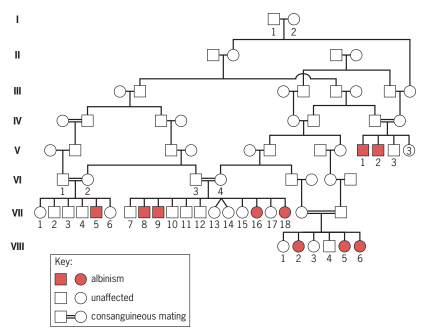
Inbreeding occurs when mates are related to each other by virtue of common ancestry. A mating between relatives is often referred to as a **consanguineous mating**, from Latin words meaning "of the same blood." In human populations, these types of matings are rare, with the incidence depending on cultural and ethnic traditions and on geography. In many cultures, marriages between close relatives—for example, between siblings or half siblings—are expressly forbidden, and marriages between more distant relatives, though allowed, must be approved by civil or religious authorities before they can occur. These restrictions exist because inbreeding tends to produce more diseased and debilitated children than matings between unrelated individuals. This tendency, as we now know, arises from an increased chance for the children of a consanguineous mating to be homozygous for a harmful recessive allele. In some cultures, however, consanguineous matings have been accepted and even encouraged. In ancient Egypt, for example, the royal line was perpetuated by brother–sister marriages, presumably to preserve the "purity" of the royal blood. Similar practices existed in Polynesia until relatively recent times.

The occurrence of consanguineous matings in human populations has helped in the analysis of genetic conditions caused by recessive alleles. In fact, the very first

gene to be identified in humans was brought to light by observing a greater frequency of recessive homozygotes in the children of first cousins; for more information, see A Milestone in Genetics: Garrod's Inborn Errors of Metabolism in the Student Companion site. Many of the classic studies in human genetics were based on the analysis of consanguineous matings in socially closed groups-for example, the Amish, a religious sect scattered in small communities in the eastern and midwestern United States. **Figure 4.17** shows an Amish pedigree in which 10 individuals have albinism. The affected individuals are all descendants of two people (I-1 and I-2) who had immigrated from Europe. The consanguineous matings in the pedigree are indicated by double lines connecting the mates. All the affected individuals come from such matings. Thus, this pedigree shows how inbreeding brings out a recessive condition, which geneticists can then analyze.

The effects of inbreeding are also evident in experimental species where it is possible to arrange matings between relatives. For example, animals such as rats, mice, and guinea pigs can be mated brother to sister, generation after generation, to create an **inbred line**. Although

these lines are genetically quite pure—that is, they do not segregate different alleles of particular genes—they often are less vigorous than lines maintained by matings between unrelated individuals. We refer to this loss of vigor as **inbreeding depression**. In plants where self-fertilization is possible, very highly inbred lines can be created by repeated self-fertilization over several generations. Each line would be expected to



■ FIGURE 4.17 Albinism in the offspring of consanguineous marriages in an Amish community from the Midwestern United States. Consanguineous marriages are indicated by double lines between the mates. The individuals with albinism, who are homozygous for a recessive allele, all come from consanguineous matings.

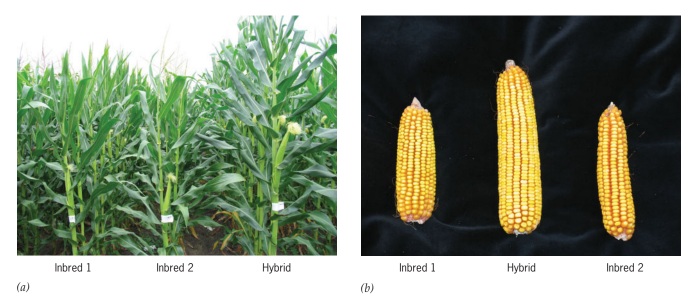
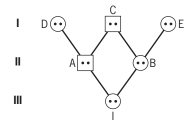


FIGURE 4.18 (a) Inbred varieties of maize and the hybrid produced by crossing them. The inbred plants are shorter and less robust than the hybrid plant. (b) Cobs from inbred plants are considerably smaller than cobs from hybrid plants.

be homozygous for different alleles that were present in the founding population of plants. **Figure 4.18** shows the result of this process in maize. The inbred plants are short and produce small ears with few kernels. By comparison, the plants generated by crossing the two inbred strains are tall and produce large ears with many kernels. These plants are expected to be heterozygous for many genes. Their robustness is a phenomenon called hybrid vigor, or **heterosis**. This term was introduced in 1914 by George Shull, a pioneering plant breeder who began the practice of crossing inbred strains to produce uniformly high-yielding, heterozygous offspring. Shull's technique has since become standard in the plant breeding industry.

GENETIC ANALYSIS OF INBREEDING

Matings between full siblings, between half siblings, and between first cousins are all examples of inbreeding. When such matings occur, we speak of the offspring as being *inbred*. Inbred individuals differ from the offspring of unrelated parents in one important way: the two copies of a gene they carry may be identical to each other by virtue of common ancestry—that is, because the genes have descended from a gene that was present in an ancestor of the inbred individual. To understand this concept, let's consider a simple pedigree that illustrates a mating between half siblings.



The two dots in each individual represent the two copies of a particular gene, and the lines that connect individuals show how genes have passed from parent to offspring. This way of drawing a pedigree is different from the one we have used previously. It clarifies how each parent contributes genes to its offspring, and it allows us to trace the descent of a particular gene through multiple generations.

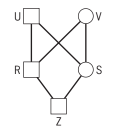
The two individuals in Generation II, labeled A and B, are half siblings. These individuals had a common father, C, but different mothers (D and E). The mating between A and B produced an offspring, I, who is inbred. Notice that I inherits one gene copy from A and one copy from B. However, both of these copies might have

originated in C, the common father of A and B. Thus, the two gene copies in I might be identical to each other by descent from one of the gene copies that was present in C. This possibility of *identity by descent* is the important consequence of inbreeding. Any individual whose gene copies are identical by descent must be homozygous for a particular allele of that gene. Thus, consanguineous matings are expected to produce relatively more homozygotes than matings between unrelated individuals, which, as we have seen, is one of the conspicuous effects of inbreeding.

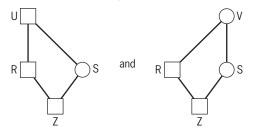
In the pedigree we are considering, C is referred to as the *common ancestor* of I because two paths of descent from C converge in I, the inbred individual. The two paths are $C \rightarrow A \rightarrow I$ and $C \rightarrow B \rightarrow I$, and together they form what geneticists call an *inbreeding loop*. This loop shows how a particular gene copy in C can be passed down both sides of the pedigree to produce two identical gene copies in I.

The fundamental determination in any analysis of inbreeding is to calculate the probability that two gene copies in an individual are identical by descent. Intuitively, this probability should increase with the intensity of inbreeding. Thus, the offspring of a mating between full siblings should have a greater probability of identity by descent than the offspring of a mating between half siblings. The effort to measure inbreeding intensity began with the pioneering work of the American geneticist Sewall Wright. In 1921 Wright discovered a mathematical quantity he called the **inbreeding coefficient**. Wright's investigations—too complicated to be discussed here—involved an analysis of correlations between the individuals in a pedigree. In these investigations, he discovered how to calculate the inbreeding coefficient and used it to measure the intensity of inbreeding. Then, in the 1940s, another American, Charles Cotterman, showed that Wright's inbreeding coefficient was equivalent to the probability of identity by descent. Thus, we can define the inbreeding coefficient, symbolized by the letter F, as the probability that two gene copies in an individual are identical by descent from a common ancestor.

To calculate the inbreeding coefficient, we follow the procedures developed by Wright and Cotterman. First, we identify the common ancestor(s) of the inbred individual. A common ancestor is connected to the inbred individual through both of that individual's parents. In the pedigree we are considering, I has only one common ancestor; however, in other types of pedigrees, an inbred individual might have more than one common ancestor. For example, the offspring of a mating between full siblings has two common ancestors:



In this case, both of Z's grandparents (U and V) are common ancestors. Two genetic paths descend from each grandparent and converge in Z. Thus, the pedigree for full-sib mating has two distinct inbreeding loops:

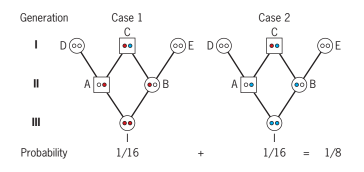


The second step in calculating the inbreeding coefficient is to count the number of individuals (n) in each inbreeding loop defined by a common ancestor. In the pedigree for mating between half siblings, there is one inbreeding loop and it has three individuals. (We do not count the inbred individual itself.) Thus, for the pedigree with

half-sib mating, n = 3. In the pedigree for full-sib mating there are two inbreeding loops, each with three individuals; thus, for each of these loops, n = 3.

The third step in the procedure to calculate the inbreeding coefficient is to compute the quantity $(1/2)^n$ for each inbreeding loop and then sum the results. The sum we obtain is the inbreeding coefficient, *F*, of the inbred individual—that is, the probability that its two gene copies are identical to each other by descent from a common ancestor. For the offspring of a mating between half siblings, we obtain $F = (1/2)^3 = 1/8$. For the offspring of a mating between full siblings, we obtain $F = (1/2)^3 + (1/2)^3 = 1/4$. Thus, the inbreeding coefficient of the offspring of full-sib mating is greater than the inbreeding coefficient of the offspring of half-sib mating, as expected.

The factor $(1/2)^n$ that we compute for each inbreeding loop is the probability that *either* of the two gene copies in the common ancestor of that loop produces two identical gene copies in the inbred individual. To understand this probability, let's focus on the mating between half siblings. We must consider two cases, labeled 1 and 2 in the following illustration.



In Case 1, the chance that the gene copy on the left (shown in red) in the common ancestor C is transmitted to the daughter A is 1/2; once in A, the chance that this gene copy is transmitted to I is 1/2. Thus, the probability that the "left" gene copy in C makes its way down to I through A is $(1/2) \times (1/2) = 1/4$. Similarly, the chance that the "left" gene copy makes its way down to I through B is $(1/2) \times (1/2) = 1/4$. Altogether, then, the probability that the "left" gene copy in C produces two identical gene copies in I, one transmitted through A and the other through B, is $(1/4) \times (1/4) = 1/16$. By similar reasoning in Case 2, we find the probability that the "right" gene copy (shown in blue) in C produces two identical gene copies in I to be 1/16. Thus, the probability that *either* the "left" or the "right" gene copies in C will produce two identical gene copies in I is (1/16) + (1/16) = 1/8, which, as we have seen, is $(1/2)^3$. The procedure of calculating the factor $(1/2)^n$ is therefore a shortcut to find the probability that either of the gene copies in a particular common ancestor will give rise to two identical gene copies in the inbred individual.

This method of calculating inbreeding coefficients works for most pedigrees. However, when a common ancestor is itself inbred, the method needs to be modified. We multiply the factor $(1/2)^n$ for the common ancestor by the term $[1 + F_{CA}]$, where F_{CA} is the inbreeding coefficient of the common ancestor. For example, in this pedigree

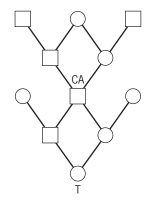
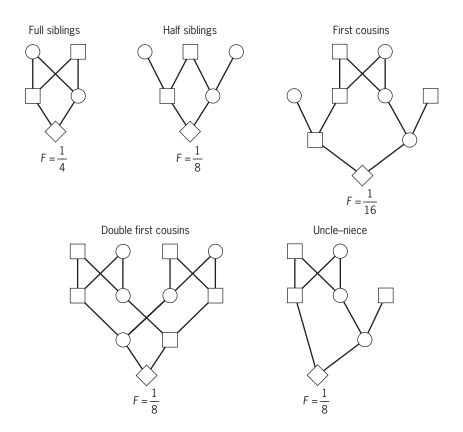


FIGURE 4.19 Values of the inbreeding coefficient, *F*, for different pedigrees.



the inbreeding coefficient of T is $F_T = (1/2)^3 \times [1 + F_{CA}]$, and because $F_{CA} = (1/2)^3 = 1/8$, we conclude that $F_T = (1/8) \times [1 + (1/8)] = 9/64$. The modifying term $[1 + F_{CA}]$ accounts for the possibility that the "left" and "right" gene copies in CA are already identical by descent. To test your ability to apply this theory, try Solve It: Compound Inbreeding.

Wright and Cotterman defined the inbreeding coefficient as an accurate measure of inbreeding intensity. **Figure 4.19** presents values of this coefficient for the offspring of different types of consanguineous matings.

One use of the inbreeding coefficient is to explain the increased frequency of recessive disorders among the offspring of consanguineous matings. In the human population, for example, the incidence of phenylketonuria (PKU) among the offspring of unrelated parents is about 1/10,000; among the offspring of first-cousin marriages, it is about 7/10,000. The difference between these frequencies, 6/10,000, is the effect of inbreeding with F = 1/16. For the offspring of closer relatives, we would expect a greater difference in the frequencies of PKU. For example, the offspring of half siblings have an inbreeding coefficient of 1/8, twice that of the offspring of first cousins. Because the effect of inbreeding is proportional to F, we would expect the incidence of PKU among the offspring of half siblings to be twice the inbreeding effect seen with the offspring of first cousins, plus the incidence of PKU in the general population. Thus, the predicted frequency of PKU among the offspring of half siblings is $2 \times (0.0006) + 0.0001 = 0.0013$. Among the offspring of full siblings, the predicted frequency is $4 \times (0.0006) + 0.0001 = 0.0025$ (because they have an inbreeding coefficient four times that of the offspring of first cousins).

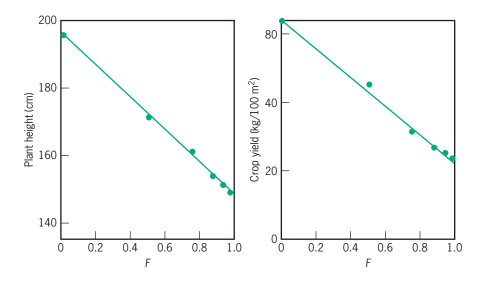
Another use of the inbreeding coefficient is to measure the decline in a complex phenotype, such as plant height or crop yield. Such traits are influenced by large numbers of genes. **Figure 4.20** shows data collected from inbred strains of maize that were obtained through a program of repeated self-fertilization. Seed was saved at each stage of the inbreeding process, and at the end, maize plants were grown from the seed in test plots to study two traits, plant height and crop yield. As Figure 4.20 shows, both of these traits declined linearly as a function of the inbreeding coefficient. The simplest explanation for this linear decline is that recessive alleles of different genes were made homozygous as the inbreeding proceeded—that is, in proportion

Solve It!

Compound Inbreeding

Two unrelated individuals mate to produce two offspring, A and B. These offspring then mate to produce an offspring, C, which mates to two different individuals to produce one offspring from each mating. These offspring then mate with each other to produce an individual in which the inbreeding effect has been compounded. What is the inbreeding coefficient of this last individual?

To see the solution to this problem, visit the Student Companion site. ■ **FIGURE 4.20** Inbreeding decline in plant height and crop yield in maize. The intensity of inbreeding is measured by the inbreeding coefficient, *F*.



to the value of F—and that these homozygotes manifested lower values for the traits. Thus, an increase in the incidence of deleterious recessive homozygotes is the basis for inbreeding depression.

MEASURING GENETIC RELATIONSHIPS

The inbreeding coefficient can also be used to measure the closeness of genetic relationships. Full siblings are obviously more closely related than half siblings. Are uncle and niece more closely related than half siblings? Are half siblings more closely related than first cousins? Are half siblings more closely related than double first cousins? To answer these questions, we must determine the fraction of genes that two relatives share by virtue of common ancestry.

For regular relatives—that is, relatives that are not themselves inbred—we can calculate the fraction of genes that are shared by imagining that the relatives have mated and produced an offspring. Obviously, because this offspring is inbred, we can calculate its inbreeding coefficient according to the usual procedure. Then, to determine the fraction of genes that the two relatives share, we simply multiply the offspring's inbreeding coefficient by 2. The result is sometimes called the **coefficient of relationship**. For full siblings, the inbreeding coefficient of an imaginary offspring is 1/4; thus, the coefficient of relationship of full siblings (or the fraction of genes they share) is $2 \times (1/4) = 1/2$. By similar reasoning, the coefficient of relationship of half siblings is 1/4, that of first cousins is 1/8, and that of double first cousins is 1/4. For uncle and niece, the coefficient of relationship is 1/4. Thus, half siblings, double first cousins, and uncle and niece are equivalently related because each shares the same fraction of their genes, 1/4. Siblings, by comparison, are more closely related because they share only one-eighth of their genes.

KEY POINTS

- Inbreeding increases the frequency of homozygotes and decreases the frequency of heterozygotes.
- The effects of inbreeding are proportional to the inbreeding coefficient, which is the probability that two gene copies in an individual are identical by descent from a common ancestor.
- The coefficient of relationship is the fraction of genes that two individuals share by virtue of common ancestry.

Basic Exercises

Illustrate Basic Genetic Analysis

- A researcher has discovered a new blood-typing system for humans. The system involves two antigens, P and Q, each determined by a different allele of a gene named N. The alleles for these antigens are about equally frequent in the general population. If the N^p and N^Q alleles are codominant, what antigens should be detected in the blood of N^p N^Q heterozygotes?
- **Answer:** Both the P and the Q antigens should be detected because codominance implies that both of the alleles in heterozygotes will be expressed.
- **2.** Flower color in a garden plant is under the control of a gene with multiple alleles. The phenotypes of the homo-zygotes and heterozygotes of this gene are as follows:

Homozygotes

WW	red
ww	pure white

- $w^s w^s$ white stippled with red
- $w^p w^p$ white with regular red patches

Heterozygotes

W	with any other allele	red
w^p	with either w^s or w	white with regular red patches
w ^s w		white stippled with red

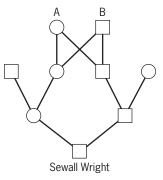
Arrange the alleles in a dominance hierarchy.

- **Answer:** W is dominant to all the other alleles, w^p is dominant to w^s and w, and w^s is dominant to w. Thus, the dominance hierarchy is $W > w^p > w^s > w$.
- **3.** Two independently discovered strains of mice are homozygous for a recessive mutation that causes the eyes to be small; the phenotypes of the two strains are indistinguishable. The mutation in one strain is called *little eye*, and the mutation in the other is called *tiny eye*. A third strain is heterozygous for a dominant mutation that eliminates the eyes altogether; the mutation in this strain is called *Eyeless*. How would you determine if the *little eye*, *tiny eye*, and *Eyeless* mutations are alleles of the same gene?
- **Answer:** The procedure to determine if two recessive mutations are alleles of the same gene is to cross their respective homozygotes to obtain hybrid progeny and then evaluate the phenotype of the hybrids. If the phenotype is mutant, the mutations are alleles of the same gene; if it is wild-type, they are not alleles. In this case, we should therefore cross *little eye* mice with *tiny eye* mice and look at their off-spring. If the offspring have small eyes, the two mutations

are alleles of the same gene; if they have eyes of normal size, the two mutations are alleles of different genes. For a dominant mutation such as *Eyeless*, no test of allelism is possible. Thus, we cannot determine if *Eyeless* is an allele of either the *little eye* or the *tiny eye* mutation.

- **4.** Distinguish between incomplete penetrance and variable expressivity.
- **Answer:** Incomplete penetrance occurs when an individual with the genotype for a trait does not express that trait at all. Variable expressivity occurs when a trait is manifested to different degrees in a set of individuals with the genotype for that trait.
- 5. In a species of fly, the wild-type eye color is red. In a mutant strain homozygous for the w mutation, the eye color is pure white; in another mutant strain homozygous for the y mutation, the eye color is yellow. Homozygous white mutants were crossed to homozygous yellow mutants, and the offspring all had red eyes. When these offspring were intercrossed, they produced three classes of progeny: 92 red, 33 yellow, and 41 pure white. (a) From the results of these crosses, how many genes control eye color? Explain. (b) If the answer to (a) is greater than one, is any one mutant gene epistatic to any other mutant gene?
- **Answer:** To answer (a), we note that the F_1 flies all had red—that is, wild-type—eyes. The w and y mutations are therefore not alleles of the same gene, and we conclude that at least two genes must control eye color in this species. To answer (b), we note that in the F_2 flies, the phenotypic segregation ratio departs from the 9:3:3:1 ratio expected for two genes assorting independently. The F_2 consists of only three classes, which, moreover, appear in the ratio of 9 red:4 white:3 yellow. Evidently, the ww homozygotes cause the flies to have white eyes regardless of what alleles of the y gene are present. Thus, the w mutant should be considered epistatic to the y mutant.
- 6. Sewall Wright, the discoverer of the inbreeding coefficient, was the offspring of a marriage between first cousins. Draw the pedigree of Dr. Wright's family and identify his common ancestors and the inbreeding loops they define. Then calculate Dr. Wright's inbreeding coefficient.

Answer: A pedigree for a first-cousin marriage is:



In this pedigree there are two common ancestors, A and B, each defining an inbreeding loop that terminates in the inbred individual. One loop is on the left side of the pedigree, the other on the right. Not counting the inbred individual, each of the loops contains five people. Thus, assuming that the common ancestors are not affected by prior inbreeding, the inbreeding coefficient of the offspring of the first-cousin marriage (Dr. Wright) is $(1/2)^5 + (1/2)^5 = 1/16$.

Testing Your Knowledge

Integrate Different Concepts and Techniques

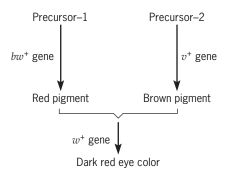
- 1. A geneticist has obtained two true-breeding strains of mice, each homozygous for an independently discovered recessive mutation that prevents the formation of hair on the body. One mutant strain is called *naked*, and the other is called *hairless*. To determine whether the two mutations are alleles, the geneticist crosses *naked* and *hairless* mice with each other. All the offspring are phenotypically wild-type; that is, they have hairs all over their bodies. After intercrossing these F₁ mice, the geneticist observes 115 wild-type mice and 85 mutant mice in the F₂. Are the *naked* and *hairless* mutations alleles? How would you explain the segregation of wild-type and mutant mice in the F₂.
- **Answer:** The *naked* and *hairless* mutations are not alleles because the F_1 hybrids are phenotypically wild-type. Thus, *naked* and *hairless* are mutations of two different genes. To explain the phenotypic ratio in the F_2 , let's first adopt symbols for these mutations and their dominant wild-type alleles:
 - n = naked mutation, N = wild-type allele
 - b = hairless mutation, H = wild-type allele

With these symbols, the genotypes of the true-breeding parental strains are nn HH (naked) and NN hh (hair*less*). The F_1 hybrids produced by crossing these strains are therefore Nn Hh. When these hybrids are intercrossed, we expect many different genotypes to appear in the offspring. However, each recessive allele, when homozygous, prevents the formation of hair on the body. Thus, only mice that are genotypically N- H- will develop hair; all the othershomozygous *nn* or homozygous *bb*, or homozygous for both recessive alleles-will fail to develop body hair. We can predict the frequencies of the wild and mutant phenotypes if we assume that the naked and hairless genes assort independently. The frequency of mice that will be N- H- is $(3/4) \times$ (3/4) = 9/16 = 0.56 (by the Multiplicative Rule of Probability), and the frequency of mice that will be either *nn* or *bb* (or both) is $(1/4) + (1/4) - [(1/4) \times (1/4)] = 7/16 = 0.44$ (by the Additive Rule of Probability). Thus, in a sample of 200 F, progeny, we expect $200 \times 0.56 = 112$ to be wild-type and $200 \times 0.44 = 88$ to be mutant. The observed frequencies of 115 wild-type and 85 mutant mice are close to these expected numbers, suggesting that the hypothesis of two independently assorting genes for body hair is indeed correct.

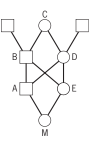
2. In fruit flies a recessive mutation, *w*, causes the eyes to be white, another recessive mutation, *v*, causes them to be vermilion, and a third recessive mutation, *bw*, causes them to be brown. The wild-type eye color is dark red. Hybrids

produced by crossing any two homozygous mutants have dark red eyes, and all the doubly homozygous mutant combinations have white eyes. How many genes do these three mutations define? If the dark red color of wild-type eyes is due to the accumulation of two different pigments, one red and the other brown, which gene controls the expression of which pigment? Can the genes be ordered into a pathway for pigment accumulation?

Answer: The three mutations define three different genes because when any two homozygous mutations are crossed, the offspring have wild-type eye color. The w mutation prevents the expression of all pigment because flies homozygous for it have neither red nor brown pigment in their eyes; the v mutation prevents the expression of brown pigment because flies homozygous for it have vermilion (bright red) eyes; and the *bw* mutation prevents the expression of red pigment because flies that are homozygous for it have brown eyes. Thus, the wild-type v gene controls the expression of brown pigment, the wild-type bw gene controls the expression of red pigment, and the wild-type w gene is necessary for the expression of both pigments. We can summarize these findings by proposing that each pigment is expressed in a different pathway and that the functioning of these pathways depends on the wild-type w gene.



3. In the following pedigree, calculate the inbreeding coefficient of M.



Answer: M has three common ancestors, B, C, and D, because two lines of descent from each of these individuals ultimately converge in M. There are four distinct inbreeding loops (common ancestor underlined):

(1) A B <u>C</u> D E	(n = 5)
(2) A D <u>C</u> B E	(n = 5)

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **4.1** What blood types could be observed in children born to a woman who has blood type M and a man who has blood type MN?
- **4.2** In rabbits, coloration of the fur depends on alleles of the gene *c*. From information given in the chapter, what phenotypes and proportions would be expected from the following crosses: (a) $c^+c^+ \times cc$; (b) $c^+c \times c^+c$; (c) $c^+c^b \times c^+c^{cb}$; (d) $cc^{cb} \times cc$; (e) $c^+c^b \times c^+c$; (f) $c^bc \times cc$?
- 4.3 In mice, a series of five alleles determines fur color. In order of dominance, these alleles are: A^Y yellow fur but homozygous lethal; A^L, agouti with light belly; A⁺, agouti (wild-type); a^t, black and tan; and a, black. For each of the following crosses, give the coat color of the parents and the phenotypic ratios expected among the progeny: (a)A^YA^L×A^YA^L; (b)A^Ya×A^La^t; (c)a^ta×A^Ya; (d)A^La^t×A^LA^L; (e) A^LA^L × A^YA⁺; (f) A⁺a^t × a^ta; (g) a^ta × aa; (h) A^YA^L × A⁺A^ta^t; and (i) A^YA^L × A^YA⁺
- 4.4 In several plants, such as tobacco, primrose, and red clover, combinations of alleles in eggs and pollen have been found to influence the reproductive compatibility of the plants. Homozygous combinations, such as S¹S¹, do not develop because S¹ pollen is not effective on S¹- stigmas. However, S¹ pollen is effective on S²S³ stigmas. What progeny might be expected from the following crosses (seed parent written first): (a) S¹S² × S²S³; (b) S¹S² × S³S⁴; (c) S⁴S⁵ × S⁴S⁵; and (d) S³S⁴ × S⁵S⁶?
- 4.5 From information in the chapter about the ABO blood types, what phenotypes and ratios are expected from the following matings: (a) I^AI^A × I^BI^B; (b) I^AI^B × ii; (c) I^Ai × I^Bi; and (d) I^Ai × ii?
- **4.6** A woman with type O blood gave birth to a baby, also with type O blood. The woman stated that a man with type AB blood was the father of the baby. Is there any merit to her statement?
- **4.7** Another woman with type AB blood gave birth to a baby with type B blood. Two different men claim to be the father. One has type A blood, the other type B blood. Can the genetic evidence decide in favor of either?
- **4.8 (co)** The flower colors of plants in a particular population may be blue, purple, turquoise, light-blue, or white. A

3) A <u>B</u> E	(n = 3)
4) A <u>D</u> E	(n = 3)

To calculate the inbreeding coefficient of M, F_M , we raise 1/2 to the power *n* for each of the loops and sum the results:

$$F_M = (1/2)^5 + (1/2)^5 + (1/2)^3 + (1/2)^3 = 5/16$$

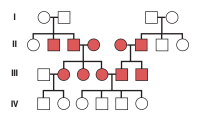
series of crosses	between different me	embers of the popu-
lation produced	the following results	:

Cross	Parents	Progeny
1	purple $ imes$ blue	all purple
2	purple \times purple	76 purple, 25 turquoise
3	blue \times blue	86 blue, 29 turquoise
4	purple \times turquoise	49 purple, 52 turquoise
5	purple \times purple	69 purple, 22 blue
6	purple \times blue	50 purple, 51 blue
7	purple \times blue	54 purple, 26 blue, 25 turquoise
8	turquoise $ imes$ turquoise	all turquoise
9	purple \times blue	49 purple, 25 blue, 23 light-blue
10	light-blue $ imes$ light-blue	60 light-blue, 29 turquoise, 31 white
11	turquoise \times white	all light-blue
12	white \times white	all white
13	$\text{purple} \times \text{white}$	all purple

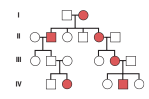
How many genes and alleles are involved in the inheritance of flower color? Indicate all possible genotypes for the following phenotypes: (a) purple; (b) blue; (c) turquoise; (d) light-blue; (e) white.

- **4.9** A woman who has blood type O and blood type M marries a man who has blood type AB and blood type MN. If we assume that the genes for the A-B-O and M-N blood-typing systems assort independently, what blood types might the children of this couple have, and in what proportions?
- **4.10** A Japanese strain of mice has a peculiar, uncoordinated gait called waltzing, which is due to a recessive allele, v. The dominant allele V causes mice to move in a coordinated fashion. A mouse geneticist has recently isolated another recessive mutation that causes uncoordinated movement. This mutation, called *tango*, could be an allele of the *waltzing* gene, or it could be a mutation in an entirely different gene. Propose a test to determine whether the *waltzing* and *tango* mutations are alleles, and if they are, propose symbols to denote them.

4.11 Congenital deafness in humans is inherited as a recessive condition. In the following pedigree, two deaf individuals, each presumably homozygous for a recessive mutation, have married and produced four children with normal hearing. Propose an explanation.



- **4.12** In the fruit fly, recessive mutations in either of two independently assorting genes, *brown* and *purple*, prevent the synthesis of red pigment in the eyes. Thus, homozygotes for either of these mutations have brownish-purple eyes. However, heterozygotes for both of these mutations have dark red, that is, wild-type eyes. If such double heterozygotes are intercrossed, what kinds of progeny will be produced, and in what proportions?
- **4.13** The dominant mutation *Plum* in the fruit fly also causes brownish-purple eyes. Is it possible to determine by genetic experiments whether *Plum* is an allele of the *brown* or *purple* genes?
- **4.14** From information given in the chapter, explain why mice with yellow coat color are not true-breeding.
- **4.15** A couple has four children. Neither the father nor the mother is bald; one of the two sons is bald, but neither of the daughters is bald.
 - (a) If one of the daughters marries a nonbald man and they have a son, what is the chance that the son will become bald as an adult?
 - (b) If the couple has a daughter, what is the chance that she will become bald as an adult?
- **4.16** The following pedigree shows the inheritance of ataxia, a rare neurological disorder characterized by uncoordinated movements. Is ataxia caused by a dominant or a recessive allele? Explain.



4.17 Chickens that carry both the alleles for rose comb (R) and pea comb (P) have walnut combs, whereas chickens that lack both of these alleles (that is, they are genotypically rr pp) have single combs. From the information about interactions between these two genes given in the chapter, determine the phenotypes and proportions expected from the following crosses:

(a) *RR Pp* × *rr Pp*(b) *rr PP* × *Rr Pp*

(c)
$$Rr Pp \times Rr pp$$

(d) $Rr pp \times rr pp$

- **4.18** Rose-comb chickens mated with walnut-comb chickens produced 15 walnut-, 14 rose-, 5 pea-, and 6 single-comb chicks. Determine the genotypes of the parents.
- **4.19** Summer squash plants with the dominant allele *C* bear white fruit, whereas plants homozygous for the recessive allele *c* bear colored fruit. When the fruit is colored, the dominant allele *G* causes it to be yellow; in the absence of this allele (that is, with genotype *gg*), the fruit color is green. What are the F_2 phenotypes and proportions expected from intercrossing the progeny of *CC GG* and *cc gg* plants? Assume that the *C* and *G* genes assort independently.
- **4.20** The white Leghorn breed of chickens is homozygous for the dominant allele *C*, which produces colored feathers. However, this breed is also homozygous for the dominant allele *I* of an independently assorting gene that inhibits coloration of the feathers. Consequently, Leghorn chickens have white feathers. The white Wyandotte breed of chickens has neither the allele for color nor the inhibitor of color; it is therefore genotypically α *ii*. What are the F₂ phenotypes and proportions expected from intercrossing the progeny of a white Leghorn hen and a white Wyandotte rooster?
- **4.21** Fruit flies homozygous for the recessive mutation *scarlet* have bright red eyes because they cannot synthesize brown pigment. Fruit flies homozygous for the recessive mutation *brown* have brownish-purple eyes because they cannot synthesize red pigment. Fruit flies homozygous for both of these mutations have white eyes because they cannot synthesize either type of pigment. The *brown* and *scarlet* mutations assort independently. If fruit flies that are heterozygous for both of these mutations are intercrossed, what kinds of progeny will they produce, and in what proportions?
- **4.22** Consider the following hypothetical scheme of determination of coat color in a mammal. Gene A controls the conversion of a white pigment P_0 into a gray pigment P_1 ; the dominant allele A produces the enzyme necessary for this conversion, and the recessive allele *a* produces an enzyme without biochemical activity. Gene B controls the conversion of the gray pigment P_1 into a black pigment P_2 ; the dominant allele *B* produces the active enzyme for this conversion, and the recessive allele *b* produces an enzyme without activity. The dominant allele C of a third gene produces a polypeptide that completely inhibits the activity of the enzyme produced by gene A; that is, it prevents the reaction $P_0 \rightarrow P_1$. Allele *c* of this gene produces a defective polypeptide that does not inhibit the reaction $P_0 \rightarrow P_1$. Genes A, B, and C assort independently, and no other genes are involved. In the F, of the cross AA bb CC \times aa BB cc, what is the expected phenotypic segregation ratio?
- **4.23** What F_2 phenotypic segregation ratio would be expected for the cross described in the preceding problem if the dominant allele, *C*, of the third gene produced a product that completely inhibited the activity of the enzyme produced by

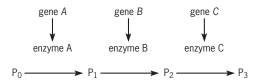
gene *B*—that is, prevented the reaction $P_1 \rightarrow P_2$ rather than inhibiting the activity of the enzyme produced by gene *A*?

- **4.24 (c)** The Micronesian Kingfisher, *Halcyon cinnamomina*, has a cinnamon-colored face. In some birds, the color continues onto the chest, producing one of three patterns: a circle, a shield, or a triangle; in other birds, there is no color on the chest. A male with a colored triangle was crossed with a female that had no color on her chest, and all their offspring had a colored shield on the chest. When these offspring were intercrossed, they produced an F_2 with a phenotypic ratio of 3 circle:6 shield:3 triangle:4 no color. (a) Determine the mode of inheritance for this trait and indicate the genotypes of the birds in all three generations. (b) If a male without color on his chest is mated to a female with a colored shield on her chest and the F_1 segregate in the ratio of 1 circle:2 shield:1 triangle, what are the genotypes of the parents and their progeny?
- **4.25** In a species of tree, seed color is determined by four independently assorting genes: *A*, *B*, *C*, and *D*. The recessive alleles of each of these genes (*a*, *b*, *c*, and *d*) produce abnormal enzymes that cannot catalyze a reaction in the biosynthetic pathway for seed pigment. This pathway is diagrammed as follows:

White precursor
$$\xrightarrow{A}$$
 Yellow \xrightarrow{B} Orange \xrightarrow{C} Red
D

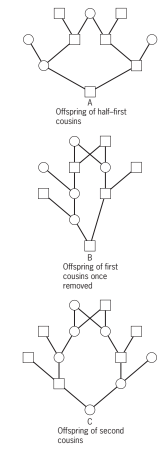
When both red and blue pigments are present, the seeds are purple. Trees with the genotypes *Aa Bb Cc Dd* and *Aa Bb Cc dd* were crossed.

- (a) What color are the seeds in these two parental genotypes?
- (b) What proportion of the offspring from the cross will have white seeds?
- (c) Determine the relative proportions of red, white, and blue offspring from the cross.
- **4.26** Multiple crosses were made between true-breeding lines of black and yellow Labrador retrievers. All the F₁ progeny were black. When these progeny were intercrossed, they produced an F₂ consisting of 91 black, 39 yellow, and 30 chocolate. (a) Propose an explanation for the inheritance of coat color in Labrador retrievers. (b) Propose a biochemical pathway for coat color determination and indicate how the relevant genes control coat coloration.
- **4.27** Two plants with white flowers, each from true-breeding strains, were crossed. All the F_1 plants had red flowers. When these F_1 plants were intercrossed, they produced an F_2 consisting of 177 plants with red flowers and 142 with white flowers. (a) Propose an explanation for the inheritance of flower color in this plant species. (b) Propose a biochemical pathway for flower pigmentation and indicate which genes control which steps in this pathway.
- **4.28** Consider the following genetically controlled biosynthetic pathway for pigments in the flowers of a hypothetical plant:

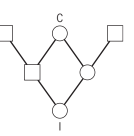


Assume that gene A controls the conversion of a white pigment, P_0 , into another white pigment, P_1 ; the dominant allele A specifies an enzyme necessary for this conversion, and the recessive allele *a* specifies a defective enzyme without biochemical function. Gene B controls the conversion of the white pigment, P_1 , into a pink pigment, P_2 ; the dominant allele, *B*, produces the enzyme necessary for this conversion, and the recessive allele, b, produces a defective enzyme. The dominant allele, C, of the third gene specifies an enzyme that converts the pink pigment, P₂, into a red pigment, P₃; its recessive allele, *c*, produces an altered enzyme that cannot carry out this conversion. The dominant allele, D, of a fourth gene produces a polypeptide that completely inhibits the function of enzyme C; that is, it blocks the reaction $P_2 \rightarrow P_3$. Its recessive allele, d, produces a defective polypeptide that does not block this reaction. Assume that flower color is determined solely by these four genes and that they assort independently. In the F_2 of a cross between plants of the genotype AA bb CC DD and plants of the genotype aa BB cc dd, what proportion of the plants will have (a) red flowers? (b) pink flowers? (c) white flowers?

4.29 In the following pedigrees, what are the inbreeding coefficients of A, B, and C?



- **4.31** Mabel and Frank are half siblings, as are Tina and Tim. However, these two pairs of half sibs do not have any common ancestors. If Mabel marries Tim and Frank marries Tina and each couple has a child, what fraction of their genes will these children share by virtue of common ancestry? Will the children be more or less closely related than first cousins?
- **4.32** Suppose that the inbreeding coefficient of I in the following pedigree is 0.25. What is the inbreeding coefficient of I's common ancestor, C?



4.33 A randomly pollinated strain of maize produces ears that are 24 cm long, on average. After one generation of self-fertilization, the ear length is reduced to 20 cm. Predict the ear length if self-fertilization is continued for one more generation.

Genomics on the Web at http://www.ncbi.nlm.nih.gov

Coat color in mammals is controlled by many different genes.

- In the mouse, the A^Y mutation, a dominant allele of the a gene, makes the coat yellow instead of agouti; in homozygous condition, this mutation is lethal. Can you find a description of the a gene and its A^Y allele in the mouse genomics database? What is the official name of this gene?
- 2. Albinism in mice is caused by recessive mutations in a gene called *Tyr*, also symbolized *c*. This gene encodes the enzyme tyrosinase, which catalyzes a step in the production of melanin pigment from the amino acid tyrosine. Can you find a

description of this gene in the mouse genomics database? Do you suspect that this gene is related, in an evolutionary sense, to the gene that, when mutant, causes albinism in rabbits?

3. Do humans have a gene related to the *Tyr* gene of mice? If they do, what condition might this gene, when mutant, be associated with?

Hint: At the web site under Popular Resources, click on Gene. Then search for A<Y> or for Tyr.

The Chromosomal Basis of Mendelism



CHAPTER OUTLINE

- Chromosomes
- The Chromosome Theory of Heredity
- Sex-Linked Genes in Humans
- Sex Chromosomes and Sex Determination
- Dosage Compensation of X-Linked Genes

Sex, Chromosomes, and Genes

What causes organisms to develop as males or females? Why are there only two sexual phenotypes? Is the sex of an organism determined by its genes? These and related questions have intrigued geneticists since the rediscovery of Mendel's work at the beginning of the twentieth century.

The discovery that genes play a role in the determination of sex emerged from a fusion between two previously distinct scientific disciplines, genetics—the study of heredity—and cytology—the study of cells. Early in the twentieth century, these disciplines were brought together through a friendship between two remarkable



The fruit fly, Drosophila melanogaster.

American scientists, Thomas Hunt Morgan and Edmund Beecher Wilson. Morgan was the geneticist and Wilson the cytologist.

As the cytologist, Wilson was interested in the behavior of chromosomes. These structures would prove to be important for sex determination in many species, including our own. Wilson was one of the first to investigate differences in the chromosomes of the two sexes. Through careful study, he and his colleagues showed that these differences were confined to a special pair of chromosomes called sex chromosomes. Wilson found that the behavior of these chromosomes during meiosis could account for the inheritance of sex.

As the geneticist, Morgan was interested in the identification of genes. He focused his research on the fruit fly, *Drosophila melanogaster*, and rather quickly discovered a gene that gave different phenotypic ratios in males and females. Morgan hypothesized that this gene was located on one of the sex chromosomes, and one of his students, Calvin Bridges, eventually proved this hypothesis to be correct. Morgan's discovery that genes reside on chromosomes was a great achievement. The abstract genetic factors postulated by Mendel were finally localized on visible structures within cells. Geneticists could now explain the Principles of Segregation and Independent Assortment in terms of meiotic chromosome behavior.

The discovery that specific genes determine the sex of an organism came much later, only after another scientific discipline, molecular biology, had joined forces with genetics and cytology. Through their combined efforts, cytologists, geneticists, and molecular biologists identified specific sex-determining genes by studying rare individuals in which the sexual phenotype was inconsistent with the sex chromosomes that were present. Today, researchers in all three fields are earnestly trying to figure out how these genes control sexual development.

Chromosomes

Each species has a characteristic set of chromosomes.

of the nineteenth century by a German cytologist, W. Waldeyer. Subsequent investigations with many different organisms established that chromosomes are characteristic of the nuclei of all cells. They are best seen by applying dyes to dividing cells; during division, the material in a chromosome is packed into a small volume, giving it the appearance of a tightly organized cylinder. During the interphase between cell divisions, chromosomes are not so easily seen, even with the best of dyes. Interphase chromosomes are loosely coiled, forming thin threads that are distributed throughout the nucleus. Consequently, when dyes are applied, the whole nucleus is stained and individual chromosomes cannot be identified. This diffuse network of threads is called **chromatin**. Some regions of the chromatin stain more darkly than others, suggesting an underlying difference in organization. The light regions are called the **euchromatin** (from the Greek word for "true"), and the dark regions are called the **heterochromatin** (from the Greek word for "different"). We will explore the functional significance of these different types of chromatin in Chapter 19.

Chromosomes were discovered in the second half

CHROMOSOME NUMBER

Within a species, the number of chromosomes is almost always an even multiple of a basic number. In humans, for example, the basic number is 23; mature eggs and sperm have this number of chromosomes. Most other types of human cells have twice as many (46), although a few kinds, such as some liver cells, have four times (92) the basic number.

The **haploid**, or basic, chromosome number (n) defines a set of chromosomes called the *haploid genome*. Most somatic cells contain two of each of the chromosomes in this set and are therefore **diploid (2n)**. Cells with four of each chromosome are **tetraploid (4n)**, those with eight of each are **octoploid (8n)**, and so on.

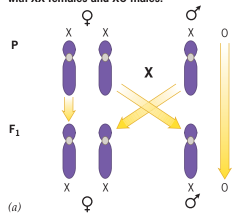
The basic number of chromosomes varies among species. Chromosome number is unrelated to the size or biological complexity of an organism, with most species containing between 10 and 40 chromosomes in their genomes (Table 5.1). The muntjac, a tiny Asian deer, has only three chromosomes in its genome, whereas some species of ferns have many hundreds.

SEX CHROMOSOMES

In some animal species such as grasshoppers, females have one more chromosome than males (**Figure 5.1a**). This extra chromosome, originally observed in other insects, is called the **X chromosome**. Females of these species have two X chromosomes, and males have only one; thus, females are cytologically XX and males are XO, where the "O" denotes the absence of a chromosome. During meiosis in the female, the two X chromosomes pair and then separate, producing eggs that contain a single X chromosome. During meiosis in the male, the solitary X chromosome moves independently of all the other chromosome. Thus, when sperm and eggs unite, two kinds of zygotes are produced: XX, which develop into females, and XO, which develop into males. Because each of these types is equally likely, the reproductive mechanism preserves a 1:1 ratio of males to females in these species.

In many other animals, including humans, males and females have the same number of chromosomes (**■** Figure 5.1*b*). This numerical equality is due to the presence of a chromosome in the male, called the **Y** chromosome, which pairs with the X during meiosis. The Y chromosome is morphologically distinguishable from the X chromosome. In humans, for example, the Y is much shorter than the X, and its centromere is located closer to one of the ends (**■** Figure 5.2). The material common to the human X and Y chromosomes is limited, consisting mainly of short segments

Inheritance of sex chromosomes in animals with XX females and XO males.



Inheritance of sex chromosomes in animals with XX females and XY males.

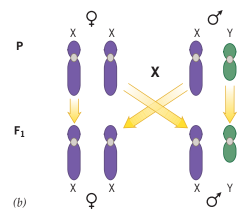


FIGURE 5.1 Inheritance of sex chromosomes in animals. (a) XX female/XO male animals, such as some grasshoppers. (b) XX female/XY male animals, such as humans and *Drosophila*.

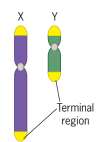
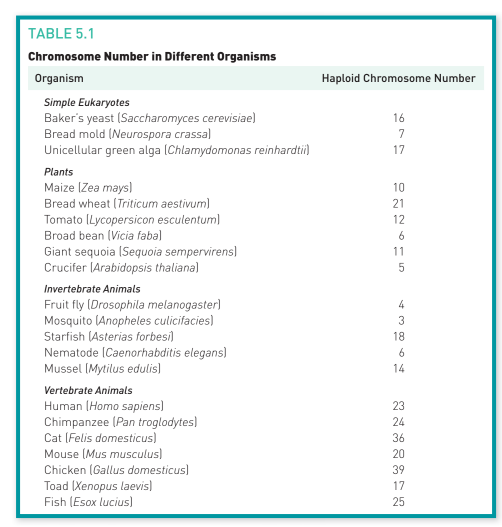


FIGURE 5.2 Human X and Y chromosomes. The terminal regions are common to both sex chromosomes.

KEY POINTS



near the ends of the chromosomes. During meiosis in the male, the X and Y chromosomes separate from each other, producing two kinds of sperm, X-bearing and Y-bearing; the frequencies of the two types are approximately equal. XX females produce only one kind of egg, which is X-bearing. If fertilization were to occur randomly, approximately half the zygotes would be XX and the other half would be XY, leading to a 1:1 sex ratio at conception. However, in humans, Y-bearing sperm have a fertilization advantage because they are lighter and move faster, and the zygotic sex ratio is about 1.3:1. During development, the excess of males is diminished by differential viability of XX and XY embryos, and at birth, males are only slightly more numerous than females (sex ratio 1.07:1). By the age of reproduction, the excess of males is essentially eliminated and the sex ratio is very close to 1:1.

The X and Y chromosomes are called **sex chromosomes.** All the other chromosomes in the genome are called **autosomes.** Sex chromosomes were discovered in the first few years of the twentieth century through the work of the American cytologists C. E. McClung, N. M. Stevens, W. S. Sutton, and E. B. Wilson. This discovery coincided closely with the emergence of Mendelism and stimulated research on the possible relationships between Mendel's principles and the meiotic behavior of chromosomes.

- Individual chromosomes become visible during cell division; between divisions they form a diffuse network of fibers called chromatin.
- Diploid somatic cells have twice as many chromosomes as haploid gametes.
- Sex chromosomes are different between the two sexes, whereas autosomes are the same.

The Chromosome Theory of Heredity

Studies on the inheritance of a sex-linked trait in *Drosophila* provided the first evidence that the meiotic behavior of chromosomes is the basis for Mendel's Principles of Segregation and Independent Assortment. By 1910 many biologists suspected that genes were situated on chromosomes, but they did not have definitive proof. Researchers needed to find a gene that could be unambiguously linked to a chromosome. This goal required that the gene be defined by a mutant allele and that the chromosome be morphologically distinguishable. Furthermore, the pattern of gene transmission had to reflect the chromosome's behavior during reproduction. All these

requirements were fulfilled when the American biologist Thomas H. Morgan discovered a particular eye color mutation in the fruit fly, *Drosophila melanogaster*. Morgan began experimentation with this species of fly in about 1909. It was ideally suited for genetics research because it reproduced quickly and prolifically and was inexpensive to rear in the laboratory. In addition, it had only four pairs of chromosomes, one being a pair of sex chromosomes—XX in the female and XY in the male. The X and Y chromosomes were morphologically distinguishable from each other and from each of the autosomes. Through careful experiments, Morgan was able to show that the eye color mutation was inherited along with the X chromosome, suggesting that a gene for eye color was physically situated on that chromosome. Later, one of his students, Calvin B. Bridges, obtained definitive proof for this Chromosome Theory of Heredity.

EXPERIMENTAL EVIDENCE LINKING THE INHERITANCE OF GENES TO CHROMOSOMES

Morgan's experiments commenced with his discovery of a mutant male fly that had white eyes instead of the red eyes of wild-type flies. When this male was crossed to wild-type females, all the progeny had red eyes, indicating that white was recessive to red. When these progeny were intercrossed with each other, Morgan observed a

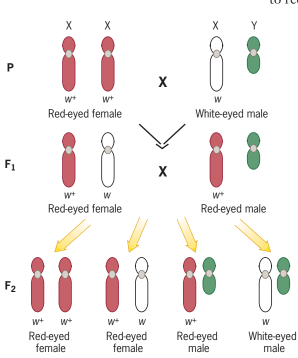
peculiar segregation pattern: all of the daughters, but only half of the sons, had red eyes; the other half of the sons had white eyes. This pattern suggested that the inheritance of eye color was linked to the sex chromosomes. Morgan proposed that a gene for eye color was present on the X chromosome, but not on the Y, and that the white and red phenotypes were due to two different alleles, a mutant allele denoted w and a wild-type allele denoted w^+ .

Morgan's hypothesis is diagrammed in **Figure 5.3**. The wild-type females in the first cross are assumed to be homozygous for the w^+ allele. Their mate is assumed to carry the mutant w allele on its X chromosome and neither of the alleles on its Y chromosome. An organism that has only one copy of a gene is called a **hemizygote**. Among the progeny from the cross, the sons inherit an X chromosome from their mother and a Y chromosome from their father; because the maternally inherited X carries the w^+ allele, these sons have red eyes. The daughters, in contrast, inherit an X chromosome from the mother and a X with w from the father. However, because w^+ is dominant to w, these heterozygous F₁ females also have red eyes.

When the F_1 males and females are intercrossed, four genotypic classes of progeny are produced, each representing a different combination of sex chromosomes. The XX flies, which are female, have red eyes because at least one w^+ allele is present. The XY flies, which are male, have either red or white eyes, depending on which X chromosome is

inherited from the heterozygous F_1 females. Segregation of the w and w^+ alleles in these females is therefore the reason half the F_2 males have white eyes.

Morgan carried out additional experiments to confirm the elements of his hypothesis. In one (\blacksquare Figure 5.4*a*), he crossed F₁ females assumed to be heterozygous for the eye color gene to mutant white males. As he expected, half the progeny



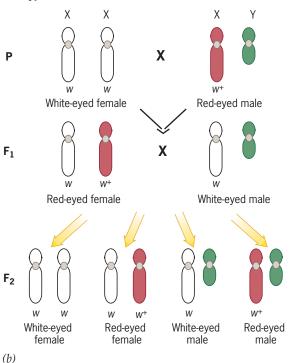
■ FIGURE 5.3 Morgan's experiment studying the inheritance of white eyes in *Drosophila*. The transmission of the mutant condition in association with sex suggested that the gene for eye color was present on the X chromosome but not on the Y chromosome.

Х Х Υ χ F_1 Х Red-eyed female White-eyed male F₂ W^+ W W⁺ W w Red-eyed White-eyed Red-eyed White-eyed female female male male *(a)*

Cross between a heterozygous female and a hemizygous

mutant male.

Cross between a homozygous mutant female and a hemizygous wild-type male.



of each sex had white eyes, and the other half had red eyes. In another experiment (**Figure 5.4b**), he crossed white-eyed females to red-eyed males. This time, all the daughters had red eyes, and all the sons had white eyes.

When he intercrossed these progeny, Morgan observed the expected segregation: half the progeny of each sex had white eyes, and the other half had red eyes. Thus, Morgan's hypothesis that the gene for eye color was linked to the X chromosome withstood additional experimental testing. ■ FIGURE 5.4 Experimental tests of Morgan's hypothesis that the gene for eye color in *Drosophila* is X-linked. (a) Experiment in which heterozygous females were crossed to white-eyed males. (b) Experiment in which white-eyed females were crossed to wild-type males.

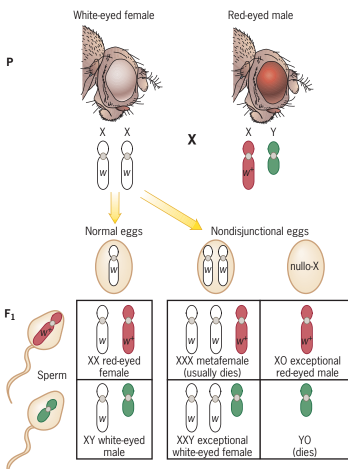
NONDISJUNCTION AS PROOF OF THE CHROMOSOME THEORY

Morgan showed that a gene for eye color was on the X chromosome of *Drosophila* by correlating the inheritance of that gene with the transmission of the X chromosome during reproduction. However, as noted earlier, it was one of his students, C. B. Bridges, who secured proof of the chromosome theory by showing that exceptions to the rules of inheritance could also be explained by chromosome behavior.

Bridges performed one of Morgan's experiments on a larger scale. He crossed white-eyed female *Drosophila* to red-eyed males and examined many F_1 progeny. Although as expected, nearly all the F_1 flies were either red-eyed females or white-eyed males, Bridges found a few exceptional flies—white-eyed females and red-eyed males. He crossed these exceptions to determine how they might have arisen. The exceptional males all proved to be sterile; however, the exceptional females were fertile, and when crossed to normal red-eyed males, they produced many progeny, including large numbers of white-eyed daughters and red-eyed sons. Thus, the exceptional F_1 females, though rare in their own right, were prone to produce many exceptional progeny.

Bridges explained these results by proposing that the exceptional F_1 flies were the result of abnormal X chromosome behavior during meiosis in the females of the P generation. Ordinarily, the X chromosomes in these females should *disjoin*, or separate from each other, during meiosis. Occasionally, however, they might fail to separate, producing an egg with two X chromosomes or an egg with no X chromosome at all. Fertilization of such abnormal eggs by normal sperm would produce zygotes with an abnormal number of sex chromosomes. **■** Figure 5.5 illustrates the possibilities.

If an egg with two X chromosomes (usually called a diplo-X egg; genotype $X^{w}X^{w}$) is fertilized by a Y-bearing sperm, the zygote will be $X^{w}X^{w}Y$. Since each of the



■ FIGURE 5.5 X chromosome nondisjunction is responsible for the exceptional progeny that appeared in Bridges' experiment. Nondisjunctional eggs that contain either two X chromosomes or no X chromosome unite with normal sperm that contain either an X chromosome or a Y chromosome to produce four types of zygotes. The XXY zygotes develop into white-eyed females, the XO zygotes develop into red-eyed, sterile males, and the YO zygotes die. Some of the XXX zygotes develop into sickly, red-eyed females, but most of them die.

X chromosomes in this zygote carries a mutant w allele, the resulting fly will have white eyes. If an egg without an X chromosome (usually called a nullo-X egg) is fertilized by an X-bearing sperm (X⁺), the zygote will be X⁺O. (Once again, "O" denotes the absence of a chromosome.) Because the single X in this zygote carries a w^+ allele, the zygote will develop into a red-eyed fly. Bridges inferred that XXY flies were female and that XO flies were male. The exceptional white-eyed females that he observed were therefore $X^w X^w Y$, and the exceptional red-eyed males were X^+O . Bridges confirmed the chromosome constitutions of these exceptional flies by direct cytological observation. Because the XO animals were male, Bridges concluded that in *Drosophila* the Y chromosome has nothing to do with the determination of the sexual phenotype. However, because the XO males were always sterile, he realized that this chromosome must be important for male sexual function.

Bridges recognized that the fertilization of abnormal eggs by normal sperm could produce two additional kinds of zygotes: $X^{w}X^{w}X^{+}$, arising from the union of a diplo-X egg and an X-bearing sperm, and YO, arising from the union of a nullo-X egg and a Y-bearing sperm. The $X^{w}X^{w}X^{+}$ zygotes develop into females that are red-eyed, but weak and sickly. These "metafemales" can be distinguished from XX females by a syndrome of anatomical abnormalities, including ragged wings and etched abdomens. Generations of geneticists have inappropriately called them "superfemales"—a term coined by Bridges—even though there is nothing super about them. The YO zygotes turn out to be completely inviable; that is, they die. In *Drosophila*, as in most other organisms with sex chromosomes, at least one X chromosome is needed for viability.

Bridges' ability to explain the exceptional progeny that came from these crosses showed the power of the chromosome theory. Each of the exceptions was due to anomalous chromosome behavior during meiosis. Bridges called the anomaly **nondisjunction** because it involved a failure of the chromosomes to disjoin during one of the meiotic divisions. This failure could result from faulty chromosome movement, imprecise or incomplete pairing, or centromere malfunction. From Bridges' data, it is impossible to specify the exact cause. However, Bridges did note that the exceptional XXY females go on to produce a high frequency of exceptional progeny, presumably because their sex chromosomes can disjoin in different ways: the X chromosomes can disjoin from each other, or either X can disjoin from the Y. In the latter case, a diplo- or nullo-X egg is produced because the X that does not disjoin from the Y is free to move to either pole during the first meiotic division. When fertilized by normal sperm, these abnormal eggs will

Bridges observed the effects of chromosome nondisjunction that had occurred during meiosis in females. We should note, however, that with appropriate experiments the effects of nondisjunction during meiosis in males can also be studied. Test your understanding of Bridges' experiment by working through Solve It: Sex Chromosome Nondisjunction.

produce exceptional zygotes.

These early studies with *Drosophila*—primarily the work of Morgan and his students (see A Milestone in Genetics: Morgan's Fly Room in the Student Companion site)—greatly strengthened the view that all genes were located on chromosomes and that Mendel's principles could be explained by the transmissional properties of chromosomes during reproduction. This idea, called the **Chromosome Theory of Heredity**, stands as one of the most important achievements in biology. Since its formulation in the early part of the twentieth century, the Chromosome Theory of Heredity has provided a unifying framework for all studies of inheritance.

THE CHROMOSOMAL BASIS OF MENDEL'S PRINCIPLES OF SEGREGATION AND INDEPENDENT ASSORTMENT

Mendel established two principles of genetic transmission: (1) the alleles of a single gene segregate from each other, and (2) the alleles of two different genes assort independently. The finding that genes are located on chromosomes made it possible to explain these principles (as well as exceptions to them) in terms of the meiotic behavior of chromosomes.

The Principle of Segregation

During the first meiotic division, homologous chromosomes pair. One of the homologues comes from the mother, the other from the father. If the mother was homozygous for an allele, A, of a gene on this chromosome, and the father was homozygous for a different allele, a, of the same gene, the offspring must be heterozygous, that is, Aa. In the anaphase of the first meiotic division, the paired chromosomes separate and move to opposite poles of the cell. One carries allele A and the other allele a. This physical separation of the two chromosomes segregates the alleles from each other; eventually, they will reside in different daughter cells. Mendel's Principle of Segregation (**True Figure 5.6**) is therefore based on the separation of homologous chromosomes during the anaphase of the first meiotic division.

The Principle of Independent Assortment

The Principle of Independent Assortment (**Figure 5.7**) is also based on this anaphase separation. To understand the relationship, we need to consider genes on two different pairs of chromosomes. Suppose that a heterozygote $Aa \ Bb$ was produced by mating an $AA \ BB$ female to an $aa \ bb$ male; also, suppose that the two genes are on different

Solve It!

Sex Chromosome Nondisjunction

A researcher crossed white-eyed males and red-eved females, each from truebreeding strains of Drosophila. The vast majority of the offspring, both males and females, had red eyes and were normal in other respects. However, some exceptional flies were observed: (a) several white-eyed males that proved to be sterile, (b) several red-eyed females with ragged wings and etched abdomens, and (c) one white-eved female. If the gene for eye color is on the X chromosome (but not on the Y chromosome), in which parent(s) did nondisjunction of the sex chromosomes occur to produce the exceptional offsprina?

To see the solution to this problem, visit the Student Companion site.

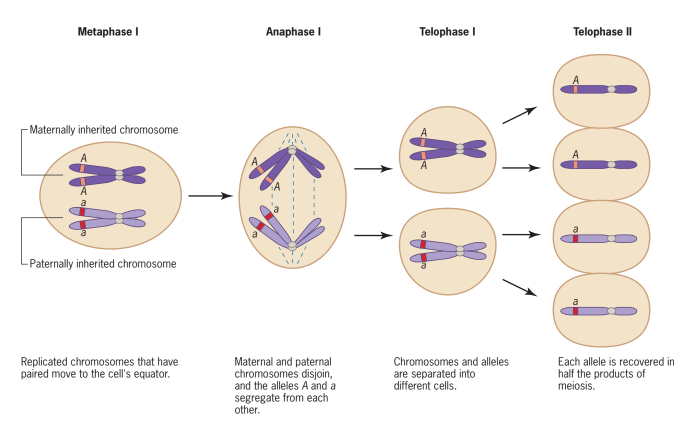


FIGURE 5.6 Mendel's Principle of Segregation and meiotic chromosome behavior. The segregation of alleles corresponds to the disjunction of paired chromosomes in the anaphase of the first meiotic division.

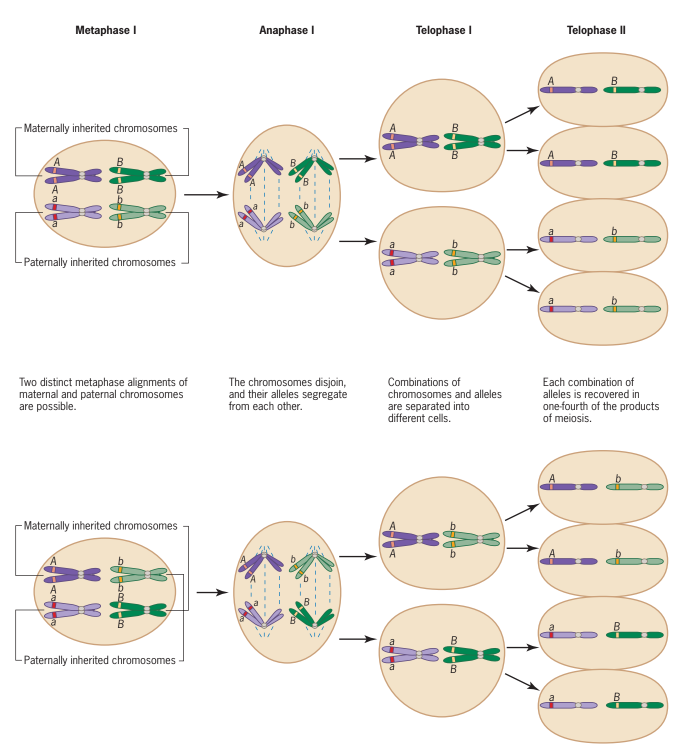


FIGURE 5.7 Mendel's Principle of Independent Assortment and meiotic chromosome behavior. Alleles on different pairs of chromosomes assort independently in the anaphase of the first meiotic division because maternally and paternally inherited chromosomes have aligned randomly on the cell's equator.

chromosomes. During the prophase of meiosis I, the chromosomes with the A and a alleles will pair, as will the chromosomes with the B and b alleles. At metaphase, the two pairs will take up positions on the meiotic spindle in preparation for the upcoming anaphase separation. Because there are two pairs of chromosomes, there are two distinguishable metaphase alignments:

$$\frac{A}{a} \frac{B}{b}$$
 or $\frac{A}{a} \frac{b}{B}$

Each of these alignments is equally likely. Here the space separates different pairs of chromosomes, and the bar separates the homologous members of each pair. During anaphase, the alleles above the bars will move to one pole, and the alleles below them will move to the other. When disjunction occurs, there is therefore a 50 percent chance that the A and B alleles will move together to the same pole and a 50 percent chance that they will move to opposite poles. Similarly, there is a 50 percent chance that the *a* and b alleles will move to the same pole and a 50 percent chance that they will move to opposite poles. At the end of meiosis, when the chromosome number is finally reduced, half the gametes should contain a parental combination of alleles (A B or a b), and half should contain a new combination (A b or a B). Altogether, there will be four types of gametes, each one-fourth of the total. This equality of gamete frequencies is a result of the independent behavior of the two pairs of chromosomes during the first meiotic division. Mendel's Principle of Independent Assortment is therefore a statement about the random alignment of different pairs of chromosomes at metaphase. In Chapter 7, we will see that genes on the same pair of chromosomes do not assort independently. Instead, because they are physically linked to each other, they tend to travel together through meiosis, violating the Principle of Independent Assortment. To test your understanding of the chromosomal basis of independent assortment, work through Problem-Solving Skills: Tracking X-Linked and Autosomal Inheritance.

PROBLEM-SOLVING SKILLS

Tracking X-Linked and Autosomal Inheritance

THE PROBLEM

In *Drosophila*, one of the genes controlling wing length is located on the X chromosome. A recessive mutant allele of this gene makes the wings miniature—hence, its symbol *m*; the wild-type allele of this gene, *m*⁺, makes the wings long. One of the genes controlling eye color is located on an autosome. A recessive mutant allele of this gene makes the eyes brown—hence, its symbol *bw*; the wildtype allele of this gene, *bw*⁺, makes the eyes red. Miniature-winged, red-eyed females from one true-breeding strain were crossed to normal-winged, brown-eyed males from another true-breeding strain. (a) Predict the phenotypes of the F₁ flies. (b) If these flies are intercrossed with one another, what phenotypes will appear in the F₂, and in what proportions?

FACTS AND CONCEPTS

- 1. Male and female offspring from a cross may show different phenotypes if the trait is X-linked.
- **2.** A male inherits its X chromosome from its mother, whereas a female inherits one of its X chromosome from its father.
- **3.** X-linked and autosomal genes assort independently.
- **4.** When genes assort independently, we multiply the probabilities associated with the components of the complete genotype.

ANALYSIS AND SOLUTION

- **a.** The parents in the initial cross were m/m; bw^+/bw^+ females and m^+/Y ; bw/bw males. In the F₁, the females will be m/m^+ ; bw/bw^+ and because both mutant alleles are recessive, they will have long wings and red eyes. The F₁ males will be m/Y; bw/bw^+ , and because they are hemizygous for the recessive X-linked mutation, they will have miniature wings; however, because they carry the dominant autosomal allele bw^+ , they will have red eyes.
- **b.** To obtain the F_2 phenotypes and their proportions, let's subdivide the problem into two parts: an X-linked part and an autosomal part. For the X-linked part, crossing the $F_1 m/m^+$ females to their m/Y brothers will produce four classes of offspring— (1) m/m females with miniature wings, (2) m/m^+ females with long wings, (3) m/Y males with miniature wings, and (4) m^+/Y males with long wings, and each class should be 1/4 of the total. For the autosomal part, crossing the $F_1 bw/bw^+$ females to their *bw/bw*⁺ brothers will produce three classes of offspring— (1) bw^+/bw^+ flies with red eyes, (2) bw/bw^+ flies with red eyes, and (3) bw/bw flies with brown eyes, and the phenotypic ratio will be 3 red: 1 brown. To combine the results of the X-linked and autosomal parts of the problem, we construct a 2 x 4 table of phenotypic frequencies. The two autosomal phenotypes and the four X-linked phenotypes define the rows and columns of the table, and the values within the cells are the frequencies of the combined phenotypes, obtained by multiplying the frequencies in the margins.

		X-Linked Phenotypes			
		Miniature Female (1/4)	Normal Female (1/4)	Miniature Male (1/4)	Normal Male (1/4)
Autosomal	Red (3/4)	3/16	3/16	3/16	3/16
Phenotypes	Brown (1/4)	1/16	1/16	1/16	1/16

For further discussion visit the Student Companion site.

KEY POINTS

- Genes are located on chromosomes.
 - The disjunction of chromosomes during meiosis is responsible for the segregation and independent assortment of genes.
 - Nondisjunction during meiosis leads to abnormal numbers of chromosomes in gametes and, ultimately, in zygotes.

Sex-Linked Genes in Humans

X- and Y-linked genes have been studied in humans.

The development of the chromosome theory depended on the discovery of the *white* eye mutation in *Drosophila*. Subsequent analysis demonstrated that this mutation was a recessive allele of an X-linked gene. Although some of

us might credit this important episode in the history of genetics to extraordinarily good luck, Morgan's discovery of the *white* eye mutation was not so remarkable. Such mutations are among the easiest to detect because they show up immediately in hemizygous males. In contrast, autosomal recessive mutations show up only after two mutant alleles have been brought together in a homozygote—a much more unlikely event.

In humans too, recessive X-linked traits are much more easily identified than are recessive autosomal traits. A male needs only to inherit one recessive allele to show an X-linked trait; however, a female needs to inherit two—one from each of her parents. Thus, the preponderance of people who show X-linked traits are male.

HEMOPHILIA, AN X-LINKED BLOOD-CLOTTING DISORDER

People with **hemophilia** are unable to produce a factor needed for blood clotting; the cuts, bruises, and wounds of hemophiliacs continue to bleed and, if not stopped by transfusion with clotting factor, can cause death. The principal type of hemophilia in humans is due to a recessive X-linked mutation, and nearly all the individuals who have it are male. These males have inherited the mutation from their hetero-zygous mothers. If they reproduce, they transmit the mutation to their daughters, who usually do not develop hemophilia because they inherit a wild-type allele from their mothers. Affected males never transmit the mutant allele to their sons. Other blood-clotting disorders are found in both males and females because they are due to mutations in autosomal genes.

The most famous case of X-linked hemophilia occurred in the Russian imperial family at the beginning of the twentieth century (**Figure 5.8**). Czar Nicholas and Czarina Alexandra had four daughters and one son, and the son, Alexis, suffered from hemophilia. The X-linked mutation responsible for Alexis's disease was transmitted to him by his mother, who was a heterozygous carrier. Czarina Alexandra was a granddaughter of Queen Victoria of Great Britain, who was also a carrier. Pedigree records show that Victoria transmitted the mutant allele to three of her nine children: Alice, who was Alexandra's mother; Beatrice, who had two sons with the disease; and Leopold, who had the disease himself. The allele that Victoria carried evidently arose as a new mutation in her germ cells, or in those of her mother, father, or a more distant maternal ancestor.

Throughout history hemophilia has been a fatal disease. Most of the people who have had it have died before the age of 20. Today, due to the availability of effective and relatively inexpensive treatments, hemophiliacs live long, healthy lives.

COLOR BLINDNESS, AN X-LINKED VISION DISORDER

In humans, color perception is mediated by light-absorbing proteins in the specialized cone cells of the retina in the eye. Three such proteins have been identified—one to absorb blue light, one to absorb green light, and one to absorb red light. Color

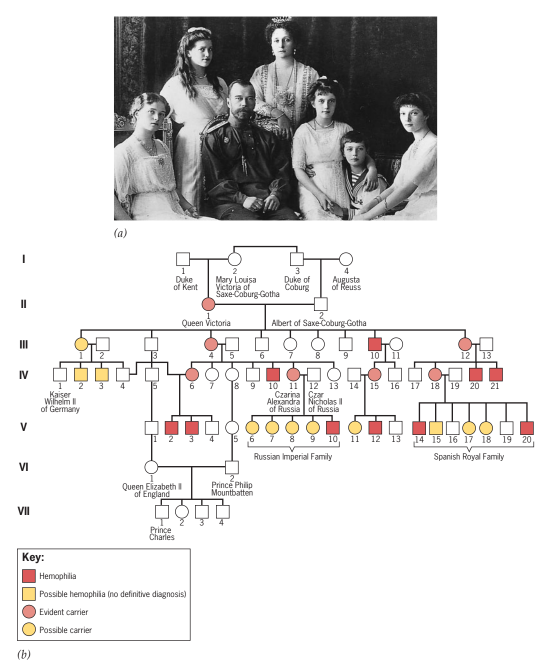


FIGURE 5.8 Royal hemophilia. (a) The Russian imperial family of Czar Nicholas II. (b) X-linked hemophilia in the royal families of Europe. Through intermarriage, the mutant allele for hemophilia was transmitted from the British royal family to the German, Russian, and Spanish royal families.

blindness may be caused by an abnormality in any of these receptor proteins. The classic type of color blindness, involving faulty perception of red and green light, follows an X-linked pattern of inheritance. About 5 to 10 percent of human males are red-green color blind; however, a much smaller fraction of females, less than 1 percent, has this disability, suggesting that the mutant alleles are recessive. Molecular studies have shown that there are two distinct genes for color perception on the X chromosome; one encodes the receptor for green light, and the other encodes the receptor for red light. Detailed analyses have demonstrated that these two receptors are structurally very similar, probably because the genes encoding them evolved from an ancestral color-receptor gene. A third gene for color perception, the one encoding the receptor for blue light, is located on an autosome.

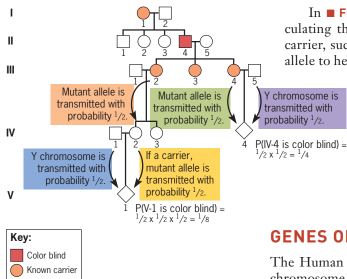
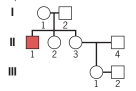


FIGURE 5.9 Analysis of a pedigree showing the segregation of X-linked color blindness.

Solve It! Calculating the Risk for Hemophilia

In this pedigree, II-1 is affected with X-linked hemophilia. If III-1 and III-2 have a child, what is the risk that the child will have hemophilia?



To see the solution to this problem, visit the Student Companion site.

KEY POINTS

In **Figure 5.9** color blindness is used to illustrate the procedures for calculating the risk of inheriting a recessive X-linked condition. A heterozygous carrier, such as III-4 in the figure, has a 1/2 chance of transmitting the mutant allele to her children. However, the risk that a particular child will be color blind

is only 1/4 since the child must be a male in order to manifest the trait. The female labeled IV-2 in the pedigree could be a carrier of the mutant allele for color blindness because her mother was. This uncertainty about the genotype of IV-2 introduces another factor of 1/2 in the risk of having a color-blind child; thus, the risk for her child is $1/4 \times 1/2 = 1/8$. Test your ability to perform this kind of analysis by working through Solve It: Calculating the Risk for Hemophilia.

GENES ON THE HUMAN Y CHROMOSOME

The Human Genome Project has identified 397 possible genes on the human Y chromosome, but fewer than 100 of them seem to be functional. By comparison, it has identified more than 1000 genes on the human X chromosome. Prior to the work of the Human Genome Project, little was known about the genetic makeup of the Y chromosome. Only a handful of Y-linked traits had been detected, even though transmission from father to son should make such traits easy to identify in conventional pedigree analysis. The results of the Human Genome Project have provided one possible explanation for the apparent lack of Y-linked traits. Several of the genes on the human Y chromosome seem to be required for male fertility. Obviously, a mutation in such a gene will interfere with a man's ability to reproduce; hence, that mutation will have little or no chance of being transmitted to the next generation.

GENES ON BOTH THE X AND Y CHROMOSOMES

Some genes are present on both the X and Y chromosomes, mostly near the ends of the short arms (see Figure 5.2). Alleles of these genes do not follow a distinct X- or Y-linked pattern of inheritance. Instead, they are transmitted from mothers and fathers to sons and daughters alike, mimicking the inheritance of an autosomal gene. Such genes are therefore called **pseudoautosomal genes**. In males, the regions that contain these genes seem to mediate pairing between the X and Y chromosomes.

• Disorders such as hemophilia and color blindness, which are caused by recessive X-linked mutations, are more common in males than in females.

- In humans the Y chromosome carries fewer genes than the X chromosome.
- In humans pseudoautosomal genes are located on both the X and Y chromosomes.

Sex Chromosomes and Sex Determination

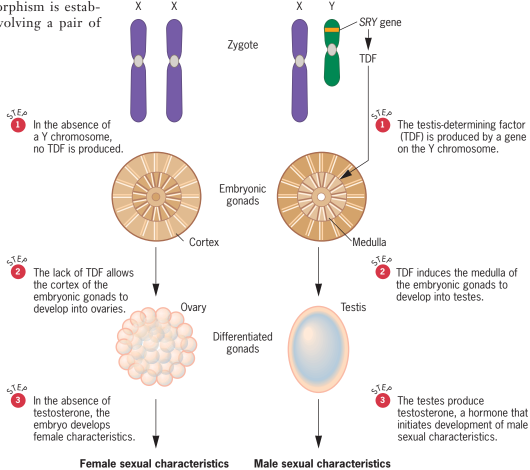
In some organisms, chromosomes—in particular, the sex chromosomes—determine male and female phenotypes. In the animal kingdom, sex is perhaps the most conspicuous phenotype. Animals with distinct males and females are sexually dimorphic. Sometimes this dimorphism is established by environmental factors. In one species of turtles, for example, sex is determined by

temperature. Eggs that have been incubated above 30°C hatch into females, whereas eggs that have been incubated at a lower temperature hatch into males.

In many other species, sexual dimorphism is established by genetic factors, often involving a pair of sex chromosomes.

SEX DETERMINATION IN HUMANS

The discovery that human females are XX and that human males are XY suggested that sex might be determined by the number of X chromosomes or by the presence or absence of a Y chromosome. As we now know, the second hypothesis is correct. In humans and other placental mammals, maleness is due to a dominant effect of the Y chromosome (**Figure 5.10**). The evidence for this fact comes from the study of individuals with an abnormal number of sex chromosomes. XO animals develop as females, and XXY animals develop as males. The dominant effect of the Y chromosome is manifested early in development, when it directs the primordial gonads to develop into testes. Once the testes have formed, they secrete testosterone, a hormone that stimu-

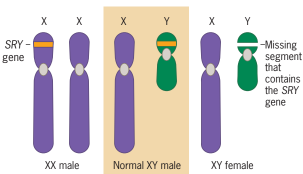


lates the development of male secondary sexual characteristics.

Researchers have shown that the **testis-determining factor (TDF)** is the product of a gene called *SRY* (for *sex-determining region Y*), which is located just outside the pseudoautosomal region in the short arm of the Y chromosome. The discovery of *SRY* was made possible by the identification of unusual individuals whose sex was inconsistent with their chromosome constitution—XX males and XY females (**Figure 5.11**). Some of the XX males were found to carry a small piece of the Y chromosome inserted into one of the X chromosomes. This piece evidently carried a gene responsible for maleness. Some of the XY females

were found to carry an incomplete Y chromosome. The part of the Y chromosome that was missing corresponded to the piece that was present in the XX males; its absence in the XY females apparently prevented them from developing testes. These complementary lines of evidence showed that a particular segment of the Y chromosome was needed for male development. Molecular analyses subsequently identified the *SRY* gene in this male-determining segment. Additional research has shown that an *SRY* gene is present on the Y chromosome of the mouse, and that like the human *SRY* gene—it triggers male development.

After the testes have formed, testosterone secretion initiates the development of male sexual characteristics. Testosterone is a hormone that binds to receptors in many kinds of cells. Once bound, the hormone–receptor complex transmits a signal to the nucleus, instructing the cell in how to differentiate. The concerted differentiation of many types of cells leads to the development of distinctly male characteristics such as heavy musculature, beard, and deep voice. If the testosterone signaling system fails, these characteristics do not appear and the individual develops as a female. One reason for failure is an inability to make the testosterone receptor ■ FIGURE 5.10 The process of sex determination in humans. Male sexual development depends on the production of the testisdetermining factor (TDF) by a gene on the Y chromosome. In the absence of this factor, the embryo develops as a female.



■ FIGURE 5.11 Evidence localizing the gene for the testis-determining factor (TDF) to the short arm of the Y chromosome in normal males. The TDF is the product of the *SRY* gene. In XX males, a small region containing this gene has been inserted into one of the X chromosomes, and in XY females, it has been deleted from the Y chromosome.

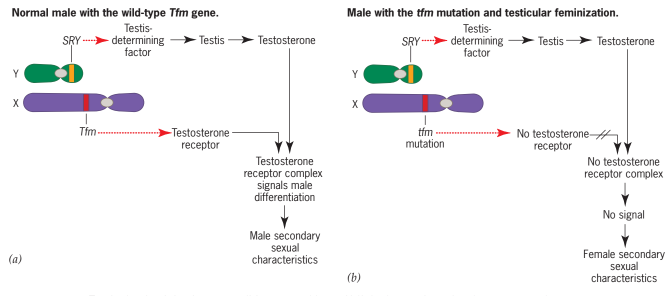


FIGURE 5.12 Testicular feminization, a condition caused by an X-linked mutation, *tfm*, that prevents the production of the testosterone receptor. (a) Normal male. (b) Feminized male with the *tfm* mutation.

(**Figure 5.12**). XY individuals with this biochemical deficiency initially develop as males—testes are formed and testosterone is produced. However, the testosterone has no effect because it cannot transmit the developmental signal inside its target cells. Individuals lacking the testosterone receptor therefore acquire female sexual characteristics. They do not, however, develop ovaries and are therefore sterile. This syndrome, called *testicular feminization*, results from a mutation in an X-linked gene, *Tfm*, which encodes the testosterone receptor. The *tfm* mutation is transmitted from mothers to their hemizygous XY offspring (who are phenotypically female) in a typical X-linked pattern.

SEX DETERMINATION IN DROSOPHILA

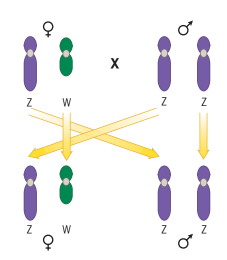
The Y chromosome in *Drosophila*—unlike that in humans—plays no role in sex determination. Instead, the sex of the fly is determined by the ratio of X chromosomes to autosomes. This mechanism was first demonstrated by Bridges in 1921 through an analysis of flies with unusual chromosome constitutions.

Normal diploid flies have a pair of sex chromosomes, either XX or XY, and three pairs of autosomes, usually denoted AA; here, each A represents one haploid set of autosomes. In complex experiments, Bridges contrived flies with abnormal numbers of chromosomes (Table 5.2). He observed that whenever the ratio of X's to A's was 1.0 or greater, the fly was female, and whenever it was 0.5 or less, the fly was male. Flies with an X:A ratio between 0.5 and 1.0 developed characteristics of both sexes; thus, Bridges called them *intersexes*. In none of these flies did the Y chromosome have any effect on the sexual phenotype. It was, however, required for male fertility.

SEX DETERMINATION IN OTHER ANIMALS

In both *Drosophila* and humans, males produce two kinds of gametes, X-bearing and Y-bearing. For this reason, they are referred to as the **heterogametic sex**; in these species females are the **homogametic sex**. In birds, butterflies, and some reptiles, this situation is reversed (**True 5.13**). Males are homogametic (usually denoted ZZ)

TABLE 5.2				
Ratio of X Chromosomes to Autosomes and the Corresponding Phenotype in Drosophila				
X Chromosomes (X) and Sets of Autosomes (A) X:A Ratio Phenotype				
1X 2A	0.5	Male		
2X 2A	1.0	Female		
3X 2A	1.5	Metafemale		
4X 3A	1.33	Metafemale		
4X 4A	1.0	Tetraploid female		
3X 3A	1.0	Triploid female		
3X 4A	0.75	Intersex		
2X 3A	0.67	Intersex		
2X 4A	0.5	Tetraploid male		
1X 3A	0.33	Metamale		

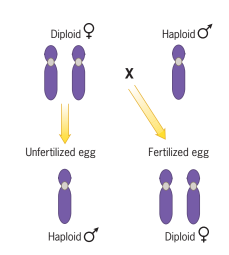


■ FIGURE 5.13 Sex determination in birds. The female is heterogametic (ZW), and the male is homogametic (ZZ). The sex of the offspring is determined by which of the sex chromosomes, Z or W, is transmitted by the female.

and females are heterogametic (ZW). However, little is known about the mechanism of sex determination in the Z–W sex chromosome system.

In honeybees, sex is determined by whether the animal is haploid or diploid (**Figure 5.14**). Diploid embryos, which develop from fertilized eggs, become females; haploid embryos, which develop from unfertilized eggs, become males. Whether or not a given female will mature into a reproductive form (queen) depends on how she was nourished as a larva. In this system, a queen can control the ratio of males to females by regulating the proportion of unfertilized eggs that she lays. Because this number is small, most of the progeny are female, albeit sterile, and serve as workers for the hive. In a haplo-diplo system of sex determination, eggs are produced through meiosis in the queen, and sperm are produced through mitosis in the male. This system ensures that fertilized eggs will have the haploid number.

Some wasps also have a haplo-diplo method of sex determination. In these species diploid males are sometimes produced, but they are always sterile. Detailed genetic analysis in one species, *Bracon hebetor*; has indicated that the diploid males are homo-zygous for a sex-determining locus, called X; diploid females are always heterozygous for this locus. Evidently, the sex locus in *Bracon* has many alleles; crosses between unrelated males and females therefore almost always produce heterozygous diploid females. However, when the mates are related, there is an appreciable chance that their offspring will be homozygous for the sex locus, in which case they develop into sterile males.



■ FIGURE 5.14 Sex determination in honeybees. Females, which are derived from fertilized eggs, are diploid, and males, which are derived from unfertilized eggs, are haploid.

- In humans sex is determined by a dominant effect of the SRY gene on the Y chromosome; the
 product of this gene, the testis-determining factor (TDF), causes a human embryo to develop
 into a male.
- In Drosophila, sex is determined by the ratio of X chromosomes to sets of autosomes (X:A); for $X:A \le 0.5$, the fly develops as a male, for $X:A \ge 1.0$, it develops as a female, and for 0.5 < X: A < 1.0, it develops as an intersex.
- In honeybees, sex is determined by the number of chromosome sets; haploid embryos develop into males and diploid embryos develop into females.

KEY POINTS

Dosage Compensation of X-Linked Genes

Different mechanisms adjust for the unequal dosage of X-linked genes in male and female animals.

Animal development is usually sensitive to an imbalance in the number of genes. Normally, each gene is present in two copies. Departures from this condition, either up or down, can cause abnormal phenotypes, and sometimes even death. It is therefore puzzling that so many species should have a sex-determination system based

on females with two X chromosomes and males with only one. In these species, how is the numerical difference of X-linked genes accommodated? *A priori*, three mechanisms may compensate for this difference: (1) each X-linked gene could work twice as hard in males as it does in females, or (2) one copy of each X-linked gene could be inactivated in females, or (3) each X-linked gene could work half as hard in females as it does in males. Extensive research has shown that all three mechanisms are utilized, the first in *Drosophila*, the second in mammals, and the third in the nematode *Caenorhabditis elegans*. These mechanisms are discussed in detail in Chapter 19; here we provide brief descriptions of the dosage compensations systems in *Drosophila* and mammals.

HYPERACTIVATION OF X-LINKED GENES IN MALE DROSOPHILA

In *Drosophila*, dosage compensation of X-linked genes is achieved by an increase in the activity of these genes in males. This phenomenon, called *hyperactivation*, involves a complex of different proteins that binds to many sites on the X chromosome in males and triggers a doubling of gene activity (see Chapter 19). When this protein complex does not bind, as is the case in females, hyperactivation of X-linked genes does not occur. In this way, total X-linked gene activity in males and females is approximately equalized.

INACTIVATION OF X-LINKED GENES IN FEMALE MAMMALS

In placental mammals, dosage compensation of X-linked genes is achieved by the inactivation of one of the female's X chromosomes. This mechanism was first proposed in 1961 by the British geneticist Mary Lyon, who inferred it from studies on mice. Subsequent research by Lyon and others has shown that the inactivation event occurs when the mouse embryo consists of a few thousand cells. At this time, each cell makes an independent decision to silence one of its X chromosomes. The chromosome to be inactivated is chosen at random; once chosen, however, it remains inactivated in all the descendants of that cell. Thus, female mammals are genetic mosaics containing two types of cell lineages; the maternally inherited X chromosome is inactivated in roughly half of these cells, and the paternally inherited X is inactivated in the other half. A female that is heterozygous for an X-linked gene is therefore able to show two different phenotypes. One of the best examples of this phenotypic mosaicism comes from the study of fur coloration in cats and mice (**Figure 5.15**). In both of these species, the X chromosome carries a gene for pigmentation of the fur. Females heterozygous for different alleles of this gene show patches of light and dark fur. The light patches express one allele, and the dark patches express the other. In cats, where one allele produces black pigment and the other produces orange pigment, this patchy phenotype is called tortoiseshell. Each patch of fur defines a clone of pigment-producing cells, or melanocytes, that were derived by mitosis from a precursor cell present at the time of X chromosome inactivation.

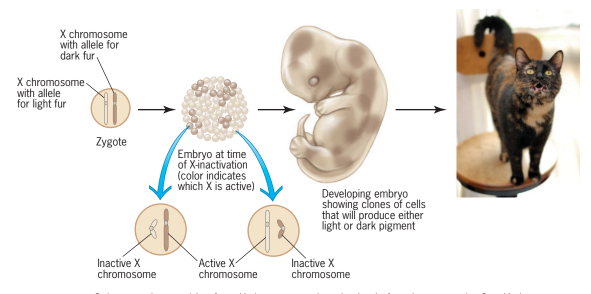


FIGURE 5.15 Color mosaics resulting from X chromosome inactivation in female mammals. One X chromosome in the zygote carries the allele for dark fur color, and the other X chromosome carries the allele for light fur color. In each cell of the early embryo, one of the two X chromosomes is randomly inactivated. Whichever X chromosome is chosen remains inactive in all the descendants of that cell. Thus, the developing embryo comes to consist of clones of cells that express only one of the fur color alleles. This genetic mosaicism produces the patches of light and dark fur that are characteristic of tortoiseshell cats.

An X chromosome that has been inactivated does not look or act like other chromosomes. Chemical analyses show that its DNA is modified by the addition of numerous methyl groups. In addition, it condenses into a darkly staining structure called a **Barr body** (**■** Figure 5.16), after the Canadian geneticist Murray Barr, who first observed it. This structure becomes attached to the inner surface of the nuclear membrane, where it replicates out of step with the other chromosomes in the cell. The inactivated X chromosome remains in this altered state in all the somatic tissues. However, in the germ tissues it is reactivated, perhaps because two copies of some X-linked genes are needed for the successful completion of oogenesis. The molecular mechanism of X-inactivation is discussed in Chapter 19.

Cytological studies have identified humans with more than two X chromosomes (see Chapter 6). For the most part, these people are phenotypically normal females, apparently because all but one of their X chromosomes is inactivated.

Often all the inactivated X's congeal into a single Barr body. These observations suggest that cells may have a limited amount of some factor needed to prevent X-inactivation. Once this factor has been used to keep one X chromosome active, all the others quietly succumb to the inactivation process.

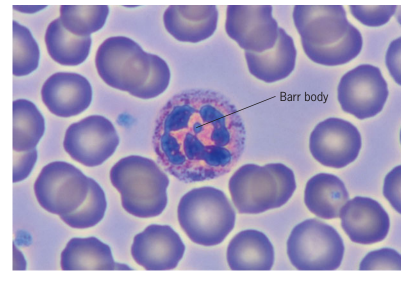


FIGURE 5.16 Barr body in a human female cell.

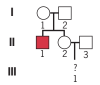
- In Drosophila, dosage compensation for X-linked genes is achieved by hyperactivating the single X chromosome in males.
- In mammals, dosage compensation for X-linked genes is achieved by inactivating one of the two X chromosomes in females.

KEY POINTS

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. A mutant *Drosophila* male with prune-colored eyes was crossed to a wild-type female with red eyes. All the F_1 off-spring of both sexes had red eyes. When these offspring were intercrossed, three different classes of F_2 flies were produced: females with red eyes, males with red eyes, and males with prune eyes. The males and females were equally frequent in the F_2 , and among the males, the two eye color classes were equally frequent. Do these results suggest that the *prune* mutation is on the X chromosome?
- Answer: The results of these crosses are consistent with the hypothesis that the *prune* mutation is on the X chromosome. According to this hypothesis, the male in the first cross must have been hemizygous for the *prune* mutation; his mate must have been homozygous for the wild-type allele of the *prune* gene. Among the F_1 , the daughters must have been heterozygous for the mutation and the wildtype allele, and the sons must have been hemizygous for the wild-type allele. When the F_1 flies were intercrossed, they produced daughters that inherited the wild-type allele from their fathers-these flies must therefore have had red eves—and they produced sons that inherited either the mutant allele or the wild-type allele from their mothers, with each of these possibilities being equally likely. Thus, according to the hypothesis, among the F₂, all the daughters and half the sons should have red eyes, and half the sons should have prune eyes, which is what was observed.
- 2. The following pedigree shows the inheritance of hemophilia in a human family. (a) What is the probability that II-2 is a carrier of the allele for hemophilia? (b) What is the probability that III-1 will be affected with hemophilia?

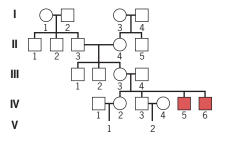


- Answer: (a) II-2 has an affected brother, which indicates that her mother was a carrier. Her chance of also being a carrier is therefore simply the probability that her mother transmitted the mutant allele to her, which is 1/2. (b) The chance that III-1 will be affected depends on three events: (1) that II-2 is a carrier, (2) that II-2 transmits the mutant allele, if she carries it, and (3) that II-3 transmits a Y chromosome. Each of these events is associated with a probability of 1/2. Thus, the probability that III-1 will be affected is $(1/2) \times$ $(1/2) \times (1/2) = 1/8$.
- **3.** How do the chromosomal mechanisms of sex determination differ between humans and *Drosophila*?
- **Answer:** In humans, sex is determined by a dominant effect of the Y chromosome. In the absence of a Y chromosome, the individual develops as a female; in its presence, the individual develops as a male. In *Drosophila*, sex is determined by the ratio of X chromosomes to sets of autosomes. When the X:A ratio is greater than or equal to one, the individual develops as a female; when the X:A ratio is less than or equal to 0.5, it develops as a male; in between these limits, the individual develops as an intersex.
- 4. How do the mechanisms that compensate for different doses of the X chromosome in the two sexes differ between humans and *Drosophila*?
- **Answer:** In humans, one of the two X chromosomes in an XX female is inactivated in the somatic cells early in development. In *Drosophila*, the single X chromosome in a male is hyperactivated so that its genes are as active as the double dose of X-linked genes present in an XX female.

Testing Your Knowledge

Integrate Different Concepts and Techniques

1. The Lesch-Nyhan syndrome is a serious metabolic disorder affecting about one in 50,000 males in the population of the United States. A class of molecules called purines, which are biochemical precursors of DNA, accumulate in the nervous tissues and joints of people with the Lesch-Nyhan syndrome. This biochemical abnormality is caused by a deficiency for the enzyme hypoxanthine phosphoribosyltransferase (HPRT), which is encoded by a gene located on the X chromosome. Individuals deficient for this enzyme are unable to control their movements and unwillingly engage in self-destructive behavior such as biting and scratching themselves. The males labeled IV-5 and IV-6 in the following pedigree have the Lesch-Nyhan syndrome. What are the risks that V-1 and V-2 will inherit this disorder?



- **Answer:** We know that III-3 must be a heterozygous carrier of the mutant allele (*b*) because two of her sons are affected. However, because she herself does not show the mutant phenotype, we know that her other X chromosome must carry the wild-type allele (H). Given that III-3 is genotypically *Hb*, there is a one-half chance that she passed the mutant allele to her daughter (IV-2). If she did, there is a one-half chance that IV-2 will transmit this allele to her child (V-1), and there is a one-half chance that this child will be a male. Thus, the risk that V-1 will have the Lesch-Nyhan syndrome is $(1/2) \times (1/2) \times (1/2) =$ 1/8. For V-2, the risk of inheriting the Lesch-Nyhan syndrome is essentially zero. This child's father (IV-3) does not have the mutant allele, and even if he did, he would not transmit it to a son. The child's mother comes from outside the family and is very unlikely to be a carrier because the trait is rare in the general population. Thus, V-2 has virtually no chance of suffering from the Lesch-Nyhan syndrome.
- 2. A geneticist crossed *Drosophila* females that had white eyes and ebony bodies to wild-type males, which had red eyes and gray bodies. Among the F_1 , all the daughters had red eyes and gray bodies, and all the sons had white eyes and gray bodies. These flies were intercrossed to produce F_2 progeny, which were classified for eye and body color and then counted. Among 384 total progeny, the geneticist obtained the following results:

Eye Color	Body Color	Males	Females	
white	ebony	20	21	
white	gray	70	73	
red	ebony	28	25	
red	gray	76	71	

How would you explain the inheritance of eye color and body color?

Answer: The results in the F_1 tell us that both mutant phenotypes are caused by recessive alleles. Furthermore, because the males and females have different eye color phenotypes, we know that the eye color gene is X-linked and that the body color gene is autosomal. In the F_2 , the two genes assort independently, as we would expect for genes located on different chromosomes. In the following table, we show the genotypes of the different classes of flies in this experiment, using w for the *white* mutation and e for the *ebony* mutation; the wild-type alleles are denoted by plus signs. Following the convention of *Drosophila* geneticists, we write the sex chromosomes (X and Y) on the left and the autosomes on the right. A question mark in a genotype indicates that either the wild-type or mutant alleles could be present.

Phen	otypes	Genotypes		
Eye Color	Body Color	Males	Females	
white	ebony	w/Y e/e	w/w e/e	
white	gray	w/Y +/?	w/w +/?	
red	ebony	+/Y e/e	+/w e/e	
red	gray	+/Y +/?	+/w +/?	

- 3. In 1906, the British biologists L. Doncaster and G. H. Raynor reported the results of breeding experiments with the currant moth, Abraxas. This moth exists in two color forms in Great Britain. One, called grossulariata, has large black spots on its wings; the other, called lacticolor, has much smaller black spots. Doncaster and Raynor crossed lacticolor females with grossulariata males and found that all the F₁ progeny were grossulariata. They then intercrossed the F_1 moths to produce an F_2 , which consisted of two types of females (grossulariata and lacticolor) and one type of males (grossulariata). Doncaster and Raynor also testcrossed the F_1 moths. Grossulariata F_1 females crossed to lacticolor males produced lacticolor females and grossulariata males-the first grossulariata males ever seen; and grossulariata F1 males crossed to lacticolor females produced four kinds of offspring: grossulariata males, grossulariata females, lacticolor males, and lacticolor females. Propose an explanation for the results of these experiments.
- Answer: The inheritance of the grossulariata and lacticolor phenotypes is obviously linked to sex. In moths, however, females are heterogametic (ZW) and males are homogametic (ZZ). Thus, we can hypothesize that lacticolor females are hemizygous for a recessive allele (l) on the Z chromosome and that grossulariata males are homozygous for a dominant allele (L) on this chromosome. When the two types of moths are crossed, they produce grossulariata females that are hemizygous for the dominant allele (L) and grossulariata males that are heterozygous for the two alleles (Ll). An intercross between these F_1 moths produces grossulariata (L) and lacticolor (l) females, each hemizygous for a different allele, and grossulariata males that are either homozygous LL or heterozygous Ll. The hypothesis that the spotting pattern in Abraxas is controlled by a gene on the Z chromosome also explains the results of the testcrosses with the F₁ grossulariata animals. Grossulariata F₁ females, which are hemizygous for the dominant allele L, when crossed to homozygous lllacticolor males produce hemizygous l lacticolor females and heterozygous Ll grossulariata males. Grossulariata F₁ males, which are Ll heterozygotes, when crossed to hemizygous l lacticolor females produce heterozygous Ll grossulariata males, hemizygous L grossulariata females, homozygous *ll* lacticolor males, and hemizygous *l* lacticolor females. Unfortunately, at the time Doncaster and Raynor reported their work, the sex chromosome constitution of Abraxas was not known. Consequently, they did not make the conceptual link between the inheritance of wing spotting and transmission of the sex chromosomes. Had they done so, T. H. Morgan's demonstration of sex linkage in Drosophila might today appear to have been an afterthought.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **5.1** What are the genetic differences between male- and female-determining sperm in animals with heterogametic males?
- **5.2** A male with singed bristles appeared in a culture of *Drosophila*. How would you determine if this unusual phenotype was due to an X-linked mutation?
- **5.3** In grasshoppers, rosy body color is caused by a recessive mutation; the wild-type body color is green. If the gene for body color is on the X chromosome, what kind of progeny would be obtained from a mating between a homozygous rosy female and a hemizygous wild-type male? (In grasshoppers, females are XX and males are XO.)
- **5.4** In the mosquito *Anopheles culicifacies*, *golden* body (*go*) is a recessive X-linked mutation, and *brown* eyes (*bw*) is a recessive autosomal mutation. A homozygous XX female with golden body is mated to a homozygous XY male with brown eyes. Predict the phenotypes of their F_1 offspring. If the F_1 progeny are intercrossed, what kinds of progeny will appear in the F_2 , and in what proportions?
- **5.5** What are the sexual phenotypes of the following genotypes in *Drosophila*: XX, XY, XXY, XXX, XO?
- **5.6** In humans, a recessive X-linked mutation, g, causes greendefective color vision; the wild-type allele, G, causes normal color vision. A man (a) and a woman (b), both with normal vision, have three children, all married to people with normal vision: a color-defective son (c), who has a daughter with normal vision (f); a daughter with normal vision (d), who has one color-defective son (g) and two normal sons (h); and a daughter with normal vision (e), who has six normal sons (i). Give the most likely genotypes for the individuals (a to i) in this family.
- **5.7** If a father and son both have defective color vision, is it likely that the son inherited the trait from his father?
- **5.8** A normal woman, whose father had hemophilia, marries a normal man. What is the chance that their first child will have hemophilia?
- **5.9** A man with X-linked color blindness marries a woman with no history of color blindness in her family. The daughter of this couple marries a normal man, and their daughter also marries a normal man. What is the chance that this last couple will have a child with color blindness? If this couple has already had a child with color blindness, what is the chance that their next child will be color blind?
- **5.10** A man who has color blindness and type O blood has children with a woman who has normal color vision and type AB blood. The woman's father had color blindness. Color blindness is determined by an X-linked gene, and blood type is determined by an autosomal gene.

- (a) What are the genotypes of the man and the woman?
- (b) What proportion of their children will have color blindness and type B blood?
- (c) What proportion of their children will have color blindness and type A blood?
- (d) What proportion of their children will be color blind and have type AB blood?
- **5.11** A *Drosophila* female homozygous for a recessive X-linked mutation that causes vermilion eyes is mated to a wild-type male with red eyes. Among their progeny, all the sons have vermilion eyes, and nearly all the daughters have red eyes; however, a few daughters have vermilion eyes. Explain the origin of these vermilion-eyed daughters.
- **5.12 ••** In *Drosophila*, vermilion eye color is due to a recessive allele (v) located on the X chromosome. Curved wings is due to a recessive allele (cu) located on one autosome, and ebony body is due to a recessive allele (e) located on another autosome. A vermilion male is mated to a curved, ebony female, and the F₁ males are phenotypically wild-type. If these males were backcrossed to curved, ebony females, what proportion of the F₂ offspring will be wild-type males?
- **5.13** A *Drosophila* female heterozygous for the recessive X-linked mutation w (for *white* eyes) and its wild-type allele w^+ is mated to a wild-type male with red eyes. Among the sons, half have white eyes and half have red eyes. Among the daughters, nearly all have red eyes; however, a few have white eyes. Explain the origin of these white-eyed daughters.
- **5.14** \bigcirc In *Drosophila*, a recessive mutation called *chocolate* (c) causes the eyes to be darkly pigmented. The mutant phenotype is indistinguishable from that of an autosomal recessive mutation called *brown* (*bw*). A cross of chocolate-eyed females to homozygous brown males yielded wild-type F₁ females and darkly pigmented F₁ males. If the F₁ flies are intercrossed, what types of progeny are expected, and in what proportions? (Assume the double mutant combination has the same phenotype as either of the single mutants alone.)
- **5.15** Suppose that a mutation occurred in the *SRY* gene on the human Y chromosome, knocking out its ability to produce the testis-determining factor. Predict the phenotype of an individual who carried this mutation and a normal X chromosome.
- **5.16** A woman carries the testicular feminization mutation (*tfm*) on one of her X chromosomes; the other X carries the wild-type allele (*Tfm*). If the woman marries a normal man, what fraction of her children will be phenotypically female? Of these, what fraction will be fertile?
- **5.17** Would a human with two X chromosomes and a Y chromosome be male or female?
- **5.18 (processive alb) (c) (**

is located on the X chromosome and on a homologous segment of the Y chromosome. Give the genotypes and phenotypes of the offspring from the following crosses:

- **5.19** Predict the sex of *Drosophila* with the following chromosome compositions (A = haploid set of autosomes):
 - (a) 4X 4A
 - (b) 3X 4A
 - (c) 2X 3A
 - (d) 1X 3A
 - (e) 2X 2A
 - (f) 1X 2A
- **5.20** In chickens, the absence of barred feathers is due to a recessive allele. A barred rooster was mated with a nonbarred hen, and all the offspring were barred. These F_1 chickens were intercrossed to produce F_2 progeny, among which all the males were barred; half the females were barred and half were nonbarred. Are these results consistent with the hypothesis that the gene for barred feathers is located on one of the sex chromosomes?
- **5.21** A *Drosophila* male carrying a recessive X-linked mutation for yellow body is mated to a homozygous wild-type female with gray body. The daughters of this mating all have uniformly gray bodies. Why aren't their bodies a mosaic of yellow and gray patches?
- **5.22** What is the maximum number of Barr bodies in the nuclei of human cells with the following chromosome compositions:
 - (a) XY
 - (b) XX
 - (c) XXY
 - (d) XXX
 - (e) XXXX
 - (f) XYY
- **5.23** Males in a certain species of deer have two nonhomologous X chromosomes, denoted X₁ and X₂, and a Y chromosome. Each X chromosome is about half as large as the Y

chromosome, and its centromere is located near one of the ends; the centromere of the Y chromosome is located in the middle. Females in this species have two copies of each of the X chromosomes and lack a Y chromosome. How would you predict the X and Y chromosomes to pair and disjoin during spermatogenesis to produce equal numbers of male- and female-determining sperm?

5.24 • A breeder of sun conures (a type of bird) has obtained two true-breeding strains, A and B, which have red eyes instead of the normal brown found in natural populations. In Cross 1, a male from strain A was mated to a female from strain B, and the male and female offspring all had brown eyes. In Cross 2, a female from strain A was mated to a male from strain B, and the male offspring had brown eyes and the female offspring had red eyes. When the F₁ birds from each cross were mated brother to sister, the breeder obtained the following results:

Phenotype	Proportion in F ₂ of Cross 1	Proportion in F ₂ of Cross 2
Brown male	6/16	3/16
Red male	2/16	5/16
Brown female	3/16	3/16
Red female	5/16	5/16

Provide a genetic explanation for these results.

5.25 In 1908 F. M. Durham and D. C. E. Marryat reported the results of breeding experiments with canaries. Cinnamon canaries have pink eyes when they first hatch, whereas green canaries have black eyes. Durham and Marryat crossed cinnamon females with green males and observed that all the F_1 progeny had black eyes, just like those of the green strain. When the F_1 males were crossed to green females, all the male progeny had black eyes, whereas all the female progeny had either black or pink eyes, in about equal proportions. When the F_1 males were crossed to cinnamon females, four classes of progeny were obtained: females with black eyes, females with pink eyes, males with black eyes, and males with pink eyes—all in approximately equal proportions. Propose an explanation for these findings.

Genomics on the Web at http://www.ncbi.nlm.nih.gov

Both humans and mice have X and Y sex chromosomes. In each species the Y is smaller than the X and has fewer genes.

- 1. What are the sizes of the human X and Y chromosomes in nucleotide pairs? How many genes does each of these chromosomes contain?
- **2.** How do the sizes of the mouse sex chromosomes compare with those of the human sex chromosomes?
- **3.** The *SRY* gene responsible for sex determination in humans is located in the short arm of the Y chromosome, near but not

in the pseudoautosomal region. Can you find its homologue, *Sry*, on the Y chromosome of the mouse?

Hint: At the web site, click on Genomes and Maps, and then under Quick Links, access the Map Viewer feature. Click on the species whose genome you want to see, and then click on one of the sex chromosomes. Use the Search function to find the *Sry* gene on the mouse's Y chromosome.



Variation in Chromosome Number and Structure

CHAPTER OUTLINE

- Cytological Techniques
- Polyploidy
- Aneuploidy
- Rearrangements of Chromosome Structure

Chromosomes, Agriculture, and Civilization

The cultivation of wheat originated some 10,000 years ago in the Middle East. Today, wheat is the principal food crop for more than a billion people. It is grown in diverse environments, from Norway to Argentina. More than 17,000 varieties have been developed, each adapted to a different locality. The total wheat production of the world is 60 million metric tons annually, accounting for more than 20 percent of the food calories consumed by the entire human population. Wheat is clearly an important agricultural crop and, some would argue, a mainstay of civilization.

Modern cultivated wheat, *Triticum aestivum*, is a hybrid of at least three different species. Its progenitors were low-yielding grasses that grew in Syria, Iran, Iraq, and Turkey. Some of these grasses appear to have been cultivated by the ancient peoples of this region. Although we do not know the exact course of events, two of the grasses apparently interbred, producing a species that excelled as a crop plant. Through human cultivation, this hybrid species was selectively improved, and then it, too, interbred with a third species, yielding a triple-hybrid that was even better suited for agriculture. Modern wheat is descended from these triple-hybrid plants.

What made the triple-hybrid wheats so superior to their ancestors? They had larger grains, they were more easily

harvested, and they grew in a wider range of conditions. We now understand the chromosomal basis for these improvements. Triple-hybrid wheat contains the chromosomes of each of its progenitors. Genetically, it is an amalgamation of the genomes of three different species.



Wheat field.

Cytological Techniques

Geneticists study chromosome number and structure by staining dividing cells with certain dyes and then examining them with a microscope. The analysis of

Geneticists use stains to identify specific chromosomes and to analyze their structures.

stained chromosomes is the main activity of the discipline called cytogenetics.

Cytogenetics had its roots in the research of several nineteenth-century European biologists who discovered chromosomes and observed their behavior during mitosis, meiosis, and fertilization. This research blossomed during the twentieth century, as microscopes improved and better procedures for preparing and staining chromosomes were developed. The demonstration that genes reside on chromosomes boosted interest in this research and led to important studies on chromosome number and structure. Today, cytogenetics has significant applied aspects, especially in medicine, where it is used to determine whether disease conditions are associated with chromosome abnormalities.

ANALYSIS OF MITOTIC CHROMOSOMES

Researchers perform most cytological analyses on dividing cells, usually cells in the middle of mitosis. To enrich for cells at this stage, they have traditionally used rapidly growing material such as animal embryos and plant root tips. However, the development of cell-culturing techniques has made it possible to study chromosomes in other types of cells (**Figure 6.1**). For example, human white blood cells can be collected from peripheral blood, separated from the nondividing red blood cells, and put into culture. The white cells are then stimulated to divide by chemical treatment, and midway through division a sample of the cells is prepared for cytological analysis. The usual procedure is to treat the dividing cells with a chemical that disables the mitotic spindle. The effect of this interference is to trap the chromosomes in mitosis, when they are most easily seen. Mitotically arrested cells are then swollen by immersion in a hypotonic solution that causes the cells to take up water by osmosis. The contents of each cell are diluted by the additional water, so that when the cells are squashed on a microscope slide, the chromosomes are spread out in an uncluttered fashion. This technique greatly facilitates subsequent analysis, especially if the chromosome number is large. For many years it was erroneously thought that human cells contained 48 chromosomes. The correct number, 46, was determined only after the swelling technique was used to separate the chromosomes within individual mitotic cells. For more details, see A Milestone in Genetics: Tjio and Levan Count Human Chromosomes Correctly in the Student Companion site.

Until the late 1960s and early 1970s, chromosome spreads were usually stained with Feulgen's reagent, a purple dye that reacts with the sugar molecules in DNA, or with aceto-carmine, a deep red dye. Because these types of dyes stain the chromosomes

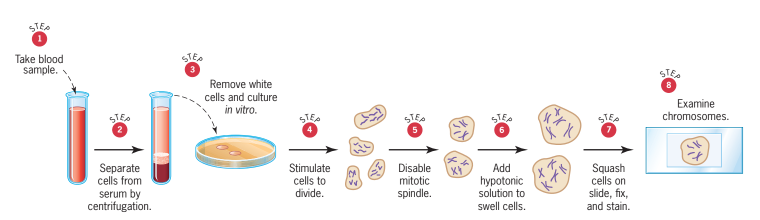


FIGURE 6.1 Preparation of cells for cytological analysis.

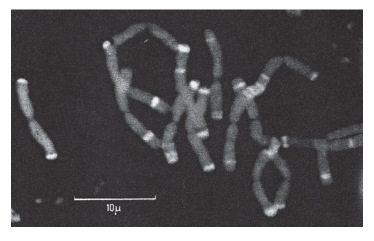


FIGURE 6.2 Metaphase chromosomes of the plant *Allium carinatum*, stained with quinacrine.

uniformly, they do not allow a researcher to distinguish one chromosome from another unless the chromosomes are very different in size or in the positions of their centromeres. Today, cytogeneticists use dyes that stain chromosomes differentially along their lengths. *Quinacrine*, a chemical relative of the antimalarial drug quinine, was one of the first of these more discriminating reagents. Chromosomes that have been stained with quinacrine show a characteristic pattern of bright bands on a darker background. However, because quinacrine is a fluorescent compound, the bands appear only when the chromosomes are exposed to ultraviolet (UV) light. Ultraviolet irradiation causes some of the quinacrine molecules that have inserted into the chromosome to emit energy. Parts of the chromosome shine brightly, whereas other parts remain dark. This bright-dark banding pattern is highly reproducible and is also specific for each chromosome (**■** Figure 6.2). Thus with quinacrine banding, cytogeneticists can identify particular chromosomes in a cell, and they can also determine if a chromosome is structurally abnormal—for example, if it is missing certain bands.

Excellent nonfluorescent staining techniques have also been developed. The most popular of these uses *Giemsa* stain, a mixture of dyes named after its inventor, Gustav Giemsa. Like quinacrine, Giemsa creates a reproducible pattern of bands on each chromosome (**Figure 6.3**). It is still not clear why chromosomes show bands when they are stained with quinacrine or Giemsa. It may be that these types of dyes react preferentially with certain DNA sequences, or with the proteins associated with them, and that these target DNA sequences are distributed in a characteristic way within each chromosome.

The most advanced technique used by cytogeneticists today is called **chromosome painting.** With this technique, colorful chromosome images are created by treating chromosome spreads with fluorescently labeled DNA fragments that have been isolated and characterized in the laboratory. Such a fragment may, for instance, come from a particular gene. The DNA fragment is chemically labeled with a fluorescent dye in the laboratory and then applied to chromosomes that have been spread on a glass slide. Under the right conditions, the DNA fragment will bind to chromosomal DNA that is complementary to it in sequence. This binding, in effect, labels the chromosomal DNA with the fluorescent dye that is present in the DNA fragment.

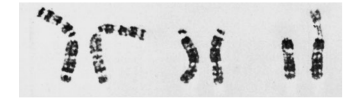
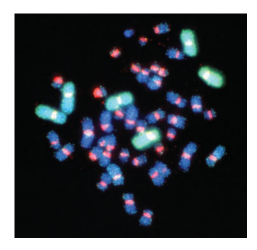


FIGURE 6.3 Metaphase chromosomes of the deerlike Asian muntjak stained with Giemsa.

Because of the specific nature of the interaction between the DNA fragment and the complementary DNA in the chromosomes, we often call the DNA fragment a *probe*. It seeks out and binds to its complement in the large mass of chromosomal DNA in a cell. After the probe has bound, the chromosome spreads are irradiated with light of an appropriate wavelength. The resulting bands or dots of color reveal where the complementary DNA sequence—the target of the probe—is located in the chromosomes. **■** Figure 6.4 shows human chromosomes that have been analyzed with this technique. The chromosomes were simultaneously painted with two different human DNA fragments, each labeled with a dye that fluoresces a different color. One of the fragments binds nonspecifically to the centromeres of each of the chromosomes, and when stimulated to fluoresce, it appears pink. The other fragment binds only to a few of the chromosomes, and when stimulated to fluoresce, it appears bright green. These few chromosomes therefore stand out among all the chromosomes in the spread. Figure 2.7 shows human chromosomes that have been painted with a panel of probes made from human DNA fragments. Each of the pairs of chromosomes has a characteristic pattern of bands. Thus, each pair can be uniquely identified using this technique.

THE HUMAN KARYOTYPE

Diploid human cells contain 46 chromosomes—44 autosomes and two sex chromosomes, which are XX in females and XY in males. At mitotic metaphase, each of the 46 chromosomes consists of two identical sister chromatids. When stained appropriately, each of the duplicated chromosomes can be recognized by its size, shape, and banding pattern. For cytological analysis, well-stained metaphase spreads are photographed, and then each of the chromosome images is cut out of the picture, matched with its partner to form homologous pairs, and arranged from largest to smallest on a chart (**■ Figure 6.5**). The largest autosome is number 1, and the smallest is number 21. (For historical reasons, the second smallest chromosome has been designated number 22.) The X chromosome is intermediate in size, and the Y chromosome is about the same size as chromosome 22. This chart of chromosome cutouts is called a **karyotype** (from the Greek word meaning "kernel," a reference to the contents of the nucleus). A skilled researcher can use a karyotype to identify abnormalities in chromosome number and structure.



■ FIGURE 6.4 Chromosome painting. Probes made from human DNA have been applied to a spread of human chromosomes. Each probe has been labeled with a dye that fluoresces a different color (pink or bright green) to reveal where DNA sequences complementary to these probes are located in the chromosomes. The pink probe targets DNA in the centromere of every chromosome, whereas the bright green probe targets DNA in only three pairs of chromosomes.

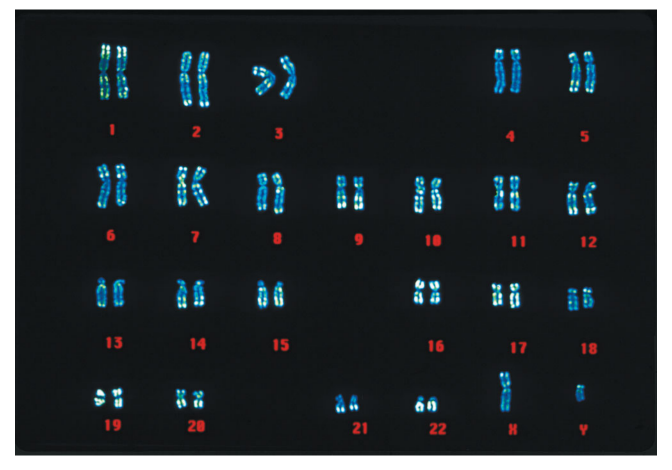
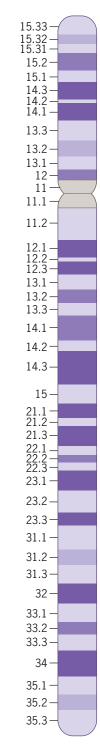


FIGURE 6.5 The karyotype of a human male stained to reveal bands on each of the chromosomes. The autosomes are numbered from 1 to 22. The X and Y are the sex chromosomes.



■ FIGURE 6.6 The ideogram of human chromosome 5. Regions within each arm are numbered consecutively starting at the centromere. Bands within each region are denoted by numbers following a decimal point.

KEY POINTS

Before the banding and painting techniques were available, it was difficult to distinguish one human chromosome from another. Cytogeneticists could only arrange the chromosomes into groups according to size, classifying the largest as group A, the next largest as group B, and so forth. Although they could recognize seven different groups, within these groups it was nearly impossible to identify a particular chromosome. Today—as a result of the banding and painting techniques—we can routinely identify each of the chromosomes. The banding and painting techniques also make it possible to distinguish each arm of a chromosome and to investigate specific regions within them. The centromere divides each chromosome into long and short arms. The short arm is denoted by the letter p (from the French word *petite*, meaning "small") and the long arm by the letter q (because it follows "p" in the alphabet). Thus, for example, a cytogeneticist can refer specifically to the short arm of chromosome 5 simply by writing "5p." Within each arm, specific regions are denoted by numbers, starting at the centromere (**Figure 6.6**). Thus, in the short arm of chromosome 5, we have region 5p11, which is closest to the centromere, followed by regions 5p12, 5p13, 5p14, and 5p15, which is farthest from the centromere. Within each region, individual bands are denoted by numbers following a decimal point; for example, 13.1, 13.2, and 13.3 refer to the three bands that make up region 5p13. The pattern of bands within the chromosome is called an *ideogram*.

CYTOGENETIC VARIATION: AN OVERVIEW

The phenotypes of many organisms are affected by changes in the number of chromosomes in their cells; sometimes even changes in part of a chromosome can be significant. These numerical changes are usually described as variations in the *ploidy* of the organism (from the Greek word meaning "fold," as in "twofold"). Organisms with complete, or normal, sets of chromosomes are said to be euploid (from the Greek words meaning "good" and "fold"). Organisms that carry extra sets of chromosomes are said to be *polyploid* (from the Greek words meaning "many" and "fold"), and the level of polyploidy is described by referring to a basic chromosome number, usually denoted n. Thus, diploids, with two basic chromosome sets, have 2n chromosomes; triploids, with three sets, have 3n; tetraploids, with four sets, have 4n; and so forth. Organisms in which a particular chromosome, or chromosome segment, is under- or overrepresented are said to be aneuploid (from the Greek words meaning "not," "good," and "fold"). These organisms therefore suffer from a specific genetic imbalance. The distinction between aneuploidy and polyploidy is that aneuploidy refers to a numerical change in part of the genome, usually just a single chromosome, whereas polyploidy refers to a numerical change in a whole set of chromosomes. Aneuploidy implies a genetic imbalance, but polyploidy does not.

Cytogeneticists have also catalogued various types of structural changes in the chromosomes of organisms. For example, a piece of one chromosome may be fused to another chromosome, or a segment within a chromosome may be inverted with respect to the rest of that chromosome. These structural changes are called *rearrangements*. Because some rearrangements segregate irregularly during meiosis, they can be associated with aneuploidy. In the sections that follow, we consider all these cytogenetic variations—polyploidy, aneuploidy, and chromosome rearrangements.

- Cytogenetic analysis usually focuses on chromosomes in dividing cells.
- Dyes such as quinacrine and Giemsa create banding patterns that are useful in identifying individual chromosomes within a cell.
- A karyotype shows the duplicated chromosomes of a cell arranged for cytogenetic analysis.

q

Polyploidy

Polyploidy, the presence of extra chromosome sets, is fairly common in plants but very rare in animals. One-half of all known plant genera contain poly-

Extra sets of chromosomes in an organism can affect, the organism's appearance and fertility.

ploid species, and about two-thirds of all grasses are polyploids. Many of these species reproduce asexually. In animals, where reproduction is primarily by sexual means, polyploidy is rare, probably because it interferes with the sex-determination mechanism.

One general effect of polyploidy is that cell size is increased, presumably because there are more chromosomes in the nucleus. Often this increase in size is correlated with an overall increase in the size of the organism. Polyploid species tend to be larger and more robust than their diploid counterparts. These characteristics have a practical significance for humans, who depend on many polyploid plant species for food. These species tend to produce larger seeds and fruits, and therefore provide greater yields in agriculture. Wheat, coffee, potatoes, bananas, strawberries, and cotton are all polyploid crop plants. Many ornamental garden plants, including roses, chrysanthemums, and tulips, are also polyploid (**■** Figure 6.7).

STERILE POLYPLOIDS

In spite of their robust physical appearance, many polyploid species are sterile. Extra sets of chromosomes segregate irregularly in meiosis, leading to grossly unbalanced (that is, aneuploid) gametes. If such gametes unite in fertilization, the resulting zygotes almost always die. This inviability among the zygotes explains why many polyploid species are sterile.

As an example, let's consider a triploid species with three identical sets of n chromosomes. The total number of chromosomes is therefore 3n. When meiosis occurs,



(c)

FIGURE 6.7 Polyploid plants with agricultural or horticultural significance: (a) Chrysanthemum (tetraploid), (b) strawberry (octoploid), (c) cotton (tetraploid), (d) banana (triploid).

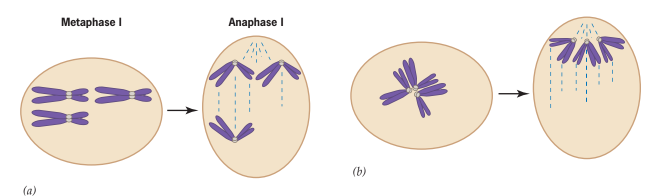
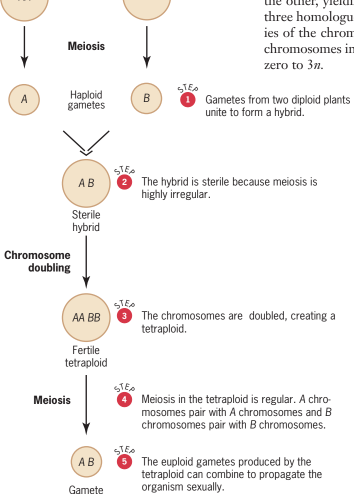


FIGURE 6.8 Meiosis in a triploid. (*a*) Univalent formation. Two of the three homologues synapse, leaving a univalent free to move to either pole during anaphase. (*b*) Trivalent formation. All three homologues synapse, forming a trivalent, which may move as a unit to one pole during anaphase. However, other anaphase disjunctions are possible.

each chromosome will try to pair with its homologues (**Tigure 6.8**). One possibility is that two homologues will pair completely along their length, leaving the third without a partner; this solitary chromosome is called a **univalent**. Another possibility is that all three homologues will synapse, forming a **trivalent** in which each member is partially paired with each of the others. In either case, it is difficult to predict how the chromosomes will move during anaphase of the first meiotic division. The more likely event is that two of the homologues will move to one pole and one homologue will move to the other, yielding gametes with one or two copies of the chromosome. However, all three homologues might move to one pole, producing gametes with zero or three copies of the chromosome. Because this segregational uncertainty applies to each trio of chromosomes in the cell, the total number of chromosomes in a gamete can vary from zero to 3n.



Diploid

plant

BΒ

Diploid plant

AA

FIGURE 6.9 Origin of a fertile tetraploid by hybridization between two diploids and subsequent doubling of the chromosomes.

Zygotes formed by fertilization with such gametes are almost certain to die; thus, most triploids are completely sterile. In agriculture and horticulture, this sterility is circumvented by propagating the species asexually. The many methods of asexual propagation include cultivation from cuttings (bananas), grafts (Winesap, Gravenstein, and Baldwin apples), and bulbs (tulips). In nature, polyploid plants can also reproduce asexually. One mechanism is **apomixis**, which involves a modified meiosis that produces unreduced eggs; these eggs then form seeds that germinate into new plants. The dandelion, a highly successful polyploid weed, reproduces in this way.

FERTILE POLYPLOIDS

The meiotic uncertainties that occur in triploids also occur in tetraploids with four identical chromosome sets. Such tetraploids are therefore also sterile. However, some tetraploids are able to produce viable progeny. Close examination shows that these species contain two distinct sets of chromosomes and that each set has been duplicated. Thus, fertile tetraploids seem to have arisen by chromosome duplication in a hybrid that was produced by a cross of two different, but related, diploid species; most often these species have the same or very similar chromosome numbers. **Figure 6.9** shows a plausible mechanism for the origin of such a tetraploid. Two diploids, denoted A and B, are crossed to produce a hybrid that receives one set of chromosomes from each of the parental species. Such a hybrid will probably be sterile because the A and B chromosomes cannot pair with each other. However, if the chromosomes in this hybrid are duplicated, meiosis will proceed in reasonably good order. Each of the A and B chromosomes will be able to pair with a perfectly homologous partner. Meiotic segregation can therefore produce gametes with a complete set of A and B chromosomes. In fertilization, these "diploid" gametes will unite to form tetraploid zygotes, which will survive because each of the parental sets of chromosomes will be balanced.

This scenario of hybridization between different but related species followed by chromosome doubling has evidently occurred many times during plant evolution. In some cases, the process has occurred repeatedly, generating complex polyploids with distinct chromosome sets. One of the best examples is modern bread wheat, *Triticum aestivum* (\blacksquare Figure 6.10). This important crop species is a hexaploid containing three different chromosome sets, each of which has been duplicated. There are seven chromosomes in each set, for a total of 21 in the gametes and 42 in the somatic cells. Thus, as we noted at the beginning of this chapter, modern wheat seems to have been formed by two hybridization events. The first involved two diploid species that combined to form a tetraploid, and the second involved a combination between this tetraploid and another diploid, to produce a hexaploid. Cytogeneticists have identified primitive cereal plants in the Middle East that may have participated in this evolutionary process. In 2010, much of the DNA in the wheat genome was sequenced. This genome is very large, roughly five times the size of the human genome. Analysis of all these DNA sequences will help us to understand wheat's evolutionary history.

Because chromosomes from different species are less likely to interfollowed by fere with each other's segregation during meiosis, polyploids arising from hybridizations between different species have a much greater chance of being fertile than do polyploids arising from the duplication of chromosomes in a single species. Polyploids created by hybridization between different species are called **allopolyploids** (from the Greek prefix for "other"); in these polyploids, the contributing genomes are qualitatively different. Polyploids created by chromosome duplication within a species are called **autopolyploids** (from the Greek prefix for "self"); in these polyploids, a single genome has been multiplied to create extra chromosome sets.

Chromosome doubling is a key event in the formation of polyploids. One possible mechanism for this event is for a cell to go through mitosis without going through cytokinesis. Such a cell will have twice the usual number of chromosomes. Through subsequent divisions, it could then give rise to a polyploid clone of cells, which might contribute either to the asexual propagation of the organism or to the formation of gametes. In plants it must be remembered that the germ line is not set aside early in development, as it is in animals. Rather, the reproductive tissues differentiate only after many cycles of cell division. If the chromosomes were accidentally doubled during one of these cell divisions, the reproductive tissues that would ultimately develop might be polyploid. Another possibility is for meiosis to be altered in such a way that unreduced gametes (with twice the normal number of chromosomes) are produced. If such gametes participate in fertilization, polyploid zygotes will be formed. These zygotes may then develop into mature organisms, which, depending on the nature of the polyploidy, may be able to produce gametes themselves. Enhance your understanding of these possibilities by working through Solve It: Chromosome Pairing in Polyploids.

TISSUE-SPECIFIC POLYPLOIDY AND POLYTENY

In some organisms, certain tissues become polyploid during development. This polyploidization is probably a response to the need for multiple copies of each chromosome and the genes it carries. The process that produces such polyploid cells, called **endomitosis**, involves chromosome duplication, followed by separation of the resulting sister chromatids. However, because there is no accompanying cell division, extra chromosome

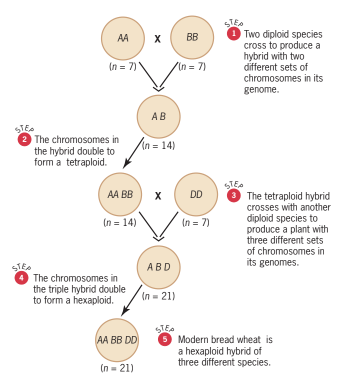


FIGURE 6.10 Origin of hexaploid wheat by sequential hybridization of different species. Each hybridization event is followed by doubling of the chromosomes.

Solve It!

Chromosome Pairing in Polyploids

There are six chromosomes in the gametes of plant species A and nine chromosomes in the gametes of plant species B. A cross between these two species produced sterile hybrids in which no chromosome pairing could be observed in the microspore mother cells of the anthers. However, the A imes B hybrid genotype could be propagated vegetatively by rooting cuttings from the plants. One of these cuttings grew into a robust plant that happened to be fertile, and when the microspore mother cells of this plant were examined cytologically, 15 bivalents were observed. This fertile plant was then backcrossed to species A, and the microspore mother cells of the offspring were examined cytologically. (a) Explain the origin of the robust plant that was fertile. (b) How many bivalents would you expect to see in the microspore mother cells of its backcross offspring? (c) How many unpaired chromosomes (univalents) would you expect to see in these offspring?

To see the solution to this problem, visit the Student Companion site.

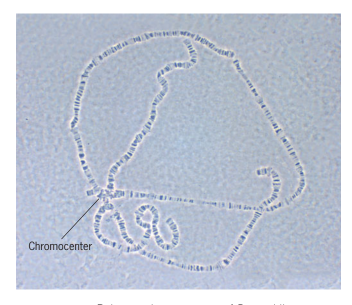


FIGURE 6.11 Polytene chromosomes of Drosophila.

sets accumulate within a single nucleus. In the human liver and kidney, for example, one round of endomitosis produces tetraploid cells.

Sometimes polyploidization occurs without the separation of sister chromatids. In these cases, the duplicated chromosomes pile up next to each other, forming a bundle of strands that are aligned in parallel. The resulting chromosomes are said to be polytene, from the Greek words meaning "many threads." The most spectacular examples of polytene chromosomes are found in the salivary glands of Drosophila larvae. Each chromosome undergoes about nine rounds of replication, producing a total of about 500 copies in each cell. All the copies pair tightly, forming a thick bundle of chromatin fibers. This bundle is so large that it can be seen under low magnification with a dissecting microscope. Differential coiling along the length of the bundle causes variation in the density of the chromatin. When dves are applied to these chromosomes, the denser chromatin stains more deeply, creating a pattern of dark and light bands (**Figure 6.11**). This pattern is highly reproducible, permitting detailed analysis of chromosome structure.

The polytene chromosomes of *Drosophila* show two additional features:

- 1. Homologous Polytene Chromosomes Pair. Ordinarily, we think of pairing as a property of meiotic chromosomes; however, in many insect species the somatic chromosomes also pair—probably as a way of organizing the chromosomes within the nucleus. When *Drosophila* polytene chromosomes pair, the large chromatin bundles become even larger. Because this pairing is precise—point-for-point along the length of the chromosome—the two homologues come into perfect alignment. Thus, the banding patterns of each are exactly in register, so much so that it is almost impossible to distinguish the individual members of a pair.
- 2. All the Centromeres of *Drosophila* Polytene Chromosomes Congeal into a Body Called the Chromocenter. Material flanking the centromeres is also drawn into this mass. The result is that the chromosome arms seem to emanate out of the chromocenter. These arms, which are banded, consist of euchromatin, that portion of the chromosome that contains most of the genes; the chromocenter consists of heterochromatin, a gene-poor material that surrounds the centromere. Unlike the euchromatic chromosome arms, this centric heterochromatin does not become polytene. Thus, compared to the euchromatin, it is vastly underreplicated.

In the 1930s C. B. Bridges published detailed drawings of the polytene chromosomes (**Figure 6.12**). Bridges arbitrarily divided each of the chromosomes

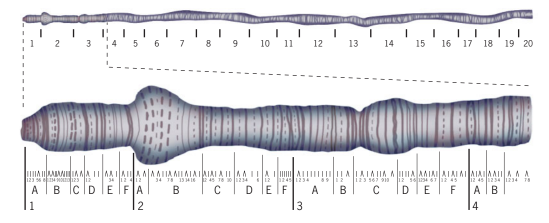


FIGURE 6.12 Bridges' polytene chromosome maps. (*Top*) Banding pattern of the polytene X chromosome. The chromosome is divided into 20 numbered sections. (*Bottom*) Detailed view of the left end of the polytene X chromosome showing Bridges' system for denoting individual bands.

into sections, which he numbered; each section was then divided into subsections, which were designated by the letters A to F. Within each subsection, Bridges enumerated all the dark bands, creating an alphanumeric directory of sites along the length of each chromosome. Bridges' alphanumeric system is still used today to describe the features of these remarkable chromosomes.

The polytene chromosomes of *Drosophila* are trapped in the interphase of the cell cycle. Thus, although most cytological analyses are performed on mitotic chromosomes, the most thorough and detailed analyses are performed on polytenized interphase chromosomes. Such chromosomes are found in many species within the insect order Diptera, including flies and mosquitoes. Unfortunately, humans do not have polytene chromosomes; thus, the high-resolution cytological analysis that is possible for *Drosophila* is not possible for our own species.

- Polyploids contain extra sets of chromosomes.
- Many polyploids are sterile because their multiple sets of chromosomes segregate irregularly in meiosis.
- Polyploids produced by chromosome doubling in interspecific hybrids may be fertile if their constituent genomes segregate independently.
- In some somatic tissues—for example, the salivary glands of Drosophila larvae—successive
 rounds of chromosome replication occur without intervening cell divisions and produce large
 polytene chromosomes that are ideal for cytogenetic analysis.

Aneuploidy

Aneuploidy describes a numerical change in part of the genome, usually a change in the dosage of a single chromosome. Individuals that have an extra chromo-

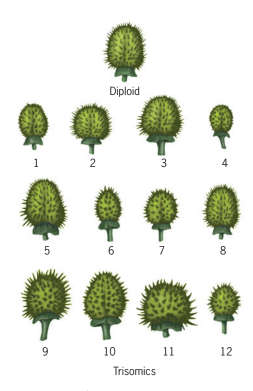
The under- or overrepresentation of a chromosome or a chromosome segment can affect a phenotype.

some, are missing a chromosome, or have a combination of these anomalies are said to be aneuploid. This definition also includes pieces of chromosomes. Thus, an individual in which a chromosome arm has been deleted is also considered to be aneuploid.

Aneuploidy was originally studied in plants, where it was shown that a chromosome imbalance usually has a phenotypic effect. The classic study was one by Albert Blakeslee and John Belling, who analyzed chromosome anomalies in Jimson weed, *Datura stramonium*. This diploid species has 12 pairs of chromosomes, for a total of 24 in the somatic cells. Blakeslee collected plants with altered phenotypes and discovered that in some cases the phenotypes were inherited in an irregular way. These peculiar mutants were apparently caused by dominant factors that were transmitted primarily through the female. By examining the chromosomes of the mutant plants, Belling found that in every case an extra chromosome was present. Detailed analysis established that the extra chromosome was different in each mutant strain. Altogether there were 12 different mutants, each corresponding to a triplication of one of the *Datura* chromosomes (**■ Figure 6.13**). Such triplications are called **trisomies**. The transmissional irregularities of these mutants were due to anomalous chromosome behavior during meiosis.

Belling also discovered the reason for the preferential transmission of the trisomic phenotypes through the female. During pollen tube growth, an euploid pollen in particular, pollen with n + 1 chromosomes—does not compete well with euploid pollen. Consequently, trisomic plants almost always inherit their extra chromosome from the female parent. Belling's work with *Datura* demonstrated that each chromosome must be present in the proper dosage for normal growth and development.

Since Belling's work, aneuploids have been identified in many species, including our own. An organism in which a chromosome, or a piece of a chromosome, is underrepresented is referred to as a *hypoploid* (from the Greek prefix for "under"). An organism in which a chromosome or chromosome segment is overrepresented is referred to as a *hyperploid* (from the Greek prefix for "over"). Each of these terms covers a wide range of abnormalities.



■ **FIGURE 6.13** Seed capsules of normal and trisomic *Datura stramonium*. Each of the 12 trisomies is shown.

KEY POINTS

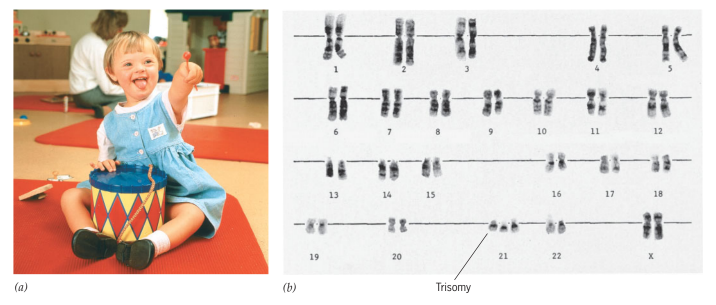


FIGURE 6.14 Down syndrome. (*a*) A young girl with Down syndrome. (*b*) Karyotype of a child with Down syndrome, showing trisomy for chromosome 21 (47, XX, +21).

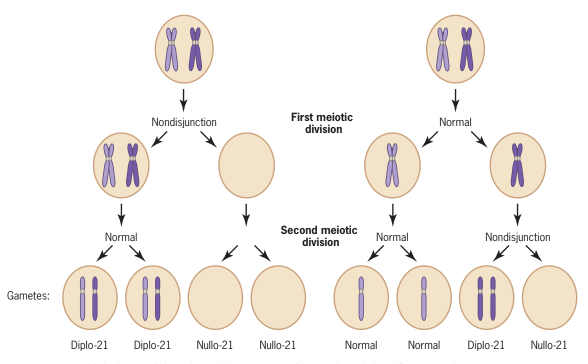
TRISOMY IN HUMANS

The best-known and most common chromosome abnormality in humans is **Down** syndrome, a condition associated with an extra chromosome 21 (**Figure 6.14a**). This syndrome was first described in 1866 by a British physician, Langdon Down, but its chromosomal basis was not clearly understood until 1959. People with Down syndrome are typically short in stature and loose-jointed, particularly in the ankles; they have broad skulls, wide nostrils, large tongues with a distinctive furrowing, and stubby hands with a crease on the palm. Impaired mental abilities require that they be given special education and care. The life span of people with Down syndrome is much shorter than that of other people. Down syndrome individuals also almost invariably develop Alzheimer's disease, a form of dementia that is fairly common among the elderly. However, people with Down syndrome develop this disease in their fourth or fifth decade of life, much sooner than other people.

The extra chromosome 21 in Down syndrome is an example of a trisomy. **Figure 6.14b** shows the karyotype of a female Down patient. Altogether, there are 47 chromosomes, including two X chromosomes as well as the extra chromosome 21. The karyotype of this individual is therefore written 47, XX, +21.

Trisomy 21 can be caused by chromosome nondisjunction in one of the meiotic cell divisions (**Figure 6.15**). The nondisjunction event can occur in either parent, but it seems to be more likely in females. In addition, the frequency of nondisjunction increases with maternal age. Thus, among mothers younger than 25 years old, the risk of having a child with Down syndrome is about 1 in 1500, whereas among mothers 40 years old, it is 1 in 100. This increased risk is due to factors that adversely affect meiotic chromosome behavior as a woman ages. In human females, meiosis begins in the fetus, but it is not completed until after the egg is fertilized. During the long time prior to fertilization, the meiotic cells are arrested in the prophase of the first division. In this suspended state, the chromosomes may become unpaired. The longer the time in prophase, the greater the chance for unpairing and subsequent chromosome non-disjunction. Older females are therefore more likely than younger females to produce aneuploid eggs.

Trisomies for chromosomes 13 and 18 have also been reported. However, these are rare, and the affected individuals show serious phenotypic abnormalities and are short-lived, usually dying within the first few weeks after birth. Another viable trisomy



■ FIGURE 6.15 Meiotic nondisjunction of chromosome 21 and the origin of Down syndrome. Nondisjunction at meiosis I does not produce normal gametes; the gametes either carry two copies of chromosome 21 (diplo-21) or no copy of this chromosome (nullo-21). Nondisjunction at meiosis II produces a gamete with two identical sister chromosomes (diplo-21) and a gamete lacking chromosome 21 (nullo-21).

that has been observed in humans is the triplo-X karyotype, 47, XXX. These individuals survive because two of the three X chromosomes are inactivated, reducing the dosage of the X chromosome so that it approximates the normal level of one. Triplo-X individuals are female and are phenotypically normal, or nearly so; sometimes they exhibit a slight mental impairment and reduced fertility.

The 47, XXY karyotype is also a viable trisomy in humans. These individuals have three sex chromosomes, two X's and one Y. Phenotypically, they are male, but they can show some female secondary sexual characteristics and are usually sterile. In 1942 H. F. Klinefelter described the abnormalities associated with this condition, now called **Klinefelter syndrome**; these include small testes, enlarged breasts, long limbs, knock-knees, and underdeveloped body hair. The XXY karyotype can originate by fertilization of an exceptional XX egg with a Y-bearing sperm or by fertilization of an X-bearing egg with an exceptional XY sperm. The XXY karyotype accounts for about three-fourths of all cases of Klinefelter syndrome. Other cases involve more complex karyotypes such as XXYY, XXXY, XXXYY, XXXYY, XXXYY, and XXXXXY. All individuals with Klinefelter syndrome have one or more Barr bodies in their cells, and those with more than two X chromosomes usually have some degree of mental impairment.

The 47, XYY karyotype is another viable trisomy in humans. These individuals are male, and except for a tendency to be taller than 46, XY men, they do not show a consistent syndrome of characteristics. All the other trisomies in humans are embry-onic lethals, demonstrating the importance of correct gene dosage. Unlike *Datura*, in which each of the possible trisomies is viable, humans do not tolerate many types of chromosomal imbalance (see Table 6.1).

MONOSOMY

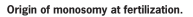
Monosomy occurs when one chromosome is missing in an otherwise diploid individual. In humans, there is only one viable monosomic, the 45, X karyotype.

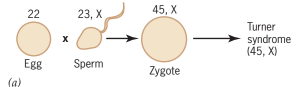
TABLE 6.1

Aneuploidy Resulting from Nondisjunction in Humans

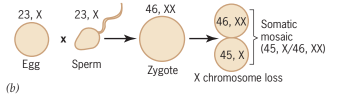
Karyotype	Chromosome Formula	Clinical Syndrome	Estimated Frequency at Birth	Phenotype
47, +21	2 n + 1	Down	1/700	Short, broad hands with palmar crease, short stature, hyperflexibility of joints, mental retardation, broad head with round face, open mouth with large tongue, epicanthal fold.
47, +13	2 <i>n</i> + 1	Patau	1/20,000	Mental deficiency and deafness, minor muscle seizures, cleft lip and/or palate, cardiac anomalies, posterior heel prominence.
47, +18	2n + 1	Edward	1/8000	Congenital malformation of many organs, low-set, malformed ears, receding mandible, small mouth and nose with general elfin appearance, mental deficiency, horseshoe or double kidney, short sternum; 90 percent die within first six months after birth.
45, X	2 n – 1	Turner	1/2500 female births	Female with retarded sexual development, usually sterile, short stature, webbing of skin in neck region, cardiovascu- lar abnormalities, hearing impairment.
47, XXY	2 n + 1	Klinefelter	1/500 male births	Male, subfertile with small testes, developed breasts, feminine-pitched voice, knock-knees, long limbs.
48, XXXY	2 <i>n</i> + 2			
48, XXYY	2 n + 2			
49, XXXXY	2 n + 3			
50, XXXXXY	2 n + 4			
47, XXX	2 n + 1	Triplo-X	1/700	Female with usually normal genitalia and limited fertility, slight mental retardation.

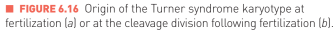
These individuals have a single X chromosome as well as a diploid complement of autosomes. Phenotypically, they are female, but because their ovaries are rudimentary, they are almost always sterile. 45, X individuals are usually short in stature; they have webbed necks, hearing deficiencies, and significant cardiovascular abnormalities. Henry H. Turner first described the condition in 1938; thus, it is now called **Turner syndrome**. 45, X individuals can originate from eggs or sperm that lack a sex chromosome or from the loss of a sex chromosome in mitosis sometime after fertilization (**Turner 6.16**). This latter possibility is supported by the finding that many Turner individuals are *somatic mosaics*. These people have two types of cells in their bodies; some are 45, X and others are 46, XX. This karyotypic mosaicism evi-





Origin of monosomy in the cleavage division following fertilization.





dently arises when an X chromosome is lost during the development of a 46, XX zygote. All the descendants of the cell in which the loss occurred are 45, X. If the loss occurs early in development, an appreciable fraction of the body's cells will be aneuploid and the individual will show the features of Turner syndrome. If the loss occurs later, the aneuploid cell population will be smaller and the severity of the syndrome is likely to be reduced. For a discussion of procedures used to detect aneuploidy in human fetuses, see the Focus on Amniocentesis and Chorionic Biopsy.

XX/XO chromosome mosaics also occur in *Drosophila*, where they produce a curious phenotype. Because sex in this species is determined by the ratio of X chromosomes to autosomes, such flies are part female and part male. XX cells develop in the female direction, and XO cells develop in the male direction. Flies with both male and female structures are called **gynandromorphs** (from Greek words meaning "woman," "man," and "form").

People with the 45, X karyotype have no Barr bodies in their cells, indicating that the single X chromosome that is present is not

FOCUS ON

AMNIOCENTESIS AND CHORIONIC BIOPSY

he Andersons, a couple living in Minneapolis, were expecting their first baby. Neither Donald nor Laura Anderson knew of any genetic abnormalities in their families, but because of Laura's age-38-they decided to have the fetus checked for aneuploidy. Laura's physician performed a procedure called **amniocentesis.** A small amount of fluid was removed from the cavity surrounding the developing fetus by inserting a needle into Laura's abdomen (**Figure 1**). This cavity, called the amniotic sac, is enclosed by a membrane. To prevent discomfort during the procedure, Laura was given a local anesthetic. The needle was guided into position by following an ultrasound scan, and some of the amniotic fluid was drawn out. Because this fluid contains nucleated cells sloughed off from the fetus, it is possible to determine the fetus's karyotype. Usually the fetal cells are purified from the amniotic fluid by centrifugation, and then the cells are cultured for several days to a few weeks. Cytological analysis of these cells will reveal if the fetus is aneuploid. Additional tests may be performed on the fluid recovered from the amniotic sac to detect other sorts of abnormalities, including neural tube defects and some kinds of mutations. The results of all these tests may take up to three weeks. In Laura's case, no abnormalities of any sort were detected, and 20 weeks after the amniocentesis, she gave birth to a healthy baby girl.

Chorionic biopsy provides another way of detecting chromosomal abnormalities in the fetus. The chorion is a fetal membrane that interdigitates with the uterine wall, eventually forming the placenta. The minute chorionic projections into the uterine tissue are called *villi* (singular, villus). At 10–11 weeks of gestation, before the placenta has developed, a sample of chorionic villi can be obtained by passing a hollow plastic tube into the uterus through the cervix. This tube can be guided by an ultrasound scan, and when the tube is in place, a tiny bit of material can be drawn up into the tube by aspiration. The recovered material usually consists of a mixture of maternal and fetal tissue. After these tissues are separated by dissection, the fetal cells can be analyzed for chromosomal abnormalities. Chorionic biopsy can be performed earlier than amniocentesis (10–11 weeks gestation versus 14–16 weeks), but it is not as reliable. In addition, it seems to be associated with a slightly greater chance of miscarriage than amniocentesis does, perhaps 2 to 3 percent. For these reasons, it tends to be used only in pregnancies where there is a strong reason to expect a genetic abnormality. In routine pregnancies, such as Laura Anderson's, amniocentesis is the preferred procedure.

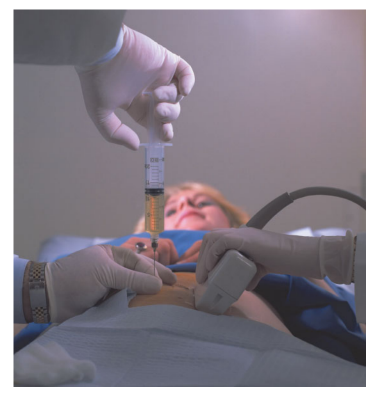


FIGURE 1 A physician taking a sample of fluid from the amniotic sac of a pregnant woman for prenatal diagnosis of a chromosomal or biochemical abnormality.

inactivated. Why, then, should Turner patients, who have the same number of active X chromosomes as normal XX females, show any phenotypic abnormalities at all? The answer probably involves a small number of genes that remain active on both of the X chromosomes in normal 46, XX females. These noninactivated genes are apparently needed in double dose for proper growth and development. The finding that at least some of these special X-linked genes are also present on the Y chromosome would explain why XY males grow and develop normally. In addition, the X chromosome that has been inactivated in 46, XX females is reactivated during oogenesis, presumably because two copies of some X-linked genes are required for normal ovarian function. 45, X individuals, who have only one copy of these genes, cannot meet this requirement and are therefore sterile.

Curiously, the cognate of the XO Turner karyotype in the mouse exhibits no anatomical abnormalities. This finding implies that the mouse homologues of the human genes that are involved in Turner syndrome need only be present in one copy for normal growth and development. To investigate the origin of the XO Turner karyotype, work through the exercise in Problem-Solving Skills: Tracing Sex Chromosome Nondisjunction.

PROBLEM-SOLVING SKILLS

Tracing Sex Chromosome Nondisjunction

THE PROBLEM

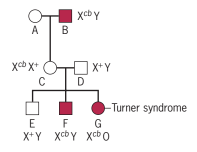
A color-blind man married a normal woman. Their daughter, who was phenotypically normal, married a normal man and the couple produced three children: a normal boy, a color-blind boy, and a color-blind girl with Turner syndrome. Explain the origin of the color-blind girl with Turner syndrome.

FACTS AND CONCEPTS

- 1. Color blindness is caused by a recessive X-linked mutation, cb.
- **2.** Turner syndrome is due to monosomy of the X chromosome (genotype XO).
- **3.** Monosomy can arise from chromosome nondisjunction during mitosis or meiosis.
- **4.** Mitotic nondisjunction in an XX individual can create a mosaic of XO and XX cells.

ANALYSIS AND SOLUTION

To start the analysis, let's diagram the pedigree and label all the people in it. In addition, because we know that color blindness is due to a recessive X-linked mutation, we can write down the genotypes of most of the people in the pedigree.



The color-blind man, B, is a key figure in this pedigree because he must have transmitted an X chromosome carrying the cb mutation to his daughter C, who is the mother of the child in question. C is not color blind herself, so she must be heterozygous for the mutant allele—that is, her genotype is $X^{cb} X^+$. Likewise, her husband, D, is not color blind, so he must have the genotype $X^+ Y$. The genotypes of the couple's first two children are also known with certainty. The last child, G, has Turner syndrome, which implies that she has just one sex chromosome—an X. Because this girl is color blind, her genotype is presumably X^{cb} O. This genotype could have been created by fertilization of an egg containing the X^{cb} chromosome by a sperm that lacked a sex chromosome. In this scenario, there must have been nondisjunction of the sex chromosomes during meiosis in G's father. Another possibility is that the X^{cb}-bearing egg was fertilized by a sperm that carried an X chromosome and this chromosome was lost during one of the early divisions in the embryo. On this second hypothesis, G would be a somatic mosaic of XO and XX cells (see Figure 6.16b). However, this explanation does not square with the observation that G is color blind, for if G were a somatic mosaic, her XX cells would have to be X^{cb} X⁺, and some of these cells would be expected to have formed normal photoreceptor cells in her retinas, thereby giving her normal color vision. The fact that G is color blind indicates that she does not have X^{cb} X⁺ cells in her retinas—or probably anywhere else in her body. Sex chromosome nondisjunction during meiosis in G's father is therefore the more plausible explanation for her color-blind, Turner phenotype.

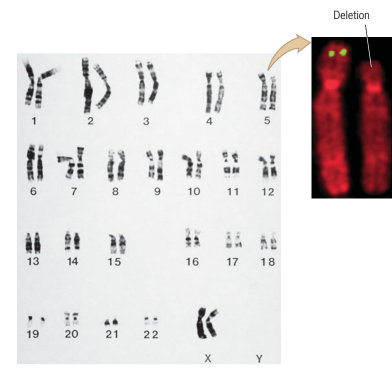
For further discussion visit the Student Companion site.

DELETIONS AND DUPLICATIONS OF CHROMOSOME SEGMENTS

A missing chromosome segment is referred to either as a **deletion** or as a **deficiency**. Large deletions can be detected cytologically by studying the banding patterns in stained chromosomes, but small ones cannot. In a diploid organism, the deletion of a chromosome segment makes part of the genome hypoploid. This hypoploidy may be associated with a phenotypic effect, especially if the deletion is large. A classic example is the *cri-du-chat* **syndrome** (from the French words for "cry of the cat") in humans (**True Figure 6.17**). This condition is caused by a deletion in the short arm of chromosome 5. The size of the deletion varies. Individuals heterozygous for the deletion and a normal chromosome have the karyotype 46 del(5)(p14), where the terms in parentheses indicate that bands in region 14 of the short arm (p) of one of the chromosomes 5 is missing. These individuals may be severely impaired, mentally as well as physically; their plaintive, catlike crying during infancy gives the syndrome its name.

An extra chromosome segment is referred to as a **duplication**. The extra segment can be attached to one of the chromosomes, or it can exist as a new and separate chromosome, that is, as a "free duplication." In either case, the effect is the same: The organism is hyperploid for part of its genome. As with deletions, this hyperploidy can be associated with a phenotypic effect.

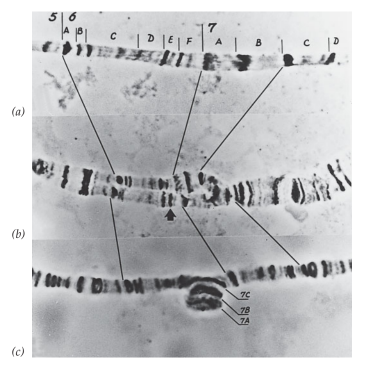




■ **FIGURE 6.17** Karyotype of a female with the *cri-du-chat* syndrome, 46, XX del(5)(p14). One of the chromosomes 5 has a deletion in its short arm. The inset shows the two chromosomes 5 labeled with a fluorescent gene-specific probe. The chromosome on the left has bound the probe because it carries this particular gene, whereas the chromosome on the right has not bound the probe because the gene, and material around it, is deleted.

Deletions and duplications are two types of aberrations in chromosome structure. Large aberrations can be detected by examination of mitotic chromosomes that have been stained with banding agents such as quinacrine or Giemsa. However, small aberrations are difficult to detect in this way and are usually identified by other genetic and molecular techniques. The best organism for studying deletions and duplications is *Drosophila*, where the polytene chromosomes afford an unparalleled opportunity for detailed cytological analysis. **Figure 6.18***b* shows a deletion in one of two paired homologous chromosomes in a *Drosophila* salivary gland. Because the two chromosomes have separated slightly, we can see that a small region is missing in the lower one.

Duplicated segments can also be recognized in polytene chromosomes. **Figure 6.18** shows a tandem duplication of a segment in the middle of the X chromosome of Drosophila. Because tandem copies of this segment pair with each other, the chromosome appears to have a knot in its middle. The Bar eye mutation in Drosophila is associated with a tandem duplication (**Figure 6.19**). This dominant X-linked mutation alters the size and shape of the compound eyes, transforming them from large, spherical structures into narrow bars. In the 1930s C. B. Bridges analyzed X chromosomes carrying the Bar mutation and found that the 16A region, which apparently contains a gene for eye shape, had been tandemly duplicated. Tandem triplications of 16A were also observed, and in these cases the compound eye was extremely small-a phenotype referred to as double-bar. The severity of the mutant eve phenotype is therefore related to the number of copies of the 16A region—clear evidence for the importance of gene dosage in determining a phenotype. Many other tandem duplications have been found in Drosophila, where polytene chromosome analysis makes their detection relatively easy. Today, molecular techniques have made it possible to detect very small tandem duplications in a wide variety of organisms. For example, the genes that encode the hemoglobin proteins have been tandemly duplicated in mammals (Chapter 19). Gene duplications appear to be relatively common and provide a significant source of variation for evolution.



■ FIGURE 6.18 Polytene chromosomes showing (a) the normal structure of regions 6 and 7 in the middle of the *Drosophila* X chromosome, (b) a heterozygote with a deletion of region 6F-7C in one of the chromosomes (arrow), and (c) an X chromosome showing a reverse tandem duplication of region 6F-7C. In (b) the prominent bands in regions 7A and 7C are present in the upper chromosome but absent in the lower one, indicating that the lower chromosome has undergone a deletion. In (c) the duplicated sequence reads 7C, 7B, 7A, 7A, 7B, 7C from left to right.

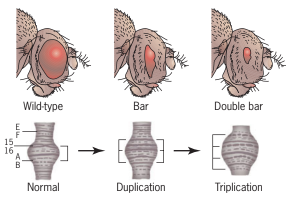


FIGURE 6.19 Effects of duplications for region 16A of the X chromosome on the size of the eyes in *Drosophila*.

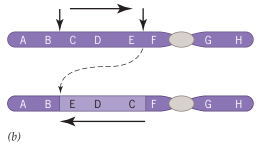
KEY POINTS

- In a trisomy, such as Down syndrome in humans, three copies of a chromosome are present; in a monosomy, such as Turner syndrome in humans, only one copy of a chromosome is present.
 - Aneuploidy may involve the deletion or duplication of a chromosome segment.

Rearrangements of Chromosome Structure

A chromosome may become rearranged internally, or it may become joined to another chromosome.

Pericentric inversion—includes centromere. A B C D E F G H A B C D G F E H (a) Paracentric inversion—excludes centromere.



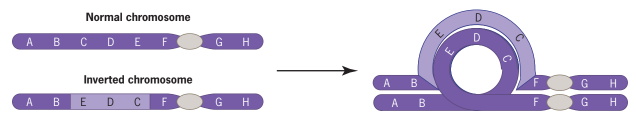
■ FIGURE 6.20 Pericentric and paracentric inversions. The chromosome has been broken at two points, and the segment between them has been inverted. A pericentric inversion (a) changes the size of the chromosome arms because the centromere is included within the inversion. By contrast, a paracentric inversion (b) does not because it excludes the centromere In nature there is considerable variation in the number and structure of chromosomes, even among closely related organisms. For example, *Drosophila melanogaster* has four pairs of chromosomes, including a pair of sex chromosomes, two pairs of large, metacentric autosomes—chromosomes with the centromere in the

middle-and a pair of small, dotlike autosomes. Drosophila virilis, which is not too distantly related, has a pair of sex chromosomes, four pairs of acrocentric autosomes-chromosomes with the centromere near one end-and a pair of dotlike autosomes. Thus, even in the same genus, species can have different chromosome arrangements. These differences imply that over evolutionary time, segments of the genome are rearranged. In fact, the observation that chromosome rearrangements can be found as variants within a single species suggests that the genome is continuously being reshaped. These rearrangements may change the position of a segment within a chromosome, or they may bring together segments from different chromosomes. In either case, the order of the genes is altered. Cytogeneticists have identified many kinds of chromosome rearrangements. Here we consider two types: inversions, which involve a switch in the orientation of a segment within a chromosome, and translocations, which involve the fusion of segments from different chromosomes. In humans, chromosome rearrangements have a medical significance because some of them are involved in predisposing individuals to develop certain types of cancer. We consider these kinds of rearrangements, and their connection to cancer, in Chapter 21.

INVERSIONS

An inversion occurs when a chromosome segment is detached, flipped around 180°, and reattached to the rest of the chromosome; as a result, the order of the segment's genes is reversed. Such rearrangements can be induced in the laboratory by X-irradiation, which breaks chromosomes into pieces. Sometimes the pieces reattach, but in the process a segment gets turned around and an inversion occurs. There is also evidence that inversions are produced naturally through the activity of transposable elements—DNA sequences capable of moving from one chromosomal position to another (Chapter 17). Sometimes, in the course of moving, these elements break a chromosome into pieces and the pieces reattach in an aberrant way, producing an inversion. Inversions may also be created by the reattachment of chromosome fragments generated by mechanical shear, perhaps as a result of chromosome entanglement within the nucleus. No one really knows what fraction of naturally occurring inversions is caused by each of these mechanisms.

Cytogeneticists distinguish between two types of inversions based on whether or not the inverted segment includes the chromosome's centromere (**Figure 6.20**). **Pericentric** inversions include the centromere, whereas **paracentric** inversions do not. The consequence is that a pericentric inversion may change the relative lengths of the two arms of the chromosome, whereas a paracentric inversion has no such effect. Thus, if an acrocentric chromosome acquires an inversion with a breakpoint in each



■ FIGURE 6.21 Pairing between normal and inverted chromosomes

of the chromosome's arms (that is, a pericentric inversion), it can be transformed into a metacentric chromosome. However, if an acrocentric chromosome acquires an inversion in which both of the breaks are in the chromosome's long arm (that is, a paracentric inversion), the morphology of the chromosome will not be changed. Hence, with the use of standard cytological methods, pericentric inversions are much easier to detect than paracentric inversions.

An individual in which one chromosome is inverted but its homologue is not is said to be an inversion heterozygote. During meiosis, the inverted and noninverted chromosomes pair point-for-point along their length. However, because of the inversion, the chromosomes must form a loop to allow for pairing in the region where their genes are in reversed order. **Figure 6.21** shows this pairing configuration; only

one of the chromosomes is looped, and the other conforms around it. In practice, either the inverted or noninverted chromosome can form the loop to maximize pairing between them. However, near the ends of the inversion, the chromosomes are stretched, and there is a tendency for some de-synapsis. We consider the genetic consequences of inversion heterozygosity in Chapter 7.

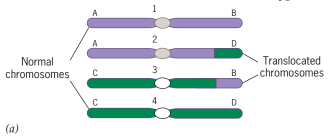
TRANSLOCATIONS

A **translocation** occurs when a segment from one chromosome is detached and reattached to a different (that is, nonhomologous) chromosome. The genetic significance is that genes from one chromosome are transferred to another.

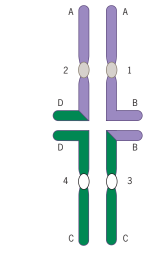
When pieces of two nonhomologous chromosomes are interchanged without any net loss of genetic material, the event is referred to as a *reciprocal translocation*. **Figure 6.22a** shows a reciprocal translocation between two large autosomes. These chromosomes have interchanged pieces of their right arms. During meiosis, these translocated chromosomes would be expected to pair with their untranslocated homologues in a cruciform, or crosslike, pattern (**Figure 6.22b**). The two translocated chromosomes face each other opposite the center of the cross, and the two untranslocated chromosomes do likewise; to maximize pairing, the translocated and untranslocated chromosomes alternate with each other, forming the arms of the cross. This pairing configuration is diagnostic of a translocation heterozygote. Cells in which the translocated chromosomes are homozygous do not form a cruciform pattern. Instead, each of the translocated chromosomes pairs smoothly with its structurally identical partner.

Because cruciform pairing involves four centromeres, which may or may not be coordinately distributed to opposite poles in the first meiotic division, chromosome disjunction in translocation heterozygotes is a somewhat uncertain process, prone to produce aneuploid gametes. Altogether there are three possible disjunctional events, illustrated in **Figure 6.23**. This simplified figure shows only

Structure of chromosomes in translocation heterozygote.

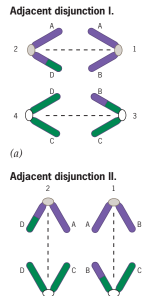


Pairing of chromosomes in translocation heterozygote.



(b)

■ FIGURE 6.22 Structure and pairing behavior of a reciprocal translocation between chromosomes. In (b) pairing occurs during the prophase of meiosis I, after the chromosomes have been duplicated.



Centromeres 1 and 3 go to one pole and centromeres 2 and 4 go to the other pole, producing aneuploid gametes.

Centromeres 1 and 2 go to one pole and centromeres 3 and 4 go to the other pole, producing aneuploid gametes.

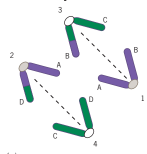
Centromeres 2 and 3 go to one pole

and centromeres 1 and

4 go to the other pole, producing euploid gametes.

Alternate disjunction.

(b)



(c) FIGURE 6.23 Types of disjunction in a translocation heterozygote during meiosis I. For simplicity, only one sister chromatid of each duplicated chromosome is shown. (a) One form of adjacent disjunction in which homologous centromeres go to opposite poles during anaphase. (b) Another form of adjacent disjunction in which homologous centromeres go to the same pole during anaphase. (c) Alternate disjunction in which homologous centromeres go to opposite poles during anaphase. one of the two sister chromatids of each chromosome. In addition, each of the centromeres is labeled to keep track of chromosome movements; the two white centromeres are homologous (that is, derived from the same chromosome pair), as are the two gray centromeres.

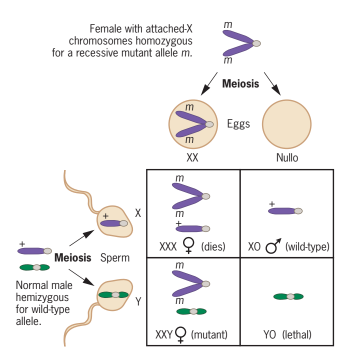
If centromeres 2 and 4 move to the same pole, forcing 1 and 3 to the opposite pole, all the resulting gametes will be aneuploid—because some chromosome segments will be deficient for genes, and others will be duplicated (Figure 6.23*a*). Similarly, if centromeres 1 and 2 move to one pole and 3 and 4 to the other, only aneuploid gametes will be produced (Figure 6.23*b*). Each of these cases is referred to as *adjacent disjunction* because centromeres that were next to each other in the cruciform pattern moved to the same pole. When the centromeres that move to the same pole are from different chromosomes (that is, they are *heterologous*), the disjunction is referred to as adjacent I (Figure 6.23*a*); when the centromeres that move to the same pole are from the same chromosome (that is, they are *homologous*), the disjunction is referred to as adjacent II (Figure 6.23*b*). Another possibility is that centromeres 1 and 4 move to the same pole, forcing 2 and 3 to the opposite pole. This case, called *alternate disjunction*, produces only euploid gametes, although half of them will carry only translocated chromosomes (Figure 6.23*c*).

The production of aneuploid gametes by adjacent disjunction explains why translocation heterozygotes have reduced fertility. When such gametes fertilize a euploid gamete, the resulting zygote will be genetically unbalanced and therefore will be unlikely to survive. In plants, aneuploid gametes are themselves often inviable, especially on the male side, and fewer zygotes are produced. Translocation heterozygotes are therefore characterized by low fertility. Investigate this effect by working through Solve It: Pollen Abortion in Translocation Heterozygotes.

COMPOUND CHROMOSOMES AND ROBERTSONIAN TRANSLOCATIONS

Sometimes one chromosome fuses with its homologue, or two sister chromatids become attached to each other, forming a single genetic unit. A **compound chromosome** can exist stably in a cell as long as it has a single functional centromere; if there are two centromeres, each may move to a different pole during division, pulling the compound chromosome apart. A compound chromosome may also be formed by the union of homologous chromosome segments. For example, the right arms of the two second chromosomes in *Drosophila* might detach from their left arms and fuse at the centromere, creating a compound half-chromosome. Cytogeneticists sometimes call this structure an **isochromosome** (from the Greek prefix for "equal"), because its two arms are equivalent. Compound chromosomes differ from translocations in that they involve fusions of homologous chromosome segments. Translocations, by contrast, always involve fusions between nonhomologous chromosomes.

The first compound chromosome was discovered in 1922 by Lillian Morgan, the wife of T. H. Morgan. This compound was formed by fusing the two X chromosomes in *Drosophila*, creating double-X or *attached-X chromosomes*. The discovery was made through genetic experimentation rather than cytological analysis. Lillian Morgan crossed females homozygous for a recessive X-linked mutation to wild-type males. From such a cross, we would ordinarily expect all the daughters to be wild-type and all the sons to be mutant. However, Morgan observed just the opposite: all the daughters were mutant and all the sons were wild-type. Further work established that the X chromosomes in the mutant females had become attached to each other. **Figure 6.24** illustrates the genetic significance of this attachment. The attached-X females produce two kinds of eggs, diplo-X and nullo-X, and their mates produce two kinds of sperm, X-bearing and Y-bearing. The union of these gametes in all possible ways produces two kinds of viable progeny: mutant XXY females, which



Solve It!

Pollen Abortion in Translocation Heterozygotes

In many plant species, aneuploid pollen are inviable. Suppose that one such plant is heterozygous for a reciprocal translocation between two large chromosomes. If adjacent I, adjacent II, and alternate disjunction in this translocation heterozygote occur with equal frequencies, what fraction of the pollen would you expect to abort?

To see the solution to this problem, visit the Student Companion site.

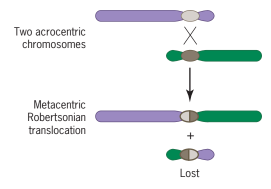
FIGURE 6.24 Results of a cross between a normal male and a female with attached-X chromosomes.

inherit the attached-X chromosomes from their mothers and a Y chromosome from their fathers; and phenotypically wild-type XO males, which inherit an X chromosome from their fathers and no sex chromosome from their mothers. Because the Y chromosome is needed for fertility, these XO males are sterile. Lillian Morgan was able to propagate the attached-X chromosomes by backcrossing XXY females to wild-type XY males from another stock. Because the sons of this cross inherited a Y chromosome from their mothers, they were fertile and could be crossed to their XXY sisters to establish a stock in which the attached-X chromosomes were permanently maintained in the female line.

Nonhomologous chromosomes can also fuse at their centromeres, creating a structure called a **Robertsonian translocation** (**Translocation** (**Translocation**)) (**Translocation** (**Translocation**)) (**Translocation** (**Translocation**)) (**Translocation**) (**Translocati**

Chromosomes can also fuse end-to-end to form a structure with two centromeres. If one of the centromeres is inactivated, the chromosome fusion will be stable. Such a fusion evidently occurred in the evolution of our own species. Human chromosome 2, which is a metacentric, has arms that correspond to two different acrocentric chromosomes in the genomes of the great apes. Detailed cytological analysis has shown that the ends of the short arms of these two chromosomes apparently fused to create human chromosome 2.

- An inversion reverses the order of genes in a segment of a chromosome.
- A translocation interchanges segments between two nonhomologous chromosomes.
- Compound chromosomes result from the fusion of homologous chromosomes or from the fusion of the arms of homologous chromosomes.
- Robertsonian translocations result from the fusion of nonhomologous chromosomes.



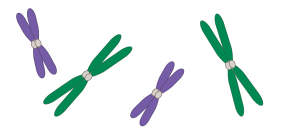
[■] FIGURE 6.25 Formation of a metacentric Robertsonian translocation by exchange between two nonhomologous acrocentric chromosomes.

KEY POINTS

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. A species has two pairs of chromosomes, one long and the other short. Draw the chromosomes at metaphase of mitosis. Show each chromatid. Are homologous chromosomes paired?
- **Answer:** Mitotic metaphase in this species would look something like the accompanying picture. Because each chromosome is duplicated, it consists of two sister chromatids. However, because the picture shows mitosis rather than meiosis, homologous chromosomes are not paired.



2. Plant species A shows 10 bivalents of chromosomes at metaphase of meiosis I; plant species B shows 14 bivalents at this stage. The two species are crossed, and the chromosomes in the offspring are doubled. (a) How many bivalents

will be seen at metaphase of meiosis I in the offspring? (b) Is the offspring expected to be fertile or sterile?

- **Answer:** (a) The offspring is a composite of the chromosomes of the two parents. In species A, the basic chromosome number is 10; in species B, it is 14. The basic chromosome number in the offspring is therefore 10 + 14 = 24, and with the chromosomes having been doubled, this is the number of bivalents that should be seen at metaphase of meiosis I. (b) The offspring is an allotetraploid and should therefore be fertile.
- **3.** What are the karyotypes of (a) a female with Down syndrome, (b) a male with trisomy 13, (c) a female with Turner syndrome, (d) a male with Klinefelter syndrome, (e) a male with a deletion in the short arm of chromosome 11?
- Answer: (a) 47, XX, +21, (b) 47, XY, +13, (c) 45, X, (d) 47, XXY, (e) 46, XY del(11p).
- **4.** What kind of pairing configuration would be seen in prophase of meiosis I in (a) an inversion heterozygote, (b) a translocation heterozygote?

Answer: (a) Loop configuration, (b) cross configuration.

Testing Your Knowledge

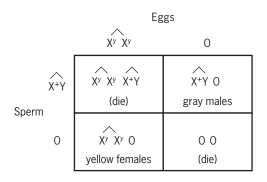
Integrate Different Concepts and Techniques

- 1. A *Drosophila* geneticist has obtained females that carry attached-X chromosomes homozygous for a recessive mutation (*y*) that causes the body to be yellow instead of gray. In one experiment, she crosses some of these females to ordinary wild-type males, and in another, she crosses these females to wild-type males that have their X and Y chromosomes attached to each other; that is, they carry a compound XY chromosome. Predict the phenotypes of the progeny from these two crosses and indicate which, if any, will be sterile.
- **Answer:** To predict the phenotypes of the progeny, we need to know their genotypes. The easiest way to determine these genotypes is to diagram the kinds of zygotes produced by each cross.

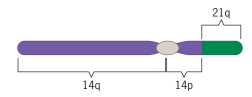
First, we consider the cross between the yellowbodied attached-X females and the ordinary wild-type males. The females produce two kinds of gametes, XX and nullo. The males also produce two kinds of gametes, X and Y. When these are combined in all possible ways, four types of zygotes are produced; however, only two types are viable. The XXY zygotes will develop into yellow-bodied females—like their mothers except that they carry a Y chromosome—and the XO zygotes will develop into gray-bodied males—like their fathers except that they lack a Y chromosome. The extra Y chromosome in the females will have no effect on fertility, but the missing Y chromosome in the males will cause them to be sterile.

		Eggs				
		X ^y X ^y	0			
0	Χ+	(die)	X ⁺ 0 gray males			
Sperm	Y	$\widehat{\chi^{y}\chi^{y}} Y$ yellow females	YO (die)			

Now we consider the cross between the yellow-bodied attached-X females and the males with a compound XY chromosome. Both sexes produce two kinds of gametes—for the females, the same as above, and for the males, either XY or nullo. When these are united in all possible ways, we find that two types of zygotes will be viable: yellow-bodied females with attached-X chromosomes and gray-bodied males with a compound XY chromosome. Both types of these viable progeny will be fertile.

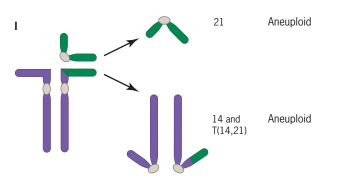


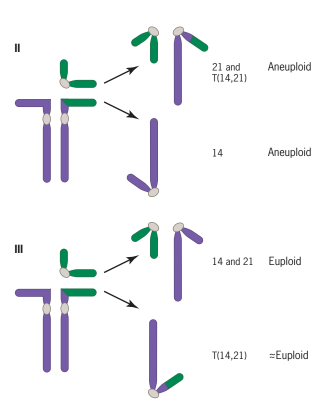
2. A phenotypically normal man carries a translocated chromosome that contains the entire long arm of chromosome 14, part of the short arm of chromosome 14, and most of the long arm of chromosome 21:



The man also carries a normal chromosome 14 and a normal chromosome 21. If he marries a cytologically (and phenotypically) normal woman, is there any chance that the couple will produce phenotypically abnormal children?

Answer: Yes, the couple could produce children with Down syndrome as a result of meiotic segregation in the cytologically abnormal man. During meiosis in this man, the translocated chromosome, T(14, 21), will synapse with the normal chromosomes 14 and 21, forming a trivalent. Disjunction from this trivalent will produce six different types of sperm, four of which are aneuploid.





Fertilization of an egg containing one chromosome 14 and one chromosome 21 by any of the aneuploid sperm will produce an aneuploid zygote as shown in the accompanying table. Although trisomy or monosomy for chromosome 14 and monosomy for chromosome 21 are all lethal conditions, trisomy for chromosome 21 is not. Thus, it is possible for the couple to give birth to a child with Down syndrome.

Disjunction	Sperm	Zygote	Condition	Outcome
Ι	21	14, 21, 21	monosomy 14	dies
	14, T(14, 21)	14, 14, T(14, 21), 21	trisomy 14	dies
II	14	14, 14, 21	monosomy 21	dies
	T(14, 21), 21	14, T(14, 21), 21, 21	trisomy 21	Down
III	14, 21	14, 14, 21, 21	euploid	normal
	T(14, 21)	14, T(14, 21), 21	≈euploid	normal

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **6.1** In the human karyotype, the X chromosome is approximately the same size as seven of the autosomes (the so-called C group of chromosomes). What procedure could be used to distinguish the X chromosome from the other members of this group?
- **6.2** In humans, a cytologically abnormal chromosome 22, called the "Philadelphia" chromosome because of the city in which it was discovered, is associated with chronic leukemia. This chromosome is missing part of its long arm. How would you denote the karyotype of an individual

who had 46 chromosomes in his somatic cells, including one normal 22 and one Philadelphia chromosome?

- **6.3** During meiosis, why do some tetraploids behave more regularly than triploids?
- 6.4 **•** The following table presents chromosome data on four species of plants and their F₁ hybrids:

Species or F ₁ Hybrid	Root Tip Chromosome Number	Number of Bivalents	Number of Univalents
A	20	10	0
В	20	10	0
С	10	5	0
D	10	5	0
$A \times B$	20	0	20
$A \times C$	15	5	5
$\mathbf{A} \times \mathbf{D}$	15	5	5
$\mathbf{C} \times \mathbf{D}$	10	0	10

Meiosis I Metaphase

- (a) Deduce the chromosomal origin of species A.
- (b) How many bivalents and univalents would you expect to observe at meiotic metaphase I in a hybrid between species C and species B?
- (c) How many bivalents and univalents would you expect to observe at meiotic metaphase I in a hybrid between species D and species B?
- **6.5** A plant species A, which has seven chromosomes in its gametes, was crossed with a related species B, which has nine. The hybrids were sterile, and microscopic observation of their pollen mother cells showed no chromosome pairing. A section from one of the hybrids that grew vigorously was propagated vegetatively, producing a plant with 32 chromosomes in its somatic cells. This plant was fertile. Explain.
- **6.6** A plant species X with n = 5 was crossed with a related species Y with n = 7. The F₁ hybrid produced only a few pollen grains, which were used to fertilize the ovules of species Y. A few plants were produced from this cross, and all had 19 chromosomes. Following self-fertilization, the F₁ hybrids produced a few F₂ plants, each with 24 chromosomes. These plants were phenotypically different from either of the original species and were highly fertile. Explain the sequence of events that produced these fertile F₂ hybrids.
- **6.7** Identify the sexual phenotypes of the following genotypes in humans: XX, XY, XO, XXX, XXY, XYY
- **6.8** If nondisjunction of chromosome 21 occurs in the division of a secondary oocyte in a human female, what is the chance that a mature egg derived from this division will receive two number 21 chromosomes?

- **6.9** A *Drosophila* female homozygous for a recessive X-linked mutation causing yellow body was crossed to a wild-type male. Among the progeny, one fly had sectors of yellow pigment in an otherwise gray body. These yellow sectors were distinctly male, whereas the gray areas were female. Explain the peculiar phenotype of this fly.
- **6.10** The *Drosophila* fourth chromosome is so small that flies monosomic or trisomic for it survive and are fertile. Several genes, including *eyeless* (*ey*), have been located on this chromosome. If a cytologically normal fly homozygous for a recessive eyeless mutation is crossed to a fly monosomic for a wild-type fourth chromosome, what kinds of progeny will be produced, and in what proportions?
- **6.11** A woman with X-linked color blindness and Turner syndrome had a color-blind father and a normal mother. In which of her parents did nondisjunction of the sex chromosomes occur?
- 6.12 **••** In humans, Hunter syndrome is known to be an X-linked trait with complete penetrance. In family A, two phenotypically normal parents have produced a normal son, a *daughter* with Hunter and Turner syndromes, and a son with Hunter syndrome. In family B, two phenotypically normal parents have produced two phenotypically normal daughters and a *son* with Hunter and Klinefelter syndromes. In family C, two phenotypically normal parents have produced a phenotypically normal daughter, a *daughter* with Hunter syndrome, and a son with Hunter syndrome. For each family, explain the origin of the child indicated in italics.
- **6.13** Although XYY men are phenotypically normal, would they be expected to produce more children with sex chromosome abnormalities than XY men? Explain.
- **6.14** In a *Drosophila* salivary chromosome, the bands have a sequence of 1 2 3 4 5 6 7 8. The homologue with which this chromosome is synapsed has a sequence of 1 2 3 6 5 4 7 8. What kind of chromosome change has occurred? Draw the synapsed chromosomes.
- **6.15** Other chromosomes have sequences as follows: (a) 1 2 5 6 7 8; (b) 1 2 3 4 4 5 6 7 8; (c) 1 2 3 4 5 8 7 6. What kind of chromosome change is present in each? Illustrate how these chromosomes would pair with a chromosome whose sequence is 1 2 3 4 5 6 7 8.
- **6.16** In plants translocation heterozygotes display about 50 percent pollen abortion. Why?
- **6.17** One chromosome in a plant has the sequence A B C D E F, and another has the sequence M N O P Q R. A reciprocal translocation between these chromosomes produced the following arrangement: A B C P Q R on one chromosome and M N O D E F on the other. Illustrate how these translocated chromosomes would pair with their normal counterparts in a heterozygous individual during meiosis.
- 6.18 In *Drosophila*, the genes *bw* and *st* are located on chromosomes 2 and 3, respectively. Flies homozygous for *bw*

mutations have brown eyes, flies homozygous for *st* mutations have scarlet eyes, and flies homozygous for *bw* and *st* mutations have white eyes. Doubly heterozygous males were mated individually to homozygous *bw*; *st* females. All but one of the matings produced four classes of progeny: wild-type, and brown-, scarlet- and white-eyed. The single exception produced only wild-type and white-eyed progeny. Explain the nature of this exception.

- **6.19** A phenotypically normal boy has 45 chromosomes, but his sister, who has Down syndrome, has 46. Suggest an explanation for this paradox.
- **6.20** Distinguish between a compound chromosome and a Robertsonian translocation.
- **6.21** A yellow-bodied *Drosophila* female with attached-X chromosomes was crossed to a white-eyed male. Both of the parental phenotypes are caused by X-linked recessive mutations. Predict the phenotypes of the progeny.
- **6.22** A man has attached chromosomes 21. If his wife is cytologically normal, what is the chance their first child will have Down syndrome?
- **6.23** Analysis of the polytene chromosomes of three populations of *Drosophila* has revealed three different banding sequences in a region of the second chromosome:

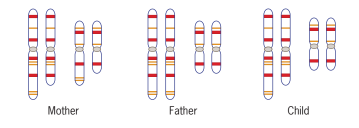
Population	Banding Sequence	
P1	1 2 3 4 5 6 7 8 9 10	
P2	1 2 3 9 8 7 6 5 4 10	
P3	1 2 3 9 8 5 6 7 4 10	

Explain the evolutionary relationships among these populations.

- **6.24** Each of six populations of *Drosophila* in different geographic regions had a specific arrangement of bands in one of the large autosomes:
 - (a) 12345678
 - (b) 12263478
 - (c) 15432678
 - (d) 14322678
 - (e) 16223478
 - (f) 154322678

Assume that arrangement (a) is the original one. In what order did the other arrangements most likely arise, and what type of chromosomal aberration is responsible for each change?

6.25 The following diagram shows two pairs of chromosomes in the karyotypes of a man, a woman, and their child. The man and the woman are phenotypically normal, but the child (a boy) suffers from a syndrome of abnormalities, including poor motor control and severe mental impairment. What is the genetic basis of the child's abnormal phenotype? Is the child hyperploid or hypoploid for a segment in one of his chromosomes?



- 6.26 a male mouse that is heterozygous for a reciprocal translocation between the X chromosome and an autosome is crossed to a female mouse with a normal karyotype. The autosome involved in the translocation carries a gene responsible for coloration of the fur. The allele on the male's translocated autosome is wild-type, and the allele on its nontranslocated autosome is mutant; however, because the wild-type allele is dominant to the mutant allele, the male's fur is wild-type (dark in color). The female mouse has light color in her fur because she is homozygous for the mutant allele of the color-determining gene. When the offspring of the cross are examined, all the males have light fur and all the females have patches of light and dark fur. Explain these peculiar results.
- 6.27 In *Drosophila*, the autosomal genes *cinnabar* (*cn*) and *brown* (*bw*) control the production of brown and red eye pigments, respectively. Flies homozygous for *cinnabar* mutations have bright red eyes, flies homozygous for *brown* mutations have brown eyes, and flies homozygous for mutations in both of these genes have white eyes. A male homozygous for mutations in the *cn* and *bw* genes has bright red eyes because a small duplication that carries the wild-type allele of *bw* (*bw*⁺) is attached to the Y chromosome. If this male is mated to a karyotypically normal female that is homozygous for the *cn* and *bw* mutations, what types of progeny will be produced?
- **6.28** In *Drosophila*, vestigial wing (vg), hairy body (b), and eyeless (ey) are recessive mutations on chromosomes 2, 3, and 4, respectively. Wild-type males that had been irradiated with X rays were crossed to triply homozygous recessive females. The F₁ males (all phenotypically wild-type) were then testcrossed to triply homozygous recessive females. Most of the F₁ males produced eight classes of progeny in approximately equal proportions, as would be expected if the vg, b, and ey genes assort independently. However, one F₁ male produced only four classes of offspring, each approximately one-fourth of the total: (1) wild-type, (2) eyeless, (3) vestigial, hairy, and (4) vestigial, hairy, eyeless. What kind of chromosome aberration did the exceptional F₁ male carry, and which chromosomes were involved?
- **6.29** Cytological examination of the sex chromosomes in a man has revealed that he carries an insertional translocation. A small segment has been deleted from the Y chromosome and inserted into the short arm of the X chromosome; this segment contains the gene responsible for male differentiation (*SRY*). If this man marries a karyotypically normal woman, what types of progeny will the couple produce?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

1. Many crop plants are polyploid. What progress has been made in sequencing the polyploid genomes of soybean (*Glycine max*), wheat (*Triticum aestivum*), and potato (*Solanum tuberosum*)?

Hint: At the web site, click on Genomes and Maps, then on Genome Project, and finally on Plant Genomes. Find each species and read about ongoing DNA sequencing efforts.

2. When triplicated, chromosome 21, the smallest of the autosomes in the human genome, causes Down syndrome. How many nucleotide pairs are present in this chromosome? How many genes does it contain?

Hint: Use Map Viewer to find chromosome 21 and then determine its size and gene content.

3. The gene for amyloid precursor protein, APP, is located on human chromosome 21. This protein appears to play an important role in the etiology of Alzheimer's disease. Locate

the *APP* gene on the ideogram of human chromosome 21. In what band does it lie?

Hint: Search for APP using the "Find in This View" function. Click on the highlighted gene name to find more information about it.

- 4. Chromosome 21 as well as a few other chromosomes in the human genome have secondary constrictions as well as a primary constriction, which is situated at the centromere. The material distal to the secondary constriction—that is, going away from the centromere toward the nearest end of the chromosome—is called a satellite. Find the secondary constriction and the satellite on the ideogram of chromosome 21.
- **5.** Secondary constrictions on some chromosomes contain genes for ribosomal RNA. Is this true for human chromosome 21?

Hint: Use the Map Viewer function to examine the ideogram of chromosome 21. Search for ribosomal RNA genes using the "Find in This View" function.

Linkage, Crossing Over, and Chromosome Mapping in Eukaryotes



CHAPTER OUTLINE

- Linkage, Recombination, and Crossing Over
- Chromosome Mapping
- Cytogenetic Mapping
- Linkage Analysis in Humans
- Recombination and Evolution

The World's First Chromosome Map

The modern picture of chromosome organization emerged from a combination of genetic and cytological studies. T. H. Morgan laid the foundation for these studies when he demonstrated that the gene for *white* eyes in *Drosophila* was located on the X chromosome. Soon afterward Morgan's students showed that other genes were X-linked, and eventually they were able to locate each of these genes on a map of the chromosome. This map was a straight line, and each gene was situated at a particular point, or locus, on it [**Figure 7.1**]. The structure of the map therefore implied that a chromosome was simply a linear array of genes.

The procedure for mapping chromosomes was invented by Alfred H. Sturtevant, an undergraduate working in Morgan's laboratory. One night in 1911 Sturtevant put aside his algebra homework in order to evaluate some experimental data. Before the sun rose the next day, he had constructed the world's first chromosome map. How was Sturtevant able to determine the map locations of individual genes? No microscope was powerful enough to see genes, nor was any measuring device accurate enough to obtain the distances between them. In fact, Sturtevant did not use any sophisticated instruments in his work. Instead, he relied completely on the analysis of data from experimental crosses with *Drosophila*. His method was simple and elegant, and exploited a phenomenon that regularly occurs during meiosis. This methodology laid the foundation for all subsequent efforts to study the organization of genes in chromosomes.



Linkage between genes was first discovered in experiments with sweet peas.

Linkage, Recombination, and Crossing Over

Genes that are on the same chromosome travel through meiosis together; however, alleles of chromosomally linked genes can be recombined by crossing over. Sturtevant based his mapping procedure on the principle that genes on the same chromosome should be inherited together. Because such genes are physically attached to the same structure, they should travel as a unit through meiosis. This phenomenon is called

linkage. The early geneticists were unsure about the nature of linkage, but some of them, including Morgan and his students, thought that genes were attached to one another much like beads on a string. Thus, these researchers clearly had a linear model of chromosome organization in mind.

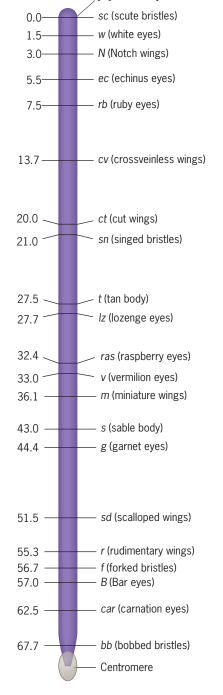
The early geneticists also knew that linkage was not absolute. Their experimental data demonstrated that genes on the same chromosome could be separated as they went through meiosis and that new combinations of genes could be formed. However, this phenomenon, called **recombination**, was difficult to explain by simple genetic theory.

One hypothesis was that during meiosis, when homologous chromosomes paired, a physical exchange of material separated and recombined genes. This idea was inspired by the cytological observation that chromosomes could be seen in pairing configurations that suggested they had switched pieces with each other. At the switch points, the two homologues were crossed over, as if each had been broken and then reattached to its partner. A crossover point was called a **chiasma** (plural, **chiasmata**), from the Greek word meaning "cross." Geneticists began to use the term *crossing over* to describe the process that created the chiasmata—that is, the actual process of exchange between paired chromosomes. They considered recombination—the separation of linked genes and the formation of new gene combinations—to be a result of the physical event of crossing over.

EARLY EVIDENCE FOR LINKAGE AND RECOMBINATION

Some of the first evidence for linkage came from experiments performed by W. Bateson and R. C. Punnett (**Figure 7.2**). These researchers crossed varieties of sweet peas that differed in two traits, flower color and pollen length. Plants with red flowers and long pollen grains were crossed to plants with white flowers and short pollen grains. All the F₁ plants had red flowers and long pollen grains, indicating that the alleles for these two phenotypes were dominant. When the F_1 plants were self-fertilized, Bateson and Punnett observed a peculiar distribution of phenotypes among the offspring. Instead of the 9:3:3:1 ratio expected for two independently assorting genes, they obtained a ratio of 24.3:1.1:1:7.1. We can see the extent of the disagreement between the observed results and the expected results at the bottom of Figure 7.2. Among the 803 F₂ plants that were examined, the classes that resembled the original parents (called the parental classes) are significantly overrepresented and the two other (nonparental) classes are significantly underrepresented. For such obvious discrepancies, it hardly seems necessary to calculate a chi-square statistic to test the hypothesis that the two traits, flower color and pollen grain length, have assorted independently. Clearly they have not. Nevertheless, we have included the chi-square calculation in Figure 7.2 just to show how much the observed results are out of line with the expected results. The chi-square value is enormous—much greater than 7.8, which is the critical value for a chi-square distribution with three degrees of freedom (see Table 3.2). Consequently, we must reject the hypothesis that the genes for flower color and pollen grain length have assorted independently.

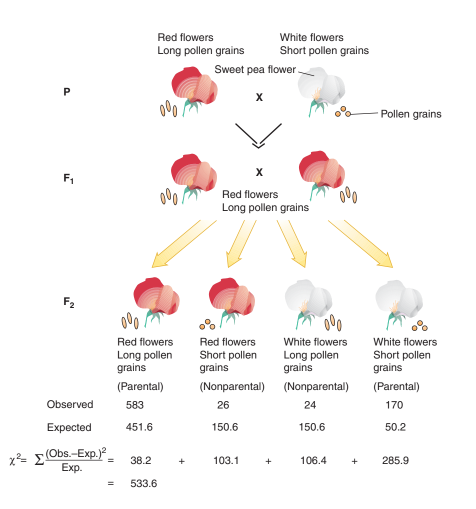
Bateson and Punnett devised a complicated explanation for their results, but it turned out to be wrong. The correct explanation for the lack of independent assortment in the data is that the genes for flower color and pollen length are located on the same chromosome—that is, they are linked. This explanation is diagrammed in



y (yellow body)

FIGURE 7.1 A map of genes on the X chromosome of *Drosophila melanogaster*.

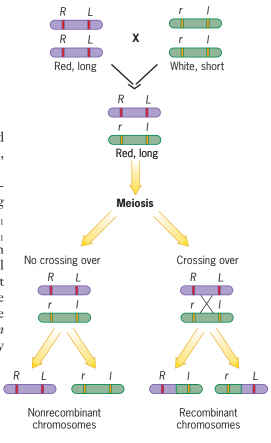
FIGURE 7.2 Bateson and Punnett's experiment with sweet peas. The results in the F_2 indicate that the genes for flower color and pollen length do not assort independently.



■ Figure 7.3. The alleles of the flower color gene are R (red) and r (white), and the alleles of the pollen length gene are L (long) and l **P** (short); the R and L alleles are dominant. (Note here that for historical reasons, the allele symbols are derived from the dominant rather than the recessive phenotypes.) Because the flower color and pollen length genes are linked, we expect the doubly heterozygous F_1 plants to produce two kinds of gametes, R L and r l. However, F_1 once in a while a crossover will occur between the two genes and their alleles will be recombined, producing two other kinds of gametes, R l and r L. The frequency of these two types of recombinant gametes should, of course, depend on the frequency of crossing over between the two genes.

Bateson and Punnett might have come up with this explanation if they had performed a testcross instead of an intercross in the F_1 . With a testcross the offspring would directly reveal the types of gametes produced by the doubly heterozygous F_1 plants. **Figure 7.4** presents the analysis of such a testcross. Doubly heterozygous F_1 sweet peas were crossed with plants homozygous for the recessive alleles of both genes. Among 1000 progeny scored, 920 resemble one or the other of the parental strains and the remaining 80 are recombinant. The frequency of the recombinant progeny produced by the heterozygous F_1 plants is therefore 80/1000 = 0.08. Because this is a testcross, 0.08 is also the frequency of recombinant gametes produced by the heterozygous F_1 plants. We can use this frequency, usually called the *recombination frequency*, to measure the intensity of linkage between genes. Genes that are tightly

FIGURE 7.3 Hypothesis of linkage between the genes for flower color and pollen length in sweet peas. In the F_1 plants the two dominant alleles, R and L, of the genes are situated on the same chromosome; their recessive alleles, r and l, are situated on the homologous chromosome.



F₁ Gametes

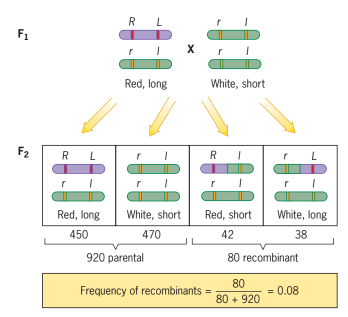


FIGURE 7.4 A testcross for linkage between genes in sweet peas. Because the recombinant progeny in the F_2 are 8 percent of the total, the genes for flower color and pollen length are rather tightly linked.

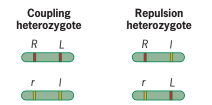


FIGURE 7.5 Coupling and repulsion linkage phases in double heterozygotes.

linked seldom recombine, whereas genes that are loosely linked recombine often. Here the recombination frequency is fairly low. This implies that crossing over between the two genes is a rather rare event.

For any two genes, the recombination frequency never exceeds 50 percent. This upper limit is obtained when genes are on different chromosomes; 50 percent recombination is, in fact, what we mean when we say that the genes assort independently. For example, let's assume that genes A and B are on different chromosomes and that an AA BB individual is crossed to an *aa bb* individual. From this cross the *Aa Bb* offspring are then testcrossed to the double recessive parent. Because the A and B genes assort independently, the F_2 will consist of two classes (*Aa Bb* and *aa bb*) that are phenotypically like the parents in the original cross and two classes (*Aa bb* and *aa Bb*) that are phenotypically recombinant. Furthermore, each F_2 class will occur with a frequency of 25 percent (see Figure 5.7). Thus, the total frequency of recombinant progeny from a testcross involving two genes on different chromosomes will be 50 percent. A frequency of recombination less than 50 percent implies that the genes are linked on the same chromosome.

Crosses involving linked genes are usually diagrammed to show the **linkage phase**—the way in which the alleles are arranged in heterozygous individuals (**Figure 7.5**). In Bateson and Punnett's sweet pea experiment,

the heterozygous F_1 plants received two dominant alleles, R and L, from one parent and two recessive alleles, r and l, from the other. Thus, we write the genotype of these plants R L/r l, where the slash (/) separates alleles inherited from different parents. Another way of interpreting this symbolism is to say that the alleles on the left and right of the slash entered the genotype on different homologous chromosomes, one from each parent. Whenever the dominant alleles are all on one side of the slash, as in this example, the genotype has the *coupling* linkage phase. When the dominant and recessive alleles are split on both sides of the slash, as in R l/r L, the genotype has the *repulsion* linkage phase. These terms provide us with a way of distinguishing between the two kinds of double heterozygotes.

CROSSING OVER AS THE PHYSICAL BASIS OF RECOMBINATION

Recombinant gametes are produced as a result of crossing over between homologous chromosomes. This process involves a physical exchange between the chromosomes, as diagrammed in **Figure 7.6**. The exchange event occurs during the prophase of the first meiotic division, when duplicated chromosomes have paired. Although four homologous chromatids are present, forming what is called a **tetrad**, only two chromatids cross over at any one point. Each of these chromatids breaks at the site of the

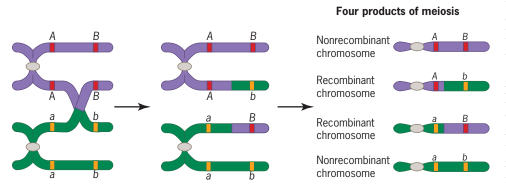


FIGURE 7.6 Crossing over as the basis of recombination between genes. An exchange between paired chromosomes during meiosis produces recombinant chromosomes at the end of meiosis.

crossover, and the resulting pieces reattach to produce the recombinants. The other two chromatids are not recombinant at this site. Each crossover event therefore produces two recombinant chromatids among a total of four.

Notice that only two chromatids are involved in an exchange at any one point. However, the other two chromatids may cross over at a different point. Thus, there is a possibility for multiple exchanges in a tetrad of chromatids (**Figure 7.7**). There may, for example, be two, three, or even four separate exchanges—customarily called double, triple, or quadruple crossovers. (We consider the genetic significance of these in a later section of this chapter.) Note,

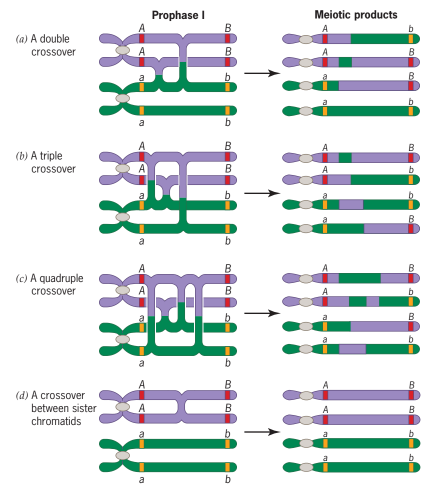


FIGURE 7.7 Consequences of multiple exchanges between chromosomes and exchange between sister chromatids during prophase I of meiosis.

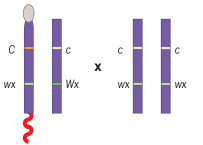
however, that an exchange between sister chromatids does not produce genetic recombinants because the sister chromatids are identical.

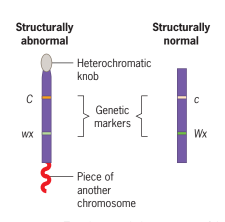
What is responsible for the breakage of chromatids during crossing over? The breaks are caused by enzymes acting on the DNA within the chromatids. Enzymes are also responsible for repairing these breaks—that is, for reattaching chromatid fragments to each other. We consider the molecular details of this process in Chapter 13.

EVIDENCE THAT CROSSING OVER CAUSES RECOMBINATION

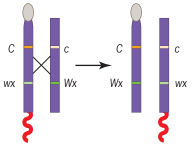
In 1931 Harriet Creighton and Barbara McClintock obtained evidence that genetic recombination was associated with a material exchange between chromosomes. Creighton and McClintock studied homologous chromosomes in maize that were morphologically distinguishable. The goal was to determine whether physical exchange between these homologues was correlated with recombination between some of the genes they carried.

Two forms of chromosome 9 were available for analysis; one was normal, and the other had cytological aberrations at each end—a heterochromatic knob at one end and a piece of a different chromosome at the other (**Trigure 7.8**). These two forms of chromosome 9 were also genetically marked to detect recombination. One marker gene controlled kernel color (C, colored; c, colorless), and the other controlled kernel texture (Wx, starchy; wx, waxy). Creighton and McClintock performed the following testcross:





They then examined the recombinant progeny for evidence of exchange between the two different forms of chromosome 9. Their results showed that the C Wx and c wx recombinants carried a chromosome with only one of the abnormal cytological markers; the other abnormal marker had evidently been lost through an exchange with the normal chromosome 9 in the previous generation:



■ FIGURE 7.8 Two forms of chromosome 9 in maize used in the experiments of Creighton and McClintock.

These findings strongly argued that recombination was caused by a physical exchange between paired chromosomes.

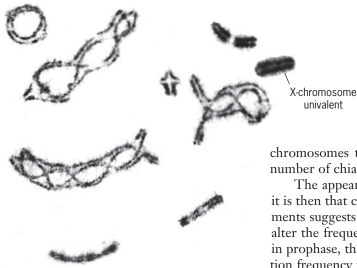


FIGURE 7.9 Diplonema of male meiosis in the grasshopper *Chorthippus parallelus*. There are eight autosomal bivalents and an X-chromosome univalent. The four smaller bivalents each have one chiasma. The remaining bivalents have two to five chiasmata.

CHIASMATA AND THE TIME OF CROSSING OVER

The cytological evidence for crossing over can be seen during late prophase of the first meiotic division when the chiasmata become clearly visible. At this time paired chromosomes repel each other slightly, maintaining close contact only at the centromere and at each chiasma (**Figure 7.9**). This partial separation makes it possible to count the chiasmata accurately. As we might expect, large

chromosomes typically have more chiasmata than small chromosomes. Thus, the number of chiasmata is roughly proportional to chromosome length.

The appearance of chiasmata late in the first meiotic prophase might imply that it is then that crossing over occurs. However, evidence from several different experiments suggests that it occurs earlier. Some of these experiments used heat shocks to alter the frequency of recombination. When the heat shocks were administered late in prophase, there was little effect, but when they were given earlier, the recombination frequency was changed. Thus, the event responsible for recombination, namely, crossing over, occurs rather early in the meiotic prophase. Additional evidence comes from molecular studies on the time of DNA synthesis. Although almost all the DNA is synthesized during the interphase that precedes the onset of meiosis, a small amount is made during the first meiotic prophase. This limited DNA synthesis has been interpreted as part of a process to repair broken chromatids, which, as we have discussed, is thought to be associated with crossing over. Careful timing experiments have shown that this DNA synthesis occurs in early to mid-prophase, but not later. The accumulated evidence therefore suggests that crossing over occurs in early to mid-prophase, long before the chiasmata can be seen.

What, then, are chiasmata, and what do they mean? Most geneticists believe that the chiasmata are merely vestiges of the actual exchange process. Chromatids that have experienced an exchange probably remain entangled with each other during most of prophase. Eventually, these entanglements are resolved, and the chromatids are separated by the meiotic spindle apparatus to opposite poles of the cell. Therefore, each chiasma probably represents an entanglement that was created by a crossover event earlier in prophase.

But why do these entanglements occur at all? Many geneticists believe that the entanglements created by crossing over are a way of holding the members of a bivalent together during prophase I. In some organisms, prophase I is protracted. In human females, for example, it may last as long as 40 years. Without crossovers, paired homologues might accidentally separate from each other during this long time, and homologues that have separated might not disjoin properly during the ensuing anaphase. Faulty disjunction of the chromosomes during the first meiotic division would ultimately lead to aneuploid gametes. Thus, crossing over appears to be a mechanism to hold paired homologues together so that when division does occur, the homologues are distributed appropriately to each of the daughter cells. In this way, then, the possibility for nondisjunction is minimized, and aneuploidy in the gametes is largely prevented.

KEY POINTS

- Linkage between genes is detected as a deviation from expectations based on Mendel's Principle of Independent Assortment.
- The frequency of recombination measures the intensity of linkage. In the absence of linkage, this frequency is 50 percent; for very tight linkage, it is close to zero.
- Recombination is caused by a physical exchange between paired homologous chromosomes early in prophase of the first meiotic division after chromosomes have duplicated.
- At any one point along a chromosome, the process of exchange (crossing over) involves only two of the four chromatids in a meiotic tetrad.
- Late in prophase I, crossovers become visible as chiasmata.

Chromosome Mapping

Crossing over during the prophase of the first meiotic division has two observable outcomes:

Linked genes can be mapped on a chromosome by studying how often their alleles recombine.

1

- **1.** Formation of chiasmata in late prophase.
- 2. Recombination between genes on opposite sides of the crossover point.

However, the second outcome can only be seen in the next generation, when the genes on the recombinant chromosomes are expressed.

Geneticists construct chromosome maps by counting the number of crossovers that occur during meiosis. However, because the actual crossover events cannot be seen, they cannot count them directly. Instead, they must estimate how many crossovers have taken place by counting either chiasmata or recombinant chromosomes. Chiasmata are counted through cytological analysis, whereas recombinant chromosomes are counted through genetic analysis. Before we proceed further, we must define what we mean by distance on a chromosome map.

CROSSING OVER AS A MEASURE OF GENETIC DISTANCE

Sturtevant's fundamental insight was to estimate the distance between points on a chromosome by counting the number of crossovers between them. Points that are far apart should have more crossovers between them than points that are close together. However, the number of crossovers must be understood in a statistical sense. In any particular cell, the chance that a crossover will occur between two points may be low, but in a large population of cells, this crossover will probably occur several times simply because there are so many independent opportunities for it. Thus,

the quantity that we really need to measure is the *average* number of crossovers in a particular chromosome region. Genetic map distances are, in fact, based on such averages. This idea is sufficiently important to justify a formal definition: The distance between two points on the genetic map of a chromosome is the average number of crossovers between them.

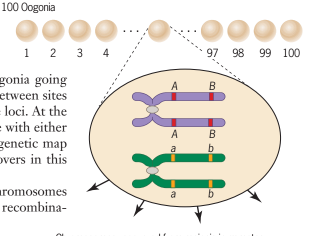
One way for us to understand this definition is to consider 100 oogonia going through meiosis (**Figure 7.10**). In some cells, no crossovers will occur between sites A and B; in others, one, two, or more crossovers will occur between these loci. At the end of meiosis, there will be 100 gametes, each containing a chromosome with either zero, one, two, or more crossovers between A and B. We estimate the genetic map distance between these loci by calculating the average number of crossovers in this sample of chromosomes. The result from the data in Figure 7.10 is 0.42.

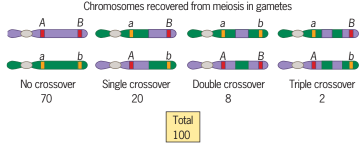
In practice, we cannot "see" each of the exchange points on the chromosomes coming out of meiosis. Instead, we infer their existence by observing the recombina-

tion of the alleles that flank them. A chromosome in which alleles have recombined must have arisen by crossing over. Counting recombinant chromosomes therefore provides a way of counting crossover exchange points.

RECOMBINATION MAPPING WITH A TWO-POINT TESTCROSS

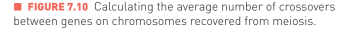
To illustrate the mapping procedure, let's consider the two-point testcross in Figure 7.11. Wild-type *Drosophila* females were mated to males homozygous for two autosomal mutations—vestigial (vg), which produces short wings, and *black* (b), which produces a black body. All the F_1 flies had long wings and gray bodies; thus, the wild-type alleles (vg^+ and b^+) are dominant. The F₁ females were then testcrossed to vestigial, black males, and the F₂ progeny were





Average number of crossovers between A and B =

$$0 \times \left(\frac{70}{100}\right) + 1 \times \left(\frac{20}{100}\right) + 2 \times \left(\frac{8}{100}\right) + 3 \times \left(\frac{2}{100}\right) = 0.42$$



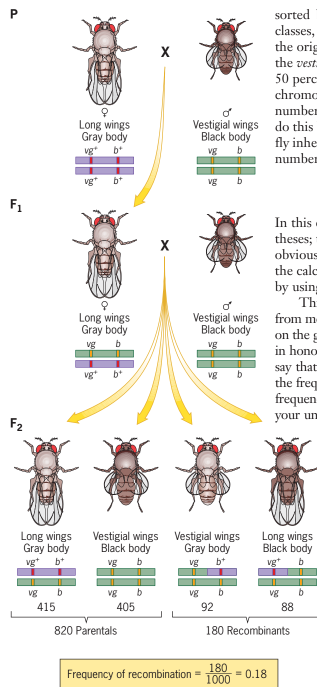


FIGURE 7.11 An experiment involving two linked genes, *vg* (*vestigial* wings) and *b* (*black* body), in *Drosophila*.

sorted by phenotype and counted. As the data show, there were four phenotypic classes, two abundant and two rare. The abundant classes had the same phenotypes as the original parents, and the rare classes had recombinant phenotypes. We know that the *vestigial* and *black* genes are linked because the recombinants are much fewer than 50 percent of the total progeny counted. These genes must therefore be on the same chromosome. To determine the distance between them, we must estimate the average number of crossovers in the gametes of the doubly heterozygous F_1 females. We can do this by calculating the frequency of recombinant F_2 flies and noting that each such fly inherited a chromosome that had crossed over once between vg and b. The average number of crossovers in the whole sample of progeny is therefore

nonrecombinants recombinants (0) \times 0.82 + (1) \times 0.18 = 0.18

In this expression, the number of crossovers for each class of flies is placed in parentheses; the other number is the frequency of that class. The nonrecombinant progeny obviously do not add any crossover chromosomes to the data, but we include them in the calculation to emphasize that we must calculate the average number of crossovers by using all the data, not just those from the recombinants.

This simple analysis indicates that, on average, 18 out of 100 chromosomes recovered from meiosis had a crossover between vg and b. Thus, vg and b are separated by 18 units on the genetic map. Sometimes geneticists call a map unit a **centiMorgan**, abbreviated cM, in honor of T. H. Morgan; 100 centiMorgans equal one Morgan (M). We can therefore say that vg and b are 18 cM (or 0.18 M) apart. Notice that the map distance is equal to the frequency of recombination, written as a percentage. Later we will see that when the frequency of recombination approaches 0.5, it underestimates the map distance. To test your understanding of the principles underlying recombination mapping, work through

the exercise in Solve It: Mapping Two Genes with Testcross Data.

RECOMBINATION MAPPING WITH A THREE-POINT TESTCROSS

We can also use the recombination mapping procedure with data from testcrosses involving more than two genes. **Figure 7.12** illustrates an experiment by C. B. Bridges and T. M. Olbrycht, who crossed wild-type *Drosophila* males to females homozygous for three recessive X-linked mutations—*scute* (*sc*) bristles, *echinus* (*ec*) eyes, and *crossveinless* (*cv*) wings. They then intercrossed the F_1 progeny to produce F_2 flies, which they classified and counted. We note that the F_1 females in this intercross carried the three recessive mutations on one of their X chromosomes and the wild-type alleles of these mutations on the other X chromosome. Furthermore, the F_1 males carried the three recessive mutations on their single X chromosome. Thus, this intercross was equivalent to a testcross with all three genes in the F_1 females present in the coupling configuration.

The \vec{F}_2 flies from the intercross comprised eight phenotypically distinct classes, two of them parental and six recombinant. The parental

classes were by far the most numerous. The less numerous recombinant classes each represented a different kind of crossover chromosome. To figure out which crossovers were involved in producing each type of recombinant, we must first determine how the genes are ordered on the chromosome.

Determining the Gene Order

There are three possible gene orders:

- **1.** *sc*—*ec*—*cv*
- **2.** *ec*—*sc*—*cv*
- **3.** *ec*—*cv*—*sc*

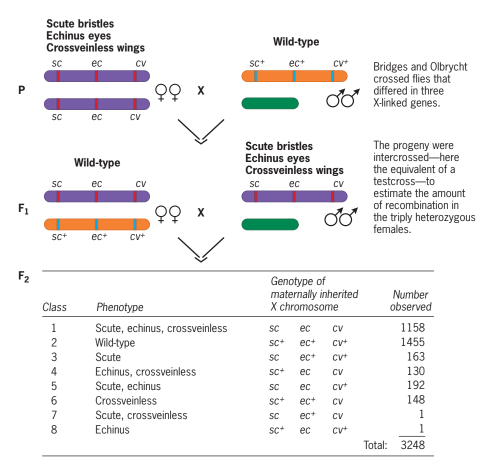


FIGURE 7.12 Bridges and Olbrycht's three-point cross with the X-linked genes *sc* (*scute* bristles), *ec* (*echinus* eyes), and *cv* (*crossveinless* wings) in *Drosophila*. Data from Bridges, C. B., and Olbrycht, T. M., 1926. *Genetics* 11: 41.

Other possibilities, such as cv-ec-sc, are the same as one of these because the left and right ends of the chromosome cannot be distinguished. Which of the orders is correct?

To answer this question, we must take a careful look at the six recombinant classes. Four of them must have come from a single crossover in one of the two regions delimited by the genes. The other two must have come from double crossing over—one exchange in each of the two regions. Because a double crossover switches the gene in the middle with respect to the genetic markers on either side of it, we have, in principle, a way of determining the gene order. Intuitively, we also know that a double crossover should occur much less frequently than a single crossover. Consequently, among the six recombinant classes, the two rare ones must represent the double crossover chromosomes.

In our data, the rare, double crossover classes are 7 (sc $ec^+ cv$) and 8 (sc⁺ $ec cv^+$), each containing a single fly (Figure 7.12). Comparing these to parental classes 1 (sc ec cv) and 2 (sc⁺ $ec^+ cv^+$), we see that the *echinus* allele has been switched with respect to *scute* and *crossveinless*. Consequently, the *echinus* gene must be located between the other two. The correct gene order is therefore (1) sc - ec - cv.

Calculating the Distances between Genes

Having established the gene order, we can now determine the distances between adjacent genes. Again, the procedure is to compute the average number of crossovers in each chromosomal region (**■** Figure 7.13).

Solve It!

Mapping Two Genes with Testcross Data

In maize, the gene for leaf color has two alleles, recessive q for green leaves and dominant G for purple leaves, and the gene for stalk height has two alleles, recessive *s* for short stalk and dominant *S* for tall stalk. A plant with green leaves and a short stalk was crossed to a plant with purple leaves and a tall stalk. All the F_1 plants from this cross had purple leaves and tall stalks. When they were backcrossed to plants with green leaves and short stalks, they produced an F_2 in which, among a total of 200 plants, four phenotypic classes were observed: (1) green leaves, short stalks, 75; (2) purple leaves, tall stalks, 79; (3) green leaves, tall stalks, 24; and (4) purple leaves, short stalks, 22. (a) What is the evidence that the genes for leaf color and stalk height are linked? (b) What is the linkage phase of the dominant and recessive alleles of these two genes in the F₁ plants? (c) Among the F₂ plants, what is the frequency of recombination? (d) What is the distance in centiMorgans between the leaf color and stalk height genes?

To see the solution to this problem, visit the Student Companion site.

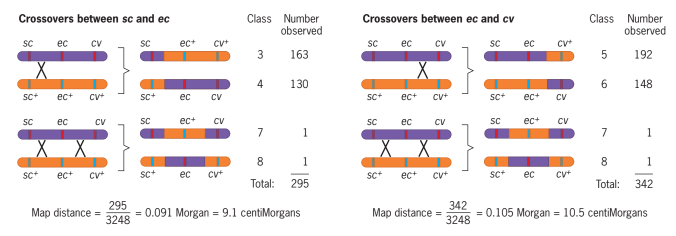


FIGURE 7.13 Calculation of genetic map distances from Bridges and Olbrycht's data. The distance between each pair of genes is obtained by estimating the average number of crossovers.

We can obtain the length of the region between *sc* and *ec* by identifying the recombinant classes that involved a crossover between these genes. There are four such classes: 3 (*sc* $ec^+ cv^+$), 4 ($sc^+ ec cv$), 7 (*sc* $ec^+ cv$), and 8 ($sc^+ ec cv^+$). Classes 3 and 4 involved a single crossover between *sc* and *ec*, and classes 7 and 8 involved two crossovers, one between *sc* and *ec* and the other between *ec* and *cv*. We can therefore use the frequencies of these four classes to estimate the average number of crossovers between *sc* and *ec*:

Class 3 Class 4 Class 7 Class 8

$$\frac{163 + 130 + 1 + 1}{\text{Total}} = \frac{295}{3248} = 0.091$$

Thus, in every 100 chromosomes coming from meiosis in the F_1 females, 9.1 had a crossover between *sc* and *ec*. The distance between these genes is therefore 9.1 map units (or, if you prefer, 9.1 centiMorgans).

In a similar way, we can obtain the distance between *ec* and *cv*. Four recombinant classes involved a crossover in this region: 5 (*sc ec cv*⁺), 6 (*sc*⁺ *ec*⁺ *cv*), 7, and 8. The double recombinants are also included here because one of their two crossovers was between *ec* and *cv*. The combined frequency of these four classes is:

Class 5 Class 6 Class 7 Class 8

$$\frac{192 + 148 + 1 + 1}{\text{Total}} = \frac{342}{3248} = 0.105$$

Consequently, ec and cv are 10.5 map units apart.

Combining the data for the two regions, we obtain the map

Map distances computed in this way are additive. Thus, we can estimate the distance between sc and cv by summing the lengths of the two map intervals between them:

$$9.1 \text{ cM} + 10.5 \text{ cM} = 19.6 \text{ cM}$$

We can also obtain this estimate by directly calculating the average number of crossovers between these genes:

Non-crossover	Single crossover	Double crossover	
classes	classes	classes	
1 and 2	3, 4, 5, and 6	7 and 8	
$(0) \times 0.805$	+ $(1) \times 0.195$	+ $(2) \times 0.0006$	= 0.196

Here the number of crossovers is given in parentheses, and its multiplier is the combined frequency of the classes with that many crossovers. In other words, each recombinant class contributes to the map distance according to the product of its frequency and the number of crossovers it represents.

Bridges and Olbrycht actually studied seven X-linked genes in their recombination experiment: *sc*, *ec*, *cv*, *ct* (*cut* wings), *v* (*vermilion* eyes), *g* (*garnet* eyes), and *f* (*forked* bristles). By calculating recombination frequencies between each pair of adjacent genes, they were able to construct a map of a large segment of the X chromosome (**True Figure 7.14**); *sc* was at one end, and *f* was at the other. Each of the seven genes that

Bridges and Olbrycht studied was, in effect, a *marker* for a particular site on the X chromosome. Summing all the map intervals between these markers, they estimated the total length of the mapped segment to be 66.8 cM. Thus, the average number of crossovers in this segment was 0.668.

Interference and the Coefficient of Coincidence

A three-point cross has an important advantage over a two-point cross: it allows the detection of double crossovers, permitting us to determine if exchanges in adjacent regions are independent of each other. For example, does a crossover in the region between sc and ec (region I on the map of the X chromosome) occur independently of a crossover in the region between ec and cv (region II)? Or does one crossover inhibit the occurrence of another nearby?

To answer these questions, we must calculate the expected frequency of double crossovers, based on the idea of independence. We can do this by multiplying the crossover frequencies for two adjacent chromosome regions. For example, in region I on Bridges and Olbrycht's map, the crossover frequency was (163 + 130 + 1 + 1)/3248 = 0.091, and in region II, it was (192 + 148 + 1 + 1)/3248 = 0.105. If we assume independence, the expected frequency of double crossovers in the interval between sc and cv would therefore be $0.091 \times 0.105 = 0.0095$. We can now compare this frequency with the observed frequency, which was 2/3248 = 0.0006. Double crossovers between sc and cv were much less frequent than expected. This result suggests that one crossover inhibited the occurrence of another nearby, a phenomenon called **interference**. The extent of the interference is customarily measured by the **coefficient of coincidence**, c, which is the ratio of the observed frequency of double crossovers to the expected frequency:

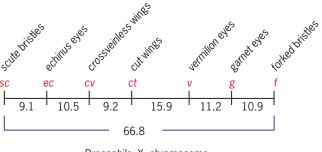
 $c = \frac{\text{observed frequency of double crossovers}}{\text{expected frequency of double crossovers}}$ $= \frac{0.0006}{0.0095} = 0.063$

The level of interference, symbolized *I*, is calculated as I = 1 - c = 0.937.

Because in this example the coefficient of coincidence is close to zero, its lowest possible value, interference was very strong (I is close to 1). At the other extreme, a coefficient of coincidence equal to one would imply no interference at all; that is, it would imply that the crossovers occurred independently of each other.

Many studies have shown that interference is strong over map distances less than 20 cM; thus, double crossovers seldom occur in short chromosomal regions. However, over long regions, interference weakens to the point that crossovers occur more or less independently. The strength of interference is therefore a function of map distance.

Once a genetic map has been constructed, it is possible to use the map to predict the results of experiments. To see how map-based predictions are made, work through the exercise in Problem-Solving Skills: Using a Genetic Map to Predict the Outcome of a Cross.



Drosophila X chromosome

FIGURE 7.14 Bridges and Olbrycht's map of seven X-linked genes in *Drosophila*. Distances are given in centiMorgans.

PROBLEM-SOLVING SKILLS

Using a Genetic Map to Predict the Outcome of a Cross

THE PROBLEM

The genes r, s, and t reside in the middle of the Drosophila X chromosome; r is 15 cM to the left of s, and t is 20 cM to the right of s. In this region, the coefficient of coincidence (c) is 0.2. A geneticist wishes to create an X chromosome that carries the recessive mutant alleles of all three genes. One stock is homozygous for r and t, and another stock is homozygous for s. By crossing the two stocks, the geneticist obtains females that are triple heterozygotes, $r s^+ t/t$ r^+ s t^+ . These females are then crossed to wild-type males. If the geneticist examines 10,000 sons from these females, how many of them will be triple mutants, *r s t*?

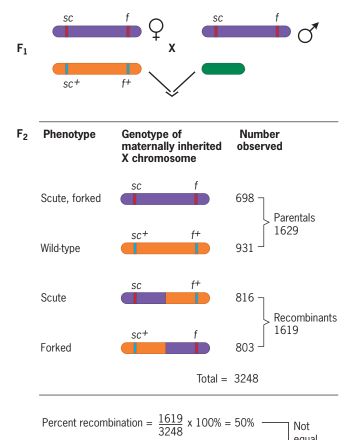
FACTS AND CONCEPTS

- 1. For small map intervals (<20 cM), the map distance equals the frequency of a single crossover in the interval.
- 2. The coefficient of coincidence equals the observed frequency of double crossovers/expected frequency of double crossovers.
- 3. The expected frequency of double crossovers is calculated on the assumption that the two crossovers occur independently.
- 4. Males inherit their X chromosome from their mothers.

ANALYSIS AND SOLUTION

Triple mutant males will be produced only if a double crossover occurs in the $r s^+ t/r^+ s t^+$ females that were crossed to wild-type males. The frequency of such double crossovers is a function of the two map distances (15 cM and 20 cM) and the level of interference, which is measured by the coefficient of coincidence (here c = 0.2). Because c = observed frequency of double crossovers/ expected frequency of double crossovers, we can solve for the observed frequency of double crossovers after a simple algebraic rearrangement: observed frequency of double crossovers = $c \times$ expected frequency of double crossovers. The expected frequency of double crossovers is calculated from the map distances assuming that crossovers in adjacent map intervals occur independently: $0.15 \times 0.20 = 0.03$. Thus, among 10,000 sons, 0.2×3 per cent should carry an X chromosome that had one crossover between the r and s genes and another crossover between the s and t genes. However, only half of these 60 sons—that is, 30—will carry the triply mutant X chromosome; the other 30 will be triply wild-type.

For further discussion visit the Student Companion site.



Map distance = 66.8 centiMorgans

equal

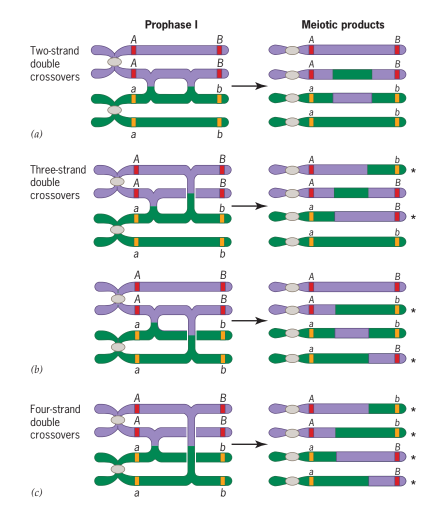
RECOMBINATION FREQUENCY AND GENETIC MAP DISTANCE

In the preceding sections, we have considered how to construct chromosome maps from data on the recombination of genetic markers. These data allow us to infer where crossovers have occurred in a sample of chromosomes. By localizing and counting these crossovers, we can estimate the distances between genes and then place the genes on a chromosome map.

This method works well as long as the genes are fairly close together. However, when they are far apart, the frequency of recombination may not reflect the true map distance (**Figure 7.15**). As an example, let's consider the genes at the ends of Bridges and Olbrycht's map of the X chromosome; sc, at the left end, was 66.8 cM away from f, at the right end. However, the frequency of recombination between sc and f was 50 percent—the maximum possible value. Using this frequency to estimate map distance, we would conclude that sc and f were 50 map units apart. Of course, the distance obtained by summing the lengths of the intervening regions on the map, 66.8 cM, is much greater.

This example shows that the true genetic distance, which depends on the average number of crossovers on a chromosome, may be much greater than the observed recombination frequency. Multiple crossovers may occur between widely separated genes, and some of these crossovers

FIGURE 7.15 A discrepancy between map distance and percent recombination. The map distance between the genes *sc* and *f* is greater than the observed percent recombination between them.

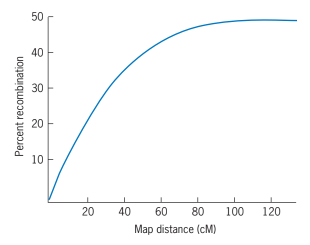


■ FIGURE 7.16 Consequences of double crossing over between two loci. Recombinant chromosomes are denoted by an asterisk. (a) Two-strand double crossovers produce only nonrecombinant chromosomes. (b) Three-strand double crossovers produce half recombinant and half nonrecombinant chromosomes. (c) Four-strand double crossovers produce only recombinant chromosomes.

may not produce genetically recombinant chromosomes (**Figure 7.16**). To see this, let's assume that a single crossover occurs between two chromatids in a tetrad, causing recombination of the flanking genetic markers. If another cross-over occurs between these same two chromatids, the flanking markers will be restored to their original configuration; the second crossover essentially cancels the effect of the first, converting the recombinant chromatids back into non-recombinants. Thus, even though two crossovers have occurred in this tetrad, none of the chromatids that come from it will be recombinant for the flanking markers.

This second example shows that a double crossover may not contribute to the frequency of recombination, even though it contributes to the average number of exchanges on a chromosome. A quadruple crossover would have the same effect. These and other multiple exchanges are responsible for the discrepancy between recombination frequency and genetic map distance. In practice, this discrepancy is small for distances less than 20 cM. Over such distances, interference is strong enough to suppress almost

all multiple exchanges, and the recombination frequency is a good estimator of the true genetic distance. For values greater than 20 cM, these two quantities diverge, principally because multiple exchanges become much more likely. **Figure 7.17** shows the mathematical relationship between recombination frequency and genetic map distance.



■ FIGURE 7.17 Relationship between frequency of recombination and genetic map distance. For values less than 20 cM, there is approximately a linear relationship between percent recombination and map distance; for values greater than 20 cM, the percent recombination underestimates the map distance.

KEY POINTS

- The genetic maps of chromosomes are based on the average number of crossovers that occur during meiosis.
- Genetic map distances are estimated by calculating the frequency of recombination between genes in experimental crosses.
- Recombination frequencies less than 20 percent estimate map distance directly; however, recombination frequencies greater than 20 percent underestimate map distance because multiple crossover events do not always produce recombinant chromosomes.

Cytogenetic Mapping

Geneticists have developed techniques to localize genes on the cytological maps of chromosomes.

Recombination mapping allows us to determine the relative positions of genes by using the frequency of crossing over as a measure of distance. However, it does not allow us to localize genes with respect to

cytological landmarks, such as bands, on chromosomes. This kind of localization requires a different procedure that involves studying the phenotypic effects of chromosome rearrangements, such as deletions and duplications. Because these types of rearrangements can be recognized cytologically, their phenotypic effects can be correlated with particular regions along the length of a chromosome. If these phenotypic effects can be associated with genes that have already been positioned on a recombination map, then the map positions of those genes can be tied to locations on the cytological map of a chromosome. This process, called cytogenetic mapping, has been most thoroughly developed in *Drosophila* genetics, where the large, banded polytene chromosomes provide researchers with extraordinarily detailed cytological maps.

LOCALIZING GENES USING DELETIONS AND DUPLICATIONS

As an example of cytogenetic mapping, let's consider ways of localizing the X-linked *white* gene of *Drosophila*, a wild-type copy of which is required for pigmentation in the eyes. This gene is situated at map position 1.5 near one end of the X chromosome. But which of the two ends is it near, and how far is it, in cytological terms, from that end? To answer these questions, we need to find the position of the *white* gene on the cytological map of the polytene X chromosome.

One procedure is to produce flies that are heterozygous for a recessive null mutation of the *white* gene (*w*) and a cytologically defined deletion (or deficiency, usually symbolized Df) for part of the X chromosome (**Tigure 7.18**). These *w*/*Df* heterozygotes provide a functional test for the location of *white* relative to the deficiency. If the *white*

gene has been deleted from the Df chromosome, then the w/Df heterozygotes will not be able to make eye pigment because they will not have a functional copy of the *white* gene on either of their X chromosomes. The eyes of the w/Df heterozygotes will therefore be white (the mutant phenotype). If, however, the *white* gene has not been deleted from the Dfchromosome, then the w/Df heterozygotes will have a functional *white* gene somewhere on that chromosome, and their eyes will be red (the wild phenotype). By looking at the eyes of the w/Df heterozygotes, we can therefore determine whether or not a specific deficiency has deleted the *white* gene. If it has, *white* must be located within the boundaries of that deficiency.

Different X chromosome deficiencies have allowed researchers to locate the *white* gene to a position near the left end of the X chromosome (**Figure 7.19**). Each deficiency was combined with a recessive *white* mutation, but only one of the deficiencies, $Df(1)w^{rfl}$, produced white eyes. Because this deficiency "uncovers" the *white* mutation, we know that the *white* gene must be located within the segment of the

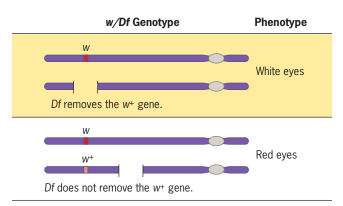
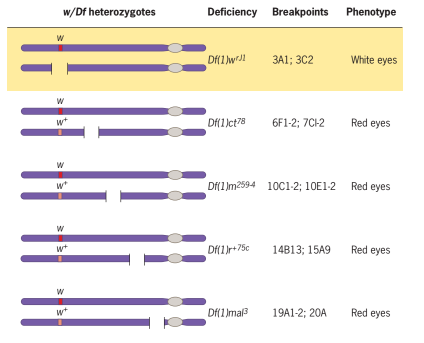


FIGURE 7.18 Principles of deletion mapping to localize a gene within a *Drosophila* chromosome. The *white* gene on the X chromosome, defined by the recessive mutation *w* which causes white eyes, is used as an example.

■ FIGURE 7.19 Localization of the *white* gene in the *Drosophila* X chromosome by deletion mapping. The deficiency breakpoints are presented using the coordinates of Bridges' cytological map of the polytene X chromosome.



The mutant eye color observed with $Df(1)w^{rJ1}$ indicates that the *white* gene is between the deficiency breakpoints in bands 3A1 and 3C2 on the X chromosome.

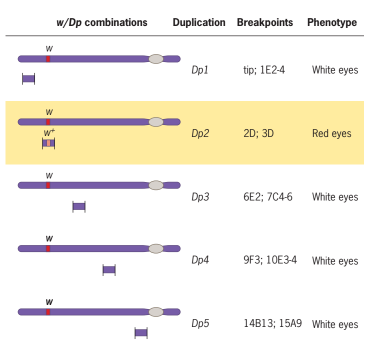
chromosome that it deletes, that is, somewhere between polytene chromosome bands 3A1 and 3C2. With smaller deficiencies, the *white* gene has been localized to polytene chromosome band 3C2, near the right boundary of $Df(1)w^{rfl}$.

We can also use duplications to determine the cytological locations of genes. The procedure is similar to the one using deletions, except that we look for a duplication that masks the phenotype of a recessive mutation. **Figure 7.20** shows an example utilizing duplications for small segments of the X chromosome that have been translocated to another chromosome. Only one of these duplications, Dp2, masks—or, as geneticists like to say, "covers"—the *white* mutation; thus, a wild-type copy of *white* must be present within it. This localizes the *white* gene somewhere between sections 2D and 3D on the polytene X chromosome, which is consistent with the results of the deletion tests already discussed.

Deletions and duplications have been extraordinarily useful in locating genes on the cytological maps of *Drosophila* chromosomes. The basic principle in *deletion mapping* is that a deletion that *uncovers* a recessive mutation must lack a wildtype copy of the mutant gene. This fact localizes that gene within the boundaries of the deletion. The basic principle in *duplication mapping* is that a duplication that *covers* a recessive mutation must contain a wild-type copy of the mutant gene. This fact localizes that gene within the boundaries of the duplication. To test your ability to localize genes using deficiencies and duplications, work through the exercise in Solve It: Cytological Mapping of a *Drosophila* Gene.

GENETIC DISTANCE AND PHYSICAL DISTANCE

The procedures for measuring genetic distance and for constructing recombination maps are based on the incidence of crossing over between paired chromosomes. Intuitively, we



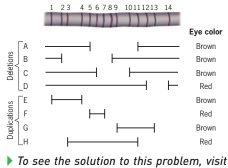
The wild-type eye color observed with *Dp2* indicates that the *white* gene is between the duplication breakpoints in regions 2D and 3D on the X chromosome.

■ FIGURE 7.20 Localization of the *white* gene in the *Drosophila* X chromosome by duplication mapping. Each duplication is a segment of the X chromosome that has been translocated to another chromosome. For simplicity, however, the other chromosome is not shown. The duplication breakpoints are presented using the coordinates of Bridges' cytological map of the polytene X chromosome.

Solve It!

Cytological Mapping of a *Drosophila* Gene

A recessive, X-linked mutation causes the eyes of Drosophila that are hemizygous or homozygous for it to be brown; the eyes of wild-type flies are red. A geneticist produced females that carried this recessive mutation on one of their X chromosomes; their other X chromosome had a cytologically defined deletion. The geneticist also produced males that carried the browneye mutation on their X chromosome; the Y chromosome in these males carried a cytologically defined duplication of a small segment of the X chromosome. The extent of each deletion and duplication is shown below in reference to a map of 14 bands within the polytene X chromosome. Each of the mutation/deletion females and mutation/duplication males was scored for eye color. From the results, locate the eye color gene in the smallest possible interval on the cytological map.



the Student Companion site.

KEY POINTS

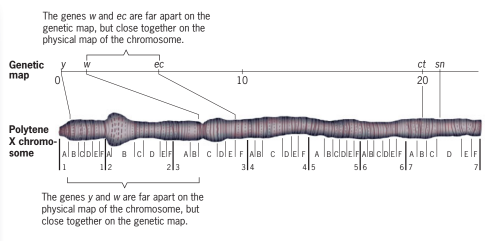


FIGURE 7.21 Left end of the polytene X chromosome of *Drosophila* and the corresponding portion of the genetic map showing the genes for *yellow* body (*y*), *white* eyes (*w*), *echinus* eyes (*ec*), *cut* wings (*ct*), and *singed* bristles (*sn*).

expect that long chromosomes should have more crossovers than short ones and that this relationship will be reflected in the lengths of their genetic maps. For the most part, our assumption is true; however, within a chromosome some regions are more prone to crossing over than others. Thus, distances on the genetic map do not correspond exactly to physical distances along the chromosome's cytological map (**■** Figure 7.21). Crossing over is less likely to occur near the ends of a chromosome and also around the centromere; consequently, these regions are condensed on the genetic map. Other regions, in which crossovers occur more frequently, are expanded.

Even though there is not a uniform relationship between genetic and physical distance, the genetic and cytological maps of a chromosome are colinear; that is, particular sites have the same order. Recombination mapping therefore reveals the true order of the genes along a chromosome. However, it does not tell us the actual physical distances between them.

- In Drosophila, genes can be localized on maps of the polytene chromosomes by combining recessive mutations with cytologically defined deletions and duplications.
- A deletion will reveal the phenotype of a recessive mutation located between its endpoints, whereas a duplication will conceal the mutant phenotype.
- Genetic and cytological maps are colinear; however, genetic distances are not proportional to cytological distances.

Linkage Analysis in Humans

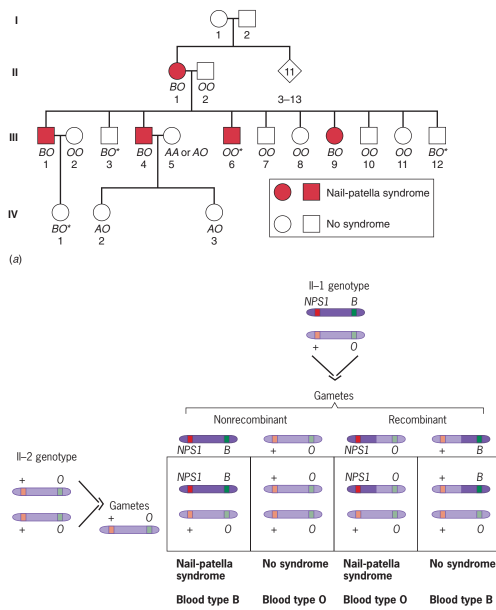
Pedigree analysis provides ways of localizing genes on human chromosomes.

To detect and analyze linkage in humans, geneticists must collect data from pedigrees. Often these data are limited or incomplete, or the information they provide is ambiguous. The task of constructing human linkage maps therefore confronts researchers with many

challenging problems. Classical studies of linkage in humans focused on pedigrees in which it was possible to follow the inheritance of two or more genes simultaneously. Today, modern molecular methods permit researchers to analyze the inheritance of

dozens of different markers in the same set of pedigrees. This multi-locus analysis has greatly increased the ability to detect linkage and to construct detailed chromosome maps. The linkage relationships that are easiest to study in humans are those between genes on the X chromosome. Such genes follow a pattern of inheritance that is readily identified. If two genes show this pattern, they must be linked. Determining linkage between autosomal genes is much more difficult. The human genome has 22 different autosomes, and a gene that does not show X-linkage could be on any one of them. Which autosome is the gene on, what other genes are linked to it, and what are the map positions of these genes? These are challenging questions for the human geneticist.

To see how linkage is detected in human pedigrees, let's examine some of the work of J. H. Renwick and S. D. Lawler. In 1955 these researchers reported evidence for linkage between the gene controlling the ABO blood groups (see Chapter 4) and a dominant mutation responsible for a rare, autosomal disorder called the nail-patella syndrome. People with this syndrome have abnormal nails and kneecaps. A portion of one of the pedigrees that Renwick and Lawler studied is shown in **Figure 7.22a**. Each individual in this pedigree was characterized for the presence or absence of the mutation for the nail-patella syndrome, denoted *NPS1*; in addition, most of the individuals were typed for the ABO blood groups.



■ FIGURE 7.22 Linkage analysis in a human pedigree. (a) A portion of a pedigree showing linkage between the ABO and nail-patella loci. Individuals affected with the nail-patella syndrome are denoted by red symbols. Where known, the genotype of the ABO locus is given underneath each symbol. Asterisks denote recombinants. (b) A Punnett square showing the genotypes produced by the couple in generation II.

The woman in generation II must represent a new occurrence of the *NPS1* mutation. Neither of her parents nor any of her 11 siblings showed the nail-patella phenotype. Among the five individuals who showed the nail-patella syndrome in this pedigree, all but one (III-6) of them had blood type B. This observation suggests that the *NPS1* mutation is genetically linked to the *B* allele of the *ABO* blood group locus. If we assume this inference to be correct, then the woman in generation II must have the genotype *NPS1* B/+O; that is, she is a repulsion heterozygote. Her husband's genotype is clearly +O/+O.

Figure 7.22 illustrates the genetic phenomena underlying this pedigree and suggests a strategy to estimate, albeit crudely, the distance between the NPS1 and ABO loci. The mating indicated in Figure 7.22*b* is essentially a testcross. The woman II-1 can produce four different kinds of gametes, two carrying recombinant chromosomes and two carrying nonrecombinant chromosomes. When these gametes are combined with the single type of gamete (+O) produced by the man II-2, four different genotypes can result. As the pedigree in Figure 7.22*a* shows, II-1 and II-2 produced all four types of children. However, only 3 (III-3, III-6, III-12, indicated by asterisks in Figure 7.22a) of their 10 children were recombinants; the other 7 were nonrecombinants. Thus, we can estimate the frequency of recombination between the NPS1 and ABO loci as 3/10 =30 percent. However, this estimate does not use all the information in the pedigree. To refine it, we can incorporate the information from the couples' three grandchildren, only one (IV-1) of whom was a recombinant. Altogether, then, 3 + 1 = 4 of the 10 + 3 = 313 offspring in the pedigree were recombinants. Thus, we conclude that the frequency of recombination between the NPS1 and ABO loci is 4/13 = 31 percent. In terms of a linkage map, we estimate that the distance between these genes is about 31 cM. Renwick and Lawler analyzed other pedigrees for linkage between the NPS1 and ABO genes. By combining all the data, they estimated the frequency of recombination to be about 10 percent. Thus, the distance between the NPS1 and ABO genes is about 10 cM.

Renwick and Lawler's study of the NPS1 and ABO loci established that these two genes are linked, but it could not identify the specific autosome that carried them. The first localization of a gene to a specific human autosome came in 1968, when R. P. Donahue and coworkers demonstrated that the Duffy blood group locus, denoted FY, is on chromosome 1. This demonstration hinged on the discovery of a variant of chromosome 1 that was longer than normal. Pedigree analysis showed that in a particular family, this long chromosome segregated with specific FY alleles. Thus, the FY locus was assigned to chromosome 1. Subsequent research has placed this locus at region 1p31 on that chromosome. Using different techniques, the NPS1 and ABO loci have been situated near the tip of the long arm of chromosome 9.

Until the early 1980s, progress in human gene mapping was extremely slow because it was difficult to find pedigrees that were segregating linked markers—say, for example, two different genetic diseases. In the 1980s, however, it became possible to identify genetic variants in the DNA itself. These variants result from differences in the DNA sequence in parts of chromosomes. For example, in one individual a particular sequence might be GAATTC on one of the DNA strands, and in another individual the corresponding DNA sequence might be GATTTC—a difference of just one nucleotide. Although we must defer to later chapters a discussion of the techniques that are used to reveal such molecular differences, here we can explore how they have helped to map human genes, including many that are involved in serious inherited diseases. If, in addition to the usual phenotypic analysis, the members of a pedigree are analyzed for the presence or absence of molecular markers in the DNA, a researcher can look for linkage between each marker and the gene under study. Then, with appropriate statistical techniques, he or she can estimate the distances between the gene and the markers that are linked to it.

This approach has allowed geneticists to map a large number of genes involved in human diseases. One of the most dramatic examples is the research that located the gene for Huntington's disease (*HD*), a debilitating and ultimately fatal neurological disorder, on chromosome 4. This effort, discussed in A Milestone in Genetics: Mapping the Gene for Huntington's Disease on the Student Companion site, analyzed large pedigrees for linkage between the *HD* gene and an array of molecular markers. Through painstaking work, the *HD* gene was mapped to within 4 cM of one of these markers. This precise localization laid the foundation for the isolation and molecular characterization of the *HD* gene itself.

Molecular markers have also made it possible to build up maps of human chromosomes from completely independent analyses. If gene A has been shown to be linked to marker x in one set of pedigrees, and gene B has been shown to be linked to marker x in another set of pedigrees, then gene A and gene B are obviously linked to each other. Thus, the analysis of these markers allows human geneticists to determine linkage relationships between genes that are not segregating in the same pedigrees.

The analysis of recombination data from pedigrees allows geneticists to construct linkage maps of chromosomes. However, except in the case of X linkage, this analysis does not tell us which chromosome is being mapped, or where a particular gene resides on the physical image of that chromosome. These challenges have been addressed by developing cytological techniques such as chromosome banding and chromosome painting (Chapter 6).

- Linkage between human genes can be detected by analyzing pedigrees.
- Pedigree analysis also provides estimates of recombination frequencies to map genes on human chromosomes.

Recombination and Evolution

Recombination is an essential feature of sexual reproduction. During meiosis, when chromosomes come together and cross over, there is an opportunity

to create new combinations of alleles. Some of these may benefit the organism by enhancing survival or reproductive ability. Over time, such beneficial combinations would be expected to spread through a population and become standard features of the genetic makeup of the species. Meiotic recombination is therefore a way of shuffling genetic variation to potentiate evolutionary change.

in evolution.

EVOLUTIONARY SIGNIFICANCE OF RECOMBINATION

We can appreciate the evolutionary advantage of recombination by comparing two species, one capable of reproducing sexually and the other not. Let's suppose that a beneficial mutation has arisen in each species. Over time, we would expect these mutations to spread. Let's also suppose that while they are spreading, another beneficial mutation occurs in a nonmutant individual within each species. In the asexual organism, there is no possibility that this second mutation will be recombined with the first, but in the sexual organism, the two mutations can be recombined to produce a strain that is better than either of the single mutants by itself. This recombinant strain will be able to spread through the whole species population. In evolutionary terms, recombination can allow favorable alleles of different genes to come together in the same organism.

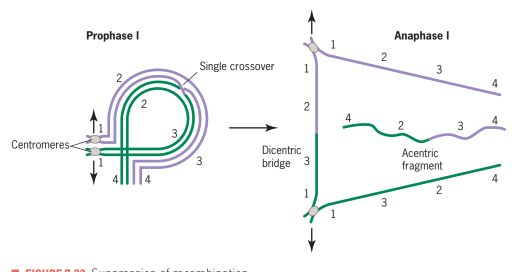
SUPPRESSION OF RECOMBINATION BY INVERSIONS

The gene-shuffling effect of recombination can be thwarted by chromosome rearrangements. Crossing over is usually inhibited near the breakpoints of a rearrangement in heterozygous condition, probably because the rearrangement disrupts chromosome pairing. Many rearrangements are therefore associated with a reduction in the frequency of recombination. This effect is most pronounced in inversion heterozygotes because the inhibition of crossing over that occurs near the breakpoints of the inversion is compounded by the selective loss of chromosomes that have undergone crossing over within the inverted region.

To see this recombination-suppressing effect, we consider an inversion in the long arm of a chromosome (**Figure 7.23**). If a crossover occurs between inverted and

KEY POINTS

Recombination—or the lack of it—plays a key role



■ FIGURE 7.23 Suppression of recombination in an inversion heterozygote. The dicentric (1 2 3 1) and acentric (4 3 2 4) chromosomes formed from the crossover chromatids are aneuploid and will cause inviability in the next generation. Consequently, the products of crossing over between the inverted and noninverted chromosomes are not recovered.

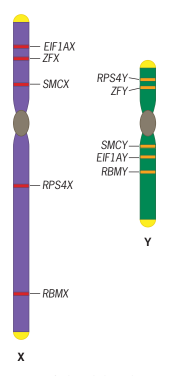


FIGURE 7.24 Order of shared genes outside the pseudoautosomal regions on the human X and Y chromosomes.

noninverted chromatids within the tetrad, it will produce two recombinant chromatids; however, both of these chromatids are likely to be lost during or after meiosis. One of the chromatids lacks a centromere-it is an acentric fragmentand will therefore be unable to move to its proper place during anaphase of the first meiotic division. The other chromatid has two centromeres and will therefore be pulled in opposite directions, forming a dicentric chromatid bridge. Eventually, this bridge will break and split the chromatid into pieces. Even if the acentric and dicentric chromatids produced by crossing over within the inversion survive meiosis, they are not likely to form viable zygotes. Both

of these chromatids are aneuploid—duplicate for some genes and deficient for others and such aneuploidy is usually lethal. These chromatids will therefore be eliminated by natural selection in the next generation. The net effect of this chromatid loss is to suppress recombination between inverted and noninverted chromosomes in heterozygotes.

Geneticists have exploited the recombination-suppressing properties of inversions to keep alleles of different genes together on the same chromosome. Let's assume, for example, that a chromosome that is structurally normal carries the recessive alleles a, b, c, d, and e. If this chromosome is paired with another structurally normal chromosome that carries the corresponding wild-type alleles a^+ , b^+ , c^+ , d^+ , and e^+ , the recessive and wild-type alleles will be scrambled by recombination. To prevent this scrambling, the chromosome with the recessive alleles can be paired with a wild-type chromosome that has an inversion. Unless double crossovers occur within the inverted region, this structural heterozygosity will suppress recombination. The multiply mutant chromosome can then be transmitted to the progeny as an intact genetic unit.

This recombination-suppressing technique has often been used in experiments with *Drosophila*, where the inverted chromosome usually carries a dominant mutation that permits it to be tracked through a whole series of crosses without cytological examination. Such a marked inversion chromosome is called a **balancer** because it allows a chromosome of interest to be kept in heterozygous condition without recombinational breakup.

Suppression of recombination by inversions seems to have played an important role in the evolution of the sex chromosomes in mammals. The evidence comes from analyses by Bruce Lahn and David Page, who studied 19 genes that are present on both the human X and Y chromosomes. These shared genes occupy different postions on the X and Y chromosomes—a finding indicating that inversions have rearranged them relative to one another during the course of evolution. In addition, the DNA sequences of the X- and Y-linked copies of these shared genes have diverged from one another to different extents. By analyzing variation in the extent of divergence, Lahn and Page have discerned four "evolutionary strata" in the human sex chromosomes-regions in which recombination has been suppressed for different lengths of evolutionary time. Lahn and Page conjecture that the X and Y chromosomes originated from a pair of autosomes sometime after the mammalian evolutionary line diverged from the line of ancient reptiles that led to dinosaurs, crocodiles, and birds. Between 240 and 320 million years ago, an inversion in what was to become the Y chromosome led to regional suppression of recombination between the X and the Y. In the lineage that ultimately led to humans, at least three additional inversions occurred, two of them sometime between 80 and 130 million years ago, and one of them between 30 and 50 million years ago. The net effect of these inversions has been to suppress recombination between most of the regions on the X and Y chromosomes. Through natural selection, functional genes have been retained on the X chromosome, but on the Y chromosome most of the genes have degenerated through the accumulation of random mutations. Thus, today the Y chromosome has many fewer functional genes than the X chromosome, and the ones that remain are arranged in a different order (**Figure 7.24**).

GENETIC CONTROL OF RECOMBINATION

It is not surprising that a process as important as recombination should be under genetic control. Studies with several organisms, including yeast and *Drosophila*, have demonstrated that recombination involves the products of many genes. Some of these gene products play a role in chromosome pairing, others catalyze the process of exchange, and still others help to rejoin broken chromatid segments. We will consider some of these activities in greater detail in Chapter 13.

One curious phenomenon, which no one has yet explained, is that there is no crossing over in *Drosophila* males. In this regard, *Drosophila* is different from most species, including our own, where crossing over occurs in both sexes. In addition, we know that the amount of recombination varies among species. Perhaps the events that lead to recombination are themselves subject to evolutionary change.

- Recombination can bring favorable mutations together.
- Chromosome rearrangements, especially inversions, can suppress recombination.
- Recombination is under genetic control.

Basic Exercises

Illustrate Basic Genetic Analysis

- An inbred strain of snapdragons with violet flowers and dull leaves was crossed to another inbred strain with white flowers and shiny leaves. The F₁ plants, which all had violet flowers and dull leaves, were backcrossed to the strain with white flowers and shiny leaves, and the following F₂ plants were obtained: 50 violet, dull; 46 white, shiny; 12 violet, shiny; and 10 white, dull. (a) Which of the four classes in the F₂ are recombinants? (b) What is the evidence that the genes for flower color and leaf texture are linked? (c) Diagram the crosses of this experiment. (d) What is the frequency of recombination between the flower color and leaf texture genes? (e) What is the genetic map distance between these genes?
- Answer: (a) The last two classes—violet, shiny, and white, dull—in the F₂ are recombinants. Neither of these combinations of phenotypes was present in the strains used in the initial cross. (b) The recombinants are 18.6 percent of the F_2 plants—much less than the 50 percent that would be expected if the flower color and leaf texture genes were unlinked. Therefore, these genes must be linked on the same chromosome in the snapdragon genome. (c) To diagram the crosses, we must first assign symbols to the alleles of the flower color and leaf texture genes: W = violet, w = white; S = dull, s = shiny; a capital letter indicates that the allele is dominant. The first cross is $WS/WS \times w s/w s$, yielding F_1 plants with the genotype W S/w s. The backcross is $WS/w s \times w s/w s$, yielding four classes of progeny: (1) W S/w s, (2) w s/w s, (3) W s/w s, and (4) w S/w s. Classes 1 and 2 are parental types, and classes 3 and 4 are recombinants. (d) The frequency of recombination is 18.6 percent. (e) The genetic map distance is estimated by the frequency of recombination as 18.6 centiMorgans.

2. What is the cytological evidence that crossing over has occurred? When and where would you look for it?

KEY POINTS

- **Answer:** Crossing over probably occurs during early to midprophase of meiosis I. However, the chromosomes are not easily analyzed in these stages, and exchanges are difficult, if not impossible, to identify by cytological methods. The best cytological evidence that crossing over has occurred is obtained from cells near the end of the prophase of meiosis I. In this stage, paired homologues repel each other slightly, and the exchanges between them are seen as chiasmata.
- **3.** A geneticist has estimated the number of exchanges that occurred during meiosis on each of 100 chromatids that were recovered in gametes. The data are as follows:

Number of Exchanges	Frequency
0	18
1	20
2	40
3	16
4	6

What is the genetic length in centiMorgans of the chromosome analyzed in this study?

Answer: The genetic length of a chromosome is the average number of exchanges on a chromatid at the end of meiosis. For the data at hand, the average is $0 \times (18/100) + 1 \times (20/100) + 2 \times (40/100) + 3 \times (16/100) + 4 \times (6/100) = 1.72$ Morgans or 172 centi-Morgans. 4. Drosophila females heterozygous for three recessive X-linked markers, y (yellow body), ct (cut wings), and m (miniature wings), and their wild-type alleles were crossed to y ct m males. The following progeny were obtained:

Phenotypic Class	Number
1. yellow, cut, miniature	30
2. wild-type	33
3. yellow	10
4. cut, miniature	12
5. miniature	8
6. yellow, cut	5
7. yellow, miniature	1
8. cut	1
	Total: $\overline{100}$

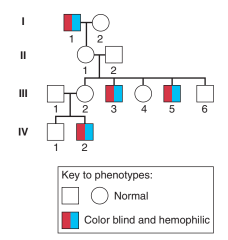
(a) Which classes are parental types? (b) Which classes represent double crossovers? (c) Which gene is in the middle of the other two? (d) What was the genotype of the heterozygous females used in the cross? (Show the correct linkage phase as well as the correct order of the markers along the chromosome.)

- **Answer:** (a) The parental classes are the most numerous; therefore, in these data, classes 1 and 2 are parental types. (b) The double crossover classes are the least numerous; therefore, in these data, classes 7 and 8 are the double crossover classes. (c) The parental classes tell us that all three mutant alleles entered the heterozygous females on the same X chromosome; the other X chromosome in these females must have carried all three wild-type alleles. The double crossover classes tell us which of the three genes is in the middle because the middle marker will be separated from each of the flanking markers by the double exchange process. In these data, the *ct* allele is separated from *y* and *m* in the double crossover classes; therefore, the *ct* gene must lie between the y and m genes. (d) The genotype of the heterozygous females used in the cross must have been $y \ ct \ m/+ + +$.
- 5. A Drosophila geneticist has conducted experiments to localize the singed (sn) bristle gene on the cytological map of the X chromosome. Males hemizygous for a recessive sn mutation were mated to females that carried various deficiencies (symbolized Df) in the X chromosome balanced over a multiply inverted X chromosome marked with the semidominant mutation for Bar (B) eyes. Thus, the crossing scheme was sn/Y males $\times Df/B$ females. The results of crosses with four different deficiencies are as follows:

Deficiency	Breakpoints	Phenotype of Non-Bar Daughters
1	2F; 3C	wild-type
2	4D; 5C	wild-type
3	6F; 7E	singed
4	7C; 8C	singed

The cytological map of the X chromosome is divided into 20 numbered sections, each subdivided into subsections A–F. Where is the *singed* gene on this cytological map?

- **Answer:** The non-Bar daughters that were examined for the singed phenotype were genotypically *Df/sn*. The *singed* mutation was "uncovered" by two of the deficiencies, 3 and 4; thus, it must lie in the deleted region on the X chromosome that is common to both—that is, in region 7C–7E.
- 6. The following pedigree shows four generations of a family described in 1928 by M. Madlener. The great-grandfather, I-1, has both color blindness and hemophilia. Letting *c* represent the allele for color blindness and *b* represent the allele for hemophilia, what are the genotypes of the man's five grandchildren? Do any of the individuals in the pedigree provide evidence of recombination between the genes for color blindness and hemophilia?



Answer: The genes for color blindness and hemophilia are X-linked. Because I-1 has both color blindness and hemophilia, his genotype must be c b. His daughter, II-1, is phenotypically normal and must therefore carry the nonmutant alleles, C and H, of these two X-linked genes. Moreover, because II-1 inherited both c and b from her father, the two nonmutant alleles that she carries must be present on the X chromosome she inherited from her mother. II-1's genotype is therefore C H/c b—that is, she is a coupling heterozygote for the two loci. III-2, the first granddaughter of I-1, is also a coupling heterozygote. We infer that she has this genotype because her son has both color blindness and hemophilia (c b), and her father is phenotypically normal (C H). Evidently, III-2 inherited the c b chromosome from her mother. Among the grandsons of

I-1, two (III-3 and III-5) of them have both hemophilia and color blindness; thus, these grandsons are genotypically c b. The other grandson (III-6) is neither color blind nor hemophilic; his genotype is therefore C H. The genotype of the remaining granddaughter (III-4) is uncertain. This woman inherited a C H chromosome from her father. However, the chromosome she inherited from her mother could be C H, c b, C b, or c H. The pedigree does not allow us to determine which of these chromosomes she received.

Testing Your Knowledge

Integrate Different Concepts and Techniques

1. R. K. Sakai, K. Akhtar, and C. J. Dubash (1985, *J. Hered.* 76:140–141) reported data from a set of testcrosses with the mosquito *Anopheles culicifacies*, a vector for malaria in southern Asia. The data involved three mutations: *bw* (*brown* eyes), *c* (*colorless* eyes), and *Blk* (*black* body). In each cross, repulsion heterozygotes were mated to mosquitoes homozygous for the recessive alleles of the genes, and the progeny were scored as having either a parental or a recombinant genotype. Are any of the three genes studied in these crosses linked? If so, construct a map of the linkage relationships.

	Repulsion	Progeny		Percent
Cross	Heterozygote	Parental	Recombinant	Recombination
1	<i>bw</i> +/+ <i>c</i>	850	503	37.2
2	bw+/+Blk	750	237	24.0
3	c+/+ Blk	629	183	22.5

Answer: In each cross, the frequency of recombination is less than 50 percent, so all three loci are linked. To place them on a linkage map, we estimate the distances between each pair of genes from the observed recombination frequencies:

Notice that the recombination frequency between bw and c (37.2 percent, from Cross 1) is substantially less than the actual distance between these genes (46.5). This shows that for widely separated genes, the recombination frequency underestimates the true map distance.

2. Singed bristles (*sn*), crossveinless wings (*cv*), and vermilion eye color (*v*) are due to recessive mutant alleles of three X-linked genes in *Drosophila melanogaster*. When a female heterozygous for each of the three genes was testcrossed with a singed, crossveinless, vermilion male, the following progeny were obtained:

The most we can say about III-4's genotype is that she carries a chromosome with the C and H alleles.

None of the four grandchildren to whom we can assign genotypes provides evidence of recombination between the genes for color blindness and hemophilia. Neither do the two great-grandchildren shown in generation IV. One of these great-grandchildren is genotypically C H; the other is genotypically c h. Thus, in the pedigree as a whole there is no evidence for recombination between the C and H genes.

Class	Phenotype	Number
1	singed, crossveinless, vermilion	3
2	crossveinless, vermilion	392
3	vermilion	34
4	crossveinless	61
5	singed, crossveinless	32
6	singed, vermilion	65
7	singed	410
8	wild-type	3
		Total: 1000

What is the correct order of these three genes on the X chromosome? What are the genetic map distances between *sn* and *cv*, *sn* and *v*, and *cv* and *v*? What is the coefficient of coincidence?

Answer: Before attempting to analyze these data, we must establish the genotype of the heterozygous female that produced the eight classes of offspring. We do this by identifying the two parental classes (2 and 7), which are the most numerous in the data. These classes tell us that the heterozygous female had the cv and v mutations on one of her X chromosomes and the sn mutation on the other. Her genotype was therefore (cv + v)/(+ sn +), with the parentheses indicating uncertainty about the gene order.

To determine the gene order, we must identify the double crossover classes among the six types of recombinant progeny. These are classes 1 and 8—the least numerous. They tell us that the *singed* gene is between *crossveinless* and *vermilion*. We can verify this by investigating the effect of a double crossover in a female with the genotype.

Two exchanges in this genotype will produce gametes that are either cv sn v or + + +, which correspond to classes 1 and 8, the observed double crossovers. Thus, the proposed gene order—cv sn v—is correct.

Having established the gene order, we can now determine which recombinant classes represent crossovers between cvand sn, and which represent crossovers between sn and v.

Crossovers between *cv* and *sn*:

Class:	3		5		1		8		
Number:	34	+	32	+	3	+	3	=	72
-									

Crossovers between *sn* and *v*:

Class: 4 6 1 8Number: 61 + 65 + 3 + 3 = 132

We determine the distances between these pairs of genes by calculating the average number of crossovers. Between cv and sn, the distance is 72/1000 = 7.2 cM, and between sn and v it is 132/1000 = 13.2 cM. We can estimate the distance between cv and v as the sum of these values: 7.2 + 13.2 = 20.4 cM. The linkage map of these three genes is therefore:

cv—7.2—*sn*—13.2—*v*

To calculate the coefficient of coincidence, we use the observed and expected frequencies of double crossovers:

 $c = \frac{\text{observed frequency}}{\text{of double crossovers}} = \frac{0.006}{0.072 \times 0.132} = 0.63$ of double crossovers

which indicates only moderate interference.

A Drosophila geneticist is studying a recessive lethal 3. mutation, l(1)r13, located on the X chromosome. This mutation is maintained in a stock with a balancer X chromosome marked with a semidominant mutation for Bar eyes (B). In homozygous and hemizygous condition, the *B* mutation reduces the eyes to narrow bars. In heterozygous condition, it causes the eyes to be kidneyshaped. Flies that are homozygous or hemizygous for the wild-type allele of B have large, spherical eyes. To maintain the l(1)r13 mutation in stock, for each generation the geneticist crosses B males to l(1)r13/B females and selects daughters with kidney-shaped eyes for crosses with their Bar-eyed brothers. The geneticist wishes to determine the cytological location of l(1)r13. To accomplish this goal, she crosses l(1)r13/B females to various males that carry duplications for short segments of the X chromosome in their genomes. Each duplication is attached to the Y chromosome. Thus, the genotype of the males used in these crosses can be represented as X/Y-Dp. The geneticist screens the progeny of each cross for the presence of non-Bar sons. From

the results shown in the following table, determine the cytological location of l(1)r13.

Dp Name	Dp Segment*	Non-Bar Sons Present
1	2D-3D	Yes
2	3A-3E	Yes
3	3D-4A	No
4	4A–4D	No
5	4B-4E	No

*The long arm of the X chromosome is divided into 20 numbered sections, starting with section 1 at the tip and ending with section 20 near the centromere. Each section is divided into six subsections, ordered alphabetically A through F. Subsection A is on the tip-side of a numbered section.

- **Answer:** The cross to maintain the lethal mutation in stock is B/Y males $\times l(1)r13/B$ females $\rightarrow B/Y$ males (Bar eyes), *l(1)r13/Y* males (die), *l(1)r13/B* females (kidney-shaped eyes), and *B*/*B* females (Bar eyes). Each generation, the *B*/*Y* males and the l(1)r13/B females are selected for crosses to perpetuate the lethal mutation. A cross to determine the cytological location of the lethal mutation can be represented as l(1)r13/B females $\times X/Y$ -Dp males $\rightarrow l(1)r13/Y$ -Dp males (if viable, non-Bar eyes), B/Y-Dp males (Bar eyes), *l(1)r13/X* females (non-Bar eyes), and *B/X* females (kidneyshaped eyes). The first class of flies-males with non-Bar eyes-provides the data on whether or not a specific duplication "covers" the lethal mutation. If it does, these males will appear among the progeny in the culture. If it does not, they will not appear. From the data, we see that two duplications, Dp 1 and Dp 2, cover the lethal mutation. Thus, the mutation must lie within the boundaries of these duplications-that is, somewhere between 2D and 3E. We can refine the lethal mutation's location by noting that the two duplications overlap from subsection 3A to subsection 3D. The mutation must therefore lie within the 3A-3D region of the X chromosome.
- 4. A woman has two dominant traits, each caused by a mutation in a different gene: cataract (an eye abnormality), which she inherited from her father, and polydactyly (an extra finger), which she inherited from her mother. Her husband has neither trait. If the genes for these two traits are 15 cM apart on the same chromosome, what is the chance that the first child of this couple will have both cataract and polydactyly?
- **Answer:** To calculate the chance that the child will have both traits, we first need to determine the linkage phase of the mutant alleles in the woman's genotype. Because she inherited the cataract mutation from her father and the polydactyly mutation from her mother, the mutant alleles must be on opposite chromosomes, that is, in the repulsion linkage phase:

For a child to inherit both mutant alleles, the woman would have to produce an egg that carried a recombinant chromosome, CP. We can estimate the probability of this event from the distance between the two genes, 15 cM,

which, because of interference, should be equivalent to 15 percent recombination. However, only half the recombinants will be *C P*. Thus, the chance that the child will inherit both mutant alleles is (15/2) percent = 7.5 percent.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- 7.1 Mendel did not know of the existence of chromosomes. Had he known, what change might he have made in his Principle of Independent Assortment?
- 7.2 From a cross between individuals with the genotypes $Cc \ Dd \ Ee \times cc \ dd \ ee$, 1000 offspring were produced. The class that was C-D-ee included 351 individuals. Are the genes c, d, and e on the same or different chromosomes? Explain.
- 7.3 If *a* is linked to *b*, and *b* to *c*, and *c* to *d*, does it follow that a recombination experiment would detect linkage between *a* and *d*? Explain.
- 7.4 Mice have 19 autosomes in their genome, each about the same size. If two autosomal genes are chosen randomly, what is the chance that they will be on the same chromosome?
- **7.5** Genes on different chromosomes recombine with a frequency of 50 percent. Is it possible for two genes on the same chromosome to recombine with this frequency?
- 7.6 If two loci are 10 cM apart, what proportion of the cells in prophase of the first meiotic division will contain a single crossover in the region between them?
- 7.7 Genes *a* and *b* are 20 cM apart. An $a^+ b^+/a^+ b^+$ individual was mated with an *a b/a b* individual.
 - (a) Diagram the cross and show the gametes produced by each parent and the genotype of the F_1 .
 - (b) What gametes can the F_1 produce, and in what proportions?
 - (c) If the F_1 was crossed to *a b/a b* individuals, what offspring would be expected, and in what proportions?
 - (d) Is this an example of the coupling or repulsion linkage phase?
 - (e) If the F₁ were intercrossed, what offspring would be expected and in what proportions?
- **7.8** Answer questions (a)–(e) in the preceding problem under the assumption that the original cross was $a^+ b/a^+ b \times a b^+/a b^+$.
- **7.9** If the recombination frequency in the previous two problems were 40 percent instead of 20 percent, what change would occur in the proportions of gametes and testcross progeny?
- **7.10** A homozygous variety of maize with red leaves and normal seeds was crossed with another homozygous variety with green leaves and tassel seeds. The hybrids were then backcrossed to the green, tassel-seeded variety, and the

following offspring were obtained: red, normal 124; red, tassel 126; green, normal 125; green, tassel 123. Are the genes for plant color and seed type linked? Explain.

- 7.11 A phenotypically wild-type female fruit fly that was heterozygous for genes controlling body color and wing length was crossed to a homozygous mutant male with black body (allele *b*) and vestigial wings (allele *vg*). The cross produced the following progeny: gray body, normal wings 126; gray body, vestigial wings 24; black body, normal wings 26; black body, vestigial wings 124. Do these data indicate linkage between the genes for body color and wing length? What is the frequency of recombination? Diagram the cross, showing the arrangement of the genetic markers on the chromosomes.
- **7.12** Another phenotypically wild-type female fruit fly heterozygous for the two genes mentioned in the previous problem was crossed to a homozygous black, vestigial male. The cross produced the following progeny: gray body, normal wings 23; gray body, vestigial wings 127; black body, normal wings 124; black body, vestigial wings 26. Do these data indicate linkage? What is the frequency of recombination? Diagram the cross, showing the arrangement of the genetic markers on the chromosomes.
- **7.13** In rabbits, the dominant allele *C* is required for colored fur; the recessive allele *c* makes the fur colorless (albino). In the presence of at least one *C* allele, another gene determines whether the fur is black (*B*, dominant) or brown (*b*, recessive). A homozygous strain of brown rabbits was crossed with a homozygous strain of albinos. The F_1 were then crossed to homozygous double recessive rabbits, yielding the following results: black 34; brown 66; albino 100. Are the genes *b* and *c* linked? What is the frequency of recombination? Diagram the crosses, showing the arrangement of the genetic markers on the chromosomes.
- **7.14** In tomatoes, tall vine (*D*) is dominant over dwarf (*d*), and spherical fruit shape (*P*) is dominant over pear shape (*p*). The genes for vine height and fruit shape are linked with 20 percent recombination between them. One tall plant (I) with spherical fruit was crossed with a dwarf, pear-fruited plant. The cross produced the following results: tall, spherical 81; dwarf, pear 79; tall, pear 22; dwarf, spherical 17. Another tall plant with spherical fruit (II) was crossed with the dwarf, pear-fruited plant, and the following results were obtained: tall, pear 21; dwarf, spherical 18; tall, spherical 5; dwarf, pear 4. Diagram these two crosses, showing the

genetic markers on the chromosomes. If the two tall plants with spherical fruit were crossed with each other, that is, $I \times II$, what phenotypic classes would you expect from the cross, and in what proportions?

- **7.15** In *Drosophila*, the genes *sr* (*stripe* thorax) and *e* (*ebony* body) are located at 62 and 70 cM, respectively, from the left end of chromosome 3. A striped female homozygous for e^+ was mated with an ebony male homozygous for sr^+ . All the offspring were phenotypically wild-type (gray body and unstriped).
 - (a) What kind of gametes will be produced by the F₁ females, and in what proportions?
 - (b) What kind of gametes will be produced by the F₁ males, and in what proportions?
 - (c) If the F₁ females are mated with striped, ebony males, what offspring are expected, and in what proportions?
 - (d) If the F₁ males and females are intercrossed, what offspring would you expect from this intercross, and in what proportions?
- **7.16** The *Drosophila*, genes *a* and *b* are located at positions 22.0 and 42.0 on chromosome 2, and genes *c* and *d* are located at positions 10.0 and 25.0 on chromosome 3. A fly homozygous for the wild-type alleles of these four genes was crossed with a fly homozygous for the recessive alleles, and the F_1 daughters were backcrossed to their quadruply recessive fathers. What offspring would you expect from this backcross, and in what proportions?
- **7.17** The *Drosophila* genes vg (*vestigial* wings) and *cn* (*cinnabar* eyes) are located at 67.0 and 57.0, respectively, on chromosome 2. A female from a homozygous strain of vestigial flies was crossed with a male from a homozygous strain of cinnabar flies. The F₁ hybrids were phenotypically wild-type (long wings and dark red eyes).
 - (a) How many different kinds of gametes could the F₁ females produce, and in what proportions?
 - (b) If these females are mated with cinnabar, vestigial males, what kinds of progeny would you expect, and in what proportions?
- **7.18** In *Drosophila*, the genes *st* (*scarlet* eyes), *ss* (*spineless* bristles), and *e* (*ebony* body) are located on chromosome 3, with map positions as indicated:

Each of these mutations is recessive to its wild-type allele $(st^+, dark red eyes; ss^+, smooth bristles; e^+, gray body)$. Phenotypically wild-type females with the genotype st ss e^+/st^+ ss^+ e were crossed with triply recessive males. Predict the phenotypes of the progeny and the frequencies with which they will occur assuming (a) no interference and (b) complete interference.

7.19 In maize, the genes *Pl* for purple leaves (dominant over *pl* for green leaves), *sm* for salmon silk (recessive to *Sm* for yellow silk), and *py* for pigmy plant (recessive to *Py* for normal-size plant) are on chromosome 6, with map positions as shown:

$$\frac{pl}{45} \quad \frac{sm}{55} \quad \frac{py}{65}$$

Hybrids from the cross $Pl sm py/Pl sm py \times pl Sm Py/pl Sm Py$ were testcrossed with pl sm py/pl sm py plants. Predict the phenotypes of the offspring and their frequencies assuming (a) no interference and (b) complete interference.

- 7.20 0 In maize, the genes Tu, j2, and gl3 are located on chromosome 4 at map positions 101, 106, and 112, respectively. If plants homozygous for the recessive alleles of these genes are crossed with plants homozygous for the dominant alleles, and the F_1 plants are testcrossed to triply recessive plants, what genotypes would you expect, and in what proportions? Assume that interference is complete over this map interval.
- **7.21** A *Drosophila* geneticist made a cross between females homozygous for three X-linked recessive mutations (*y*, *yellow* body; *ec*, *echinus* eye shape; *w*, *white* eye color) and wild-type males. He then mated the F_1 females to triply mutant males and obtained the following results:

Females	Males	Number
+ + +/y ec w	+ + +	475
y ec w/y ec w	y ec w	469
$y + +/y \ ec \ w$	y + +	8
+ ec w/y ec w	+ ec w	7
y + w/y ec w	y + w	18
+ ec +/y ec w	+ ec +	23
$+ + w/y \ ec \ w$	+ + w	0
y ec +/y ec w	<i>y ec</i> +	0

Determine the order of the three loci *y*, *ec*, and *w*, and estimate the distances between them on the linkage map of the X chromosome.

7.22 A *Drosophila* geneticist crossed females homozygous for three X-linked mutations (*y*, *yellow* body; *B*, *bar* eye shape; *v*, *vermilion* eye color) to wild-type males. The F_1 females, which had gray bodies and bar eyes with dark red pigment, were then crossed to *y* B^+ *v* males, yielding the following results:

Phenotype	Number
yellow, bar, vermilion wild-type	546
yellow bar, vermilion	244
yellow, vermilion] bar	160
yellow, bar } vermilion	50

Determine the order of these three loci on the X chromosome and estimate the distances between them.

7.23 Female *Drosophila* heterozygous for three recessive mutations *e* (*ebony* body), *st* (*scarlet* eyes), and *ss* (*spineless* bristles) were testcrossed, and the following progeny were obtained:

Phenotype	Number	
wild-type	67	
ebony	8	
ebony, scarlet	68	
ebony, spineless	347	
ebony, scarlet, spineless	78	
scarlet	368	
scarlet, spineless	10	
spineless	54	

(a) What indicates that the genes are linked?

- (b) What was the genotype of the original heterozygous females?
- (c) What is the order of the genes?
- (d) What is the map distance between *e* and *st*?
- (e) Between e and ss?
- (f) What is the coefficient of coincidence?
- (g) Diagram the crosses in this experiment.
- 7.24 Consider a female *Drosophila* with the following X chromosome genotype:

$$\frac{w \quad dor^{+}}{w^{+} \quad dor}$$

The recessive alleles w and *dor* cause mutant eye colors (white and deep orange, respectively). However, w is epistatic over *dor*; that is, the genotypes w *dor*/Y and w *dor*/w *dor* have white eyes. If there is 40 percent recombination between w and *dor*; what proportion of the sons from this heterozygous female will show a mutant phenotype? What proportion will have either red or deep orange eyes?

- **7.25** In *Drosophila*, the X-linked recessive mutations *prune* (*pn*) and *garnet* (*g*) recombine with a frequency of 0.4. Both of these mutations cause the eyes to be brown instead of dark red. Females homozygous for the *pn* mutation were crossed to males hemizygous for the *g* mutation, and the F_1 daughters, all with dark red eyes, were crossed with their brown-eyed brothers. Predict the frequency of sons from this last cross that will have dark red eyes.
- 7.26 Assume that in *Drosophila* there are three X-linked genes x, y, and z, with each mutant allele recessive to the wild-type allele. A cross between females heterozygous for these three loci and wild-type males yielded the following progeny:

Females	+ + +	1010
Males	+ + +	39
	+ + z	430
	+ y z	32
	x + +	27
	x y +	441
	x y z	31
		Total: 2010

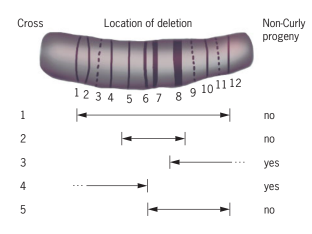
Using these data, construct a linkage map of the three genes and calculate the coefficient of coincidence.

- 7.27 In the nematode *Caenorhabditis elegans*, the linked genes *dpy* (*dumpy* body) and *unc* (*uncoordinated* behavior) recombine with a frequency *P*. If a repulsion heterozygote carrying recessive mutations in these genes is self-fertilized, what fraction of the offspring will be both dumpy and uncoordinated?
- 7.28 In the following testcross, genes *a* and *b* are 20 cM apart, and genes *b* and *c* are 10 cM apart: $a + c/+b + \times a b c/a b c$. If the coefficient of coincidence is 0.5 over this interval on the linkage map, how many triply homozygous recessive individuals are expected among 1000 progeny?
- **7.29** *Drosophila* females heterozygous for three recessive mutations, *a*, *b*, and *c*, were crossed to males homozygous for all three mutations. The cross yielded the following results:

Phenotype	Number
+ + +	75
+ + c	348
+ b c	96
<i>a</i> + +	110
a b +	306
a b c	65

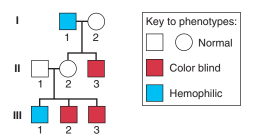
Construct a linkage map showing the correct order of these genes and estimate the distances between them.

7.30 A Drosophila second chromosome that carried a recessive lethal mutation, l(2)g14, was maintained in a stock with a balancer chromosome marked with a dominant mutation for curly wings. This latter mutation, denoted Cy, is also associated with a recessive lethal effect—but this effect is different from that of l(2)g14. Thus, l(2)g14/Cy flies survive, and they have curly wings. Flies without the Cy mutation have straight wings. A researcher crossed l(2)g14/Cy females to males that carried second chromosomes with different deletions (all homozygous lethal) balanced over the Cy chromosome (genotype Df/Cy). Each cross was scored for the presence or absence of progeny with straight wings.

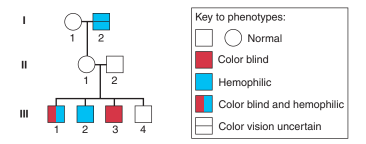


In which band is the lethal mutation l(2)g14 located?

7.31 The following pedigree, described in 1937 by C. L. Birch, shows the inheritance of X-linked color blindness and hemophilia in a family. What is the genotype of II-2? Do any of her children provide evidence for recombination between the genes for color blindness and hemophilia?



7.32 The following pedigree, described in 1938 by B. Rath, shows the inheritance of X-linked color blindness and hemophilia in a family. What are the possible genotypes of II-1? For each possible genotype, evaluate the children of II-1 for evidence of recombination between the color blindness and hemophilia genes.



- 7.33 A normal woman with a color-blind father married a normal man, and their first child, a boy, had hemophilia. Both color blindness and hemophilia are due to X-linked recessive mutations, and the relevant genes are separated by 10 cM. This couple plans to have a second child. What is the probability that it will have hemophilia? color blindness? both hemophilia and color blindness? neither hemophilia nor color blindness?
- **7.34** Two strains of maize, M1 and M2, are homozygous for four recessive mutations, *a*, *b*, *c*, and *d*, on one of the large chromosomes in the genome. Strain W1 is homozygous for the dominant alleles of these mutations. Hybrids produced by crossing M1 and W1 yield many different classes of recombinants, whereas hybrids produced by crossing M2 and W1 do not yield any recombinants at all. What is the difference between M1 and M2?
- **7.35** A *Drosophila* geneticist has identified a strain of flies with a large inversion in the left arm of chromosome 3. This inversion includes two mutations, *e* (*ebony* body) and *cd* (*cardinal* eyes), and is flanked by two other mutations, *sr* (*stripe* thorax) on the right and *ro* (*rough* eyes) on the left. The geneticist wishes to replace the *e* and *cd* mutations inside the inversion with their wild-type alleles; he plans to accomplish this by recombining the multiply mutant, inverted chromosome with a wild-type, inversion-free chromosome. What event is the geneticist counting on to achieve his objective? Explain.

Genomics on the Web at http://www.ncbi.nlm.nih.gov

Chromosome maps were first developed by T. H. Morgan and his students, who used *Drosophila* as an experimental organism.

- 1. Find the genetic map positions of the genes w (white eyes), m (miniature wings), and f (forked bristles) on the X chromosome (also denoted as chromosome 1) of Drosophila melanogaster.
- 2. Find the positions of these three genes on the cytogenetic map of the X chromosome of *D. melanogaster*.

Hint: At the web site, click on Genomes and Maps, then on Genome Project, and finally on Genomic Biology. Then under Genome resources, click on Insects. From there, open the page on *Drosophila melanogaster* and then under Sequencing Centers in the sidebar, click on FlyBase, which is the database for genomic information about *Drosophila*. On the FlyBase main page, search for each of the three genes to obtain the genetic and cytological locations.

- 3. Use the Map Viewer function on the web site to locate w, m, and f on the ideogram of the X chromosome.
- **4.** Homologous genes are genes that have been derived from a common ancestor. The *SRY* gene for sex determination in humans is located on the Y chromosome. A homologue of this gene, called *SOX3*, is located on the X chromosome. Find these two genes on the ideograms of the human sex chromosomes. In what bands do they lie?
- **5.** *RBMX* and *RBMY* are another pair of homologous genes on the human X and Y chromosomes. Locate these two genes relative to *SOX3* and *SRY*. Considering the evolutionary history of the X and Y chromosomes, what might account for the positions of these two pairs of genes on the sex chromosomes?

Hint: Search using the "Find in This View" function on the Map Viewer page of the web site.

The Genetics of Bacteria and Their Viruses

CHAPTER OUTLINE

- Viruses and Bacteria in Genetics
- The Genetics of Viruses
- The Genetics of Bacteria
- Mechanisms of Genetic Exchange in Bacteria
- The Evolutionary Significance of Genetic Exchange in Bacteria

1970s. Indeed, many physicians thought that TB might be totally eliminated. Unfortunately, they were wrong!

On November 16, 1991, a headline in the *New York Times* stated: "A Drug-Resistant TB Results in 13 Deaths in New York Prisons." Then, a prison guard in Syracuse was killed by the same drugresistant strain of *M. tuberculosis* as the prisoners, and, unfortunately,

winter months. Because TB is so contagious, families with the disease lived in almost total isolation. Their friends were afraid to visit for fear of contracting the disease. When Oscar was 14 years old, his mother died, and his life changed immediately. He quit school so that he could take care of his younger siblings while his dad worked.

Multi-Drug-Resistant Bacteria:

Oscar Peterson was a happy child, the son of Norwegian im-

migrants who moved to the Minnesota frontier at the end of the nineteenth century. However, his happy childhood was shortlived. His mother soon became very ill, with incessant coughing, chest pains, and high fevers. She had tuberculosis (TB), a

dreaded disease caused by the bacterium Mycobacterium tuber-

culosis. TB is highly contagious because *M. tuberculosis* is transmitted via aerosolized droplets produced when an infected person

coughs or sneezes. The disease was often fatal because there was

no effective treatment at the time. Fresh air was prescribed, so the

Peterson family slept with the windows open, even during the cold

A Ticking Timebomb?

Thousands of frontier families like the Petersons fought to survive the scourge of TB in the first part of the twentieth century. Then, Alexander Fleming discovered penicillin, and a revolution in the treatment of bacterial diseases followed. During the 1940s and 1950s, scientists



Mycobacterium tuberculosis, the bacterium that causes tuberculosis in humans.

this drug-resistant strain was just the tip of the iceberg. Today, many strains of M. tuberculosis are resistant to a whole battery of drugs and antibiotics. These drug-resistant strains are of two types: multi-drug-resistant (MDR) strains—those resistant to most normally prescribed antibiotics, and extensively drug-resistant (XDR) strains—those also resistant to the antibiotics used to treat MDR-TB. MDR and XDR strains of *M. tuberculosis* are present throughout the world, with especially high frequencies in prisons from New York to Siberia.

discovered an arsenal of highly effective antibiotics. As a result, the incidence of TB decreased sharply in the United States during the

The genetic basis of this multi-drug resistant TB is discussed later in this chapter (see Plasmids and Episomes) and in Chapter 17. How serious a threat does the evolution of MDR and XDR bacteria pose to human health? Dr. Lee Reichman, one of the world's leading experts on TB, has referred to MDR-*M. tuberculosis* as a "timebomb." Worldwide, 2 billion people (15 million in the United States) are infected with latent *M. tuberculosis*. Of these, 8.4 million develop active TB and 2 million die every year. On March 18, 2010, the World Health Organization reported that MDR-TB and XDR-TB have increased to record levels, with 440,000 cases worldwide in 2008. In some areas of the world, one in four individuals with TB cannot be treated effectively with standard antibiotic regimes; they are infected with MDR and XDR strains of *M. tuberculosis*. Perhaps we should initiate steps to confront the crisis of MDR- and XDR-TB now—before the "timebomb" explodes.

Viruses and Bacteria in Genetics

Bacteria and viruses have made important contributions to the science of genetics.

We live in a world along with countless bacteria and viruses. Some bacteria, like *M. tuberculosis*, are harmful; others, like those we use to make yogurt are helpful. Bacteria play important roles in the Earth's ecosytems.

They erode rock, capture energy from materials in their environments, fix atmospheric nitrogen into compounds that other organisms can use, and break down the bodies of organisms that have died. If bacteria did not carry out these functions, life as we know it would not be possible. These tiny organisms enable large, multicellular organisms like us to survive.

Geneticists began to study bacteria and their viruses in the middle of the twentieth century, years after Mendel's Principles and the Chromosome Theory of Heredity had been firmly established. To the first bacterial and viral geneticists, these tiny organisms seemed to offer the possibility of extending genetic analysis to a deeper, biochemical level—indeed, to the very molecules that make up genes and chromosomes. As we will see in this and succeeding chapters, this exciting prospect was realized. The genetic analysis of bacteria and viruses has allowed researchers to probe the chemical nature of genes and their products. All that we now call molecular biology has been founded on the study of bacteria and viruses.

For a research scientist, bacteria and viruses have several advantages compared to creatures like maize or Drosophila. First, they are small, reproduce quickly, and form large populations in just a matter of days. An experimenter can grow 10^{10} bacteria in a small culture tube; 10^{10} Drosophila, by contrast, would fill a 14 ft \times 14 ft \times 14 ft room. Second, bacteria and viruses can be grown on biochemically defined culture media. Because the constituents of the culture medium can be changed as desired, a researcher can identify the chemical needs of the organism and investigate how it processes these chemicals during its metabolism. Drugs such as antibiotics can also be added to the medium to kill bacteria selectively. This type of treatment allows a researcher to identify resistant and sensitive strains of a bacterial species-for example, to determine if M. tuberculosis cultured from a patient is resistant to a particular antibiotic. Third, bacteria and viruses have relatively simple structures and physiology. They are therefore ideal for studying fundamental biological processes. Finally, genetic variability is easy to detect among these tiny microorganisms. If we examine bacteria or viruses, we almost always find that they manifest different phenotypes and that these differences are heritable. For example, some strains of a bacterial species can grow on a biochemically defined medium containing lactose as the only energy source, whereas other strains cannot. Strains that are not able to grow on this type of medium are mutant with respect to the metabolism of lactose. The ability to obtain mutant strains of bacteria and viruses has allowed geneticists to dissect complex phenomena such as energy recruitment, protein synthesis, and cell division at the molecular level.

The advances in molecular biology during the last few decades have provided a wealth of information about the genomes of many bacteria and viruses. Today, we know the complete nucleotide sequences of the genomes of a large number of viruses and bacteria. These sequences are providing detailed information about the genetic control of metabolism in diverse microbial species and, especially, about their evolutionary relationships. We will examine some of this information in Chapter 15 (see Comparative Genomics).

In this chapter we will concentrate on a few bacteria and viruses that have played major roles in genetic analysis. These tiny organisms include the bacterium *Escherichia coli* and two viruses that infect it. We will begin our investigation with the simplest microorganisms—the viruses that infect bacteria such as *E. coli*.

- Their small size, short generation time, and simple structures have made bacteria and viruses valuable model systems for genetic studies.
- Many basic concepts of genetics were first deduced from studies of bacteria and viruses.

The Genetics of Viruses

Viruses straddle the line between the living and the nonliving. Consider, for example, a virus that causes discoloration on the leaves of tobacco plants, a condition called tobacco mosaic disease. The tobacco mosaic virus (TMV) can be crystallized and stored on a shelf for years. In this state, it exhibits none of the properties normally associated with living systems: it does not reproduce; it does not grow or develop; it does not utilize energy; and it does not respond to environmental stimuli. However, if a liquid suspension containing TMV is rubbed onto the leaf of a tobacco Viruses can only reproduce by infecting living host cells. Bacteriophages are viruses that infect bacteria. Several important genetic concepts have been discovered through studies of bacteriophages.

suspension containing TMV is rubbed onto the leaf of a tobacco plant, the viruses in the suspension infect the cells, reproduce, utilize energy supplied by the plant cells, and respond to cellular signals. Clearly, they exhibit the properties of living systems.

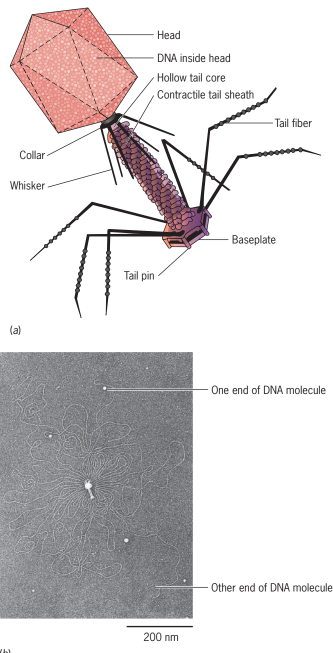
Indeed, it is the simplicity of viruses that has made them ideal research tools for genetic analysis. Questions that have been difficult to answer using more complicated eukaryote systems have often been addressed using viruses. In Chapter 9, we will discuss experiments that used viruses to demonstrate that genetic information is stored in DNA and RNA. In Chapters 10, 11, and 12, we will discuss experiments that used viruses to elucidate the mechanisms of DNA replication, transcription, and translation. In this chapter, we will focus on viruses that infect bacteria: we will discuss the organization of their genomes and the methods that geneticists have developed to analyze them.

Viruses that infect bacteria are called **bacteriophages** (from the Greek "to eat bacteria"). Among the many bacteriophages that have been identified, two have played especially important roles in the elucidation of genetic concepts. Both of these viruses infect the colon bacillus *Escherichia coli*. Bacteriophages can be categorized into two types—virulent and temperate—based on their lifestyles in infected cells. Bacteriophage T4 (phage T4) is a virulent phage; it uses the metabolic machinery of the host cell to produce progeny viruses and kills the host in the process. Bacteriophage lambda (λ), a temperate phage, is another coliphage (phage that infects *E. coli*); however, this phage can either kill the host cell like phage T4, or it can enter into a special association with the host and replicate its genome along with the host cell's genome during each cell duplication. The results of studies performed on bacteriophages T4 and lambda have established genetic paradigms that are relevant to understanding other types of viruses, such as the human immunodeficiency virus, HIV (see Chapter 17 for a discussion of HIV).

BACTERIOPHAGE T4

Bacteriophage T4 is a large virus that stores its genetic information in a doublestranded DNA molecule packaged inside a proteinaceous head (**■** Figure 8.1a). The virus is composed almost entirely of proteins and DNA—approximately half of each (**■** Figure 8.1b). The T4 chromosome is approximately 168,800 base pairs long and contains about 150 characterized genes and an equal number of uncharacterized sequences thought to be genes. The tail of the virus contains several important components. Its central hollow core provides the channel through which the phage

KEY POINTS



(b)

FIGURE 8.1 Bacteriophage T4. (a) Diagram showing the structure of bacteriophage T4 and (b) electron micrograph of a T4 bacteriophage (center) from which the DNA has been released by osmotic shock. Both ends of the linear DNA molecule are visible.

DNA is injected into the bacterium. The tail sheath functions as a small muscle that contracts and pushes the tail core through the bacterial cell wall. The six tail fibers are used to locate receptors on the host cell, and the tail pins on the baseplate then attach firmly to these receptors. All of these components must function correctly for the phage to infect an *E. coli* cell successfully.

Bacteriophage T4 is a **lytic phage**; when it infects a bacterium, it replicates and kills the host, producing about 300 progeny viruses per infected cell (**Figure 8.2**). After the phage DNA is injected into the host bacterium, it quickly (within 2 minutes) directs the synthesis of proteins that shut off the transcription, translation, and replication of bacterial genes, allowing the virus to take control of the metabolic machinery of the host. Some of the phage genes encode nucleases that degrade the host DNA. Other phage proteins initiate the replication of phage DNA. Somewhat later, the genes that encode the structural components of the virus are expressed. Thereafter, the assembly of progeny phage begins; infectious progeny phage start to accumulate in the host cell at about 17 minutes after infection. At about 25 minutes after infection, a phage-encoded enzyme called *lysozyme* degrades the bacterial cell wall and ruptures the host bacterium, releasing about 300 progeny phage per infected cell.

As mentioned above, T4 encodes nucleases that degrade the host DNA. The degradation products are then used in the synthesis of phage DNA. But how do these enzymes degrade host DNA without destroying the DNA of the virus? The answer is that T4 DNA contains an unusual base—5-hydroxymethylcytosine (HMC; cytosine with a —CH₂OH group attached to one of the atoms in the cytosine molecule)—instead of cytosine. In addition, derivatives of glucose molecules are attached to the HMC. These modifications protect T4 DNA from degradation by the nucleases that it uses to degrade the DNA of the host cell.

Genes on bacteriophage T4 chromosomes can be mapped using recombination frequencies, just as in eukaryotes. However, because viruses have a single chromosome that does not go through meiosis, the mapping procedure is somewhat different from that used for an organism like *Drosophila*. Crosses are performed by simultaneously infecting host bacteria with two different types of phage and then screening the progeny phage for recombinant genotypes. Map distances, in centiMorgans, are then calculated as the average number of crossovers that have occurred between genetic markers. For short distances, map distances are approximately equal to the percentage of recombinant chromosomes among the progeny.

There are many different kinds of mutant alleles in T4 phage. Temperature-sensitive (ts) mutations are among the most useful. Wild-type T4 can grow at temperatures ranging from about 25° to over 42° C, whereas heat-sensitive mutants can grow at 25°, but not at 42° C. Thus, ts mutants can be distinguished from wild-type phage by culturing the phage at low and high temperatures. We will

discuss these heat-sensitive mutations and other types of T4 mutants in Chapters 12 and 13.

BACTERIOPHAGE LAMBDA

Bacteriophage lambda (λ) is another coliphage that has made large contributions to genetics. Lambda is smaller than T4; however, its life cycle is more complex. The lambda genome contains about 50 genes in a double-stranded DNA molecule 48,502 base pairs long. This linear DNA molecule is packaged in the λ head (**Tigure 8.3**). Soon after it is injected into an *E. coli* cell, the λ DNA molecule is converted to a circular form, which participates in all subsequent intracellular events.

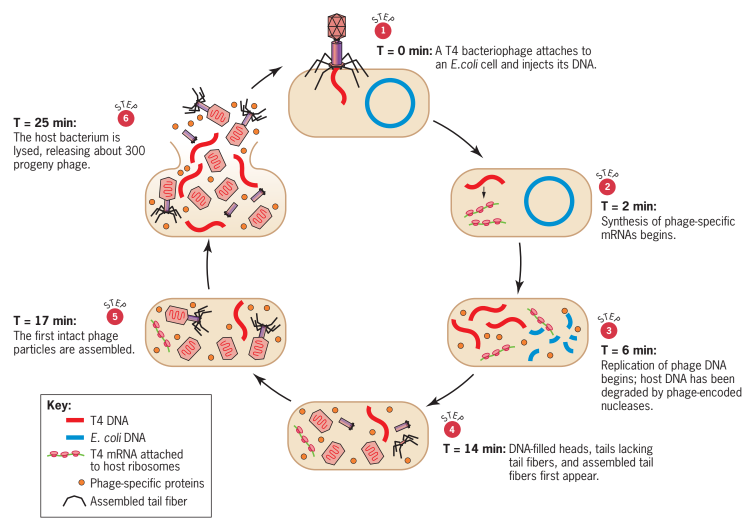


FIGURE 8.2 The life cycle of bacteriophage T4.

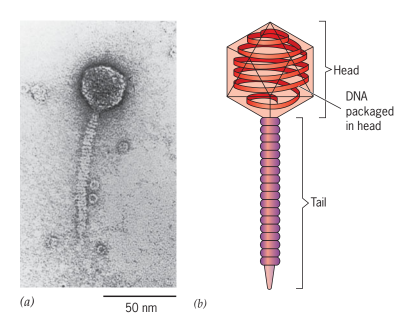


FIGURE 8.3 Bacteriophage λ . Electron micrograph (a) and diagram (b) showing the structure of bacteriophage λ .

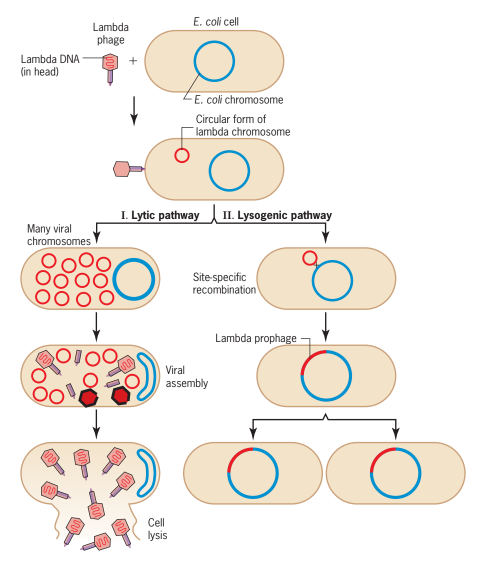


FIGURE 8.4 The life cycle of bacteriophage λ . The two intracellular states of bacteriophage lambda: lytic growth and lysogeny.

Inside the cell, the circular λ chromosome can proceed down either of two pathways (**True 8.4**). It can enter a lytic cycle, during which it reproduces and encodes enzymes that lyse the host cell, just like phage T4. Or, it can enter a **lysogenic** pathway, during which it is inserted into the chromosome of the host bacterium and thereafter is replicated along with that chromosome. In this integrated state, the λ chromosome is called a **prophage**. For this state to continue, the genes of the prophage that encode products involved in the lytic pathway—for example, enzymes involved in the replication of phage DNA, structural proteins required for phage morphogenesis, and the lysozyme that catalyzes cell lysis—must not be expressed.

Integration of the λ chromosome occurs by a site-specific recombination event between the circular λ DNA and the circular *E. coli* chromosome (**True 8.5**). This recombination occurs at specific attachment sites—*attP* on the λ chromosome and *attB* on the bacterial chromosome—and is mediated by the product of the λ *int* gene, the λ integrase. It covalently inserts the λ DNA into the chromosome of the host cell. The site-specific recombination occurs in the central region of the attachment sites where both *attP* and *attB* have the same sequence of 15 nucleotide pairs:

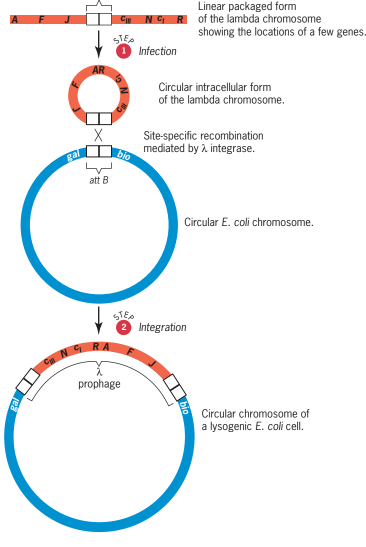
GCTTTTTTTATACTAA CGAAAAAATATGATT

With the exception of this core sequence, *attP* and *attB* have quite different sequences. Because recombination occurs within this core sequence during integration, the resulting *attB/P* and *attP/B* sites that flank the integrated prophage also both contain the 15-nucleotide-pair sequence. These structures are important because they facilitate excision of the prophage by a very similar site-specific recombination event.

About once in every 10^5 cell divisions, the λ prophage spontaneously excises from the host chromosome and enters the lytic pathway. This phenomenon is the reason the prophage is said to be in a lysogenic state, that is, one capable of causing lysis, albeit at low frequency. Excision of the λ prophage can also be induced, for example, by irradiation with ultraviolet light. The excision process is usually precise, with site-specific recombination between the core sequences in *attB/P* and *attP/B*. It produces an autonomous λ chromosome that has the original pre-integration form. Excision requires the λ integrase and the product of the λ xis gene, λ excisase. These two enzymes mediate a site-specific recombination event that is essentially the reverse of the integration event. Occasionally, excision occurs anomalously, and bacterial DNA is excised along with phage DNA. When this occurs, the resulting virus can transfer bacterial genes from one host bacterium to another. We will discuss this process later (see Mechanisms of Genetic Exchange in Bacteria).

Studies on phage λ have contributed much to our understanding of genetic phenomena. We will discuss the replication of the λ chromosome in Chapter 9. The discovery of the λ prophage (for which André Lwoff was awarded a share of the 1965 Nobel Prize in Physiology or Medicine) provided the paradigm for the proviral states of the human immunodeficiency virus (HIV) (Chapter 17) and various vertebrate RNA tumor viruses (Chapter 21).

- Viruses are obligate parasites that can reproduce only by infecting living host cells.
- Bacteriophages are viruses that infect bacteria.
- Bacteriophage T4 is a lytic phage that infects E. coli, reproduces, and lyses the host cell.
- Bacteriophage lambda (λ) can enter a lytic pathway, like T4, or it can enter a lysogenic pathway, during which its chromosome is inserted into the chromosome of the bacterium.
- In its integrated state, the λ chromosome is called a prophage, and its lytic genes are kept turned off.



att P

FIGURE 8.5 Integration of the λ DNA molecule into the chromosome of *E. coli*.

The Genetics of Bacteria

The genetic information of most bacteria is stored in a single main chromosome carrying a few thousand genes and a variable number of "mini-chromosomes" called plasmids and episomes. Plasmids are autonomously replicating, circular DNA molecules that Bacteria contain genes that mutate to produce altered phenotypes. Gene transfer in bacteria is unidirectional from donor cells to recipient cells.

KEY POINTS

carry anywhere from three genes to several hundred genes. Some bacteria contain as many as 11 different plasmids in addition to the main chromosome. Episomes are similar to plasmids, but episomes can replicate either autonomously or as part of the main chromosome—in an integrated state like the λ prophage.

Bacteria reproduce asexually by simple fission, with each daughter cell receiving one copy of the chromosome. They are monoploid but "multinucleate"; that is, the cell usually contains two or more identical copies of the chromosome. The chromosomes of bacteria do not go through the mitotic and meiotic condensation cycles that occur during cell division and gametogenesis in eukaryotes. Therefore, the recombination events—independent assortment and meiotic crossing over—that occur during sexual reproduction in eukaryotes do not occur in bacteria.

Nevertheless, recombination has been just as important in the evolution of bacteria as it has been in the evolution of eukaryotes. Indeed, processes that are akin to sexual reproduction—parasexual processes—occur in bacteria. We will consider these processes after discussing some of the types of mutants used in bacterial genetics and the unidirectional nature of gene transfer between bacteria.

MUTANT GENES IN BACTERIA

Bacteria will grow in liquid medium, often requiring aeration, or on the surface of semisolid medium containing agar. If grown on semisolid medium, each bacterium will divide and grow exponentially, producing a visible colony on the surface of the medium. The number of colonies that appear on a culture plate can be used to estimate the number of bacteria that were originally present in the suspension applied to the plate.

Each bacterial species produces colonies with a specific color and morphology. *Serratia marcescens*, for example, produces a red pigment that results in distinctive red colonies (**Figure 8.6**). Mutations in bacterial genes can change both colony color and morphology. Moreover, any mutation that slows the growth rate of the bacterium will produce small or petite colonies. Some mutations alter the morphology of the bacterium without changing colony morphology. Besides these colony color and morphology mutants, other types of mutants have been useful in genetic studies of bacteria.



■ FIGURE 8.6 Bacterial colonies. Photograph showing colonies of the bacterium *Serratia marcescens* growing on agar-containing medium. The distinctive color of the colonies results from the red pigment produced by this species.

Mutants Blocked in Their Ability to Utilize Specific Energy Sources

Wild-type *E. coli* can use almost any sugar as an energy source. However, some mutants are unable to grow on the milk sugar lactose. They grow well on other sugars but cannot grow on medium containing lactose as the sole energy source. Other mutants are unable to grow on galactose, and still others are unable to grow on arabinose. The standard nomenclature for describing these and other types of mutants in bacteria is to use three-letter abbreviations with appropriate superscripts. For phenotypes, the first letter is capitalized; for genotypes, all three letters are lowercase and italicized. Therefore, wild-type *E. coli* is phenotypically Lac⁺ (able to use lactose as an energy source) and genotypically *lac*⁺. Mutants that are unable to utilize lactose as an energy source are phenotypically Lac⁻ and genotypically *lac*⁻ (or, sometimes just *lac*).

Mutants Unable to Synthesize an Essential Metabolite

Wild-type *E. coli* can grow on medium (minimal medium) containing an energy source and some inorganic salts. These cells can synthesize all of the metabolites—amino acids, vitamins, purines, pyrimidines, and so on—they need from these substances. These wild-type bacteria are called **prototrophs**. When a mutation occurs in a gene encoding an enzyme required for the synthesis of an essential metabolite, the bacterium carrying that mutation will have a new growth requirement. It will grow if the metabolite is added to the medium, but it will not grow in the absence of the metabolite. Such mutants are called **auxotrophs**; they require auxiliary nutrients for growth. As an example, wild-type *E. coli* can synthesize tryptophan *de novo*; these cells are phenotypically Trp^+ and genotypically trp^+ . Tryptophan auxotrophs are Trp^- and trp^- .

Mutants Resistant to Drugs and Antibiotics

Wild-type *E. coli* cells are killed by antibiotics such as ampicillin and tetracycline. Phenotypically, they are Amp^s and Tet^s. The mutant alleles that make *E. coli* resistant to these antibiotics are designated *amp^r* and *tet^r*, respectively. Bacteria that contain these mutant alleles can grow on medium containing the antibiotics, whereas wild-type bacteria cannot. Thus, antibiotics can be used to select bacteria that carry genes for resistance. The resistance genes function as dominant selectable markers.

Bacteria divide rapidly and produce large populations of cells for genetic studies. Moreover, media that select specific bacterial genotypes (selective media) are relatively easy to prepare. As a result, bacteria have been used to study rare events such as mutation within genes and recombination between closely linked genes. We will discuss some of these studies in Chapter 13.

UNIDIRECTIONAL GENE TRANSFER IN BACTERIA

The recombination events that occur in bacteria involve transfers of genes from one bacterium to another, rather than the reciprocal exchanges of genes that occur during meiosis in eukaryotes. Thus, gene transfer is *unidirectional* rather than bidirectional. Recombination events in bacteria usually occur between a fragment of one chromosome (from a **donor cell**) and a complete chromosome (in a **recipient cell**), rather than between two complete chromosomes as in eukaryotes. With rare exceptions, the recipient cells become partial diploids, containing a linear piece of the donor chromosome and a complete circular recipient chromosome. As a result, *crossovers must occur in pairs* and must insert a segment of the donor chromosome into the recipient chromosome (**Tigure 8.7a**). If a single crossover (or any odd number of crossovers) occurs, it will destroy the integrity of the recipient chromosome, producing a nonviable linear DNA molecule (**Tigure 8.7b**).

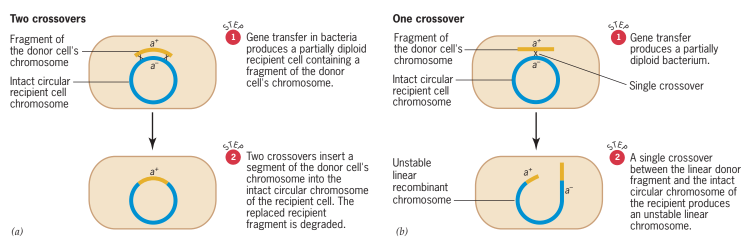


FIGURE 8.7 Recombination in bacteria. The parasexual processes that occur in bacteria produce partial diploids containing linear fragments of the donor cell's chromosome and intact circular chromosomes of the recipient cells. (a) To maintain the integrity of the circular chromosomes, crossovers must occur in pairs, inserting segments of the donor chromosomes into the chromosomes of the recipient. (b) A single crossover between a fragment of a donor chromosome and a circular recipient chromosome destroys the integrity of the circular chromosome, producing a linear DNA molecule that is unable to replicate and is subsequently degraded.

KEY POINTS • Bacteria usually contain one main chromosome.

- Wild-type bacteria are prototrophs; they can synthesize everything they need to grow and reproduce given an energy source and some inorganic molecules.
- Auxotrophic mutant bacteria require additional metabolites for growth.
- Gene transfer in bacteria is unidirectional; genes from a donor cell are transferred to a recipient cell, with no transfer from recipient to donor.

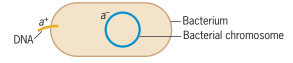
Mechanisms of Genetic Exchange in Bacteria

Bacteria exchange genetic material through three different parasexual processes.

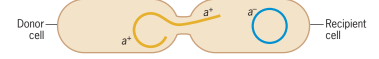
Three distinct parasexual processes occur in bacteria. The most obvious difference between these three processes is the mechanism by which DNA is transferred from one cell to

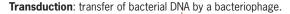
another (**Figure 8.8**). **Transformation** involves the uptake of free DNA molecules released from one bacterium (the donor cell) by another bacterium (the recipient cell). **Conjugation** involves the direct transfer of DNA from a donor cell to a recipient cell. **Transduction** occurs when bacterial genes are carried from a donor cell to a recipient cell by a bacteriophage.

Transformation: uptake of free DNA.



Conjugation: direct transfer of DNA from one bacterium to another.





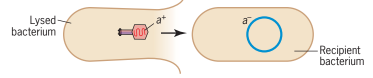


FIGURE 8.8 The three types of gene transfer in bacteria.

TABLE 8.1

Distinguishing between the Three Parasexual Processes in Bacteria

	Criterion			
Recombination Process	Cell Contact Required?	Sensitive to DNase?		
Transformation	no	yes		
Conjugation	yes	no		
Transduction	no	no		

The three parasexual processes of gene transfer—transformation, conjugation, and transduction—in bacteria can be distinguished by two simple criteria (Table 8.1). (1) Does the process require cell contact? (2) Is the process sensitive to deoxyribonuclease (DNase), an enzyme that degrades DNA? These two criteria can be tested experimentally quite easily. Sensitivity to DNase is determined simply by adding the enzyme to the medium in which the bacteria are growing. If gene transfer no longer occurs, the process involves transformation. The protein coats of bacteriophages and the walls and membranes of bacterial cells protect the donor DNA from degradation by DNase during transduction and conjugation, respectively.

A simple experiment can determine whether or not cell contact is required for bacterial gene transfer. In this experiment, bacteria with different genotypes are placed in opposite arms of a U-shaped culture tube (**Figure 8.9**). The two arms are separated by a glass filter that has pores large enough to allow DNA molecules and viruses, but not bacteria, to pass through it. If gene transfer occurs between the bacteria growing in opposite arms of the U-tube, the process cannot be conjugation, which requires direct contact between donor and recipient cells. If the observed gene transfer occurs in the presence of DNase and in the absence of cell contact, it must involve transduction.

All three parasexual processes do not occur in all bacterial species; in fact, transduction probably is the only process that occurs in all bacteria. Whether or not transformation or conjugation occurs in a species depends on whether the required genes and metabolic machinery have evolved in that species. *E. coli*, for example, does not contain genes that encode the proteins required to take up free DNA. Thus, transformation does not occur in *E. coli* growing under natural conditions. Only conjugation and transduction occur in *E. coli* cells growing in natural habitats. However, scientists have discovered how to transform *E. coli* cells in the laboratory by using chemical or physical treatments that make them permeable to DNA. In Chapter 14, we will discuss the use of artificial transformation methods to "clone" (make many copies of) foreign genes in *E. coli* cells.

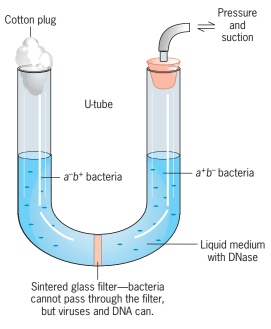
TRANSFORMATION

Frederick Griffith discovered transformation in *Streptococcus pneumoniae* (pneumococcus) in 1928. Pneumococci, like all other living organisms, exhibit genetic variability that can be recognized by the existence of different phenotypes (Table 8.2). The two phenotypic characteristics of importance in Griffith's demonstration of transformation are (1) the presence or absence of a polysaccharide (complex sugar polymer) capsule surrounding the bacterial cells, and (2) the type of capsule-that is, the specific molecular composition of the polysaccharides present in the capsule. When grown on blood agar medium in petri dishes, pneumococci with capsules form large, smooth colonies (**Figure 8.10**) and are thus designated Type S. Encapsulated pneumococci are virulent (pathogenic), causing pneumonia in mammals such as mice and humans. The virulent Type S pneumococci mutate to an avirulent (nonpathogenic) form that has no polysaccharide capsule at a frequency of about one per 10⁷ cells. When grown on blood agar medium, such nonencapsulated, avirulent pneumococci produce small, rough-surfaced colonies (Figure 8.10) and are thus designated Type R. The polysaccharide capsule is required for virulence because it protects the bacterial cell from destruction by white blood cells. When a capsule is present, it may be of several different antigenic types (Type I, II, III, and so forth), depending on the specific molecular composition of the polysaccharides and, of course, ultimately depending on the genotype of the cell.

The different capsule types can be identified immunologically. If Type II cells are injected into the bloodstream of rabbits, the immune system of the rabbits will produce antibodies that react specifically with Type II cells. Such Type II antibodies will agglutinate Type II pneumococci but not Type I or Type III pneumococci.

Griffith's unexpected discovery was that if he injected heat-killed Type IIIS pneumococci (virulent when alive) plus live Type IIR pneumococci (avirulent) into mice, many of the mice succumbed to pneumonia, and live Type IIIS cells were recovered from the carcasses (**Figure 8.11**). When mice were injected with heat-killed Type IIIS pneumococci alone, none of the mice died. The observed virulence was therefore not due to a few Type IIIS cells that survived the heat treatment. The live pathogenic pneumococci recovered from the carcasses had Type III polysaccharide capsules. This result is important because nonencapsulated Type R cells can mutate back to encapsulated Type S cells. However, when such a mutation occurs in a Type IIR cell, the resulting cell will become Type IIS, not Type IIIS. Thus, the transformation of avirulent Type IIR cells to virulent Type IIIS cells cannot be explained by mutation. Instead, some component of the dead Type IIIS cells (the "transforming principle") must have converted living Type IIR cells to Type IIIS.

Subsequent experiments by Richard Sia and Martin Dawson in 1931 showed that the phenomenon described by Griffith, now called transformation, was not mediated in any way by a living host. The same phenomenon occurred in the test tube when



■ FIGURE 8.9 The U-tube experiment with bacteria. The U-tube is used to determine whether or not cell contact is required for recombination to occur. Bacteria of different genotypes are placed in separate arms of the tube, separated by a glass filter that prevents contact between them. If recombination occurs, it cannot be due to conjugation.

TABLE 8.2

Characteristics of Streptococcus pneumoniae Strains When Grown on Blood Agar Medium

Colony Morphology					th Antiserum d Against	
Туре	Appearance	Size	Capsule	Virulence	Type IIS	Type IIIS
IIR ^a	Rough	Small	Absent	Avirulent	None	None
IIS	Smooth	Large	Present	Virulent	Agglutination	None
IIIRª	Rough	Small	Absent	Avirulent	None	None
IIIS	Smooth	Large	Present	Virulent	None	Agglutination

^a Although Type R cells are nonencapsulated, they carry genes that would direct the synthesis of a specific kind (antigenic Type II or III) of capsule if the block in capsule formation were not present. When Type R cells mutate back to encapsulated Type S cells, the capsule Type (II or III) is determined by these genes. Thus, R cells derived from Type IIS cells are designated Type IIR. When these Type IIR cells mutate back to encapsulated Type S cells, the capsuleted Type S cells, the capsules are of Type II.

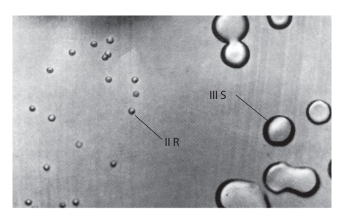


FIGURE 8.10 Colony phenotypes of the two strains of *Streptococcus pneumoniae* studied by Griffith in 1928.

live Type IIR cells were grown in the presence of heat-killed Type IIIS cells. Since Griffith's experiments demonstrated that the Type IIIS phenotype of the transformed cells was passed on to progeny cells—that is, was due to a permanent inherited change in the genotype of the cells—the demonstration of transformation set the stage for determining the chemical basis of heredity in pneumococcus. Indeed, the first proof that genetic information is stored in DNA rather than proteins was the 1944 demonstration by Oswald Avery, Colin MacLeod, and Maclyn McCarty that DNA was responsible for transformation in pneumococci. Because of its pivotal role in establishing DNA as the genetic material, we will discuss this demonstration in Chapter 9.

The mechanism of transformation has been studied in considerable detail in *S. pneumoniae*, *Bacillus subtilis*, *Haemophilus influenzae*, and *Neisseria gonorrhoeae*. The basic process is similar in all four species; however, variations in the mechanism occur in each species. *S. pneu-*

moniae and *B. subtilis* will take up DNA from any source, whereas *H. influenzae* and *N. gonorrhoeae* will only take up their own DNA or DNA from closely related species. *H. influenzae* and *N. gonorrhoeae* will only take up DNA that contains a special short nucleotide-pair sequence (11 base pairs in *Haemophilus*; 10 in *Neisseria*) that is present in about 600 copies in their respective genomes.

Even in the bacterial species that have the ability to take up DNA from their environment, not all cells can do so. Indeed, only cells that are expressing the genes that encode proteins required for the process are capable of taking up DNA. These bacteria are said to be **competent**, and the proteins that mediate the transformation process are called **competence (Com) proteins**. Bacteria develop competence during

> the late phase of their growth cycle—when cell density is high but before cell division stops. The process by which cells become competent is understood best in *B. subtilis*, where small peptides called competence pheromones are secreted by cells and accumulate at high cell density. High concentrations of the pheromones induce the expression of the genes encoding proteins required for transformation to occur.

Let's focus on the mechanism of transformation in B. subtilis (**Figure 8.12**). The competence genes are located in clusters, and each cluster is designated by a letter-for example, A, B, C. The first gene in each cluster is designated A, the second B, and so on. Thus, the protein encoded by the first gene in the fifth cluster is designated ComEA. ComEA and ComG proteins bind double-stranded DNA to the surfaces of competent cells. As the bound DNA is pulled into the cell by the ComFA DNA translocase (an enzyme that moves or "translocates" DNA), one strand of DNA is degraded by a deoxyribonuclease (an enzyme that degrades DNA), and the other strand is protected from degradation by a coating of single-stranded DNA-binding protein and RecA protein (a protein required for recombination). With the aid of RecA and other proteins that mediate recombination, the single strand of transforming DNA invades the chromosome of the recipient cell, pairing with the complementary strand of DNA and replacing the equivalent strand. The replaced recipient strand is then degraded. If the donor and recipient cells carry different alleles of a gene, the resulting recombinant double helix will have one allele in one strand and the other allele in the second strand. A DNA double helix of this type is called a heteroduplex (a "heterozygous" double helix); it will segregate into two homoduplexes when it replicates.

The DNA molecules taken up by competent cells during transformation are usually only 0.2 to 0.5 percent of the complete chromosome. Therefore, unless two genes are quite close together, they will never be present on the same molecule of transforming DNA. Double transformants for two genes (say, *a* to a^+ and *b* to b^+ , using an a^+b^+ donor and an

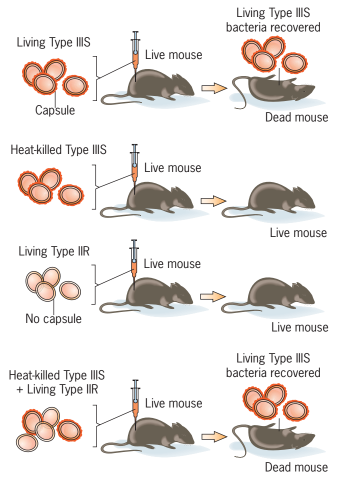


FIGURE 8.11 Griffith's discovery of transformation in *Streptococcus pneumoniae*.

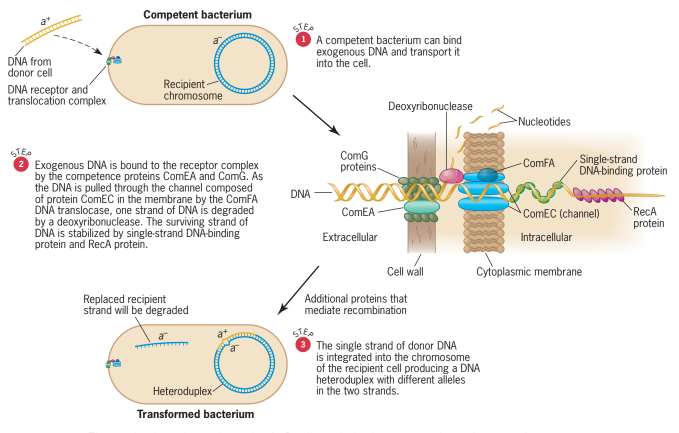


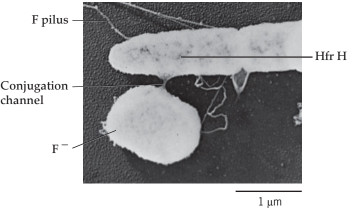
FIGURE 8.12 The mechanism of transformation in *Bacillus subtilis*. A competent bacterium contains a DNA receptor/translocation complex that can bind exogenous DNA and transport it into the cell, where it can recombine with chromosomal DNA of the recipient cell. ComEA, EC, FA, and G are competence proteins; they are synthesized only in competent cells. See the text for additional details.

a b recipient) will require two independent transformation events (uptake and integration of one DNA molecule carrying a^+ and of another molecule carrying b^+). The probability of two such independent events occurring together will equal the product of the probability of each occurring alone. If, by contrast, two genes are closely linked, they may be carried on a single molecule of transforming DNA, and double transformants may be formed at a high frequency. The frequency with which two genetic markers are cotransformed can thus be used to estimate how far apart they are on the host chromosome.

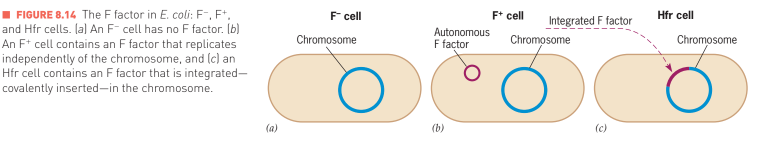
CONJUGATION

Transformation does not occur in *E. coli*—the most intensely studied bacterial species—under natural conditions. Thus, we could ask if there is any kind of gene transfer between *E. coli* cells. The answer to this question is "yes." In 1946, Joshua Lederberg and Edward Tatum discovered that *E. coli* cells transfer genes by conjugation. Their important discovery is discussed further in A Milestone in Genetics: Conjugation has proven to be an important method of genetic mapping in bacterial species where it occurs, and it is an invaluable tool in genetic research.

During conjugation, DNA is transferred from a donor cell to a recipient cell through a specialized intercellular conjugation channel, which forms between them (**Figure 8.13**). Note that the donor and recipient cells are in direct contact during conjugation; the separation observed in Figure 8.13 is the result of stretching forces during preparation for microscopy.



■ FIGURE 8.13 Conjugation in *E. coli*. This early electron micrograph by Thomas F. Anderson shows conjugation between an Hfr H cell and an F⁻ cell. Donor and recipient cells are actually in close juxtaposition during conjugation. The conjugation channel shown here has been stretched during preparation for microscopy.



Donor cells have cell-surface appendages called F pili (singular, F pilus). The synthesis of these F pili is controlled by genes present on a small circular molecule of DNA called an **F factor** (for *f*ertility *factor*). Most F factors are approximately 10⁵ nucleotide pairs in size (see Figure 8.20). Bacteria that contain an F factor are able to transfer genes to other bacteria. The F pili of a donor cell make contact with a recipient cell that lacks an F factor and attach to that cell, so that the two cells can be pulled into close contact. In the past, DNA was thought to move from a donor cell to a recipient cell through an F pilus. However, more recent experiments have shown this idea to be incorrect. The F pili are involved only in establishing cell contact, not in DNA transfer. After the F pili bring a donor cell and a recipient cell together, a conjugation channel forms between the cells, and DNA is transferred from the donor cell to the recipient cell through this channel.

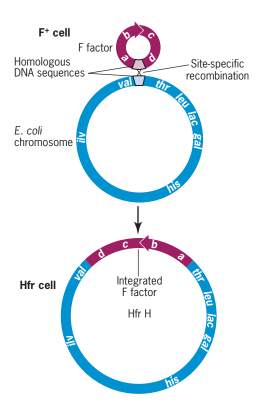
The F factor can exist in either of two states: (1) the *autonomous state*, in which it replicates independently of the bacterial chromosome, and (2) the *integrated state*, in which it is covalently inserted into the bacterial chromosome and replicates like any other segment of that chromosome (**Figure 8.14**). Genetic elements with these properties are called episomes (see Plasmids and Episomes later in this chapter). A donor cell carrying an autonomous F factor is called an F^+ cell. A recipient cell lacking an F factor is called an F^- cell. When an F^+ cell conjugates (or "mates") with an F^- recipient cell, only the F factor is transferred. Both cells (donor and recipient) become F^+ cells because the F factor is replicated during transfer, and each cell receives a copy. Thus, if a population of F^+ cells is mixed with a population of F^- cells, virtually all of the cells will acquire an F factor.

The F factor can integrate into the bacterial chromosome by site-specific recombination events (**Theorem Figure 8.15**). The integration of the F factor is mediated by short DNA sequences that are present in multiple copies in both the F factor and the bacterial chromosome. Thus, an F factor can integrate at many different sites in the bacterial chromosome. A cell that carries an integrated F factor is called an **Hfr cell** (for *b*igh-*fr*equency *r*ecombination). In its integrated state, the F factor mediates the transfer of the chromosome from the Hfr cell to a recipient (F^-) cell during conjugation. Usually, the cells separate before chromosome transfer is complete; thus, only rarely will an entire chromosome be transferred from an Hfr cell to a recipient cell.

The mechanism that transfers DNA from a donor cell to a recipient cell during conjugation appears to be the same if just the F factor is being transferred, as in $F^+ \times F^-$ matings, or if the bacterial chromosome is being transferred, as in Hfr $\times F^$ matings. Transfer is initiated at a special site called *oriT*—the *orig*in of *t*ransfer—one of three sites on the F factor at which DNA replication can be initiated. The other two sites—*oriV* and *oriS*—are used to initiate replication during cell division, not during conjugation. *oriV* is the primary origin of replication during cell fission; *oriS* is a secondary origin that performs this function when *oriV* is absent or nonfunctional.

During conjugation, one strand of the circular DNA molecule is cut at *oriT* by an enzyme, and one end is transferred into the recipient cell through the channel that forms between the conjugating cells (**■** Figure 8.16). The F factor or the Hfr chromosome containing the F factor replicates during transfer by a mechanism called *rolling-circle replication*, because the circular DNA molecule "rolls" during replication (see Chapter 10, Figure 10.30). During conjugation, one copy of the donor chromosome is synthesized in the donor cell, and the transferred strand of donor DNA is replicated in the recipient cell.

Because transfer is initiated within the integrated F factor, part of the F factor is transferred prior to the transfer of chromosomal genes in Hfr \times F⁻ matings. The



■ FIGURE 8.15 The formation of an Hfr cell by the integration of an autonomous F factor. The F factor is covalently inserted into the chromosome by site-specific recombination between homologous DNA sequences in the F factor and the chromosome.

rest of the F factor is transferred after the chromosomal genes. Thus, the recipient cell acquires a complete F factor and is converted to an Hfr cell only in rare cases when an entire Hfr chromosome is transferred.

Several of the details of conjugation were worked out using one particular Hfr strain called Hfr H (for the English microbial geneticist William Hayes, who isolated it). In this strain, the F factor is integrated near the thr (threonine) and leu (leucine) loci, as shown in Figure 8.15. In 1957 Elie Wollman and François Jacob, working at the Pasteur Institute in Paris, provided new insight into the process of conjugation by crossing Hfr H cells of genotype thr^+ leu⁺ azi^s ton^s lac⁺ gal⁺ str^s with F^- cells of genotype $thr^- leu^- azi^r ton^r$ $lac^{-}gal^{-}str^{r}$. The *thr* gene and the *leu* gene are responsible for the syntheses of the amino acids threonine and leucine, respectively. Allele pairs azis/azir, tons/tonr, and str^s/str^r control sensitivity (s) or resistance (r) to sodium azide, bacteriophage T1, and streptomycin, respectively. Alleles lac⁺ and lac^{-} and alleles gal^{+} and gal^{-} govern the ability (+) or inability (-) to utilize lactose and galactose, respectively, as energy sources.

At varying times after the Hfr H and F⁻ cells were mixed to initiate mating, samples were removed and agitated vigorously in a blender to break the conjugation bridges and separate the conjugating cells. These cells, whose mating had been so unceremoniously interrupted, were then plated on medium containing the antibiotic streptomycin but lacking the amino acids threonine and leucine. Only recombinant cells carrying the *thr*⁺ and *leu*⁺ genes of the Hfr H parent and the str^r gene of the F

Mechanisms of Genetic Exchange in Bacteria F⁺ donor cell F⁻ recipient cell F factor `Pilus Donor Recipient chromosome chromosome The F pili of the F⁺ donor cell make contact with the F⁻ recipient cell and pull the cells together. Genes on the F factor direct the synthesis of the conjugation bridge. One strand of DNA is cleaved at the origin of replication of the F factor. Origin Conjugation bridge provides channel between cells STED Rolling-circle replication transfers one strand of the F factor into the recipient cell. Replication of the F factor occurs in both cells (one strand is synthesized in each cell) during transfer. Rolling-circle replication Replication Transfer of the F factor is completed, vielding two F⁺ bacteria.

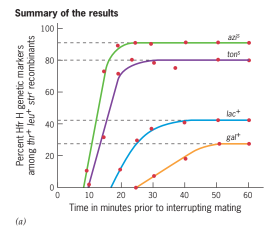
F⁺ cell

parent could grow on this selective medium. The Hfr H donor cells would be killed by the streptomycin, and the F⁻ recipient cells would not grow without threonine and leucine.

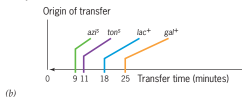
Colonies produced by $thr^+ leu^+ str^r$ recombinants were then transferred (see Chapter 13, Figure 13.15) to a series of plates containing different selective media to determine which of the other donor markers were present. The series of plates included medium containing specific supplements that allowed Wollman and Jacob to determine whether the recombinants contained donor or recipient alleles of each of the genes. Medium containing sodium azide was used to distinguish between azis and azi^r cells. Medium containing bacteriophage T1 was used to score recombinant bacteria as ton^s or ton^r. Medium containing lactose as the sole carbon source was used to determine whether recombinants were lac⁺ or lac⁻, and medium with galactose as the sole carbon source was used to identify gal^+ and gal^- recombinants.

When conjugation was interrupted prior to 8 minutes after mixing the Hfr H cells and the F^- cells, no thr⁺ leu⁺ str^r recombinants were detected. Recombinants (thr⁺ leu⁺ str^r) first appeared at about 8 1/2 minutes after mixing the Hfr H and F⁻ cells and accumulated to a maximum frequency within a few minutes. When the ■ FIGURE 8.16 Mating between an F⁺ cell and an F⁻ cell. The F factor of the donor cell is replicated during transfer from an F⁺ cell to an F^- cell. When the process is complete, each cell has a copy of the F factor.

F⁺ cell



Interpretation of the results



■ **FIGURE 8.17** Wollman and Jacob's classic interrupted mating experiment. (*a*) The frequencies of the unselected donor alleles present in *thr*⁺ *leu*⁺ *str*^r recombinants are shown as a function of the time at which mating was interrupted. (*b*) Interpretation of the results based on the linear transfer of genes from the Hfr cell to the F⁻ cell. Transfer is initiated at the origin on the F factor, and the time at which a gene is transferred to the F⁻ cell depends on its distance from the F factor.

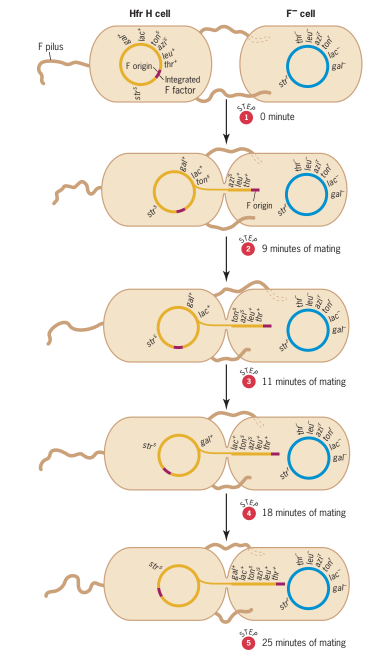


FIGURE 8.18 The interpretation of Wollman and Jacob's interrupted mating experiment. A linear transfer of genes occurs from the donor (Hfr H) cell to the recipient (F⁻) cell. Transfer begins at the origin of replication on the integrated F factor and proceeds with the sequential transfer of genes based on their location on the chromosome. The chromosome replicates during the transfer process so that the Hfr and F⁻ cells both end up with a copy of the transferred DNA.

presence of the other donor markers was examined at varying times after mixing the donor and recipient cells, donor alleles were transferred to recipient cells in a specific temporal sequence (**Figure 8.17**). The Hfr H *azi^s* gene first appeared in recombinants at about 9 minutes after mixing the Hfr and F⁻ bacteria. The *ton^s*, *lac*⁺, and *gal*⁺ markers first appeared after 11, 18, and 25 minutes of mating, respectively. These results indicated that the genes from Hfr H were being transferred to the F⁻ cells in a specific temporal order, reflecting the order of the genes on the chromosome (**Figure 8.18**).

Subsequent studies with different Hfr strains revealed that gene transfer could be initiated at different sites on the chromosome. We now know that the F factor can

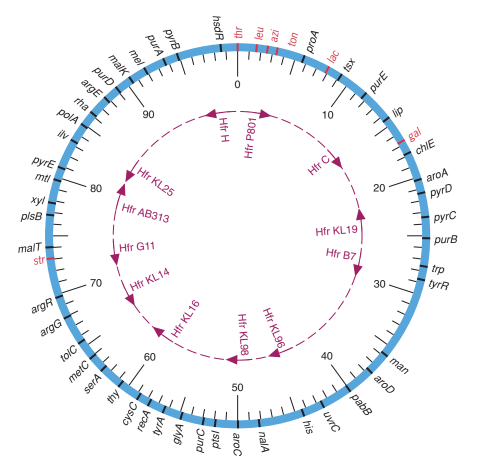


FIGURE 8.19 The circular linkage map of *E. coli*. The inner circle shows the sites of integration of the F factor in selected Hfr strains. The arrows indicate whether transfer by the Hfr's is clockwise or counterclockwise. The outer circle shows the position of selected genes. The map is divided into 100 units, where each unit is the length of DNA transferred during one minute of conjugation. The genes shown in red were used in Wollman and Jacob's famous interrupted mating experiment (see Figures 8.17 and 8.18).

integrate at many different sites in the *E. coli* chromosome and that the site of integration determines where gene transfer is initiated in each Hfr strain. Moreover, the orientation of F factor integration—either $d \ c \ b \ a$ reading clockwise or $a \ b \ c \ d$ reading clockwise (see Figure 8.15)—determines whether the transfer of genes is clockwise relative to the *E. coli* linkage map or counterclockwise (**Tegure 8.19**).

The transfer of a complete chromosome from an Hfr to an F^- cell takes about 100 minutes, and transfer appears to proceed at a fairly constant rate. Thus, the time required for transfer of genes during conjugation can be used to map genes on bacterial chromosomes. A map distance of 1 minute corresponds to the length of a chromosomal segment transferred in 1 minute of conjugation under standard conditions. The linkage map of *E. coli* is therefore divided into 100 one-minute intervals (see Figure 8.19). The zero coordinate of this circular map has been arbitrarily set at the *thrA* gene. When a new mutation is identified in *E. coli*, its location on the chromosome is first determined by conjugation mapping. To test your understanding of conjugation mapping, deduce the chromosomal locations of the genes discussed in Problem-Solving Skills: Mapping Genes Using Conjugation Data at the end of this section. More precise mapping can subsequently be done using transformation or transduction.

PLASMIDS AND EPISOMES

As previously mentioned, the genetic material of a bacterium is carried in one main chromosome plus from one to several extrachromosomal DNA molecules called plasmids.

PROBLEM-SOLVING SKILLS

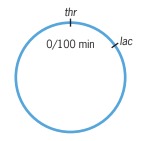
Mapping Genes Using Conjugation Data

THE PROBLEM

You have identified a mutant *E. coli* strain that cannot synthesize the amino acid tryptophan (Trp⁻). To determine the location of the *trp⁻* mutation on the *E. coli* chromosome, you have carried out interrupted mating experiments with four different Hfr strains. In all cases, the Hfr strains carried the dominant wild-type alleles of the marker genes, and the F⁻ strain carried the recessive mutant alleles of these genes. The following chart shows the time of entry in minutes (in parentheses) of the wild-type alleles of the marker genes into the Trp⁻ F⁻ strain. The marker genes are *thr*⁺, *aro*⁺, *his*⁺, *tyr*⁺, *met*⁺, *arg*⁺, and *ilv*⁺ (encoding enzymes required for the synthesis of the amino acids threonine, the aromatic amino acids phenylalanine, tyrosine, and tryptophan, histidine, tyrosine, methionine, arginine, and isoleucine plus valine, respectively), and *man*⁺, *gal*⁺, *lac*⁺, and *xyl*⁺ (required for the ability to catabolize the sugars mannose, galactose, lactose, and xylose, respectively, and use them as energy sources).

Hfr A — man ⁺ (1)	trp+ (9)	<i>aro</i> +(17)	gal+(20)	lac+ (29)	thr+ (37)
Hfr B — <i>trp</i> ⁺ (6)	man ⁺ (14)	his+(22)	tyr+ (34)	met+ (42)	arg+ (48)
Hfr C — <i>thr</i> +(3)	ilv+ (20)	xyl+(25)	arg+(33)	met+ (39)	tyr+ (47)
Hfr D — <i>met</i> ⁺ (2)	arg+ (8)	<i>xyl</i> +(16)	<i>ilv</i> + (21)	thr+ (38)	lac ⁺ (46)

On the map of the circular *E. coli* chromosome shown here, indicate (1) the relative location of each gene, (2) the position where the F factor is integrated in each of the four Hfr's, and (3) the direction of chromosome transfer for each Hfr (clockwise or counterclockwise; indicate direction with an arrow).



FACTS AND CONCEPTS

- 1. The chromosome of *E. coli* contains a circular DNA molecule.
- Chromosomal DNA is transferred from Hfr donor cells to F⁻ recipient cells by rolling-circle replication.
- **3.** Rolling-circle replication, and thus transfer of chromosomal genes, is initiated at the origin of replication on the integrated F factor.

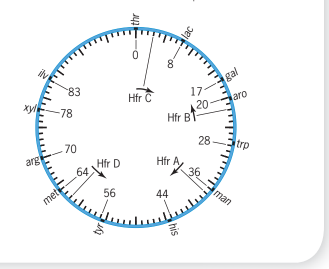
- The direction of transfer (clockwise or counterclockwise) depends on the orientation of the F factor in the Hfr chromosome.
- **5.** The F factor can integrate at many sites in the *E. coli* chromosome and in either orientation (clockwise or counterclockwise).
- **6.** The genetic map of the *E. coli* chromosome is divided into minutes, where 1 minute is the length of DNA transferred from an Hfr strain to an F⁻ strain during 1 minute of conjugation.
- Transfer of the entire chromosome from an Hfr cell to an F[−] cell takes 100 minutes; therefore, the linkage map of the complete circular chromosome totals 100 minutes.
- **8.** The *thr* locus has been arbitrarily assigned position "0" on the map of the *E. coli* chromosome, with linkage distance increasing from 0 to 100 minutes moving clockwise from *thr*.

ANALYSIS AND SOLUTION

If we examine the sequence in which genes are transferred from each Hfr strain to the ${\sf F}^-$ strain, we observe a linear sequence in each case.

Moreover, note that regardless of the sequence in which genes are transferred by different Hfr strains, the distance between adjacent genes remains the same. The distance between *man* and *trp* is 8 minutes, for example, regardless of whether Hfr strain A or B is used in the experiment. Indeed, if we combine the results obtained using the four Hfr strains and place *thr* at position 0, the data yield the following circular genetic map. The circular map is a satisfying result given that we know that the chromosomal DNA of *E. coli* is also circular.

For further discussion visit the Student Companion site.

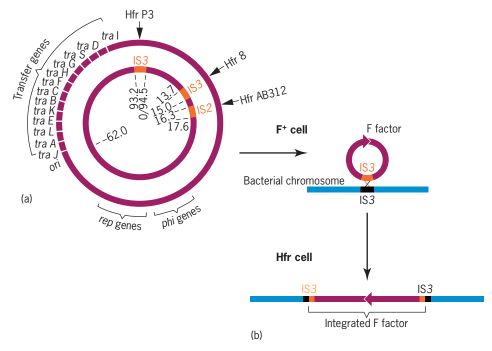


By definition, a **plasmid** is a genetic element that can replicate independently of the main chromosome in an extrachromosomal state. Most plasmids are dispensable to the host; that is, they are not required for survival of the cell in which they reside. However, under certain environmental conditions, such as when an antibiotic is present, they may be essential if they carry a gene for resistance to the antibiotic.



There are three major types of plasmids in *E. coli*: F factors, R plasmids, and Col plasmids. Fertility (F) factors were discussed earlier (see Conjugation). R plasmids (*resistance plasmids*) carry genes that make host cells resistant to antibiotics and other antibacterial drugs. Col plasmids (previously called colicinogenic factors) encode proteins that kill sensitive *E. coli* cells. There are a large number of distinct Col plasmids; however, they will not be discussed further here.

Some plasmids endow host cells with the ability to conjugate. All F⁺ plasmids, many R plasmids, and some Col plasmids have this property; we say that they are conjugative plasmids. Other R and Col plasmids do not endow cells with the ability to conjugate; we say that they are nonconjugative. The conjugative nature of many R plasmids plays an important role in the rapid spread of antibiotic and drug resistance genes through populations of pathogenic bacteria, as discussed at the beginning of this chapter. The evolution of R plasmids that make host bacteria resistant to multiple antibiotics has become a serious medical problem, and the use of antibiotics for nontherapeutic purposes has contributed to the rapid evolution of multiple drug-resistant bacteria (see On the Cutting Edge: Antibiotic-Resistant Bacteria).



■ FIGURE 8.20 IS elements mediate the integration of the F factor. (a) An abbreviated map of the structure of the F factor in *E. coli* strain K12, with distances given in kilobases (1000 nucleotide pairs). The locations of genes required for conjugative transfer (*tra* genes), replication (*rep* genes), and the inhibition of phage growth (*phi* genes) are shown, along with the positions of three IS elements. The arrows denote the specific IS element that mediated the integration of the F factor during the formation of the indicated Hfr strains. (*b*) Recombination between IS elements inserts the F factor into the bacterial chromosome, producing an Hfr.

In 1958 François Jacob and Elie Wollman recognized that the F factor and certain other genetic elements had unique properties. They defined this class of elements and called them episomes. According to Jacob and Wollman, an **episome** is a genetic element that is unessential to the host and that can replicate either autonomously or be integrated (covalently inserted) into the chromosome of the host bacterium. The terms *plasmid* and *episome* are not synonyms. Many plasmids do not exist in integrated states and are thus not episomes. Similarly, many lysogenic phage chromosomes, such as the phage λ genome, are episomes but are not plasmids.

The ability of episomes to insert themselves into chromosomes depends on the presence of short DNA sequences called insertion sequences (or IS elements). The IS elements are present in both episomes and bacterial chromosomes. These short sequences (from about 800 to about 1400 nucleotide pairs in length) are transposable; that is, they can move from one chromosome to a different chromosome (see Chapter 17). In addition, IS elements mediate recombination between otherwise nonhomologous genetic elements. The role of IS elements in mediating the integration of episomes is well documented in the case of the F factor in *E. coli*. Crossing over between IS elements in the F factor and the bacterial chromosome produces Hfr's with different origins and directions of transfer during conjugation (**Tegure 8.20**).

F' FACTORS AND SEXDUCTION

As discussed in the preceding section, an Hfr strain is produced by the integration of an F factor into the chromosome by recombination between IS elements in the chromosome and IS elements in the F factor (see Figure 8.20). Do you think that this recombination process might be reversible? Indeed, rare F⁺ cells are present in Hfr cultures, indicating that excision of the F factor does occur (by a process that is essentially the

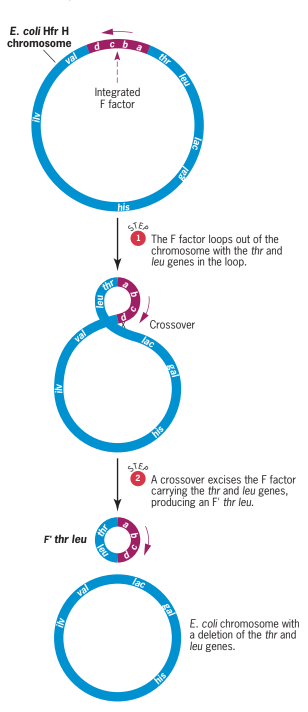
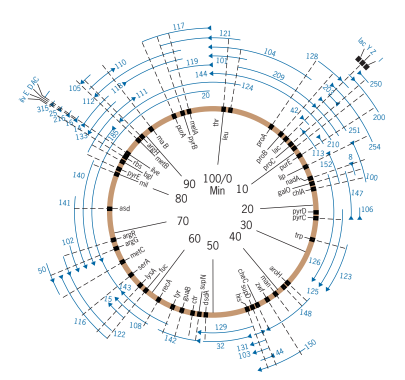


FIGURE 8.21 Formation of an F'. The anomalous excision of the F factor from an Hfr chromosome produces an F factor carrying the *E. coli thr* and *leu* genes and designated F' *thr leu*.



■ FIGURE 8.22 F's in *E. coli*. Map of the chromosome of *E. coli* K12 showing the genes present in representative F's. The F's are drawn as linear structures in order to align them with the segments of the chromosome that they contain. In reality, they are circular DNA molecules—the structures formed by joining the two ends of each F'.

reverse of the integration event shown in Figure 8.20*b*). Moreover, anomalous excision events such as the one shown in **Figure 8.21** produce autonomous F factors carrying bacterial genes. These modified F factors, called F' ("F prime") factors, were first identified by Edward Adelberg and Sarah Burns in 1959. F' factors range in size from those carrying a single bacterial gene to those carrying up to half the bacterial chromosome (**Figure 8.22**).

Transfer of F' factors to recipient (F^-) cells is called **sexduction**; it occurs by the same mechanism as F factor transfer in $F^+ \times F^-$ matings (see Figure 8.16)—with one important difference: bacterial genes incorporated into F' factors are transferred to recipient cells at a much higher frequency. The F' factors are valuable tools for genetic studies; they can be used to produce partial diploids carrying two copies of any gene or set of linked genes. Thus, sexduction can be used to determine dominance relationships between alleles and perform other genetic tests requiring two copies of a gene in the same cell.

Consider an F' thr^+leu^+ factor generated by anomalous excision of the F factor from Hfr H, as shown in Figure 8.21. Matings between F' thr^+leu^+ donor cells and thr^-leu^- recipient cells produce thr^-leu^-/F' thr^+leu^+ partial diploids. These partial diploids are unstable because the F' factor may be lost, producing thr^-leu^- haploids, or recombination may occur between the chromosome and the F', producing stable thr^+leu^+ recombinants. To examine the use of partial diploids in genetic mapping in more detail, see Solve It: How Can You Map Closely Linked Genes Using Partial Diploids?

TRANSDUCTION

Transduction—another mode of gene transfer in bacteria—was discovered by Norton Zinder and Joshua Lederberg in 1952. Zinder and Lederberg studied

auxotrophic strains of *Salmonella typhimurium* that required amino acid supplements to grow. One strain required phenylalanine, tryptophan, and tyrosine; the other required methionine and histidine. Neither strain could grow on minimal medium lacking these amino acids. However, when Zinder and Lederberg grew the strains together, rare prototrophs were produced. Moreover, when they grew the strains in medium containing DNase, but separated them in the two arms of a U-tube (see Figure 8.9), prototrophic recombinants were still produced. The insensitivity to DNase ruled out transformation as the underlying mechanism, and the fact that cell contact was not required for the appearance of the prototrophs eliminated conjugation. Subsequent experiments showed that one of the strains was infected with a virus called bacteriophage P22 and that this virus was carrying genes from one cell (the donor) to another (the recipient). The rare prototrophs that Zinder and Lederberg detected were therefore due to recombination between bacterial DNA carried by the virus and DNA in the chromosome of the recipient cell.

Later studies revealed that there are two very different types of transduction. In **generalized transduction**, a random or nearly random fragment of bacterial DNA is packaged in the phage head in place of the phage chromosome. In **specialized transduction**, a recombination event occurs between the host chromosome and the phage chromosome, producing a phage chromosome that contains a piece of bacterial DNA. Phage particles that contain bacterial DNA are called *transducing particles*. Generalized transducing particles contain only bacterial DNA. Specialized transducing particles always contain both phage and bacterial DNA.

Generalized Transduction

Generalized transducing phages can transport any bacterial gene from one cell to another—thus, the name generalized transduction. The best known generalized transducing phages are P22 in *S. typhimurium* and P1 in *E. coli*. Only about 1 to 2 percent of the phage particles produced by bacteria infected with P22 or P1 contain bacterial DNA, and only about 1 to 2 percent of the transferred DNA is incorporated into the chromosome of the recipient cell by recombination. Thus, the process is quite inefficient; the frequency of transduction for any given bacterial gene is about 1 per 10⁶ phage particles.

Specialized Transduction

Specialized transduction is characteristic of viruses that transfer only certain genes between bacteria. Bacteriophage lambda (λ) is the best-known specialized transducing phage; λ carries only the *gal* (required for the utilization of galactose as an energy source) and *bio* (essential for the synthesis of biotin) genes from one *E. coli* cell to another. Earlier in this chapter, we discussed the site-specific insertion of the λ chromosome into the *E. coli* chromosome to establish a lysogenic state (see Bacteriophage Lambda). The insertion site is between the *gal* genes and the *bio* genes on the *E. coli* chromosome (see Figure 8.5), which explains why λ only transduces these genes.

The integrated λ chromosome—the λ prophage—in a lysogenic cell undergoes rare (about one in 10⁵ cell divisions) spontaneous excision, whereupon it enters the lytic pathway. Prophage excision can also be induced, for example, by irradiating lysogenic cells with ultraviolet light. Normal excision is essentially the reverse of the site-specific integration process and yields intact circular phage and bacterial chromosomes (**■** Figure 8.23a). Occasionally, the excision is anomalous, with the crossover occurring at a site other than the original attachment site. When this happens, a portion of the bacterial chromosome is excised with the phage DNA and a portion of the phage chromosome is left in the host chromosome (**■** Figure 8.23b). These anomalous prophage excisions produce specialized transducing phage carrying either the *gal* or *bio* genes of the host. These transducing phage are denoted $\lambda dgal$ (for λ defective phage carrying *gal* genes) and λ *dbio* (λ *d* defective phage carrying *bio* genes), respectively. They are defective phage particles

Solve It!

How Can You Map Closely Linked Genes Using Partial Diploids?

Suppose that you want to determine the order of two genes (y and z) at one locus relative to a marker (x) at a nearby locus. You perform the following reciprocal crosses:

1. $x^+ y^+ z^-$ donor X $x^- y^- z^+$ recipient, and **2.** $x^- y^- z^+$ donor X $x^+ y^+ z^-$ recipient.

Note that the order of the three genes (x, y, and z) is unknown; they are arbitrarily written in alphabetic order. Assume that the mutants are all auxotrophs and that selective media can be prepared on which only prototrophic ($x^+y^+z^+$) recombinants can grow. When equal numbers of progeny are plated on selective medium, about 200 prototrophic recombinants are observed in Cross 1, whereas over 4000 are detected in Cross 2. What is the order of the three genes on the chromosome?

To see the solution to this problem, visit the Student Companion site.

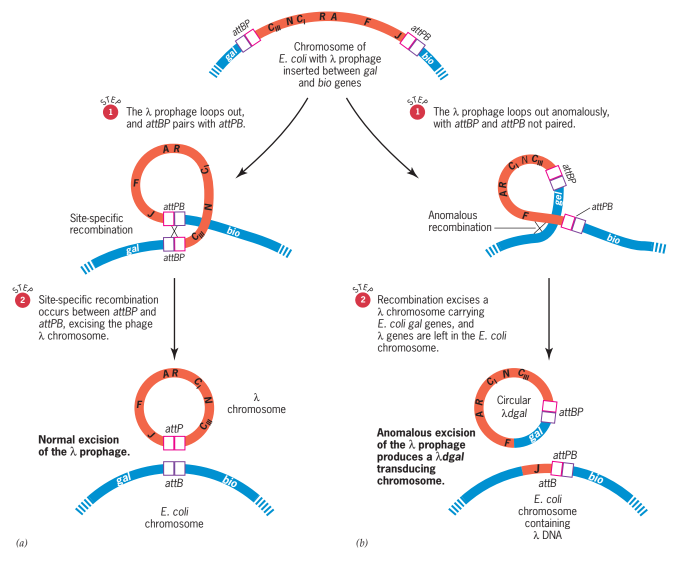


FIGURE 8.23 Lambda prophage excision. Comparison of (a) the normal excision of the λ prophage with (b) anomalous excision producing recombinant $\lambda dgal$ transducing chromosomes.

because one or more genes required for lytic or lysogenic reproduction were left in the host chromosome.

Because of the small size of the phage head, only bacterial genes located close to the prophage can be excised with the phage DNA and packaged in phage heads. Another specialized transducing phage, Φ 80, integrates near the *E. coli trp* genes (required for the synthesis of the amino acid tryptophan); this phage transduces *trp* markers. If specialized transducing particles are formed during prophage excision, as shown in Figure 8.23*b*, they should be produced only when lysogenic cells enter the lytic pathway. Indeed, transducing particles are not present in lysates produced from primary lytic infections. The frequency of transducing particles in lysates produced by induction of lysogenic cells is about one in 10⁶ progeny particles; therefore, these lysates are called *Lft* (*l*ow*f*requency *t*ransduction) lysates.

The fate of the $\lambda dgal$ and $\lambda dbio$ DNA molecules after their injection into new host cells will depend on which λ genes are missing. If genes for lytic growth are missing, but an *att* site and *int* (integrase) gene are present, the defective chromosomes will be able to integrate into the host chromosome. However, they will not be able to reproduce lytically unless a wild-type λ , acting as a "helper" phage, is present. If the *int* gene is missing, the defective phage chromosome will be able to integrate only in the

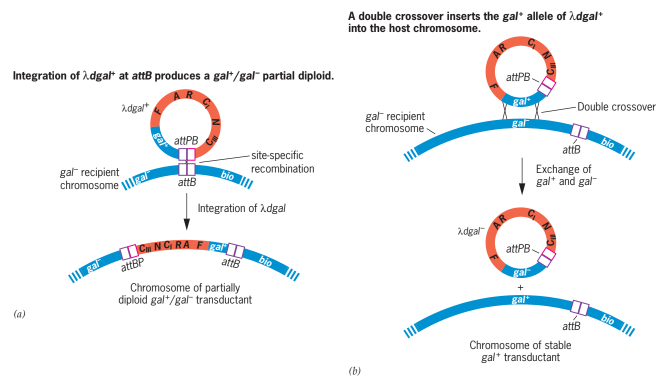


FIGURE 8.24 Recombination in *gal*⁻ recipient cells infected with $\lambda dgal^+$ transducing phage. (a) Integration of $\lambda dgal^+$ at *attB* produces an unstable *gal*⁺/*gal*⁻ partial diploid. (b) A double crossover transfers the *gal*⁺ allele from $\lambda dgal^+$ to the chromosome.

presence of a wild-type helper. If a $\lambda dgal^+$ phage infects a gal^- recipient cell, integration of the $\lambda dgal^+$ will produce an unstable gal^+/gal^- partial diploid (**■** Figure 8.24*a*), whereas rare recombination events between gal^+ in the transducing DNA and gal^- in the recipient chromosome will produce stable gal^+ transductants (**■** Figure 8.24*b*).

If the ratio of phage to bacteria is high, recipient cells will be infected with both wild-type λ *phage* and $\lambda dgal^+$; thus, these cells will be double lysogens carrying one wild-type λ prophage and one $\lambda dgal$ prophage. The resulting transductants will be gal^+/gal^- partial diploids. If the gal^+/gal^- transductants are induced with ultraviolet light, the lysates will contain about 50 percent $\lambda dgal$ particles and 50 percent λ^+ particles. Both prophages will replicate with equal efficiency using the gene products encoded by the λ^+ genome. Such lysates are called *Hft* (*bigh-frequency transduction events*; therefore, *Hft* lysates are used preferentially in transduction experiments.

 Three parasexual processes—transformation, conjugation, and transduction—occur in bacteria. These processes can be distinguished by two criteria: whether the gene transfer is inhibited by deoxyribonuclease and whether it requires cell contact.

• Transformation involves the uptake of free DNA by bacteria.

- Conjugation occurs when a donor cell makes contact with a recipient cell and then transfers DNA to the recipient cell.
- Transduction occurs when a virus carries bacterial genes from a donor cell to a recipient cell.
- Plasmids are self-replicating extrachromosomal genetic elements.
- Episomes can replicate autonomously or as integrated components of bacterial chromosomes.
- F factors that contain chromosomal genes (F' factors) are transferred to F^- cells by sexduction.

KEY POINTS

The Evolutionary Significance of Genetic Exchange in Bacteria

Genetic exchange is as important in bacteria as it is in other organisms.

Solve It!

How Do Bacterial Genomes Evolve?

A mating (Cross I) between F⁺ met⁺ ser⁺ cys⁺ str^s and F⁻ met⁻ ser⁻ cys⁻ str^r strains of E. coli resulted in all of the bacteria becoming F⁺, but produced no *met⁺ ser⁺ cys⁺* str^r prototrophic recombinants. After several generations of growth, new cultures of each strain were grown from single colonies, and the cross was repeated. This time (Cross II), met⁺ ser⁺ cys⁺ str^r recombinants were produced, but all of these recombinants were F⁻. After several additional generations of growth of the strains used in Cross II, new cultures were grown from isolated colonies, and the mating was repeated a third time (Cross III). No met⁺ ser⁺ cys⁺ str^r recombinants were produced in Cross III; instead, all the progeny that survived on medium containing streptomycin had the genotype met⁺ ser⁺ cys⁻ str^r and were phenotypically F⁺. Using a map of the E. coli chromosome, explain these results.

To see the solution to this problem, visit the Student Companion site. Mutation is the source of new genetic variation, and recombination then produces new combinations of this variation—new assortments of genes responsible for the phenotypes acted on by natural selection. In eukaryotes, recombination events—

independent assortment and crossing over—are an integral part of sexual reproduction. However, bacteria do not reproduce sexually. Nevertheless, recombination is certainly important in the evolution of bacteria, just as it is in eukaryotes. Thus, it is not surprising that mechanisms for exchanging genetic information and producing recombinant combinations of genes have evolved in bacteria. These parasexual mechanisms—transformation, conjugation, and transduction—generate new combinations of genes and allow bacteria to evolve and adapt to new environmental niches and to sudden changes in existing habitats. To test your understanding of the importance of recombination in bacteria, try Solve It: How Do Bacterial Genomes Evolve?

Although parasexual processes are beneficial to bacteria, they create serious problems for humans who are attempting to combat bacterial diseases. These parasexual processes are partially responsible for the rapid evolution and spread of plasmids that confer antibiotic and drug resistance to bacteria. The extensive use of antibiotics in agriculture and medicine has resulted in the evolution of plasmids that carry a whole battery of antibiotic-resistance genes (see On the Cutting Edge: Antibiotic-Resistant Bacteria). Indeed, the resurgence of tuberculosis in New York City in the 1990s has in large part been due to the evolution of a strain of *M. tuberculosis* that is resistant to seven different antibiotics. Moreover, the New York strain is closely related to the predominant strains of antibiotic-resistant *M. tuberculosis* in China and other regions of the world. Thus, what is beneficial to pathogenic bacteria is often harmful to humans.

- Parasexual recombination mechanisms produce new combinations of genes in bacteria.
 - Parasexual mechanisms enhance the ability of bacteria to adapt to changes in the environment.

ON THE CUTTING EDGE

ANTIBIOTIC-RESISTANT BACTERIA

n March 2010, the World Health Organization reported that multiple drug-resistant (MDR) strains of Mycobacterium tuberculosis, the bacterium that causes tuberculosis (TB), have increased to record levels. In some countries, up to one-fourth of the individuals with TB cannot be treated effectively with our best antibiotics.

KEY POINTS

Now, a new gene designated NDM-1, for New Delhi metallobeta-lactamase, has evolved that makes bacteria resistant to a major group of antibiotics, the carbapenems, that are often used to treat MDR strains of bacteria. The NDM-1 gene is located on a plasmid that is easily transferred from one bacterium to another. To date, the gene has been found in *E. coli* and *Klebsiella pneumonia*—both causing severe urinary tract infections—but there is little doubt that it will spread to other bacterial species. Unfortunately, only two antibiotics are being developed with the potential to be effective in treating NDM-1 superbugs.

What has led to the evolution of these antibiotic- and drugresistant bacteria? How have humans contributed to this potential crisis? Can we resolve this problem? If so, how? Let's start by considering the history of antibiotics and antibiotic-resistant bacteria.

Selman Waksman, a Ukrainian immigrant to the United States, discovered streptomycin in 1943. Later he named this class of antibacterial drugs *antibiotics*. The first documented treatment of a human with streptomycin involved a 21-year-old patient at the Mayo



Clinic in Rochester, Minnesota. This woman had an advanced case of tuberculosis. She began receiving experimental injections of streptomycin in 1944, and to everyone's surprise her tuberculosis was cured. Streptomycin and other antibiotics quickly became "miracle drugs." Their application to people with bacterial infections saved millions of lives.

Humans soon began using large amounts of antibiotics. In 1950, the world used 10 tons of streptomycin. By 1955, worldwide use of streptomycin had increased to 50 tons, along with about 10 tons each of chloramphenicol and tetracycline.

However, the bacteria soon began to fight back. They evolved new genes encoding products that protected them from antibiotics. The evolution of antibiotic-resistant bacteria confirmed the power of natural selection. A bacterium without an antibiotic-resistance gene is killed by the antibiotic. A bacterium with an antibiotic-resistance gene grows, divides, and produces a population of bacteria, all resistant to the antibiotic. The result was inevitable: antibiotic-resistant bacteria spread.

Some of the first studies documenting the evolution of antibiotic- and drug-resistant bacteria were performed in Japan on the four "species" of *Shigella*—*S. dysenteriae*, *S. flexneri*, *S. boydii*, and *S. sonnei*, which cause dysentery. Only 0.2 percent of the *Shigella* strains isolated from sewers and polluted rivers in 1953 were resistant to any of the antibiotics and drugs tested. Just 12 years later, the frequency of antibiotic- and drug-resistant *Shigella* strains isolated from the same places had increased to 58 percent. However, the really bad news was not that these strains were resistant to antibiotics; rather, it was that most of them were resistant to at least four of the six antibiotics and drugs—ampicillin, kanamycin, tetracycline, streptomycin, sulfanilamide, and chloramphenicol—tested. They were multi-drug-resistant strains of *Shigella*. MDR strains of other bacteria, such as *M. tuberculosis*, also began appearing.

The genes that protect bacteria from antibiotics often are present on small DNA molecules called R-plasmids (see

Plasmids and Episomes). Many R-plasmids are self-transmissible; that is, they carry genes that mediate their own transfer from one cell to another and even from one species to another. In addition, the antibiotic-resistance genes are often present on genetic elements—transposable genetic elements or "jumping genes"—that can move from one DNA molecule to another (see Chapter 17). Thus, the genes on these R-plasmids can spread rapidly through bacterial populations.

One reason that MDR strains of bacteria have evolved so rapidly is that we overuse antibiotics. All too often, antibiotics are prescribed for viral infections such as the common cold and flu. Antibiotics have no antiviral activity and should not be used to treat viral infections. In addition, antibiotics are widely used in vast amounts as "growth promoters" in animal feed, where they prevent bacterial infections that reduce growth rate. Indeed, almost half of the antibiotics produced in the United States are used as additives in animal feed. They are added at rates of 2 to 50 grams per ton of feed, and the inevitable happens antibiotic-resistant bacteria evolve. These resistant bacteria are then transmitted to humans caring for the animals, working in the meatpackaging industry, or consuming undercooked meat products.

Given the widespread evolution of MDR strains of *M. tuber-culosis*, *Staphylococcus aureus*, *Shigella dysenteriae*, and other pathogenic bacteria, perhaps we should restrict the use of some of our best antibiotics to the treatment of potentially fatal human diseases. Indeed, Denmark banned the use of penicillins and tetracyclines as animal growth promoters in the 1970s, and Sweden banned all nontherapeutic use of antibiotics, including as animal growth promoters, in 1986. The negative effects of banning the use of antibiotics in animal feed on productivity have been minimal, and in Sweden, the overall use of antibiotics has decreased by 55 percent since the ban on nontherapeutic uses began. Perhaps it is time for the United States and the rest of the world to follow Scandinavia's lead—to ban, or at least limit, the nontherapeutic use of antibiotics. Do we really need antibiotics in animal feed? And do we need them in our hand soap?

Basic Exercises

Illustrate Basic Genetic Analysis

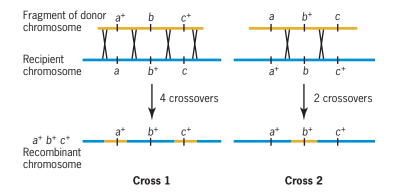
- 1. What advantages for genetic research do viruses have over cellular and multicellular organisms?
- **Answer:** The two major advantages for genetic studies that viruses have over cellular and multicellular organisms are (1) their structural simplicity and (2) their short life cycle. Viruses usually contain a single chromosome with a relatively small number of genes, and they can complete their life cycle in from about 20 minutes to a few hours.
- 2. What are the major differences between crossing over in bacteria and in eukaryotes?
- **Answer:** In bacteria, crossing over usually occurs between a fragment of the chromosome from a donor cell and an intact circular chromosome in a recipient cell (see

Figure 8.7*a*). As a result, crossovers must occur in pairs that insert segments of the donor cell's chromosome into the recipient cell's chromosome. Single crossovers, or any odd number of crossovers, will destroy the integrity of the circular chromosome and yield a linear DNA molecule in its place (see Figure 8.7*b*).

3. When grown together, two strains of *E. coli*, $a b^+$ and a^+b , are known to exchange genetic material, leading to the production of a^+b^+ recombinants. However, when these two strains are grown in opposite arms of a U-tube (see Figure 8.9), no a^+b^+ recombinants are produced. What parasexual process is responsible for the formation of the a^+b^+ recombinants when these strains are grown together?

- **Answer:** The two *E. coli* strains are exchanging information by conjugation, the only parasexual process in bacteria that requires cell contact. The glass filter separating the arms of the U-tube prevents contact between cells in these arms.
- 4. You have identified three closely linked genetic markers—*a*, *b*, and *c*—in *E. coli*. The markers are transferred from an Hfr strain to an F⁻ strain in less than 1 minute, and they are present on the chromosome in the order *a*—*b*—*c*. You perform phage P1 transduction experiments using strains of genotype $a^+ b c^+$ and $a b^+ c$. In cross 1, the donor cells are $a^+ b c^+$ and the recipient cells are $a^+ b c^+$. For both crosses, you prepare minimal medium plates on which only $a^+ b^+ c^+$ recombinants can form colonies. In which cross would you expect to observe the most $a^+ b^+ c^+$ recombinants?
- **Answer:** You would expect more $a^+ b^+ c^+$ recombinants in cross 2 because the formation of a chromosome carrying all

three wild-type markers requires only two crossovers (one pair of crossovers) in that cross, whereas four crossovers (two pairs) are required to produce an $a^+ b^+ c^+$ chromosome in cross 1. The required crossovers are shown in the following diagram.



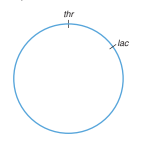
Testing Your Knowledge

Integrate Different Concepts and Techniques

1. You have identified a mutant *E. coli* strain that cannot synthesize histidine (His⁻). To determine the location of the *his*⁻ mutation on the *E. coli* chromosome, you perform interrupted mating experiments with five different Hfr strains. The following chart shows the time of entry (minutes, in parentheses) of the wild-type alleles of the first five markers (mutant genes) into the His⁻ strain.

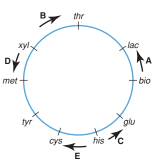
Hfr A —— <i>bio</i> (4)	glu (20)	<i>his</i> (27)	cys (37)	tyr (45)
Hfr B — xyl (6)	<i>met</i> (18)	tyr (24)	cys (32)	<i>bis</i> (42)
Hfr C —— <i>bis</i> (3)	cys (13)	tyr (21)	<i>met</i> (27)	xyl (39)
Hfr D — xyl (7)	thr (25)	<i>lac</i> (40)	<i>bio</i> (48)	<i>glu</i> (62)
Hfr E —— <i>bis</i> (4)	<i>glu</i> (11)	<i>bio</i> (27)	<i>lac</i> (35)	thr (50)

(a) On the following map of the circular *E. coli* chromosome, indicate (1) the relative location of each gene, (2) the position where the F factor is integrated in each of the five Hfr's, and (3) the direction of chromosome transfer for each Hfr (indicate direction with an arrow).



(b) To further define the location of the *his*⁻ mutation on the chromosome, you use the mutant strain as a recipient in a bacteriophage P1 transduction experiment. Which, if any, of the genes shown in the chart above would you expect to be cotransduced with the his^+ allele of your his^- mutant gene, given that phage P1 can package about 1 percent of the *E. coli* chromosomal DNA molecule? Note that the *E. coli* chromosome contains 4.6 million nucleotide pairs and that transfer of the entire chromosome during conjugation takes 100 minutes. Explain your answer.

Answer: (a) The gene order is as shown on the following map, and the sites of F factor integration and direction of transfer for each of the Hfr's are indicated by the arrowheads labeled A through E.



(b) None of the markers would be cotransduced with his^+ because phage P1 can package only 1 percent of the *E. coli* chromosome, and none of the other genes is within 1 min of *his*.

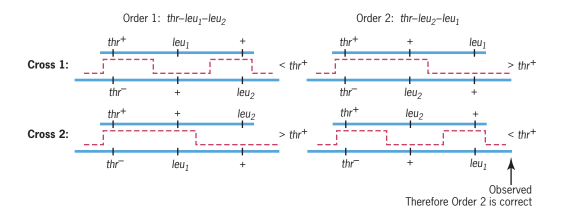
2. Reciprocal three-point transduction crosses were used to determine the order of two mutations, *leu*₁ and *leu*₂, in the *leuA* gene relative to the linked *thrA* gene of *E. coli*. In each cross, *leu*⁺ recombinants were selected on minimal medium

containing threenine but no leucine, and then tested for thr^+ or thr^- by replica plating onto plates containing no threenine. The results are given in the table below

C	ross		
Donor Markers	Recipient Markers	<i>tbr</i> Allele in <i>leu</i> ⁺ Recombinants	Percent <i>thr</i> ⁺
1. $thr^+ leu_1$	thr ⁻ leu ₂	350 thr ⁺ : 349 thr ⁻	50
2. $thr^+ leu_2$	$thr^{-}leu_{1}$	60 thr+: 300 thr-	17

What is the order of leu_1 and leu_2 relative to the outside marker *thr*?

Answer: The two crosses are diagrammed here showing the two possible orders, with the dashed red lines marking the portions of the two chromosomes that must be present in tbr^+ - leu_1^+ - leu_2^+ (+++) recombinants. Note that if order 1 is correct, the formation of +++ recombinants will require 4 crossovers (2 pairs of crossovers) in cross 1 and only 2 crossovers (1 pair) in cross 2, therefore predicting more +++ recombinants in cross 2 and fewer in cross 1. However, if order 2 is correct, there should be more +++ recombinants in cross 1 and fewer in cross 2. Since the second result was observed, the correct order is tbr- leu_2 - leu_1 .



Questions and Problems

Enhance Understanding and Develop Analytical Skills

- 8.1 By what criteria are viruses living? nonliving?
- 8.2 How do bacteriophages differ from other viruses?
- 8.3 In what ways do the life cycles of bacteriophages T4 and λ differ? In what respect are they the same?
- 8.4 How does the structure of the λ prophage differ from the structure of the λ chromosome packaged in the λ virion?
- **8.5** In what way does the integration of the λ chromosome into the host chromosome during a lysogenic infection differ from crossing over between homologous chromosomes?
- **8.6** Geneticists have used mutations that cause altered phenotypes such as white eyes in *Drosophila*, white flowers and wrinkled seeds in peas, and altered coat color in rabbits to determine the locations of genes on the chromosomes of these eukaryotes. What kinds of mutant phenotypes have been used to map genes in bacteria?
- **8.7** You have identified three mutations—*a*, *b*, and *c*—in *Streptococcus pneumoniae*. All three are recessive to their wild-type alleles a^+ , b^+ , and c^+ . You prepare DNA from a wild-type donor strain and use it to transform a strain with genotype *a b c*. You observe a^+b^+ transformants and a^+c^+ transformants, but no b^+c^+ transformants. Are these mutations closely linked? If so, what is their order on the *Streptococcus* chromosome?

- **8.8** A nutritionally defective *E. coli* strain grows only on a medium containing thymine, whereas another nutritionally defective strain grows only on medium containing leucine. When these two strains were grown together, a few progeny were able to grow on a minimal medium containing neither thymine or leucine. How can this result be explained?
- **8.9** Assume that you have just demonstrated genetic recombination (e.g., when a strain of genotype $a b^+$ is present with a strain of genotype $a^+ b$, some recombinant genotypes, $a^+ b^+$ and a b, are formed) in a previously unstudied species of bacteria. How would you determine whether the observed recombination resulted from transformation, conjugation, or transduction?
- 8.10 (a) What are the genotypic differences between F⁻ cells, F⁺ cells, and Hfr cells? (b) What are the phenotypic differences? (c) By what mechanism are F⁻ cells converted to F⁺ cells? F⁺ cells to Hfr cells? Hfr cells to F⁺ cells?
- **8.11** (a) Of what use are F' factors in genetic analysis? (b) How are F' factors formed? (c) By what mechanism does sexduction occur?
- **8.12** What are the basic differences between generalized transduction and specialized transduction?

- **8.13** What roles do IS elements play in the integration of F factors?
- **8.14** How can bacterial genes be mapped by interrupted mating experiments?
- **8.15** What does the term *cotransduction* mean? How can co-transduction frequencies be used to map genetic markers?
- 8.16 **•••** In *E. coli*, the ability to utilize lactose as a carbon source requires the presence of the enzymes β -galactosidase and β -galactoside permease. These enzymes are encoded by two closely linked genes, *lacZ* and *lacY*, respectively. Another gene, *proC*, controls, in part, the ability of *E. coli* cells to synthesize the amino acid proline. The alleles *str^r* and *str^s*, respectively, control resistance and sensitivity to streptomycin. Hfr H is known to transfer the two *lac* genes, *proC*, and *str*, in that order, during conjugation.

A cross was made between Hfr H of genotype $lacZ^$ $lacY^+$ proC⁺ str^s and an F⁻ strain of genotype $lacZ^+$ lacY⁻ proC⁻ str^s. After about 2 hours, the mixture was diluted and plated out on medium containing streptomycin but no proline. When the resulting proC⁺ str^r recombinant colonies were checked for their ability to grow on medium containing lactose as the sole carbon source, very few of them were capable of fermenting lactose. When the reciprocal cross (Hfr H $lacZ^+$ $lacY^-$ proC⁺ str^s X F⁻ $lacZ^ lacY^+$ proC⁻ str^s) was done, many of the proC⁺ str^r recombinants were able to grow on medium containing lactose as the sole carbon source. What is the order of the *lacZ* and *lacY* genes relative to proC?

8.17 An F⁺ strain, marked at 10 loci, gives rise spontaneously to Hfr progeny whenever the F factor becomes incorporated into the chromosome of the F⁺ strain. The F factor can integrate into the circular chromosome at many points, so that the resulting Hfr strains transfer the genetic markers in different orders. For any Hfr strain, the order of markers entering a recipient cell can be determined by interrupted mating experiments. From the following data for several Hfr strains derived from the same F⁺ strain, determine the order of markers in the F⁺ strain.

Hfr Strain	Markers Donated in Order
1	$-$ Z-H-E-R \rightarrow
2	$-$ O-K-S-R \rightarrow
3	$-$ K-O-W-I \rightarrow
4	$-$ Z-T-I-W \rightarrow
5	$-\operatorname{H-Z-T-I} \to$

8.18 **•••** The data in the following table were obtained from three-point transduction tests made to determine the order of mutant sites in the *A* gene encoding the α subunit of tryptophan synthetase in *E. coli. Anth* is a linked, unselected marker. In each cross, trp^+ recombi-

nants were selected and then scored for the *anth* marker $(anth^+ \text{ or } anth^-)$. What is the linear order of *anth* and the three mutant alleles of the *A* gene indicated by the data in the table?

Cross	Donor Markers	Recipient Markers	anth Allele in trp ⁺ Recombinants	% antb+
1	$antb^+ - A34$	anth ⁻ — A223	72 anth ⁺ : 332 anth ⁻	18
2	$antb^+ - A46$	anth ⁻ — A223	196 anth ⁺ : 180 anth ⁻	52
3	$antb^+ - A223$	anth ⁻ — A34	380 anth ⁺ : 379 anth ⁻	50
4	anth ⁺ — A223	anth ⁻ — A46	60 anth ⁺ : 280 anth ⁻	20

8.19 Bacteriophage P1 mediates generalized transduction in E. coli. A P1 transducing lysate was prepared by growing P1 phage on pur⁺ pro⁻ his⁻ bacteria. Genes pur, pro, and *his* encode enzymes required for the synthesis of purines, proline, and histidine, respectively. The phage and transducing particles in this lysate were then allowed to infect *pur⁻ pro⁺ bis⁺* cells. After incubating the infected bacteria for a period of time sufficient to allow transduction to occur, they were plated on minimal medium supplemented with proline and histidine, but no purines to select for *pur*⁺ transductants. *The pur*⁺ colonies were then transferred to minimal medium with and without proline and with and without histidine to determine the frequencies of each of the outside markers. Given the following results, what is the order of the three genes on the E. coli chromosome?

Genotype	Number Observed
pro+ his+	100
pro ⁻ his ⁺	22
pro+ his-	150
pro- his-	1

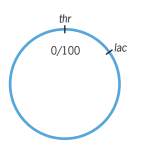
8.20 Two additional mutations in the *trp A* gene of *E. coli*, *trp A58* and *trp A487*, were ordered relative to *trp A223* and the outside marker *anth* by three-factor transduction crosses as described in Problem 8.18. The results of these crosses are summarized in the following table. What is the linear order of *anth* and the three mutant sites in the *trp A* gene?

Cross	Donor Markers	Recipient Markers	anth Allele in trp ⁺ Recombinants	% antb ⁻
1	$antb^+$ — A487	anth ⁻ — A223	72 anth ⁺ : 332 anth ⁻	82
2	anth ⁺ — A58	anth ⁻ — A223	196 anth+: 180 anth-	48
3	anth+— A223	antb ⁻ — A487	380 anth+: 379 anth-	50
4	anth+— A223	anth ⁻ — A58	60 anth ⁺ : 280 anth ⁻	80

8.21 You have identified a mutant *E. coli* strain that cannot synthesize histidine (His⁻). To determine the location of the *bis*⁻ mutation on the *E. coli* chromosome, you perform interrupted mating experiments with five different Hfr strains. The following chart shows the time of entry (minutes, in parentheses) of the wild-type alleles of the first five markers (mutant genes) into the His⁻ strain.

Hfr A ——— <i>bis</i> (1)	<i>man</i> (9)	gal (28)	<i>lac</i> (37)	thr (45)
Hfr B — <i>man</i> (15)	<i>bis</i> (23)	cys (38)	ser (42)	arg (49)
Hfr C — <i>thr</i> (3)	<i>lac</i> (11)	gal (20)	man (39)	<i>bis</i> (47)
Hfr D ——— <i>cys</i> (3)	<i>bis</i> (18)	man (26)	gal (45)	<i>lac</i> (54)
Hfr E <i>thr</i> (4)	<i>rha</i> (18)	arg (36)	ser (43)	cys (47)

On the following map of the circular *E. coli* chromosome, indicate (1) the relative location of each gene relative to *thr* (located at 0/100 Min), (2) the position where the F factor is integrated in each of the five Hfr's, and (3) the direction of chromosome transfer for each Hfr (indicate direction with an arrow or arrowhead).



8.22 Mutations *nrd* 11 (gene *nrd B*, encoding the beta subunit of the enzyme ribonucleotide reductase), *am M*69

(gene 63, encoding a protein that aids tail-fiber attachment), and *nd* 28 (*denA*, encoding the enzyme endonuclease II) are known to be located between gene 31 and gene 32 on the bacteriophage T4 chromosome. Mutations *am* N54 and *am* A453 are located in genes 31 and 32, respectively. Given the three-factor cross data in the following table, what is the linear order of the five mutant sites?

Three-Factor Cross Data

Cross	% Recombination ^a
1. am A453—am M69 × nrd 11	2.6
2. am A453—nrd 11 × am M69	4.2
3. <i>am A</i> 453— <i>am M</i> 69 × <i>nd</i> 28	2.5
4. <i>am A</i> 453— <i>nd</i> 28 × <i>am M</i> 69	3.5
5. <i>am A</i> 453— <i>nrd</i> 11 × <i>nd</i> 28	2.9
6. <i>am A</i> 453— <i>nd</i> 28 × <i>nrd</i> 11	2.1
7. am N54—am M69 × nrd 11	3.5
8. <i>am N</i> 54— <i>nrd</i> 11 × <i>am M</i> 69	1.9
9. am N54—nd 28 × am M69	1.7
10. <i>am N</i> 54— <i>am M</i> 69 × <i>nd</i> 28	2.7
11. <i>am</i> N54— <i>nd</i> 28 × <i>nrd</i> 11	2.9
12. am N54—nrd 11 × nd 28	1.9

^aAll recombination frequencies are given as $\frac{2 \text{ (wild-type progeny)}}{\text{total progeny}} \times 100.$

Genomics on the Web at http://www.ncbi.nlm.nih.gov

The *E. coli* genome was one of the first bacterial genomes sequenced. The complete nucleotide sequence (4.6 million nucleotide pairs) of the genome of *E. coli* strain K12 was published in September 1997.

- 1. How many different strains of *E. coli* have had their genomes sequenced since 1997?
- 2. Are these genomes all about the same size? If not, how much variation in size is observed between the genomes of different *E. coli* strains?
- **3.** Some *E. coli* strains, for example, 0157:H7, are more pathogenic to humans and other mammals than strains such as K12. Do these strains have larger or smaller genomes than K12? Might comparisons of the genes in the pathogenic and non-pathogenic strains provide hints as to why some strains are pathogenic and others are not?

Hint: At the NCBI web site, Genome Biology \rightarrow Entrez Genome \rightarrow Microbial Genomes \rightarrow Complete Genomes \rightarrow *Escherichia coli*.



DNA and the Molecular Structure of Chromosomes

CHAPTER OUTLINE

- Functions of the Genetic Material
- Proof That Genetic Information Is Stored in DNA
- The Structures of DNA and RNA
- Chromosome Structure in Prokaryotes and Viruses
- Chromosome Structure in Eukaryotes

Discovery of Nuclein

In 1868, Johann Friedrich Miescher, a young Swiss medical student, became fascinated with an acidic substance that he isolated from pus cells obtained from bandages used to dress human wounds. He first separated the pus cells from the bandages and associated debris, and then treated the cells with pepsin, a proteolytic enzyme that he isolated from the stomachs of pigs. After the pepsin treatment, he recovered an acidic substance that he called "nuclein." Miescher's nuclein was unusual in that it contained large amounts of both nitrogen and phosphorus, two elements known at the time to coexist only in certain types of fat. Miescher wrote a paper describing his discovery of nuclein in human pus cells and submitted it for publication in 1869. However, the editor of the journal to which the paper was sent was skeptical of the results and decided to repeat the experiments himself. As a result, Miescher's paper describing nuclein was not published until 1871, two years after its submission.

At the time, the importance of the substance that Miescher called nuclein could not have been anticipated. The existence of polynucleotide chains, the key component of the acidic material in Miescher's nuclein, was not documented until the 1940s. The role of nucleic acids in storing and transmitting genetic information was not established until 1944, and the double-helix structure of DNA was not discovered until 1953. Even in 1953, many geneticists were



Color-enhanced transmission electron micrograph of a ruptured *E. coli* cell with much of its DNA extruded.

reluctant to accept the idea that nucleic acids, rather than proteins, carried the genetic information because nucleic acids exhibited less structural variability than proteins.

Functions of the Genetic Material

In 1865, Mendel showed that "Merkmalen" (now "genes") transmitted genetic information, and in the first part of the twentieth century, their patterns of transmission were studied extensively. Although these classical genetic studies provided little insight into the

The genetic material must replicate, control the growth and development of the organism, and allow the organism to adapt to changes in the environment.

molecular nature of genes, they did demonstrate that the genetic material must perform three essential functions:

- 1. The genotypic function, **replication**. The genetic material must store genetic information and accurately transmit that information from parents to offspring, generation after generation.
- **2.** The phenotypic function, **gene expression**. The genetic material must control the development of the phenotype of the organism. That is, the genetic material must dictate the growth of the organism from the single-celled zygote to the mature adult.
- **3.** The evolutionary function, **mutation**. The genetic material must undergo changes to produce variations that allow organisms to adapt to modifications in the environment so that evolution can occur.

Other early genetic studies established a precise correlation between the patterns of transmission of genes and the behavior of chromosomes during sexual reproduction, providing strong evidence that genes are usually located on chromosomes. Thus, further attempts to discover the chemical basis of heredity focused on the molecules present in chromosomes.

Chromosomes are composed of two types of large organic molecules (macromolecules) called **proteins** and **nucleic acids**. The nucleic acids are of two types: **deoxyribonucleic acid (DNA)** and **ribonucleic acid (RNA)**. During the 1940s and early 1950s, the results of elegant experiments clearly established that the genetic information is stored in nucleic acids, not in proteins. In most organisms, the genetic information is encoded in the structure of DNA. However, in many small viruses, the genetic information is encoded in RNA.

• The genetic material must perform three essential functions: the genotypic function replication, the phenotypic function—gene expression, and the evolutionary function—mutation. KEY POINT

Proof That Genetic Information Is Stored in DNA

Several lines of indirect evidence suggested that DNA harbors the genetic information of living organisms. For example, most of the DNA of cells is located in the chromosomes, whereas RNA and proteins are also

abundant in the cytoplasm. Also, a precise correlation exists between the amount of DNA per cell and the number of sets of chromosomes per cell. Most somatic cells of diploid organisms contain twice the amount of DNA as the haploid germ cells (gametes) of the same species. The molecular composition of the DNA is the same (with rare exceptions) in all the cells of an organism, whereas the composition of both RNA and proteins is highly variable from one cell type to another. DNA is more stable than RNA or proteins. Since the genetic material must store and transmit information from parents to offspring, we might expect it to be stable, like DNA. Although these correlations strongly suggest that DNA is the genetic material, they by no means prove it.

In most organisms, the genetic information is encoded in DNA. In some viruses, RNA is the genetic material.

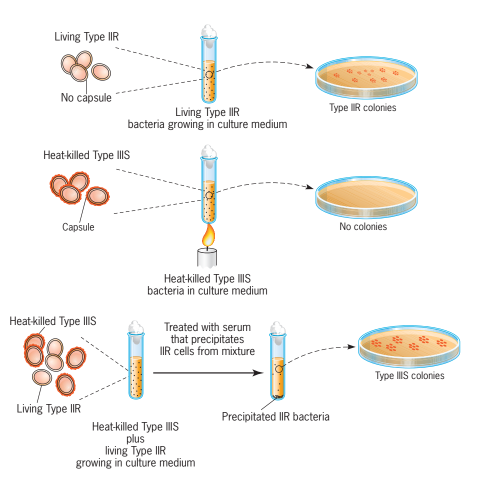


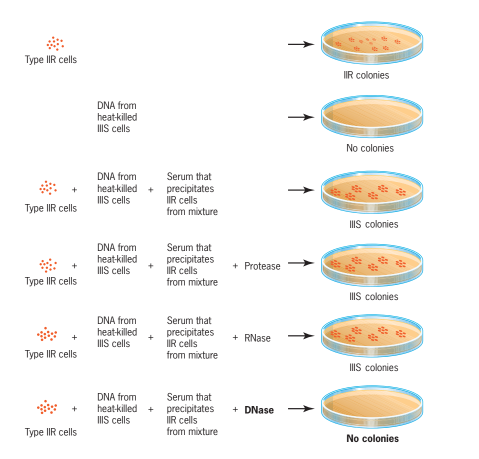
FIGURE 9.1 Sia and Dawson's demonstration of transformation in *Streptococcus pneumoniae in vitro*.

PROOF THAT DNA MEDIATES TRANSFORMATION

Frederick Griffith's discovery of transformation in *Streptococcus pneumoniae* was discussed in Chapter 8. When Griffith injected both heat-killed Type IIIS bacteria (virulent when alive) and live Type IIR bacteria (avirulent) into mice, many of the mice developed pneumonia and died, and live Type IIIS cells were recovered from their carcasses. Something from the heat-killed cells—the "transforming principle"—had converted the live Type IIR cells to Type IIIS. In 1931, Richard Sia and Martin Dawson performed the same experiment *in vitro*, showing that the mice played no role in the transformation process (**■** Figure 9.1). Sia and Dawson's experiment set the stage for Oswald Avery, Colin MacLeod, and Maclyn McCarty's demonstration that the "transforming principle" in *S. pneumoniae* is DNA. Avery and colleagues showed that DNA is the only component of the Type IIIS cells required to transform Type IIR cells to Type IIIS (**■** Figure 9.2).

But how could they be sure that the DNA was really pure? Proving the purity of any macromolecular substance is extremely difficult. Maybe the DNA preparation contained a few molecules of protein, and these contaminating proteins were responsible for the observed transformation. The most definitive experiments in Avery, MacLeod, and McCarty's proof that DNA was the transforming principle involved the use of enzymes that degrade DNA, RNA, or protein. In separate experiments, highly purified DNA from Type IIIS cells was treated with the enzymes (1) **deoxyribonuclease (DNase)**, which degrades DNA, (2) **ribonuclease (RNase)**, which degrades RNA, or (3) **proteases**, which degrade proteins; the DNA was then tested for its ability to transform Type IIR cells to Type IIIS. Only DNase treatment had any effect on the transforming activity of the DNA preparation—it eliminated all transforming activity (Figure 9.2).

Although the molecular mechanism by which transformation occurs remained unknown for many years, the results of Avery and coworkers clearly established that the genetic information in *Streptococcus* is present in DNA. Geneticists now know that the segment of DNA in the chromosome of *Streptococcus* that carries the genetic



■ FIGURE 9.2 Avery, MacLeod, and McCarty's proof that the "transforming principle" is DNA.

information specifying the synthesis of a Type III capsule is physically inserted into the chromosome of the Type IIR recipient cell during the transformation process.

PROOF THAT DNA CARRIES THE GENETIC INFORMATION IN BACTERIOPHAGE T2

Additional evidence demonstrating that DNA is the genetic material was published in 1952 by Alfred Hershey (1969 Nobel Prize winner) and Martha Chase. The results of their experiments showed that the genetic information of a particular bacterial virus (bacteriophage T2) was present in its DNA. Their results had a major impact on scientists' acceptance of DNA as the genetic material. This impact was the result of the simplicity of the Hershey–Chase experiment.

Viruses are the smallest living organisms; they are living, at least in the sense that their reproduction is controlled by genetic information stored in nucleic acids via the same processes as in cellular organisms (Chapter 8). However, viruses are acellular parasites that can reproduce only in appropriate host cells. Their reproduction is totally dependent on the metabolic machinery (ribosomes, energy-generating systems, and other components) of the host. Viruses have been extremely useful in the study of many genetic processes because of their simple structure and chemical composition (many contain only proteins and nucleic acids) and their very rapid reproduction (15 to 20 minutes for some bacterial viruses under optimal conditions).

Bacteriophage T2, which infects the common colon bacillus *Escherichia coli*, is composed of about 50 percent DNA and about 50 percent protein (**Figure 9.3**). Experiments prior to 1952 had shown that all bacteriophage T2 reproduction takes place within *E. coli* cells. Therefore, when Hershey and Chase showed that the DNA of the virus particle entered the cell, whereas most of the protein of the virus remained adsorbed to the outside of the cell, the implication was that the genetic information necessary for viral reproduction was present in DNA. The basis for the Hershey–Chase experiment is that DNA contains phosphorus but no sulfur, whereas proteins contain sulfur but virtually no phosphorus. Thus, Hershey and Chase were

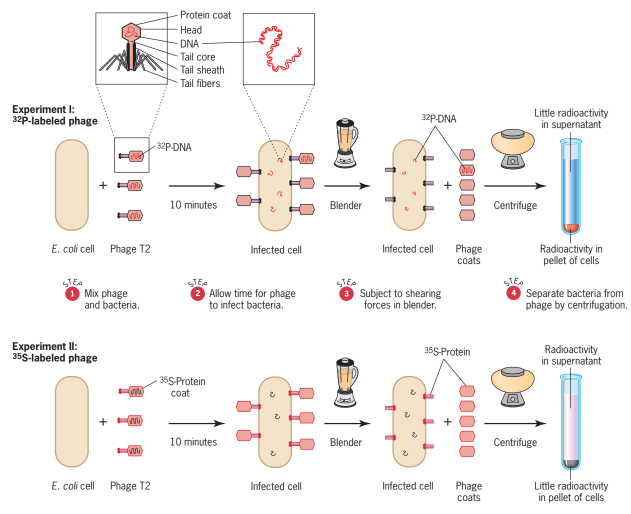
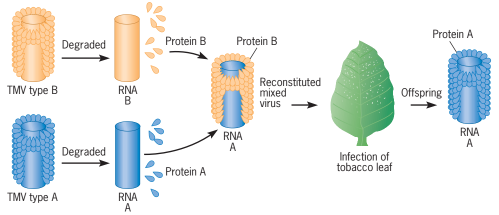


FIGURE 9.3 Hershey and Chase's demonstration that the genetic information of bacteriophage T2 resides in its DNA.

able to label specifically either (1) the phage DNA by growth in a medium containing the radioactive isotope of phosphorus, ³²P, in place of the normal isotope, ³¹P; or (2) the phage protein coats by growth in a medium containing radioactive sulfur, ³⁵S, in place of the normal isotope, ³²S (Figure 9.3).

When T2 phage particles labeled with ³⁵S were mixed with *E. coli* cells for a few minutes and the phage-infected cells were then subjected to shearing forces in a Waring blender, most of the radioactivity (and thus the proteins) could be removed from the cells without affecting progeny phage production. When T2 particles in which the DNA was labeled with ³²P were used, however, essentially all the radioactivity was found inside the cells; that is, the DNA was not subject to removal by shearing in a blender. The sheared-off phage coats were separated from the infected cells by low-speed centrifugation, which pellets (sediments) cells while leaving phage particles suspended. These results indicated that the DNA of the virus enters the host cell, whereas the protein coat remains outside the cell. Since progeny viruses are produced inside the cell, Hershey and Chase's results indicated that the genetic information directing the synthesis of both the DNA molecules and the protein coats of the progeny viruses must be present in the parental DNA. Moreover, the progeny particles were shown to contain some of the ³²P, but none of the ³⁵S of the parental phage.

There was one problem with Hershey and Chase's proof that the genetic material of phage T2 is DNA. Their results showed that a significant amount of ³⁵S (and thus protein) was injected into the host cells with the DNA. Thus, it could be argued that this small fraction of the phage proteins contained the genetic information. More recently, scientists have developed procedures by which protoplasts (cells with



■ FIGURE 9.4 The genetic material of tobacco mosaic virus (TMV) is RNA, not protein. TMV contains no DNA; it is composed of just RNA and protein.

the walls removed) of *E. coli* can be infected with pure phage DNA. Normal infective progeny phage are produced in these experiments, called **transfection** experiments, proving that the genetic material of such bacterial viruses is DNA.

PROOF THAT RNA STORES THE GENETIC INFORMATION IN SOME VIRUSES

As more and more viruses were identified and studied, it became apparent that many of them contain RNA and proteins, but no DNA. In all cases studied to date, it is clear that these RNA viruses—like all other organisms—store their genetic information in nucleic acids rather than in proteins, although in these viruses the nucleic acid is RNA. One of the first experiments that established RNA as the genetic material in RNA viruses was the so-called reconstitution experiment of Heinz Fraenkel-Conrat and coworkers, published in 1957. Their simple, but definitive, experiment was done with tobacco mosaic virus (TMV), a small virus composed of a single molecule of RNA encapsulated in a protein coat. Different strains of TMV can be identified on the basis of differences in the chemical composition of their protein coats.

Fraenkel-Conrat and colleagues treated TMV particles of two different strains with chemicals that dissociate the protein coats of the viruses from the RNA molecules and separated the proteins from the RNA. Then they mixed the proteins from one strain with the RNA molecules from the other strain under conditions that result in the reconstitution of complete, infective viruses composed of proteins from one strain and RNA from the other strain. When tobacco leaves were infected with these reconstituted mixed viruses, the progeny viruses were always phenotypically and genotypically identical to the parent strain from which the RNA had been obtained (**Figure 9.4**). Thus, the genetic information of TMV is stored in RNA, not in protein.

• The genetic information of most living organisms is stored in deoxyribonucleic acid (DNA).

• In some viruses, the genetic information is present in ribonucleic acid (RNA).

KEY POINTS

The Structures of DNA and RNA

The genetic information of all living organisms, except the RNA viruses, is stored in DNA. What is the structure of DNA, and in what form is the genetic information stored? What features of the structure of DNA facilitate the accurate transmission of genetic information from generation to generation? The answers to these questions are without doubt three of the most important facets of our understanding of the nature of life.

DNA is usually double-stranded, with adenine paired with thymine and guanine paired with cytosine. RNA is usually single-stranded and contains uracil in place of thymine.

NATURE OF THE CHEMICAL SUBUNITS IN DNA AND RNA

Nucleic acids, the major components of Miescher's nuclein, are macromolecules composed of repeating subunits called **nucleotides**. Each nucleotide is composed of (1) a phosphate group, (2) a five-carbon sugar, or pentose, and (3) a cyclic nitrogen-containing compound called a base (**Figure 9.5**). In DNA, the sugar is 2-deoxyribose (thus the name deoxyribonucleic acid); in RNA, the sugar is ribose (thus ribonucleic acid). Four different bases commonly are found in DNA: **adenine (A)**, **guanine (G)**, **thymine (T)**, and **cytosine (C)**. RNA also usually contains adenine, guanine, and cytosine but has a different base, **uracil (U)**, in place of thymine. Adenine and guanine are double-ring bases called

Nucleic acids are composed of repeating subunits called nucleotides. Each nucleotide is composed of three units.

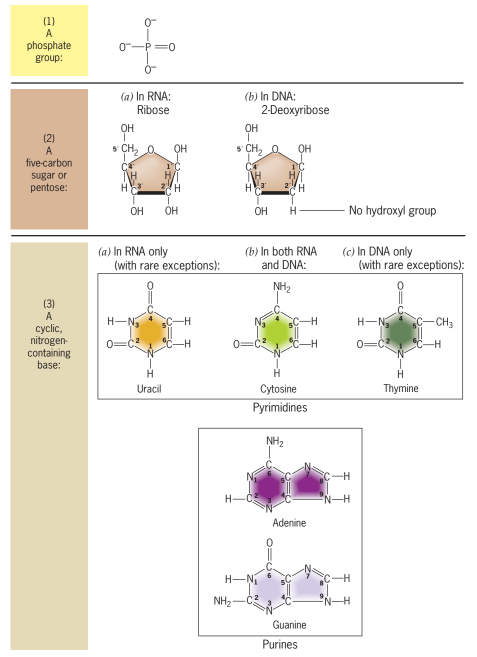


FIGURE 9.5 Structural components of nucleic acids. The standard numbering systems for the carbons in pentoses and the carbons and nitrogens in the ring structures of the bases are shown in (2) and (3), respectively. Single-ring bases are called pyrimidines, and double-ring-bases are purines.

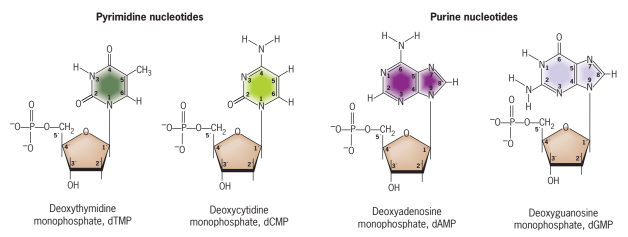


FIGURE 9.6 Structures of the four common deoxyribonucleotides present in DNA. The carbons and nitrogens in the rings of the bases are numbered 1 through 6 (pyrimidines) and 1 through 9 (purines). Therefore, the carbons in the sugars of nucleotides are numbered 1' through 5' to distinguish them from the carbons in the bases.

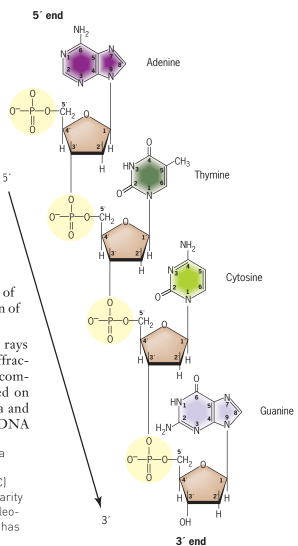
purines; cytosine, thymine, and uracil are single-ring bases called **pyrimidines**. Both DNA and RNA, therefore, contain four different subunits, or nucleotides: two purine nucleotides and two pyrimidine nucleotides (**Trigure 9.6**). In polynucleotides such as DNA and RNA, these subunits are joined together in long chains (**Trigure 9.7**). RNA usually exists as a single-stranded polymer that is composed of a long sequence of nucleotides. DNA has one additional—and very important—level of organization: it is usually a double-stranded molecule.

DNA STRUCTURE: THE DOUBLE HELIX

One of the most exciting breakthroughs in the history of biology occurred in 1953 when James Watson and Francis Crick (**Figure 9.8**) deduced the correct structure of DNA. Their double-helix model of the DNA molecule immediately suggested an elegant mechanism for the transmission of genetic information (see A Milestone in Genetics: The Double Helix on the Student Companion site). Watson and Crick's double-helix structure was based on two major kinds of evidence:

- 1. When Erwin Chargaff and colleagues analyzed the composition of DNA from many different organisms, they found that the concentration of thymine was always equal to the concentration of adenine and the concentration of cytosine was always equal to the concentration of guanine (Table 9.1). Their results strongly suggested that thymine and adenine as well as cytosine and guanine were present in DNA in some fixed interrelationship. Their data also showed that the total concentration of pyrimidines (thymine plus cytosine) was always equal to the total concentration of purines (adenine plus guanine; see Table 9.1).
- 2. When X rays are focused through fibers of purified molecules, the rays are deflected by the atoms of the molecules in specific patterns, called diffraction patterns, which provide information about the organization of the components of the molecules. These *X-ray diffraction patterns* can be recorded on X-ray-sensitive film just as patterns of light can be recorded with a camera and light-sensitive film. Watson and Crick used X-ray diffraction data on DNA

■ FIGURE 9.7 Structure of a polynucleotide chain. The tetranucleotide chain shown is a DNA chain containing the sugar 2'-deoxyribose. RNA chains contain the sugar ribose. The nucleotides in polynucleotide chains are joined by phosphodiester (C−O−P−O−C) linkages. Note that the polynucleotide shown has a 5' (top) to 3' (bottom) chemical polarity because each phosphodiester linkage joins the 5' carbon of 2'-deoxyribose in one nucleo-tide to the 3' carbon of 2'-deoxyribose in the adjacent nucleotide. Therefore, the chain has a 5' carbon terminus at the top and a 3' carbon terminus at the bottom.



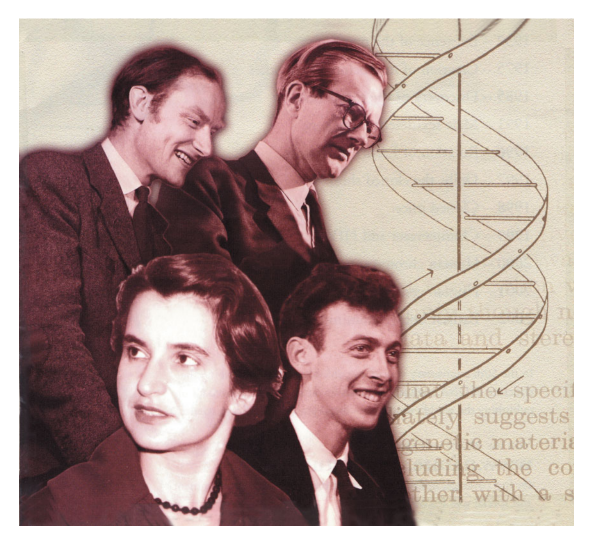


FIGURE 9.8 The four major players—Francis Crick, Maurice Wilkins, James Watson, and Rosalind Franklin (clockwise from top left)—in the discovery of the double-helix structure of DNA.

TABLE 9.1

Base Composition of DNA from Various Organisms

					Mola	Molar Ratios	
Creation	0/ Adamina	% Quenine	0/ Outoping	0/ Thursing	$\frac{A + G}{T + C}$	$\frac{A+T}{G+C}$	
Species	% Adenine	% Guanine	% Cytosine	% Thymine	I + C	0 + 0	
I. Viruses							
Bacteriophage λ	26.0	23.8	24.3	25.8	0.99	1.08	
Bacteriophage T2	32.6	18.1	16.6	32.6	1.03	1.88	
Herpes simplex	13.8	37.7	35.6	12.8	1.06	0.36	
II. Bacteria							
Escherichia coli	26.0	24.9	25.2	23.9	1.04	1.00	
Micrococcus lysodeikticus	14.4	37.3	34.6	13.7	1.07	0.39	
Ramibacterium ramosum	35.1	14.9	15.2	34.8	1.00	2.32	
III. Eukaryotes							
Saccharomyces cerevisiae	31.7	18.3	17.4	32.6	1.00	1.80	
Zea mays (corn)	25.6	24.5	24.6	25.3	1.00	1.04	
Drosophila melanogaster	30.7	19.6	20.2	29.4	1.01	1.51	
<i>Homo sapiens</i> (human)	30.2	19.9	19.6	30.3	1.01	1.53	

5'

structure (**Figure 9.9**) provided by Maurice Wilkins, Rosalind Franklin (see Figure 9.8), and their coworkers. These data indicated that DNA was a highly ordered, two-stranded structure with repeating substructures spaced every 0.34 nanometer (1 nm = 10^{-9} meter) along the axis of the molecule.

On the basis of Chargaff's chemical data, Wilkins' and Franklin's X-ray diffraction data, and inferences from model building, Watson and Crick proposed that DNA exists as a righthanded **double helix** in which the two polynucleotide chains are coiled about one another in a spiral (**I Figure 9.10**). Watson, Crick, and Wilkins shared the 1962 Nobel Prize in Physiology or Medicine for their work on the double-helix model. Unfortunately, Franklin died prematurely (age 37) in 1958, and Nobel Prizes cannot be awarded posthumously.

Each of the two polynucleotide chains in a double helix consists of a sequence of nucleotides linked together by phosphodiester bonds, joining adjacent deoxyribose moieties (Table 9.2). The two polynucleotide strands are held together in

their helical configuration by hydrogen bonding (Table 9.2) between bases in opposing strands; the resulting base pairs are stacked between the two chains perpendicular to the axis of the molecule like the steps of a spiral staircase (Figure 9.10). The base-pairing is specific: adenine is always paired with thymine, and guanine is always paired with cytosine. Thus, all base pairs consist of one purine and one pyrimidine. The specificity of

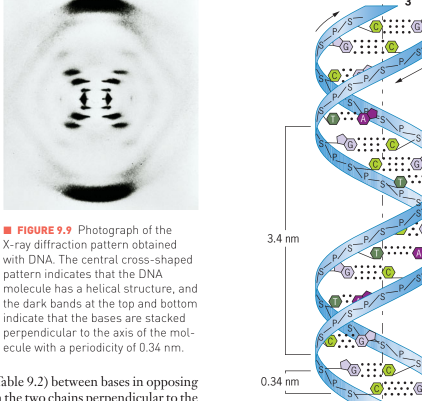
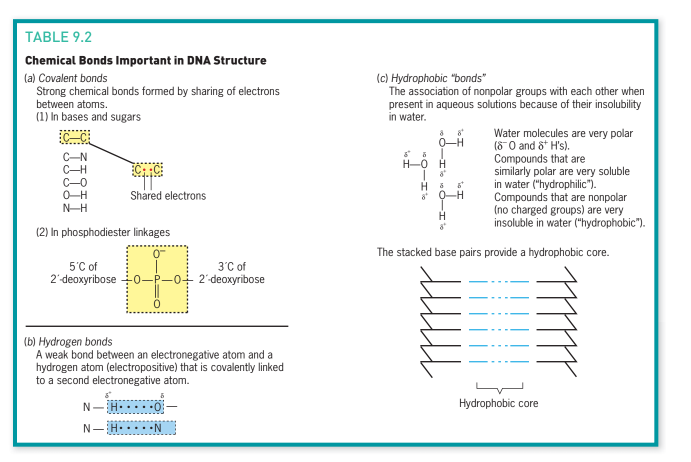
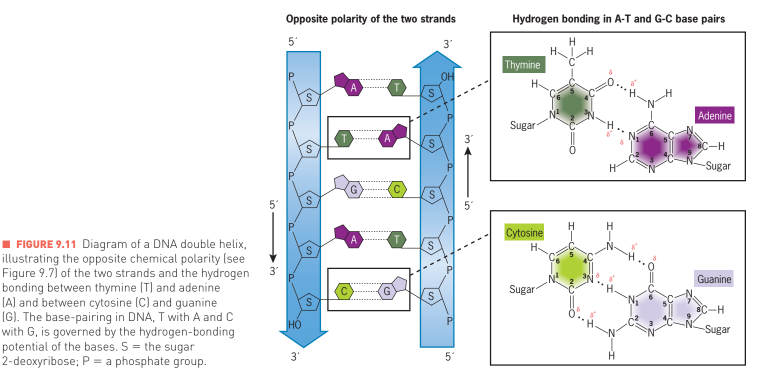


FIGURE 9.10 Diagram of the double-helix structure of DNA.





base-pairing results from the hydrogen-bonding capacities of the bases in their normal configurations (**Figure 9.11**). In their common structural configurations, adenine and thymine form two hydrogen bonds, and guanine and cytosine form three hydrogen bonds. Hydrogen bonding is not possible between cytosine and adenine or thymine and guanine when they exist in their common structural states.

Once the sequence of bases in one strand of a DNA double helix is known, the sequence of bases in the other strand is also known because of the specific base-pairing (see Problem-Solving Skills: Calculating Base Content in DNA). The two strands of a

(G). The base-pairing in DNA, T with A and C with G, is governed by the hydrogen-bonding potential of the bases. S = the sugar 2-deoxyribose; P = a phosphate group.

FIGURE 9.11 Diagram of a DNA double helix, illustrating the opposite chemical polarity (see

bonding between thymine (T) and adenine

(A) and between cytosine (C) and guanine

PROBLEM-SOLVING SKILLS

Calculating Base Content in DNA

THE PROBLEM

Double-stranded genomic DNA was isolated from the bacterium Mycobacterium tuberculosis, and chemical analysis showed that 33 percent of the bases in the DNA were guanine residues. Given this information, is it possible to determine what percent of the bases in the DNA of *M. tuberculosis* were adenine residues?

Single-stranded genomic DNA was isolated from bacteriophage Φ X174, and chemical analysis showed that 22 percent of the bases in the Φ X174 DNA were cytosines. Based on this information, is it possible to determine what percent of the bases in the DNA packaged in the Φ X174 virion were adenines?

FACTS AND CONCEPTS

- 1. In double-stranded DNA, adenine in one strand is always paired with thymine in the complementary strand, and quanine in one strand is always paired with cytosine in the other strand.
- 2. In single-stranded DNA, there is no strict base pairing. There is some base-pairing between bases within the single strands forming hairpin structures, but there is no strict A:T and G:C base pairing as in double-stranded DNA.

ANALYSIS AND SOLUTION

In the double-stranded genomic DNA of *M. tuberculosis*, every A in one strand is hydrogen-bonded to a T in the complementary strand, and every G is hydrogen-bonded to a C in the complementary strand. Thus, if 33 percent of the bases are guanines, 33 percent of the bases are cytosines. That means that 66 percent of the bases are G's and C's and 34 percent (100% - 66%) of the bases are A's and T's. Since A always pairs with T, half are A's and half are T's. Therefore, 17 percent (34% \times 1/2) of the bases in the DNA of *M. tuberculosis* are adenines.

In the single-stranded DNA of bacteriophage Φ X174, there is no strict base pairing, only the occasional pairing between bases within the single strand of DNA. As a result, one cannot predict the proportion of adenine residues in the DNA based on the proportion of cytosines. Indeed, one cannot even predict the percentage of adenines based on the percentage of thymines in single-stranded DNA, like the DNA packaged in the Φ X174 virion.

For further discussion go to the Student Companion site.

DNA double helix are thus said to be complementary. *This property, the* **complementarity** of the two strands of the double helix, makes DNA uniquely suited to store and transmit genetic information from generation to generation (Chapter 10).

The base pairs in DNA are stacked about 0.34 nm apart, with 10 base pairs per turn (360°) of the double helix (Figure 9.10). The sugar-phosphate backbones of the two complementary strands are *antiparallel* (Figure 9.11). Unidirectionally along a DNA double helix, the phosphodiester bonds in one strand go from a 3' carbon of one nucleotide to a 5' carbon of the adjacent nucleotide, whereas those in the complementary strand go from a 5' carbon to a 3' carbon. This "opposite polarity" of the complementary strands of a DNA double helix plays an important role in DNA replication, transcription, and recombination.

The stability of DNA double helices results in part from the large number of hydrogen bonds between the base pairs (even though each hydrogen bond by itself is weak, much weaker than a covalent bond) and in part from the hydrophobic bonding (or stacking forces) between adjacent base pairs (Table 9.2). The stacked nature of the base pairs is best illustrated with a space-filling diagram of DNA structure (**Figure 9.12**). The planar sides of the base pairs are relatively nonpolar and thus tend to be hydrophobic (water-insoluble). Because of this insolubility in water, the hydrophobic core of stacked base pairs contributes considerable stability to DNA molecules present in the aqueous protoplasms of living cells. The space-filling drawing also shows that the two grooves of a DNA double helix are not identical; one, the major groove, is much wider than the other, the minor groove. The difference between the major groove and the minor groove is important when one examines the interactions between DNA and proteins that regulate gene expression. Some proteins bind to the major groove; others bind to the minor groove. Test your understanding of DNA structure by answering the questions posed in Solve It: What Are Some Important Features of Double-Stranded DNA?

DNA STRUCTURE: ALTERNATE FORMS OF THE DOUBLE HELIX

The Watson–Crick double-helix structure just described is called **B-DNA**. B-DNA is the conformation that DNA takes under physiological conditions (in aqueous solutions containing low concentrations of salts). The vast majority of the DNA molecules present in the aqueous protoplasms of living cells exist in the B conformation. However, DNA is not a static, invariant molecule. On the contrary, DNA molecules exhibit considerable conformational flexibility.

The structures of DNA molecules change as a function of their environment. The exact conformation of a given DNA molecule or segment of a DNA molecule will depend on the nature of the molecules with which it is interacting. In fact, intracellular B-DNA appears to have an average of 10.4 nucleotide pairs per turn, rather than precisely 10 as shown in Figure 9.10. In high concentrations of salts or in a partially dehydrated state, DNA exists as **A-DNA**, which is a right-handed helix like B-DNA, but with 11 nucleotide pairs per turn (**Table 9.3**). A-DNA is a shorter, thicker double helix with a diameter of 2.3 nm. DNA molecules almost certainly never exist as A-DNA *in vivo*. However, the A-DNA conformation is important because DNA-RNA

TABLE 9.3							
Alternate Forms of DNA							
Helix Form	Helix Direction	Base Pairs per Turn	Helix Diameter				
А	Right-handed	11	2.3 nm				
В	Right-handed	10	1.9 nm				
Z	Left-handed	12	1.8 nm				

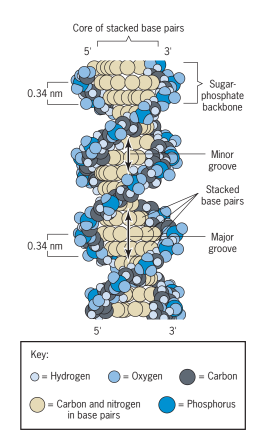


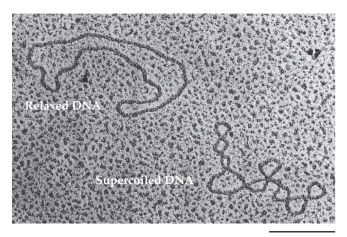
FIGURE 9.12 Space-filling diagram of a DNA double helix.

Solve It!

What Are Some Important Features of Double-Stranded DNA?

One strand of DNA in the coding region of the human *HBB* gene (encoding β -globin) begins with the nucleotide sequence 5'-ATGGTGCATCTGACTCCTGAGGAGA-AGTCT-3', where 5' and 3' designate the carbons on the 2-deoxyribose groups at the ends of the strand. Therefore, this strand of DNA has a 5' \longrightarrow 3' chemical polarity reading left to right. What is the nucleotide sequence of the complementary strand of DNA in this region of the HBB gene? What is the chemical polarity of the complementary strand? What is the length of this segment of the HBB gene when present in a cell as double-stranded DNA? How many 2-deoxyribose molecules are present in this segment of DNA? How many pyrimidine molecules are present in this segment of the HBB gene?

To see the solution to this problem, visit the Student Companion site.



0.1 μm

■ FIGURE 9.13 Comparison of the relaxed and negatively supercoiled structures of DNA. The relaxed structure is B-DNA with 10.4 base pairs per turn of the helix. The negatively supercoiled structure results when B-DNA is underwound, with less than one turn of the helix for every 10.4 base pairs.

heteroduplexes (double helices containing a DNA strand base-paired with a complementary RNA strand) or RNA-RNA duplexes exist in a very similar structure *in vivo*.

Certain DNA sequences have been shown to exist in a left-handed, double-helical form called **Z-DNA** (Z for the zigzagged path of the sugarphosphate backbones of the structure). Z-DNA was discovered by X-ray diffraction analysis of crystals formed by DNA oligomers containing alternating G:C and C:G base pairs. Z-DNA occurs in double helices that are G:C-rich and contain alternating purine and pyrimidine residues. In addition to its unique left-handed helical structure, Z-DNA (Table 9.3) differs from the A and B conformations in having 12 base pairs per turn, a diameter of 1.8 nm, and a single deep groove. The function of Z-DNA in living cells is still not clear.

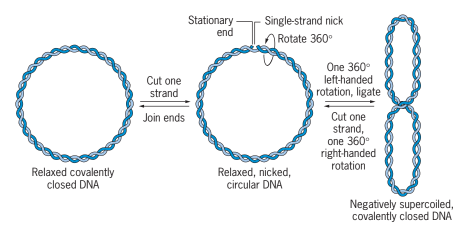
DNA STRUCTURE: NEGATIVE SUPERCOILS IN VIVO

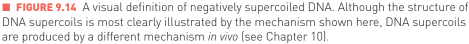
All the functional DNA molecules present in living cells display one other very important level of organization—they are supercoiled. **Supercoils** are introduced into a DNA molecule when one or both strands are cleaved and when the complementary strands at one end are rotated or twisted around each other with the other end held fixed in space—and thus not allowed to

spin. This supercoiling causes a DNA molecule to collapse into a tightly coiled structure similar to a coiled telephone cord or twisted rubber band (**Figure 9.13**, lower right). Supercoils are introduced into and removed from DNA molecules by enzymes that play essential roles in DNA replication (Chapter 10) and other processes.

Supercoiling occurs only in DNA molecules with fixed ends, ends that are not free to rotate. Obviously, the ends of the circular DNA molecules (Figure 9.13) present in most prokaryotic chromosomes and in the chromosomes of eukaryotic organelles such as mitochondria are fixed. The large linear DNA molecules present in eukaryotic chromosomes are also fixed by their attachment at intervals and at the ends to non-DNA components of the chromosomes. These attachments allow enzymes to introduce supercoils into the linear DNA molecules present in eukaryotic chromosomes, just as they are incorporated into the circular DNA molecules present in most prokaryotic chromosomes.

We can perhaps visualize supercoiling most easily by considering a circular DNA molecule. If we cleave one strand of a covalently closed, circular double helix of DNA, and rotate one end of the cleaved strand a complete turn (360°) around the complementary strand while holding the other end fixed, we will introduce one supercoil into the molecule (**Figure 9.14**). If we rotate the free end in the same





direction as the DNA double helix is wound (right-handed), a positive supercoil (overwound DNA) will be produced. If we rotate the free end in the opposite direction (left-handed), a negative supercoil (underwound DNA) will result. Although this is the simplest way to define supercoiling in DNA, it is not the mechanism by which supercoils are produced in DNA *in vivo*. That mechanism is discussed in Chapter 10.

The DNA molecules of almost all organisms, from the smallest viruses to the largest eukaryotes, exhibit **negative supercoiling** *in vivo*, and many of the biological functions of chromosomes can be carried out only when the participating DNA molecules are negatively supercoiled. (The DNA of some viruses that infect Archaea is positively supercoiled.) Considerable evidence indicates that negative supercoiling is involved in replication (Chapter 10), recombination, gene expression, and regulation of gene expression. Similar amounts of negative supercoiling exist in the DNA molecules present in bacterial chromosomes and eukaryotic chromosomes.

- DNA usually exists as a double helix, with the two strands held together by hydrogen bonds between the complementary bases: adenine paired with thymine and guanine paired with cytosine.
- The complementarity of the two strands of a double helix makes DNA uniquely suited to store and transmit genetic information.
- The two strands of a DNA double helix have opposite chemical polarity.
- RNA usually exists as a single-stranded molecule containing uracil instead of thymine.
- The functional DNA molecules in cells are negatively supercoiled.

KEY POINTS

Chromosome Structure in Prokaryotes and Viruses

Much of the information about the structure of DNA has come from studies of prokaryotes, primarily because they are less complex, both genetically and biochemically, than eukaryotes. Prokaryotes are monoploid (mono = one); they have only one set of genes (one copy of the genome). ("Monoploid" should not be conThe DNA molecules of prokaryotes and viruses are organized into negatively supercoiled domains.

fused with "haploid," which refers specifically to the reduced chromosome number in gametes.) In most viruses and prokaryotes, the single set of genes is stored in a single chromosome, which in turn contains a single molecule of nucleic acid (either RNA or DNA).

The smallest known RNA viruses have only three genes, and the complete nucleotide sequences of the genomes of many viruses are known. For example, the single RNA molecule in the genome of bacteriophage MS2 consists of 3569 nucleotides and contains 4 genes. The smallest known DNA viruses have only 9 to 11 genes. Again, the complete nucleotide sequences are known in several cases. For example, the genome of bacteriophage ϕ X174 is a single DNA molecule 5386 nucleotides in length that contains 11 genes. The largest DNA viruses, like bacteriophage T2 and the animal pox viruses, contain about 150 genes. Bacteria like *E. coli* have 2500 to 3500 genes, most of which are present in a single molecule of DNA.

In the past, prokaryotic chromosomes were often characterized as "naked molecules of DNA," in contrast to eukaryotic chromosomes with their associated proteins and complex morphology. This misconception resulted in part because (1) most of the published pictures of prokaryotic "chromosomes" were electron micrographs of isolated DNA molecules, not metabolically active or functional chromosomes, and (2) most of the published photographs of eukaryotic chromosomes were of highly condensed meiotic or mitotic chromosomes—again, metabolically inactive chromosomal states. Functional prokaryotic chromosomes, or nucleoids (nucleoids rather than

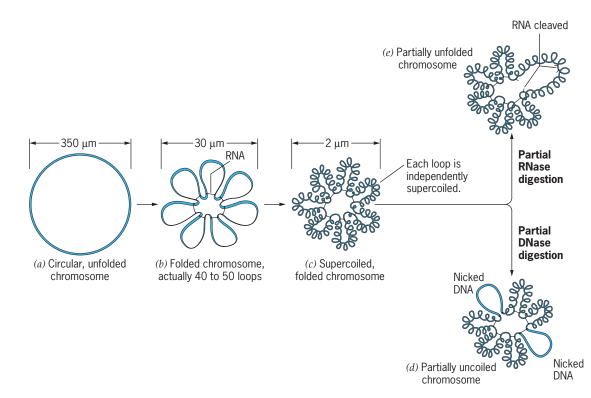


FIGURE 9.15 Diagram of the structure of the functional state of the E. coli chromosome.

nuclei because they are not enclosed in a nuclear membrane), are now known to bear little resemblance to the isolated viral and bacterial DNA molecules seen in electron micrographs, just as the metabolically active interphase chromosomes of eukaryotes have little morphological resemblance to mitotic or meiotic metaphase chromosomes.

The contour length of the circular DNA molecule present in the chromosome of the bacterium *Escherichia coli* is about 1500 μ m. Because an *E. coli* cell has a diameter of only 1 to 2 μ m, the large DNA molecule present in each bacterium must exist in a highly condensed (folded or coiled) configuration. When *E. coli* chromosomes are isolated by gentle procedures in the absence of ionic detergents (commonly used to lyse cells) and are kept in the presence of a high concentration of cations such as polyamines (small basic or positively charged proteins) or 1*M* salt to neutralize the negatively charged phosphate groups of DNA, the chromosomes remain in a highly condensed state comparable in size to the nucleoid *in vivo*. This structure, called the **folded genome**, is the functional state of a bacterial chromosome. Though smaller, the functional intracellular chromosomes of bacterial viruses are very similar to the folded genomes of bacteria.

Within the folded genome, the large DNA molecule in an *E. coli* chromosome is organized into 50 to 100 *domains* or loops, each of which is independently negatively supercoiled (**Figure 9.15**). RNA and protein are both components of the folded genome, which can be partially relaxed by treatment with either deoxyribonuclease (DNase) or ribonuclease (RNase). Because each domain of the chromosome is independently supercoiled, the introduction of single-strand "nicks" in DNA by treatment of the chromosomes with a DNase that cleaves DNA at internal sites will relax the DNA only in the nicked domains, and all unnicked loops will remain supercoiled. Destruction of the RNA connectors by RNase will unfold the folded genome partially by eliminating the organization of the DNA molecule into 50 to 100 loops. However, RNase treatment will not affect the supercoiling of the domains of the chromosome.



• The DNA molecules in prokaryotic and viral chromosomes are organized into negatively supercoiled domains.

Bacterial chromosomes contain circular molecules of DNA segregated into about 50 domains.

Chromosome Structure in Eukaryotes

Eukaryotic genomes contain levels of complexity that are not encountered in prokaryotes. In contrast to prokaryotes, most eukaryotes are diploid, having two complete sets of genes, one from each parent. As we discussed in Chapter 6, some flowering plants are polyploid; that is, they carry several copies of the genome. Although eukaryotes have only about 2 to 15 times as many genes as *E. coli*, they have orders of magnitude more DNA. Moreover, much of this DNA does not contain genes, at least not genes encoding proteins or RNA molecules.

Not only do most eukaryotes contain many times the amount of DNA in prokaryotes, but also this DNA is packaged into several chromosomes, and each chromosome is present in two (diploids) or more (polyploids) copies. Recall that the chromosome of *E. coli* has a contour length of 1500 μ m, or about 1.5 mm. Now consider that the haploid chromosome complement, or genome, of a human contains about 1000 mm of DNA (or about 2000 mm per diploid cell). Moreover, this meter of DNA is subdivided among 23 chromosomes of variable size and shape, with each chromosome containing 15 to 85 mm of DNA. In the past, geneticists had little information as to how this DNA was arranged in the chromosomes. Is there one molecule of DNA per chromosome as in prokaryotes, or are there many? If many, how are the molecules arranged relative to each other? How does the 85 mm (85,000 μ m) of DNA in the largest human chromosome get condensed into a mitotic metaphase structure that is about 0.5 μ m in diameter and 10 μ m long? What are the structures of the metabolically active interphase chromosomes? We consider the answers to some of these questions in the following sections.

CHEMICAL COMPOSITION OF EUKARYOTIC CHROMOSOMES

Interphase chromosomes are usually not visible with the light microscope. However, chemical analysis, electron microscopy, and X-ray diffraction studies of isolated **chromatin** (the complex of the DNA, chromosomal proteins, and other chromosome constituents isolated from nuclei) have provided valuable information about the structure of eukaryotic chromosomes.

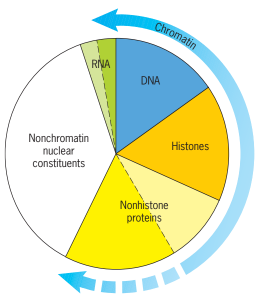
When chromatin is isolated from interphase nuclei, the individual chromosomes are not recognizable. Instead, one observes an irregular aggregate of nucleoprotein. Chemical analysis of isolated chromatin shows that it consists primarily of DNA and proteins with lesser amounts of RNA (**Figure 9.16**). The proteins are of two major classes: (1) basic (positively charged at neutral pH) proteins called **histones** and (2) a heterogeneous, largely acidic (negatively charged at neutral pH) group of proteins collectively referred to as **nonhistone chromosomal proteins**.

Histones play a major structural role in chromatin. They are present in the chromatin of all eukaryotes in amounts equivalent to the amounts of DNA. This relationship suggests that an interaction occurs between histones and DNA that is conserved in eukaryotes. The histones of all plants and animals consist of five classes of proteins. These five major histone types, called *H1*, *H2a*, *H2b*, *H3*, and *H4*, are present in almost all cell types. A few exceptions exist, most notably some sperm, where the histones are replaced by another class of small basic proteins called **protamines**.

The five histone types are present in molar ratios of approximately 1 H1:2 H2a:2 H2b:2 H3:2 H4. Four of the five types of histones are specifically complexed with DNA to produce the basic structural subunits of chromatin, small (approximately 11 nm in diameter by 6.5 nm high) ellipsoidal beads called **nucleosomes**. The histones have been highly conserved during evolution—four of the five types of histone are similar in all eukaryotes.

Most of the 20 amino acids in proteins are neutral in charge; that is, they have no charge at pH 7. However, a few are basic and a few are acidic. The histones are basic because they contain 20 to 30 percent arginine and lysine, two positively charged

Eukaryotic chromosomes contain huge molecules of DNA that are highly condensed during mitosis and meiosis. The centromeres and telomeres of eukaryotic chromosomes have unique structures.



■ FIGURE 9.16 The chemical composition of chromatin as a function of the total nuclear content. The DNA and histone contents of chromatin are relatively constant, but the amount of nonhistone proteins present depends on the procedure used to isolate the chromatin (dashed arrow).

amino acids (see Figure 12.1). The exposed $-NH_3^+$ groups of arginine and lysine allow histones to act as polycations. The positively charged side groups on histones are important in their interaction with DNA, which is polyanionic because of the negatively charged phosphate groups.

The remarkable constancy of histones H2a, H2b, H3, and H4 in all cell types of an organism and even among widely divergent species is consistent with the idea that they are important in chromatin structure (DNA packaging) and are only nonspecifically involved in the regulation of gene expression. However, as will be discussed later, chemical modifications of histones can alter chromosome structure, which, in turn, can enhance or decrease the level of expression of genes located in the modified chromatin.

In contrast, the nonhistone protein fraction of chromatin consists of a large number of heterogeneous proteins. Moreover, the composition of the nonhistone chromosomal protein fraction varies widely among different cell types of the same organism. Thus, the nonhistone chromosomal proteins probably do not play central roles in the packaging of DNA into chromosomes. Instead, they are likely candidates for roles in regulating the expression of specific genes or sets of genes.

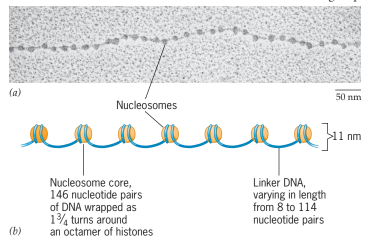
ONE LARGE DNA MOLECULE PER CHROMOSOME

A typical eukaryotic chromosome contains 1 to 20 cm (10^4 to 2×10^5 µm) of DNA. During metaphase of meiosis and mitosis, this DNA is packaged in a chromosome with a length of only 1 to 10 µm. How is all of this DNA condensed into the compact chromosomes that are present during mitosis and meiosis? Do many DNA molecules run parallel throughout the chromosome—the *multineme* or "multistrand" model—or is there just one DNA double helix extending from one end of the chromosome to the other—the *unineme* or single-strand model? (Note that "strand" here refers to the DNA double helix, not the individual polynucleotide chains of DNA.)

Considerable evidence now indicates that each chromosome contains a single, giant molecule of DNA that extends from one end through the centromere all the way to the other end of the chromosome. However, as we will discuss in the following section, this giant DNA molecule is highly condensed (coiled and folded) within the chromosome.

THREE LEVELS OF DNA PACKAGING IN EUKARYOTIC CHROMOSOMES

The largest chromosome in the human genome contains about 85 mm (85,000 μ m, or 8.5×10^7 nm) of DNA that is believed to exist as one giant molecule. This DNA molecule somehow gets packaged into a metaphase structure that is about 0.5 μ m in diameter and



■ FIGURE 9.17 Electron micrograph (a) and low-resolution diagram (b) of the beads-on-a-string nucleosome substructure of chromatin isolated from interphase nuclei. *In vivo*, the DNA linkers are probably wound between the nucleosomes forming a condensed 11-nm fiber.

kaged into a metaphase structure that is about 0.5 μ m in diameter and about 10 μ m in length—a condensation of almost 10⁴-fold in length from the naked DNA molecule to the metaphase chromosome. How does this condensation occur? What components of the chromosomes are involved in the packaging processes? Is there a universal packaging scheme? Are there different levels of packaging? Clearly, meiotic and mitotic chromosomes are more extensively condensed than interphase chromosomes. What additional levels of condensation occur in these special structures that are designed to assure the proper segregation of the genetic material during cell divisions? Are DNA sequences of genes that are not being expressed packaged differently from those of genes that are not being expressed? Let us investigate some of the evidence that establishes the existence of three different levels of packaging of DNA into chromosomes.

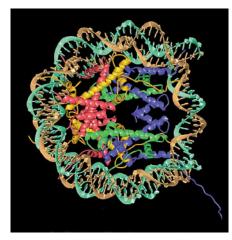
When isolated chromatin from interphase cells is examined by electron microscopy, it is found to consist of a series of ellipsoidal beads (about 11 nm in diameter and 6.5 nm high) joined by thin threads (**Figure 9.17***a*). Further evidence for a regular, periodic packaging of DNA has come from studies on the digestion of chromatin with various nucleases. Partial digestion of chromatin with these

nucleases yielded fragments of DNA in a set of discrete sizes that were integral multiples of the smallest size fragment. These results are nicely explained if chromatin has a repeating structure, supposedly the bead seen by electron microscopy (Figure 9.17*a*), within which the DNA is packaged in a nuclease-resistant form (**■** Figure 9.17*b*). This "bead" or chromatin subunit is called the **nucleosome**. According to the present concept of chromatin structure, the **linkers**, or interbead threads of DNA, are susceptible to nuclease attack.

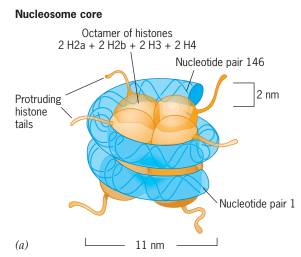
After partial digestion of the DNA in chromatin with an endonuclease (an enzyme that cleaves DNA internally), DNA approximately 200 nucleotide pairs in length is associated with each nucleosome (produced by a cleavage in each linker region). After extensive nuclease digestion, a 146-nucleotide-pair-long segment of DNA remains present in each nucleosome. This nuclease-resistant structure is called the **nucleosome core**. Its structure—essentially invariant in eukaryotes—consists of a 146-nucleotide-pair length of DNA and two molecules each of histones H2a, H2b, H3, and H4. The histones protect the segment of DNA in the nucleosome core from cleavage by endonucleases. Physical studies (X-ray diffraction and similar analyses) of nucleosome-core crystals have shown that the DNA is wound as 1.65 turns of a superhelix around the outside of the histone octamer (**Teigure 9.18a**).

The complete chromatin subunit consists of the nucleosome core, the linker DNA, and the associated nonhistone chromosomal proteins, all stabilized by the binding of one molecule of histone H1 to the outside of the structure (**Figure 9.18b**). The size of the linker DNA varies from species to species and from one cell type to another. Linkers as short as eight nucleotide pairs and as long as 114 nucleotide pairs have been reported. Evidence suggests that the complete nucleosome (as opposed to the nucleosome core) contains two full turns of DNA superhelix (a 166-nucleotide-pair length of DNA) on the surface of the histone octamer and the stabilization of this structure by the binding of one molecule of histone H1 (Figure 9.18*b*).

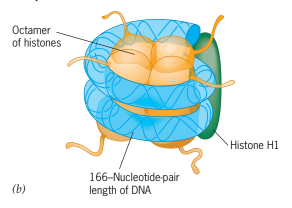
The structure of the nucleosome core has been determined with resolution to 0.28 nm by X-ray diffraction studies. The resulting high-resolution

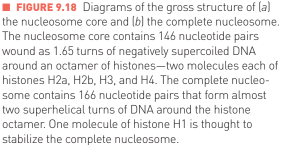


map of the nucleosome core shows the precise location of all eight histone molecules and the 146 nucleotide pairs of negatively supercoiled DNA (**Figure 9.19a** and **b**). Some of the terminal segments of the histones pass over and between the turns of the DNA superhelix to add stability to the nucleosome. The interactions

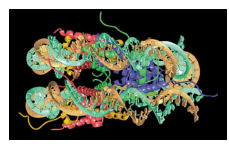


Complete nucleosome









(b)

H2a H2b

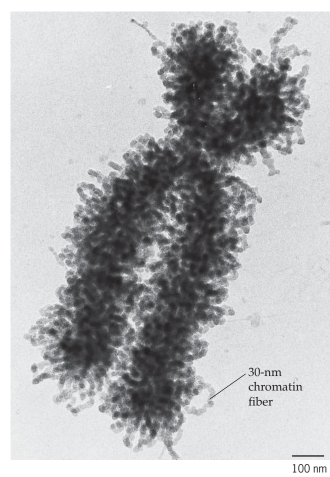
■ FIGURE 9.19 Structure of the nucleosome core based on X-ray diffraction studies with 0.28-nm resolution. The macromolecular composition of the nucleosome core is shown looking along (a) or perpendicular to (b) the axis of the superhelix. (c) Diagram of the structure of a halfnucleosome, which shows the relative positions of the DNA superhelix and the histones more clearly. The complementary strands of DNA are shown in brown and green, and histones H2a, H2b, H3, and H4 are shown in yellow, red, blue, and green, respectively.

Solve It!

How Many Nucleosomes in One Human X Chromosome?

According to the Entrez Genome Database of the National Center for Biotechnology Information, the first human X chromosome to be sequenced contained 154,913,754 nucleotide pairs. If the DNA in this chromosome is organized into nucleosomes and the average internucleosome linker DNA contains 50 nucleotide pairs, how many nucleosomes will be present in this chromosome during interphase? How many molecules of histone H3 will be present in this X chromosome?

To see the solution to this problem, visit the Student Companion site.



of a human metanl

FIGURE 9.20 Electron micrograph of a human metaphase chromosome showing the presence of 30-nm chromatin fibers. The available evidence indicates that each chromatid contains one large, highly coiled or folded 30-nm fiber.

between the various histone molecules and between the histones and DNA are seen most clearly in the structure of one-half of the nucleosome core (**Figure 9.19***c*), which contains only 73 nucleotide pairs of supercoiled DNA. Try Solve It: How Many Nucleosomes in One Human X Chromosome? to test your understanding of nucleosome structure.

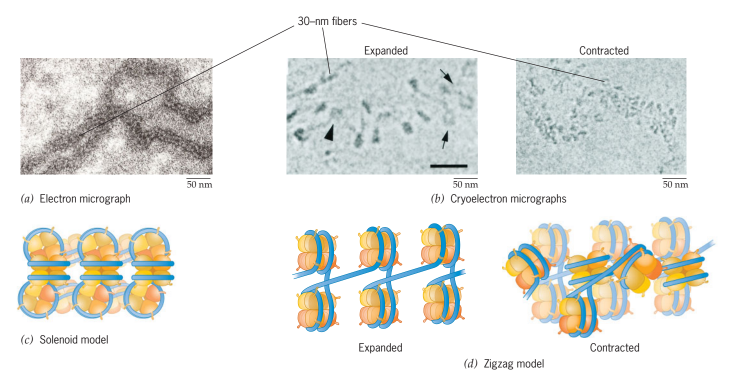
The basic structural component of eukaryotic chromatin is the nucleosome. But are the structures of all nucleosomes the same? What role(s), if any, does nucleosome structure play in gene expression and regulation of gene expression? The structure of nucleosomes in transcriptionally active regions of chromatin is known to differ from that of nucleosomes in transcriptionally inactive regions. But what are the details of this structure–function relationship? The tails of some of the histone molecules protrude from the nucleosome and are accessible to enzymes that add and remove chemical groups such as methyl ($-CH_3$) and acetyl groups. The addition of these groups can change the level of expression of genes packaged in nucleosomes containing the modified histones (see Chapter 19).

Electron micrographs of isolated metaphase chromosomes show masses of tightly coiled or folded lumpy fibers (**Figure 9.20**). These **chromatin fibers** have an average diameter of 30 nm. When the structures seen by light and electron microscopy during earlier stages of meiosis are compared, it becomes clear that the light microscope simply permits one to see those regions where these 30-nm fibers are tightly packed or condensed. Indeed, when interphase chromatin is isolated using very gentle procedures, it also consists of 30-nm fibers (**Figure 9.21a**). However, the structure of these fibers seems to be quite variable and depends on

the procedures used. When observed by cryoelectron microscopy (microscopy using quickly frozen chromatin rather than fixed chromatin), the 30-nm fibers show less tightly packed "zigzag" structures (**Figure 9.21b**).

What is the substructure of the 30-nm fiber seen in chromosomes? The two most popular models are the solenoid model (**Figure 9.21c**) and the zigzag model (**Figure 9.21d**). In vivo, the nucleosomes clearly interact with one another to condense the 11-nm nucleosomes into 30-nm chromatin fibers. Whether these have solenoid structures or zigzag structures, or both, depending on the conditions, is still uncertain. What is certain is that chromatin structure is not static; chromatin can expand and contract in response to chemical modifications of histone H1 and the histone tails that protrude from the nucleosomes.

Metaphase chromosomes are the most condensed of normal eukaryotic chromosomes. Clearly, the role of these highly condensed chromosomes is to organize and package the giant DNA molecules of eukaryotic chromosomes into structures that will facilitate their segregation to daughter nuclei without the DNA molecules of different chromosomes becoming entangled and, as a result, being broken during the anaphase separation of the daughter chromosomes. As we noted in the preceding section, the basic structural unit of the metaphase chromosome is the 30-nm chromatin fiber. However, how are these 30-nm fibers further condensed into the observed metaphase structure? Unfortunately, there is still no clear answer to this question. There is evidence that the gross structure of metaphase chromosomes is not dependent on histones. Electron micrographs of isolated metaphase chromosomes from which the histones have been removed reveal a scaffold, or central core, which is surrounded by a huge pool or halo of DNA (**Figure 9.22**). This chromosome scaffold must be composed of nonhistone chromosomal proteins. Note the absence of any apparent ends of DNA molecules in the micrograph shown in Figure 9.22; this finding again supports the concept of one giant DNA molecule per chromosome.



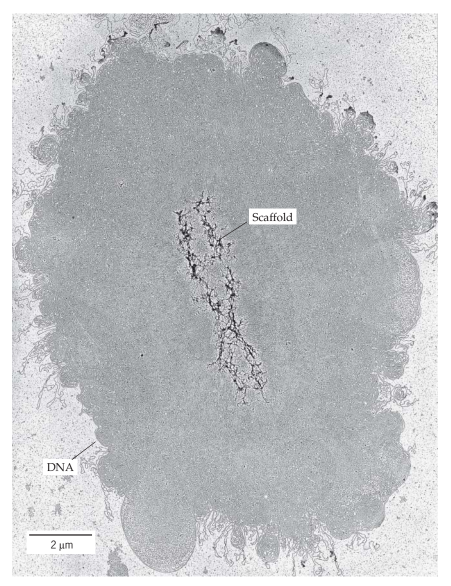
■ FIGURE 9.21 Electron micrograph (a) and cryoelectron micrographs (b) of the 30-nm chromatin fibers in eukaryotic chromosomes. The structure of 30-nm chromatin fibers seems to vary based on the procedures used to isolate and photograph them. (c) According to one popular model, the 30-nm fiber is produced by coiling the 11-nm nucleosome fiber into a solenoid structure with six nucleosomes per turn. (d) However, when chromatin is visualized after cryopreservation (quick freezing) without fixation, it exhibits a zigzag structure whose density—expanded versus contracted—varies with ionic strength and with chemical modifications of the histone molecules.

In summary, at least three levels of condensation are required to package the 10^3 to $10^5 \,\mu\text{m}$ of DNA in a eukaryotic chromosome into a metaphase structure a few microns long (**Tigure 9.23**).

- 1. The first level of condensation involves packaging DNA as a negative supercoil into nucleosomes, to produce the 11-nm-diameter interphase chromatin fiber. This clearly involves an octamer of histone molecules, two each of histones H2a, H2b, H3, and H4.
- **2.** The second level of condensation involves an additional folding or supercoiling of the 11-nm nucleosome fiber, to produce the 30-nm chromatin fiber. Histone H1 is involved in this supercoiling of the 11-nm nucleosome fiber to produce the 30-nm chromatin fiber.
- **3.** Finally, nonhistone chromosomal proteins form a scaffold that is involved in condensing the 30-nm chromatin fiber into the tightly packed metaphase chromosomes. This third level of condensation appears to involve the separation of segments of the giant DNA molecules present in eukaryotic chromosomes into independently supercoiled domains or loops. The mechanism by which this third level of condensation occurs is not known.

CENTROMERES AND TELOMERES

As we discussed in Chapter 2, the two homologous chromosomes (each containing two sister chromatids) of each chromosome pair separate to opposite poles of the meiotic spindle during anaphase I of meiosis. Similarly, during anaphase II of meiosis and the single anaphase of mitosis, the sister chromatids of each chromosome move to opposite spindle poles and become daughter chromosomes. These anaphase movements



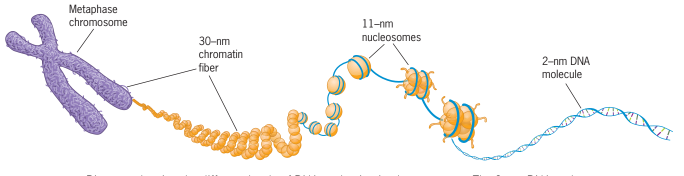
■ FIGURE 9.22 Electron micrograph of a human metaphase chromosome from which the histones have been removed. A huge pool of DNA surrounds a central "scaffold" composed of nonhistone chromosomal proteins. Note that the scaffold has roughly the same shape as the metaphase chromosome prior to removal of the histones. Also note the absence of ends of DNA molecules in the halo of DNA surrounding the scaffold.

depend on the attachment of spindle microtubules to specific regions of the chromosomes, the centromeres. Because all centromeres perform the same basic function, it is not surprising that the centromeres of different chromosomes of a species contain similar structural components.

The centromere of a metaphase chromosome can usually be recognized as a constricted region (see Figure 9.20). In fact, the production of two functional centromeres is a key step in the transition from metaphase to anaphase, and a functional centromere must be present on each daughter chromosome to avoid the deleterious effects of nondisjunction. Acentric chromosomal fragments are usually lost during mitotic and meiotic divisions.

The structures of centromeres in multicellular plants and animals vary widely from species to species. The one feature that they have in common is the presence of specific DNA sequences that are repeated many times, frequently in long tandem arrays. Other DNA sequences are often found embedded within these tandem arrays. Each centromere of human chromosomes, for example, contains 5000 to 15,000 copies of a 171 base-pair-long sequence called the alpha (sometimes "alphoid") satellite sequence (**Figure 9.24**). (Satellite sequences form distinct "satellite" bands during centrifugation in density gradients (see Chapter 10). Huntington Willard and colleagues have shown that a 450,000 base-pair segment of the centromere of the human X chromosome is sufficient for centromere function. This segment consists mostly of alpha satellite sequences but contains interspersed centromere protein (CENP) binding sites called CENP-B boxes. Both components are essential for centromere function.

It has been known for several decades that the **telomeres** (from the Greek terms *telos* and *meros*, meaning "end" and "part," respectively), or ends of eukaryotic chromosomes, have unique properties. Hermann J. Muller, who introduced the term *telomere*



■ FIGURE 9.23 Diagram showing the different levels of DNA packaging in chromosomes. The 2-nm DNA molecule is first condensed into 11-nm nucleosomes, which are further condensed into 30-nm chromatin fibers. The 30-nm fibers are then segregated into supercoiled domains or loops via their attachment to chromosome scaffolds composed of nonhistone chromosomal proteins.

in 1938, demonstrated that *Drosophila* chromosomes without natural ends—produced by breaking chromosomes with X rays—were not transmitted to progeny. In a classical study of maize chromosomes, Barbara McClintock demonstrated that new ends of broken chromosomes are sticky and tend to fuse with each other. In contrast, the natural ends of normal (unbroken) chromosomes are stable and show no tendency to fuse with other broken or native ends. McClintock's results indicated that telomeres must have special structures different from the ends produced by breakage of chromosomes.

Another reason for postulating that telomeres have unique structures is that the known mechanisms of replication of linear DNA molecules do not permit duplication of both strands of DNA at the ends of the molecules (Chapter 10). Thus, telomeres must have unique structures that facilitate their replication, or there must be some special replication enzyme that resolves this enigma. Whatever their structure, telomeres must provide at least three important functions. They must (1) prevent deoxyribonucleases from degrading the ends of the linear DNA molecules, (2) prevent fusion of the ends with other DNA molecules, and (3) facilitate replication of the ends of the linear DNA molecules without loss of material.

The telomeres of eukaryotic chromosomes have unique structures that include short nucleotide sequences present as tandem repeats. Although the sequences vary somewhat in different species, the basic repeat unit has the pattern 5' $T_{1-4}A_{0-1}G_{1-8}$ ' in all but a few species. For example, the repeat sequence in humans and other vertebrates is TTAGGG, that of the protozoan *Tetrahymena thermophila* is TTGGGG, and that of the plant *Arabidopsis thaliana* is TTTAGGG. In most species, additional repetitive DNA sequences are present adjacent to telomeres; these are referred to as telomere-associated sequences.

In vertebrates, the TTAGGG repeat is highly conserved; it has been identified in more than 100 species, including mammals, birds, reptiles, amphibians, and fishes. The number of copies of this basic repeat unit in telomeres varies from species to species, from chromosome to chromosome within a species, and even on the same chromosome in different cell types. In normal (noncancerous) human somatic cells, telomeres usually contain 500 to 3000 TTAGGG repeats and gradually shorten with age. In contrast, the telomeres of germ-line cells and cancer cells do not shorten with age (see Telomere Length and Aging in Humans in Chapter 10).

The telomeres of a few species are not composed of short tandem repeats of the type described earlier. In *D. melanogaster*; for example, telomeres are composed of two specialized DNA sequences that can move from one location in the genome to other locations. Because of their mobility, such sequences are called *transposable genetic elements* (see Chapter 17).

Most telomeres terminate with a G-rich single-stranded region of the DNA strand with the 3' end (a so-called 3' overhang). These overhangs are short (12 to 16 bases) in ciliates such as *Tetrahymena*, but they are quite long (50 to 500 bases) in humans. The guanine-rich repeat sequences of telomeres have the ability to form hydrogen-bonded structures distinct from those produced by Watson–Crick base-pairing in DNA. Oligonucleotides that contain tandem telomere repeat sequences form these special structures in solution, but whether they exist *in vivo* remains unknown.

The telomeres of humans and a few other species have been shown to form structures called **t-loops**, in which the single strand at the 3' terminus invades an upstream telomeric repeat (TTAGGG in mammals) and pairs with the complementary strand, displacing the equivalent strand (**Televenergy Probability**). The DNA in these t-loops is protected from degradation and/or modification by DNA repair processes by a telomere-specific protein complex called **shetterin**. Shelterin is composed of six different proteins, three of which bind specifically to telomere repeat sequences. TRF1 and TRF2 bind to double-stranded repeat sequences. Subunits TIN2 and TPP1 tether POT1 to DNA-bound TRF1 and TRF2, and the TRF2-associated protein Rap1 helps regulate telomere length. Shelterin is present in sufficient quantities in most cells to coat all the single- and double-stranded telomere repeat sequences in the chromosome complement.

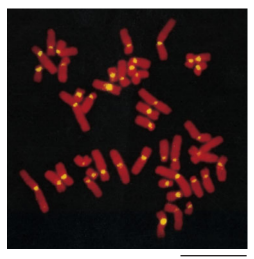
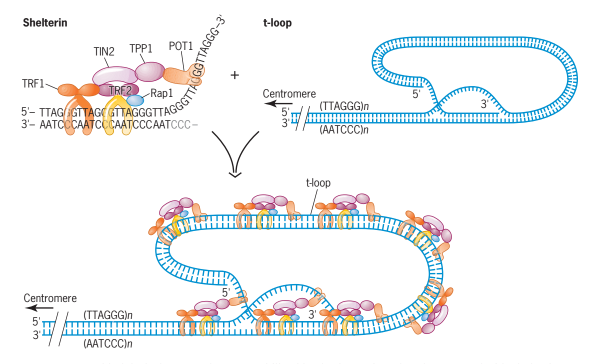




FIGURE 9.24 The location of alpha satellite DNA sequences (yellow) in the centromeres of human chromosomes (red). See Appendix C: *In Situ* Hybridization.



■ FIGURE 9.25 Model of a human telomere stabilized by the formation of a t-loop coated with shelterin. The 3'-terminus forms a t-loop by invading an upstream telomere repeat and pairing with the complementary strand. Shelterin contains six protein subunits, along with some associated proteins (not shown). TRF1 and TRF2 are telomere repeat-binding factors 1 and 2; they bind specifically to double-stranded repeat sequences. Protein POT1 binds specifically to single-stranded TTAGGG repeats displaced by the invading 3'-terminus of the telomeric DNA. TIN2 and TPP1 tether POT1 to TRF1 and TRF2, and the TRF2-associated Rap1 helps regulate telomere length.

To date, t-loops have been identified in the telomeres of vertebrates, the ciliate *Oxytricha fallax*, the protozoan *Trypanosoma brucei*, and the plant *Pisum sativum* (peas). Thus, they are probably important components of the telomeres of most species.

REPEATED DNA SEQUENCES

The centromeres and telomeres discussed in this chapter contain DNA sequences that are repeated many times. Indeed, the chromosomes of eukaryotes contain many DNA sequences that are repeated in the haploid chromosome complement, sometimes as many as a million times. DNA containing such repeated sequences, called **repetitive DNA**, is a major component (15 to 80 percent) of eukaryotic genomes.

The first evidence for repetitive DNA came from centrifugation studies of eukaryotic DNA. When the DNA of a prokaryote, such as E. coli, is isolated, fragmented, and centrifuged at high speeds for long periods of time in a 6M cesium chloride (CsCl) solution, the DNA will form a single band in the centrifuge tube at the position where its density is equal to the density of the CsCl solution (Chapter 10). For E. coli, this band will form at a position where the CsCl density is equal to the density of DNA containing about 50 percent A:T and 50 percent G:C base pairs. DNA density increases with increasing G:C content. The extra hydrogen bond in a G:C base pair results in a tighter association between the bases and thus a higher density than for A:T base pairs. The centrifugation of DNAs from eukaryotes to equilibrium conditions in such CsCl solutions usually reveals the presence of one large main band of DNA and one to several small bands. These small bands of DNA are called satellite bands (from the Latin word *satelles*, meaning "an attendant" or "subordinate") and the DNAs in these bands are often referred to as satellite DNAs. For example, the genome of Drosophila virilis, a distant relative of Drosophila melanogaster; contains three distinct satellite DNAs, each composed of a repeating sequence of seven base pairs. Other satellite DNAs in eukaryotes have long repetitive sequences.

Much of what we know about the types of repeated DNA sequences in the chromosomes of various eukaryotic species resulted from DNA renaturation experiments. The two strands of a DNA double helix are held together by a large number of relatively weak hydrogen bonds between complementary bases. When DNA molecules in aqueous solution are heated to near 100°C, these bonds are broken and the complementary strands of DNA separate. This process is called **denaturation**. If the complementary single strands of DNA are cooled slowly under the right conditions, the complementary base sequences will find each other and will re-form base-paired double helices. This reformation of double helices from the complementary single strands of DNA is called **renaturation**.

If a DNA sequence is repeated many times, denaturation will yield a large number of complementary single strands that will renature rapidly, faster than the rate of renaturation of sequences that are present only once in the genome. Indeed, the rate of DNA renaturation is directly proportional to copy number (the number of copies of the sequence in the genome)-the higher the copy number, the faster the rate and the less time required for renaturation. Mathematical analyses of the rates of renaturation of DNA sequences in eukaryotic genomes provided strong evidence for the presence of different classes of repeated DNA sequences, or repetitive DNA, in eukaryotic chromosomes. The recent genome sequencing projects have provided additional information about the different types of repetitive DNA sequences in eukaryotic genomes, and ongoing sequencing projects are providing information about the sequence variability that occurs in human populations (see On the Cutting Edge: The 1000 Genomes Project). The locations of different DNA sequences in chromosomes can be determined directly by procedures similar to the renaturation experiments described here. With this procedure, called *in situ* hybridization, labeled strands of DNA form double helices with denatured DNA still present in chromosomes (see Appendix C: In Situ Hybridization).

The most highly repeated sequences in eukaryotic genomes do not encode proteins. Indeed, they are not even transcribed. Other less repetitive sequences encode proteins, such as ribosomal proteins and the muscle proteins actin and myosin that are needed in large amounts and are each encoded by several genes. The genes that specify ribosomal RNAs are also multicopy genes because cells need large amounts of ribosomal RNA to produce the ribosomes required for protein synthesis.

The most prevalent of the repeated DNA sequences are **transposable genetic elements**, DNA sequences that can move from one location in a chromosome to another or even to a different chromosome (Chapter 17), or inactive sequences derived from transposable elements. In *D. melanogaster*, about 90 different families of transposable elements have been characterized and been given interesting names such as *hobo*, *pogo*, and *gypsy* that suggest their mobility. A much larger proportion—between 40 and 50 percent—of the human genome contains transposable elements or sequences derived from them. As much as 80 percent of the corn genome may consist of transposable genetic elements or their derivatives. These repetitive transposable elements are discussed in more detail in Chapters 15 and 17.

- Each eukaryotic chromosome contains one giant molecule of DNA packaged into 11-nm ellipsoidal beads called nucleosomes.
- The condensed chromosomes that are present in mitosis and meiosis and carefully isolated interphase chromosomes are composed of 30-nm chromatin fibers.
- At metaphase, the 30-nm fibers are segregated into domains by scaffolds composed of nonhistone chromosomal proteins.
- The centromeres (spindle-fiber-attachment regions) and telomeres (termini) of chromosomes have unique structures that facilitate their functions.
- Eukaryotic genomes contain repeated DNA sequences, with some sequences present a million times or more.

KEY POINTS

ON THE CUTTING EDGE

THE 1000 GENOMES PROJECT

ow much DNA sequence variation exists among people from a particular population? from different populations? These questions are being addressed by the 1000 Genomes Project, a venture launched in 2008 by an international consortium of scientists. The Project's goal is to sequence at least 2500 genomes from people representing ancestral groups from all over the world (**Table 1**). This collection of genome sequences will provide detailed information about human genetic diversity. Where in the genome do we find sequence differences? How frequent are they? Are genomes from the same ancestral group closer in sequence than genomes from different ancestral groups?

The nearly complete sequences of the genomes—about 3 billion nucleotide pairs—of a few individuals are now available, and

TABLE 1

The 1000 Genomes Project: Worldwide Distribution of Selected Genomes

500 Genomes of European Ancestry: 100 genomes from each of the following locations: Utah, United States; Toscani, Italy; England and Scotland; Finland; and Spain.

500 Genomes of East Asian Ancestry: 100 genomes from each of the following locations: Beijing, China; Tokyo, Japan; South China; Xishaungbanna, China; and Ho Chi Minh City, Vietnam.

500 Genomes of West African Ancestry: 100 genomes from each of the following locations: Ibadan, Nigeria; Webuye, Kenya; Western Gambia; Navrongo, Ghana; and Blantyre, Malawi.

500 Genomes of American or African American Ancestry: 70 genomes from each of the following locations: Medellin, Columbia; Lima, Peru; Puerto Rico; and Los Angeles (with Mexican ancestry); plus 79 genomes from Barbados; 80 genomes from Jackson, Mississippi (with African ancestry); and 61 genomes from the southwestern regions of the United States (with African ancestry).

500 Genomes of South Asian Ancestry: 100 genomes from each of the following locations: Assam, India; Calcutta, India; Hyderabad, India; Bombay, India; and Lahore, Pakistan.



new technologies have made sequencing much faster and less expensive, making the goals of the 1000 Genomes Project achievable.

We already know quite a bit about certain types of variation in the human genome, especially the short DNA sequences that are present as tandem repeats in chromosomes. These sequences exhibit highly variable copy number, making them invaluable in personal identification cases—that is, in identifying or distinguishing individuals. We will discuss the use of these variable sequences, a process called DNA profiling (originally DNA Fingerprinting), in forensic and paternity cases, and in identification of otherwise unidentifiable bodies after explosions, crashes, or other tragedies, in Chapter 16.

The 1000 Genomes Project will focus on other types of genetic variation, for example, single *n*ucleotide *p*olymorphisms (SNPs), insertions and deletions, and large structural changes in the DNA. The Project hopes to identify the majority of the sequence variants in the human genome that occur at a frequency of at least 1 percent.

Three pilot projects were carried out in 2008–2009 to assess the feasibility of the plan and to decide how to best achieve the overall goals. In 2008, major portions of the genomes from 180 people were sequenced, and the genomes of two three-member families (mother, father, and child) were sequenced nearly to completion. Then in 2009, the sequences of a thousand gene-rich regions from the genomes of 900 individuals were obtained. Primed by the success of this work, the main project got underway in 2009 and 2010 with efforts to sequence 2500 genomes from a total of 22 different populations. All the data accumulated by the Project are available for everyone to see on a web site maintained by the National Center for Biotechnology Information.

What might we do with this DNA sequence information? One use will be to study the genetic relationships among different human populations so that we can better understand who we are and where we came from. Another use will be to correlate particular sequence variants with alleles that influence our susceptibility to disease—heart disease, cancer, dementia, rheumatoid arthritis, behavioral disorders, and many other types of illnesses. Thus, the long-term significance of the Project is that it will enhance our understanding of the genetic basis of human health.

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. What differences in the chemical structures of DNA and protein allow scientists to label one or the other of these macromolecules with a radioactive isotope?
- **Answer:** DNA contains phosphorus (the common isotope is ³¹P) but no sulfur; DNA can be labeled by growing cells on medium containing the radioactive isotope of phosphorus, ³²P. Proteins contain sulfur (the common isotope is ³²S) but usually little or no phosphorus; proteins can be labeled

by growing cells on medium containing the radioactive isotope of sulfur, ³⁵S.

- 2. If the sequence of one strand of a DNA double helix is ATCG, what is the sequence of the other strand?
- **Answer:** Because the two strands of a double helix are complementary—adenine always paired with thymine and guanine always paired with cytosine—the sequence of the

second strand can be deduced from the sequence of the first strand. For ATCG, the double helix will have the following structure:

ATCG TÄGC

- **3.** How should the sequence of the complementary strand in the double helix in Exercise 2 be written as a single strand of DNA?
- **Answer:** Remember that the two strands of a DNA double helix have opposite chemical polarity; one strand has $5' \rightarrow 3'$ polarity, and the other has $3' \rightarrow 5'$ polarity, when both are read in the same direction. Because the accepted convention is to write sequences starting with the 5'-terminus on the left and ending with the 3'-terminus on the right, the top strand of the double helix should be written 5'-ATCG-3' and the complementary strand, 5'-CGAT-3'. The structure of the double helix should be written:

5'-ATCG-3' 3'-TÄGC-5'

4. If a mixture of DNA and protein is shown to contain genetic information by some assay such as transformation in bacteria, how can a researcher determine whether that

genetic information is present in the DNA or the protein component?

- **Answer:** The biological specificity of enzymes provides a powerful tool for use in many investigations. The enzyme deoxyribonuclease (DNase) degrades DNA to mononucleotides, and proteases degrade proteins to smaller components. If the mixture of DNA and protein is treated with DNase and the genetic information is destroyed, it is stored in DNA. If the mixture is treated with protease and the genetic information is lost, it resides in the protein component of the mixture.
- **5.** How are the single-stranded regions of DNA at the ends of human chromosomes protected from degradation by nucleases and other enzymes?
- **Answer:** The single-stranded 3' overhangs in telomeres of human chromosomes invade telomere repeat sequences (TTAGGG) upstream from the terminus and form lariat-like structures called t-loops (see Figure 9.25). The DNA molecules in the t-loops are coated with a telomere-specific protein complex called shelterin. One of the proteins (POT1) in the shelterin complex binds specifically to the single-stranded repeat sequences in the telomeres protecting them from degradation by nucleases and other enzymes involved in repair of damaged DNA.

Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. The red alga *Polyides rotundus* stores its genetic information in double-stranded DNA. When DNA was extracted from *P. rotundus* cells and analyzed, 32 percent of the bases were found to be guanine residues. From this information, can you determine what percentage of the bases in this DNA were thymine residues? If so, what percentage? If not, why not?
- Answer: The two strands of a DNA double helix are complementary to each other, with guanine (G) in one strand always paired with cytosine (C) in the other strand and, similarly, adenine (A) always paired with thymine (T). Therefore, the concentrations of G and C are always equal, as are the concentrations of A and T. If 32 percent of the bases in double-stranded DNA are G residues, then another 32 percent are C residues. Together, G and C comprise 64 percent of the bases in *P. rotundus* DNA; thus, 36 percent of the bases are A's and T's. Since the concentration of A must equal the concentration of T, 18 percent (36% × 1/2) of the bases must be T residues.
- 2. The *E. coli* virus $\phi X174$ stores its genetic information in single-stranded DNA. When DNA was extracted from $\phi X174$ virus particles and analyzed, 21 percent of the bases were found to be G residues. From this information, can you determine what percentage of the bases in this DNA were thymine residues? If so, what percentage? If not, why not?

- **Answer:** No! The A = T and G = C relationships occur only in double-stranded DNA molecules because of their complementary strands. Since base-pairing does not occur or occurs only as limited intrastrand pairing in singlestranded nucleic acids, you cannot determine the percentage of any of the other three bases from the G content of the $\phi X174$ DNA.
- 3. If each G_1 -stage human chromosome contains a single molecule of DNA, how many DNA molecules would be present in the chromosomes of the nucleus of (a) a human egg, (b) a human sperm, (c) a human diploid somatic cell in stage G_1 , (d) a human diploid somatic cell in stage G_2 , (e) a human primary oocyte?
- Answer: A normal human haploid cell contains 23 chromosomes, and a normal human diploid cell contains 46 chromosomes, or 23 pairs of homologues. If prereplication chromosomes contain a single DNA molecule, postreplication chromosomes will contain two DNA molecules, one in each of the two chromatids. Thus, normal human eggs and sperm contain 23 chromosomal DNA molecules; diploid somatic cells contain 46 and 92 chromosomal DNA molecules at stages G_1 and G_2 , respectively; and a primary oocyte contains 92 such DNA molecules.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **9.1** (a) How did the transformation experiments of Griffith differ from those of Avery and his associates? (b) What was the significant contribution of each? (c) Why was Griffith's work not evidence for DNA as the genetic material, whereas the experiments of Avery and coworkers provided direct proof that DNA carried the genetic information?
- **9.2** A cell-free extract is prepared from Type IIIS pneumococcal cells. What effect will treatment of this extract with (a) protease, (b) RNase, and (c) DNase have on its subsequent capacity to transform recipient Type IIR cells to Type IIIS? Why?
- **9.3** How could it be demonstrated that the mixing of heatkilled Type III pneumococcus with live Type II resulted in a transfer of genetic material from Type III to Type II rather than a restoration of viability to Type III by Type II?
- **9.4** What is the macromolecular composition of a bacterial virus or bacteriophage such as phage T2?
- **9.5** (a) What was the objective of the experiment carried out by Hershey and Chase? (b) How was the objective accomplished? (c) What is the significance of this experiment?
- **9.6** How did the reconstitution experiment of Fraenkel-Conrat and colleagues show that the genetic information of tobacco mosaic virus (TMV) is stored in its RNA rather than its protein?
- **9.7** (a) What background material did Watson and Crick have available for developing a model of DNA? (b) What was their contribution to building the model?
- **9.8** (a) Why did Watson and Crick choose a double helix for their model of DNA structure? (b) Why were hydrogen bonds placed in the model to connect the bases?
- **9.9** (a) If a virus particle contained double-stranded DNA with 200,000 base pairs, how many nucleotides would be present? (b) How many complete spirals would occur on each strand? (c) How many atoms of phosphorus would be present?(d) What would be the length of the DNA configuration in the virus?
- 9.10 What are the differences between DNA and RNA?
- **9.11** RNA was extracted from TMV (tobacco mosaic virus) particles and found to contain 20 percent cytosine (20 percent of the bases were cytosine). With this information, is it possible to predict what percentage of the bases in TMV are adenine? If so, what percentage? If not, why not?
- **9.12 (constructed from cells of** *Staphylococcus afermentans* and analyzed for base composition. It was found that 37 percent of the bases are cytosine. With this information, is it possible to predict what percentage of the bases are adenine? If so, what percentage? If not, why not?
- **9.13** If one strand of DNA in the Watson–Crick double helix has a base sequence of 5'-GTCATGAC-3', what is the base sequence of the complementary strand?

- **9.14** Indicate whether each of the following statements about the structure of DNA is true or false. (Each letter is used to refer to the concentration of that base in DNA.)
 - (a) A + T = G + C(b) A = G; C = T
 - (c) A/T = C/G
 - (d) T/A = C/G
 - (e) A + G = C + T
 - (f) G/C = 1
 - (g) A = T within each single strand.
 - (h) Hydrogen bonding provides stability to the double helix in aqueous cytoplasms.
 - (i) Hydrophobic bonding provides stability to the double helix in aqueous cytoplasms.
 - (j) When separated, the two strands of a double helix are identical.
 - (k) Once the base sequence of one strand of a DNA double helix is known, the base sequence of the second strand can be deduced.
 - (l) The structure of a DNA double helix is invariant.
 - (m) Each nucleotide pair contains two phosphate groups, two deoxyribose molecules, and two bases.
- **9.15** The nucleic acids from various viruses were extracted and examined to determine their base composition. Given the following results, what can you hypothesize about the physical nature of the nucleic acids from these viruses?
 - (a) 35% A, 35% T, 15% G, and 15% C
 (b) 35% A, 15% T, 25% G, and 25% C
 - (c) 35% A, 30% U, 30% G, and 5% C
- **9.16** Compare and contrast the structures of the A, B, and Z forms of DNA.
- **9.17** The temperature at which one-half of a double-stranded DNA molecule has been denatured is called the melting temperature, T_m . Why does T_m depend directly on the GC content of the DNA?
- **9.18 ••** A diploid rye plant, *Secale cereale*, has 2n = 14 chromosomes and approximately 1.6×10^{10} bp of DNA. How much DNA is in a nucleus of a rye cell at (a) mitotic metaphase, (b) meiotic metaphase I, (c) mitotic telophase, and (d) meiotic telophase II?
- **9.19** The available evidence indicates that each eukaryotic chromosome (excluding polytene chromosomes) contains a single giant molecule of DNA. What different levels of organization of this DNA molecule are apparent in chromosomes of eukaryotes at various times during the cell cycle?
- 9.20 00 A diploid nucleus of *Drosophila melanogaster* contains about 3.4×10^8 nucleotide pairs. Assume (1) that all nuclear DNA is packaged in nucleosomes and (2) that an average internucleosome linker size is 60 nucleotide pairs. How many nucleosomes would be present in a diploid nucleus

of *D. melanogaster*? How many molecules of histone H2a, H2b, H3, and H4 would be required?

- **9.21** The relationship between the melting T_m and GC content can be expressed, in its much simplified form, by the formula $T_m = 69 + 0.41$ (% GC). (a) Calculate the melting temperature of *E. coli* DNA that has about 50% GC. (b) Estimate the % GC of DNA from a human kidney cell where $T_m = 85^{\circ}$ C.
- **9.22** Experimental evidence indicates that most highly repetitive DNA sequences in the chromosomes of eukaryotes do not produce any RNA or protein products. What does this indicate about the function of highly repetitive DNA?
- **9.23** The satellite DNAs of *Drosophila virilis* can be isolated, essentially free of main-band DNA, by density-gradient centrifugation. If these satellite DNAs are sheared into approximately 40-nucleotide-pair-long fragments and are analyzed in denaturation-renaturation experiments, how would you expect their hybridization kinetics to compare with the renaturation kinetics observed using similarly sheared main-band DNA under the same conditions? Why?
- **9.24** (a) What functions do (1) centromeres, (2) telomeres provide? (b) Do telomeres have any unique structural features? (c) What is the function of telomerase? (d) When chromosomes are broken by exposure to high-energy radiation such as X rays, the resulting broken ends exhibit a pronounced tendency to stick to each other and fuse. Why might this occur?

- **9.25** Are eukaryotic chromosomes metabolically most active during prophase, metaphase, anaphase, telophase, or interphase?
- **9.26** Are the scaffolds of eukaryotic chromosomes composed of histone or nonhistone chromosomal proteins? How has this been determined experimentally?
- **9.27** (a) Which class of chromosomal proteins, histones or nonhistones, is the more highly conserved in different eukaryotic species? Why might this difference be expected? (b) If one compares the histone and nonhistone chromosomal proteins of chromatin isolated from different tissues or cell types of a given eukaryotic organism, which class of proteins will exhibit the greater heterogeneity? Why are both classes of proteins not expected to be equally homogeneous in chromosomes from different tissues or cell types?
- **9.28** (a) If the haploid human genome contains 3×10^{9} nucleotide pairs and the average molecular weight of a nucleotide pair is 660, how many copies of the human genome are present, on average, in 1 mg of human DNA? (b) What is the weight of one copy of the human genome? (c) If the haploid genome of the small plant *Arabidopsis thaliana* contains 7.7×10^{7} nucleotide pairs, how many copies of the *A. thaliana* genome are present, on average, in 1 mg of *A. thaliana* DNA? (d) What is the weight of one copy of the *A. thaliana* genome? (e) Of what importance are calculations of the above type to geneticists?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

The available evidence indicates that each eukaryotic chromosome contains one giant DNA double helix running from one end through the centromere all the way to the other end. Of course, those DNA molecules are highly condensed into nucleosomes, 30-nm fibers, and higher order folding or coiling. Human cells contain 46 chromosomes. How large is the DNA molecule in the largest human chromosome?

- 1. Which human chromosome contains the largest DNA molecule? How large is it? How many genes does it contain?
- 2. Which human chromosome contains the smallest DNA molecule? How many base pairs does it contain? How many genes?
- **3.** Which human chromosomes contain genes encoding H1 histones? Other histone genes? How many histone genes are present in the human genome?

Hint: At the NCBI web site, Map Viewer \rightarrow *Homo sapiens* genome viewer (click on largest and smallest chromosomes shown) \rightarrow Search (Question 3).

10

Replication of DNA and Chromosomes

CHAPTER OUTLINE

- Basic Features of DNA Replication In Vivo
- DNA Replication in Prokaryotes
- Unique Aspects of Eukaryotic Chromosome Replication

Monozygotic Twins: Are They Identical?

From the day of their birth, through childhood, adolescence, and adulthood, Merry and Sherry have been mistaken for one another. When they are apart, Merry is called Sherry about half of the time, and Sherry is misidentified as Merry with equal frequency. Even their parents have trouble distinguishing them. Merry and Sherry are monozygotic ("identical") twins; they both developed from a single fertilized egg. At an early cleavage stage, the embryo split into two cell masses, and both groups of cells developed into complete embryos. Both embryos developed normally, and on April 7, 1955, identical. If the human genome contains about 20,500 genes, are the progeny replicas of all these genes exactly the same in identical twins?

A human life emerges from a single cell, a tiny sphere about 0.1 mm in diameter. That cell gives rise to hundreds of billions of cells during fetal development. At maturity, a human of average size contains about 65 trillion (65,000,000,000,000) cells. With some exceptions, each of these 65 trillion cells contains a progeny replica of each of the about 20,500 genes. Moreover, the cells of the body are not static; in some tissues, old cells are continuously being replaced by new cells. For example, in a healthy individual, the bone marrow cells produce about 2 million red

one newborn was named Merry, the other Sherry.

People often explain nearly identical phenotypes of monozygotic twins like Merry and Sherry by stating that "they contain the same genes." Of course, that is not true. To be accurate. the statement should be that identical twins contain progeny replicas of the same parental genes. But this simple colloquialism suggests that most people do, indeed, believe that the progeny replicas of a gene actually are



Four pairs of twins with their mothers at the Iowa State Fair.

blood cells per minute. Although not all of the progeny replicas of genes in the human body are identical, the process by which these genes are duplicated is very accurate. The human haploid genome contains about 3×10^{9} nucleotide pairs of DNA, all of which must be duplicated during each cell division. In this chapter, we examine how DNA replicates, and we focus on the mechanisms that ensure the fidelity of this process.

Basic Features of DNA Replication In Vivo

In humans, the synthesis of a new strand of DNA occurs at the rate of about 3000 nucleotides per minute. In bacteria, about 30,000 nucleotides are added to a nascent DNA chain per minute. Clearly, the cellular machinery responsible for DNA replication must work very fast, but, even more importantly, DNA replication occurs semiconservatively, is initiated at unique origins, and usually proceeds bidirectionally from each origin of replication.

it must work with great accuracy. Indeed, the fidelity of DNA replication is amazing, with an average of only one mistake per billion nucleotides incorporated after synthesis and the correction of mistakes during and immediately after replication. Thus, the majority of the genes of identical twins are indeed identical, but some will have changed owing to replication errors and other types of mutations (Chapter 13). Most of the key features of the mechanism by which the rapid and accurate replication of DNA occurs are now known, although many molecular details remain to be elucidated.

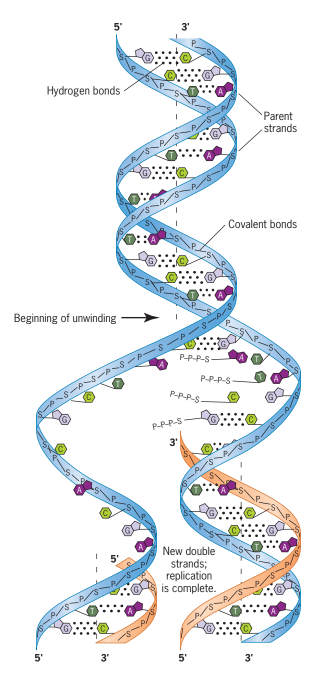
The synthesis of DNA, like the synthesis of RNA (Chapter 11) and proteins (Chapter 12), involves three steps: (1) chain initiation, (2) chain extension or elongation, and (3) chain termination. In this and the following two chapters, we examine the mechanisms by which cells carry out each of the three steps in the synthesis of these important macromolecules. First, however, we consider some key features of DNA replication.

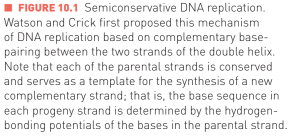
SEMICONSERVATIVE REPLICATION

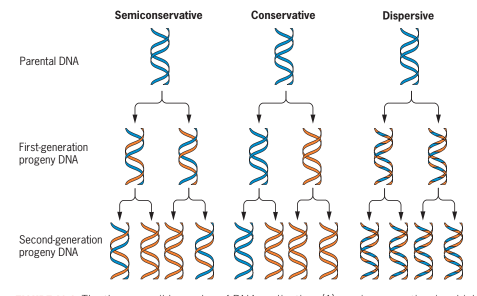
When Watson and Crick deduced the double-helix structure of DNA with its complementary base-pairing, they immediately recognized that the basepairing specificity could provide the basis for a simple mechanism for DNA duplication. Therefore, five weeks after the appearance of their paper on the double-helix structure of DNA, Watson and Crick published a paper describing a mechanism by which the double helix could replicate. They proposed that the two complementary strands of the double helix unwind and separate, and that each strand guides the synthesis of a new complementary strand (**Figure 10.1**). The sequence of bases in each parental strand is used as a template, and the base-pairing restrictions within the double helix dictate the sequence of bases in the newly synthesized strand. Adenine, for example, in the parent strand will serve as a template via its hydrogen-bonding potential for the incorporation of thymine in the nascent complementary strand. This mechanism of DNA replication is called semiconservative replication (because the parental molecule is half conserved) to distinguish it from other possible mechanisms of replication (**Figure 10.2**). In conservative replication, the parental double helix would be conserved, and a new progeny double helix would be synthesized. In dispersive replication, segments of both strands of the parental DNA molecule would be conserved and used as templates for the synthesis of complementary segments that would subsequently be joined to produce progeny DNA strands.

In 1958, Matthew Meselson and Franklin Stahl demonstrated that the chromosome of *Escherichia coli* replicates semiconservatively. Then, in 1962, John Cairns demonstrated that the *E. coli* chromosome was a single duplex of DNA. Together, the results presented by Cairns and Meselson and Stahl showed that DNA replicates semiconservatively in *E. coli*.

Meselson and Stahl grew *E. coli* cells for many generations in a medium in which the heavy isotope of nitrogen, ¹⁵N, had been substituted for the normal, light isotope, ¹⁴N. The purine and pyrimidine bases in DNA contain nitrogen. Thus, the DNA of cells grown on medium containing ¹⁵N will have a greater density (mass per unit volume) than the DNA of cells grown on medium containing ¹⁴N. Molecules with different densities can be separated







■ FIGURE 10.2 The three possible modes of DNA replication: (1) semiconservative, in which each strand of the parental double helix is conserved and directs the synthesis of a new complementary progeny strand; (2) conservative, in which the parental double helix is conserved and directs the synthesis of a new progeny double helix; and (3) dispersive, in which segments of each parental strand are conserved and direct the synthesis of new complementary strand segments that are subsequently joined to produce new progeny strands.

by **equilibrium density-gradient centrifugation.** By using this technique, Meselson and Stahl were able to distinguish between the three possible modes of DNA replication by following the changes in the density of DNA from cells grown on ¹⁵N medium and then transferred to ¹⁴N medium for various periods of time—so-called density-transfer experiments.

The density of most DNAs is about the same as the density of concentrated solutions of heavy salts such as cesium chloride (CsCl). For example, the density of 6M CsCl is about 1.7 g/cm³. *E. coli* DNA containing ¹⁴N has a density of 1.710 g/cm³. Substitution of ¹⁵N for ¹⁴N increases the density of *E. coli* DNA to 1.724 g/cm³. When a 6M CsCl solution is centrifuged at very high speeds for long periods of time, an equilibrium density gradient is formed (**■** Figure 10.3). If DNA is present in such a gradient, it will move to a position where the density of the CsCl solution is equal to its own density. Thus, if a mixture of *E. coli* DNA containing the heavy isotope of nitrogen, ¹⁵N, and *E. coli* DNA containing the normal light nitrogen isotope, ¹⁴N, is subjected to CsCl equilibrium density-gradient centrifugation, the DNA molecules will separate into two "bands," one consisting of "heavy" (¹⁵N-containing) DNA and the other of "light" (¹⁴N-containing) DNA.

Meselson and Stahl took cells that had been growing in medium containing ¹⁵N for several generations (and thus contained "heavy" DNA), washed them to remove the medium containing ¹⁵N, and transferred them to medium containing ¹⁴N. After the cells were allowed to grow in the presence of ¹⁴N for varying periods of time, the DNAs were extracted and analyzed in CsCl equilibrium density gradients. The results of their experiment (**■** Figure 10.4) are consistent only with semiconservative replication, excluding both conservative and dispersive models of DNA synthesis. All the DNA isolated from cells after one generation of growth in medium containing ¹⁴N had a density halfway between the densities of "heavy" DNA and "light" DNA. This intermediate density is usually referred to as "hybrid" density. After two generations of growth in medium containing ¹⁴N, half of the DNA was of hybrid density and half was light. These results are precisely those predicted by the Watson and Crick semiconservative mode of replication (see Figure 10.2). One generation

of semiconservative replication of a parental double helix containing ¹⁵N in medium containing only ¹⁴N would produce two progeny double helices, both of which have ¹⁵N in one strand (the "old" strand) and ¹⁴N in the other strand (the "new" strand). Such molecules would be of hybrid density.

Conservative replication would not produce any DNA molecules with hybrid density; after one generation of conservative replication of heavy DNA in light medium, half of the DNA still would be heavy and the other half would be light. If replication were dispersive, Meselson and Stahl would have observed a shift of the DNA from heavy toward light in each generation (that is, "half heavy" or hybrid after one generation, "quarter heavy" after two generations, and so forth). These possibilities are clearly inconsistent with the results of Meselson and Stahl's experiment. DNA replication was subsequently shown to occur semiconservatively in several other microorganisms. Try Solve It: Understanding the Semiconservative Replication of DNA to test your comprehension of the significance of Meselson and Stahl's results.

The semiconservative replication of eukaryotic chromosomes was first demonstrated in 1957 by the results of experiments carried out by J. Herbert Taylor, Philip Woods, and Walter Hughes on root-tip cells of the broad bean, Vicia faba. Taylor and colleagues labeled Vicia faba chromosomes by growing root tips for eight hours (less than one cell generation) in medium containing radioactive ³H-thymidine. The root tips were then removed from the radioactive medium, washed, and transferred to nonradioactive medium containing the alkaloid colchicine. It is known that colchicine binds to microtubules and prevents the formation of functional spindle fibers. As a result, daughter chromosomes do not undergo their normal anaphase separation. Thus, the number of chromosomes per nucleus will double once per cell cycle in the presence of colchicine. This doubling of the chromosome number each cell generation allowed Taylor and his colleagues to determine how many DNA duplications each cell had undergone subsequent to the incorporation of radioactive thymidine. At the first metaphase in colchicine (c-metaphase), nuclei will contain 12 pairs of chromatids (still joined at the centromeres). At the second c-metaphase, nuclei will contain 24 pairs, and so on.

Taylor and colleagues used a technique called autoradiography to examine the distribution of radioactivity in the chromosomes of cells at the first c-metaphase, the second c-metaphase, and so on. Autoradiography is a method for detecting and localizing radioactive isotopes in cytological preparations or macromolecules by exposure to a photographic emulsion that is sensitive to low-energy radiation. The emulsion contains silver halides that produce tiny black spots-often called silver grains—when they are exposed to the charged particles emitted during the decay of radioactive isotopes. Autoradiography permits a researcher to prepare an image of the localization of radioactivity in macromolecules, cells, or tissues, just as photography permits us to make a picture of what we see. The difference is that the film used for autoradiography is sensitive to radioactivity, whereas the film we use in a camera is sensitive to visible light. Autoradiography is particularly useful in studying DNA metabolism because DNA can be specifically labeled by growing cells on ³H-thymidine, a deoxyribonucleoside of thymine that contains a radioactive isotope of hydrogen (tritium). Thymidine is incorporated almost exclusively into DNA; it is not present in any other major component of the cell.

When Taylor and coworkers used autoradiography to examine the distribution of radioactivity in the *Vicia faba* chromosomes at the first c-metaphase, both

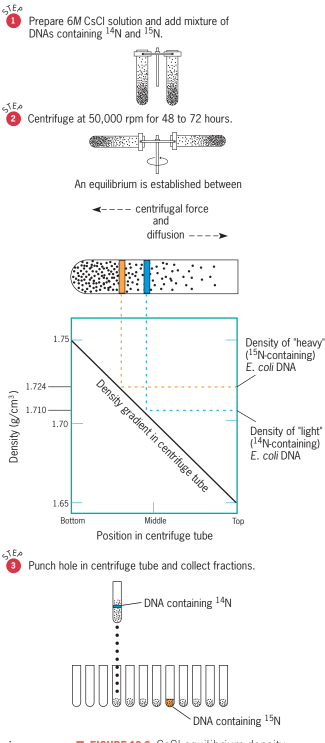
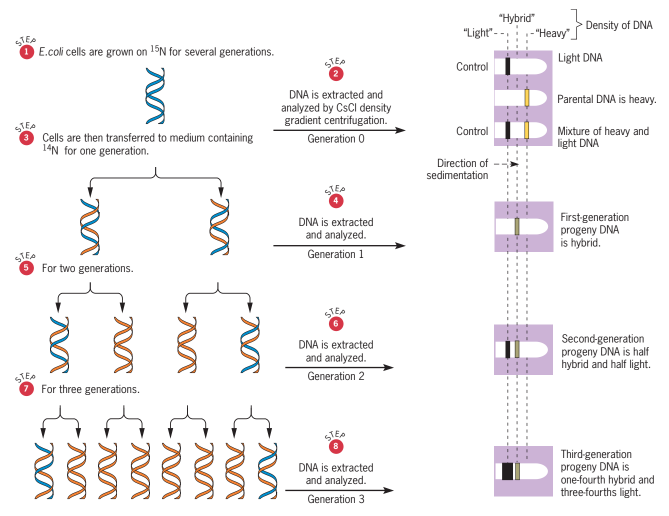
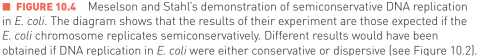


FIGURE 10.3 CsCl equilibrium densitygradient centrifugation.





Solve It!

Understanding the Semiconservative Replication of DNA

A culture of bacteria is grown for many generations in a medium in which the only available nitrogen is the heavy isotope (¹⁵N). The culture is then switched to a medium containing only ¹⁴N for one generation of growth; it is then returned to a ¹⁵N-containing medium for one final generation of growth. If the DNA from these bacteria is isolated and centrifuged to equilibrium in a CsCl density gradient, how would you predict the DNA to band in the gradient?

To see the solution to this problem, visit the Student Companion site. chromatids of each pair were radioactive (**Tigure 10.5a**). However, at the second c-metaphase, only one of the chromatids of each pair was radioactive (**Tigure 10.5b**). These are precisely the results expected if DNA replication is semiconservative, given one DNA molecule per chromosome (**Tigure 10.5c**). In 1957, Taylor and his colleagues were able to conclude that chromosomal DNA in *Vicia faba* segregated in a semiconservative manner during each cell division. The conclusion that the double helix replicated semiconservatively in the broad bean had to await subsequent evidence indicating that each chromosome contains a single molecule of DNA. Analogous experiments have subsequently been carried out with several other eukaryotes, and, in all cases, the results indicate that replication is semiconservative. Test your understanding of chromosome replication by working the problem in Problem-Solving Skills: Predicting Patterns of ³H Labeling in Chromosomes.

UNIQUE ORIGINS OF REPLICATION

John Cairns established the existence of a site of initiation or **origin of replication** on the circular chromosome of *E. coli* but provided no hint as to whether the origin was a unique site or occurred at randomly located sites in a population of replicating chromosomes. In bacterial and viral chromosomes, there is usually one unique

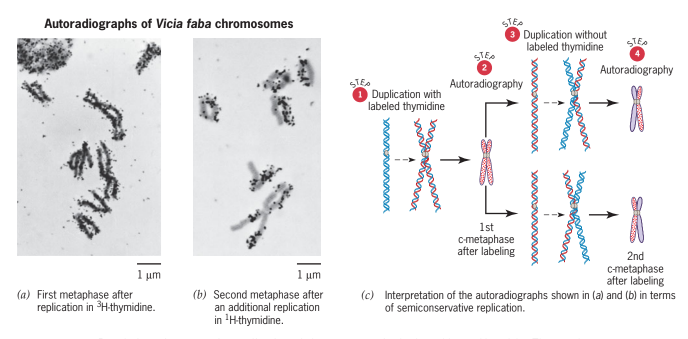


FIGURE 10.5 Proof of semiconservative replication of chromosomes in the broad bean, *Vicia faba*. The results obtained by Taylor, Woods, and Hughes (*a*, *b*) are predicted by the semiconservative replication of the DNA (*c*).

origin per chromosome, and this single origin controls the replication of the entire chromosome. In the large chromosomes of eukaryotes, multiple origins collectively control the replication of the giant DNA molecule present in each chromosome. Current evidence indicates that these multiple replication origins in eukaryotic chromosomes also occur at specific sites. Each origin controls the replication of a unit of DNA called a *replicon;* thus, most prokaryotic chromosomes contain a single replicon, whereas eukaryotic chromosomes usually contain many replicons.

The single origin of replication, called *oriC*, in the *E. coli* chromosome has been characterized in considerable detail. *oriC* is 245 nucleotide pairs long and contains two different conserved repeat sequences (**Figure 10.6**). One 13-bp sequence is present as three tandem repeats. These three repeats are rich in A:T base pairs, facilitating the formation of a localized region of strand separation referred to as the **replication bubble**. Recall that A:T base pairs are held together by only two hydrogen bonds as opposed to three in G:C base pairs (Chapter 9). Thus, the two strands of AT-rich regions of DNA come apart more easily, that is, with the input of less energy. The formation of a localized zone of denaturation is an essential first step in the replication of all double-stranded DNAs. Another conserved component of *oriC* is a 9-bp sequence that is repeated four times and is interspersed with other sequences. These four sequences are binding sites for a protein that plays a key role in the formation of the replication of DNA synthesis at origins and the proteins that are involved.

The multiple origins of replication in eukaryotic chromosomes also appear to be specific DNA sequences. In the yeast *Saccharomyces cerevisiae*, segments of chromosomal DNA that allow a fragment of circularized DNA to replicate as an independent unit (autonomously), that is, as an extrachromosomal self-replicating unit, have been identified and characterized. These sequences are called *ARS* (for *A*utonomously *Replicating Sequences*) *elements*. Their frequency in the yeast genome corresponds well with the number of origins of replication, and some have been shown experimentally to function as origins. ARS elements are about 50 base pairs in length and include a core 11-bp AT-rich sequence,

ATTTATPuTTTA TAAATAPyAAAT

PROBLEM-SOLVING SKILLS



Predicting Patterns of ³H Labeling in Chromosomes

THE PROBLEM

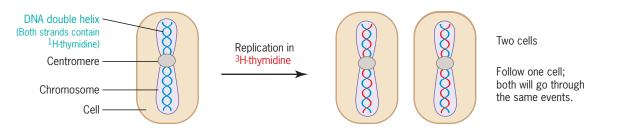
Haplopappus gracilis is a diploid plant with two pairs of chromosomes (2n = 4). A G₁-stage cell of this plant, not previously exposed to radioactivity, was placed in culture medium containing ³H-thymidine. After one generation of growth in this medium, the two progeny cells were washed with nonradioactive medium and transferred to medium containing ¹H-thymidine and colchicine. They were allowed to grow in this medium for one additional cell generation and on to metaphase of a second cell division. The chromosomes from each cell were then spread on a microscope slide, stained, photographed, and exposed to an emulsion sensitive to low-energy radiation. One of the daughter cells exhibited a metaphase plate with eight chromosomes, each with two daughter chromatids. Draw this metaphase plate showing the predicted distribution of radioactivity on the autoradiograph. Assume no crossing over!

FACTS AND CONCEPTS

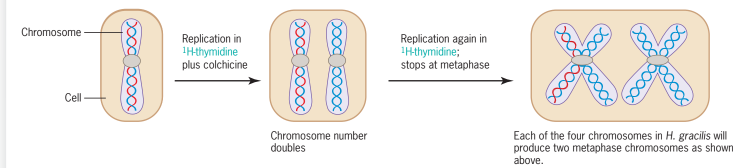
- **1.** Each G₁-stage (prereplicative) chromosome contains a single DNA double helix.
- **2.** DNA replication is semiconservative.
- **3.** Daughter chromatids remain attached to a single centromere at metaphase of mitosis.
- **4.** The centromere duplicates prior to anaphase; at that time, each chromatid becomes a daughter chromosome.
- **5.** Colchicine binds to the proteins that form the spindle fibers responsible for the separation of daughter chromosomes to the spindle poles during anaphase and prevents the formation of functional spindles. As a result, chromosome number doubles during each cell generation in the presence of colchicine.

ANALYSIS AND SOLUTION

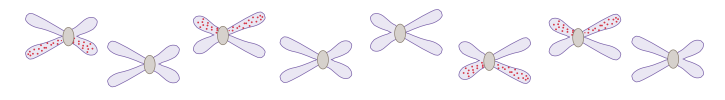
All four chromosomes will go through the same replication events. Therefore, we only need to follow one chromosome. The first replication in the presence of ³H-thymidine, but no colchicine, is shown in the following diagram with radioactive strands in red.



The second and third replications (in ¹H-thymidine and colchicine) are shown in the following diagram.



When the resulting metaphase chromosomes are subjected to autoradiography, the distribution of radioactivity (indicated by red dots) on the eight chromosomes will be as follows.



For further discussion visit the Student Companion site.

(where Pu is either of the two purines and Py is either of the two pyrimidines) and additional imperfect copies of this sequence. The ability of ARS elements to function as origins of replication is abolished by basepair changes within this conserved core sequence.

Attempts to characterize origins of replication in multicellular eukaryotes have been largely unsuccessful. Despite evidence that replication is initiated at specific sequences *in vivo* and the availability of the sequences of entire genomes, the components of a functional origin have remained elusive. There appear to be two major reasons for this failure to identify replication origins. First, the functional assays used in yeast—the ability of the origin to support the replication of a plasmid or artificial chromosome—do not yield reliable results in other eukaryotes. Sequences that support the replication of plasmids in mammalian cells, for example, often result in the initiation of replication at random or multiple sites. Second, considerable evidence now suggests that the initiation of replication involves relatively long DNA sequences—up to several thousand base pairs—in metazoans, making their origins difficult to characterize.

VISUALIZATION OF REPLICATION FORKS BY AUTORADIOGRAPHY

The gross structure of replicating bacterial chromosomes was first determined by John Cairns in 1963, again by means of autoradiography. Cairns grew E. coli cells in medium containing 3H-thymidine for varying periods of time, lysed the cells gently so as not to break the chromosomes (long DNA molecules are sensitive to shearing), and carefully collected the chromosomes on membrane filters. These filters were affixed to glass slides, coated with emulsion sensitive to β-particles (the low-energy electrons emitted during decay of tritium), and stored in the dark for a period of time to allow sufficient radioactive decay. When the films were developed, the autoradiographs (**Figure 10.7***a*) showed that the chromosomes of *E. coli* are circular structures that exist as θ -shaped intermediates during replication. The autoradiographs further indicated that the unwinding of the two complementary parental strands (which is necessary for their separation) and their semiconservative replication occur simultaneously or are closely coupled. Since the parental double helix must rotate 360° to unwind each gyre of the helix, some kind of "swivel" must exist. Geneticists now know that the required swivel is a transient single-

strand break (cleavage of one phosphodiester bond in one strand of the double helix) produced by the action of enzymes called topoisomerases.

Replication of the *E. coli* chromosome occurs bidirectionally from the unique origin of replication. Each Y-shaped structure is a **replication fork**, and the two replication forks move in opposite directions sequentially around the circular chromosome (**Figure 10.7***b*).

The bidirectional replication of the circular *E. coli* chromosome just discussed occurs during cell division. It should not be confused with rollingcircle replication, which mediates the

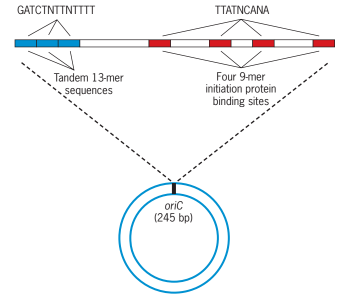
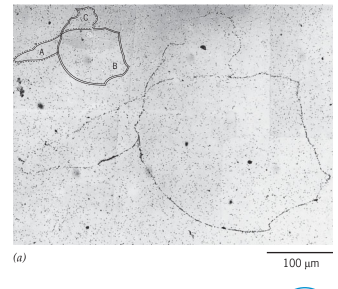
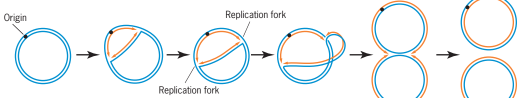


FIGURE 10.6 Structure of *oriC*, the single origin of replication in the *E. coli* chromosome.





(b) Bidirectional replication of the circular E. coli chromosome.

FIGURE 10.7 Visualization of the replication of the *E. coli* chromosome by autoradiography. (a) One of Cairns's autoradiographs of a θ -shaped replicating chromosome from a cell that had been grown for two generations in the presence of ³H-thymidine, with his interpretative diagram shown at the upper left. Radioactive strands of DNA are shown as solid lines and nonradioactive strands as dashed lines. Loops A and B have completed a second replication in ³H-thymidine; section C remains to be replicated the second time. (b) A diagram showing how Cairns's results are explained by bidirectional replication of the *E. coli* chromosome initiated at a unique origin of replication. Linear phage λ chromosome present in mature virions

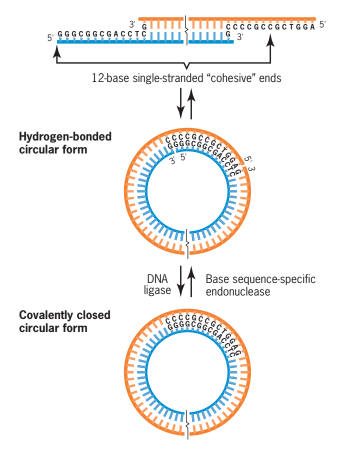


FIGURE 10.8 Three forms of the phage lambda chromosome. The conversions of the linear λ chromosome with its complementary cohesive ends to the hydrogen-bonded circular λ chromosome and then to the covalently closed circular λ chromosome are shown. The linear form of the chromosome appears to be an adaptation to facilitate its injection from the phage head through the small opening in the phage tail into the host cell during infection. Prior to replicating in the host cell, the chromosome is converted to the covalently closed circular form. Only the ends of the chromosome of the mature phage are shown; the jagged vertical line indicates that the central portion of the chromosome is 48,502 nucleotide pairs long.

transfer of chromosomes from Hfr cells to F^- cells (Chapter 8). Some viral chromosomes replicate by the rolling-circle mechanism; see the section Rolling-Circle Replication later in this chapter.

BIDIRECTIONAL REPLICATION

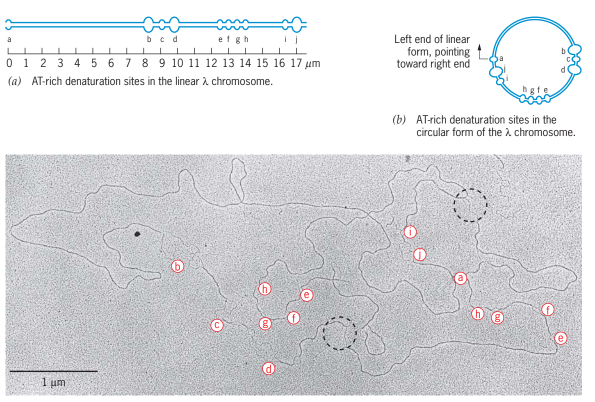
Bidirectional replication was first convincingly demonstrated by experiments with some of the small bacterial viruses that infect E. coli. Bacteriophage lambda (phage λ) contains a single linear molecule of DNA only 17.5 μ m long. The phage λ chromosome is somewhat unusual in that it has a single-stranded region, 12 nucleotides long, at the 5' end of each complementary strand (**Figure 10.8**). These single-stranded ends, called "cohesive" or "sticky" ends, are complementary to each other. The cohesive ends of a lambda chromosome can thus base-pair to form a hydrogen-bonded circular structure. One of the first events to occur after a lambda chromosome is injected into a host cell is its conversion to a covalently closed circular molecule (Figure 10.8). This conversion from the hydrogen-bonded circular form to the covalently closed circular form is catalyzed by DNA ligase, an important enzyme that seals single-strand breaks in DNA double helices. DNA ligase is required in all organisms for DNA replication, DNA repair, and recombination between DNA molecules. Like the E. coli chromosome, the lambda chromosome replicates in its circular form via θ -shaped intermediates.

The feature of the lambda chromosome that facilitated the demonstration of bidirectional replication is its differentiation into regions containing high concentrations of adenine and thymine (AT-rich regions) and regions with large amounts of guanine and cytosine (GC-rich regions). In particular, it contains a few segments with high AT content (AT-rich clusters). In the late 1960s, Maria Schnös and Ross Inman used these AT-rich clusters as physical markers to demonstrate, by means of a technique called denaturation mapping, that replication of the lambda chromosome is initiated at a unique origin and proceeds bidirectionally rather than unidirectionally.

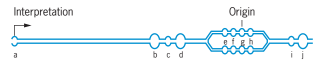
When DNA molecules are exposed to high temperature (100°C) or high pH (11.4), the hydrogen and hydrophobic bonds that hold the complementary strands together in the double-helix configuration are broken, and the two strands separate—a process called denaturation. Because AT base pairs are held together by only two hydrogen bonds, compared with three hydrogen bonds in GC base pairs, AT-rich molecules denature more easily (at lower pH or temperature) than GC-rich molecules. When lambda chromosomes are exposed to pH 11.05 for 10 minutes under the appropriate conditions, the AT-rich clusters denature to form denaturation bubbles, which are detectable by electron microscopy, whereas the GC-rich regions

remain in the duplex state (**Figure 10.9**). These denaturation bubbles can be used as physical markers whether the lambda chromosome is in its mature linear form, its circular form, or its θ -shaped replicative intermediates. By examining the positions of the branch points (Y-shaped structures) relative to the positions of the denaturation bubbles in a large number of θ -shaped replicative intermediates, Schnös and Inman demonstrated that both branch points are replication forks that move in opposite directions around the circular chromosome. **Figure 10.10** shows the results expected in Schnös and Inman's experiment if replication is (*a*) unidirectional or (*b*) bidirectional. The results clearly demonstrated that replication of the lambda chromosome is bidirectional.

Bidirectional replication from a fixed origin has also been demonstrated for several organisms with chromosomes that replicate as linear structures. Replication of the chromosome of phage T7, another small bacteriophage, begins at a unique site near one end to form an "eye" structure (**Figure 10.11***a*) and then proceeds bidirectionally until one fork reaches the nearest end. Replication of the Y-shaped



(c) AT-rich denaturation bubbles in a θ -shaped replicating λ chromosome.



(d) Diagram in linear form of the λ replicative intermediate shown in (c).

FIGURE 10.9 The use of AT-rich denaturation sites as physical markers to prove that the phage λ chromosome replicates bidirectionally rather than unidirectionally. The positions of the AT-rich denaturation bubbles are shown for the linear (a) and circular (b) forms of the λ chromosome. The electron micrograph (c) shows the positions of denaturation bubbles (labeled a-j) and replication forks (circled) in a partially replicated λ chromosome. The structure of the partially replicated chromosome in (c) is diagrammed in (d).

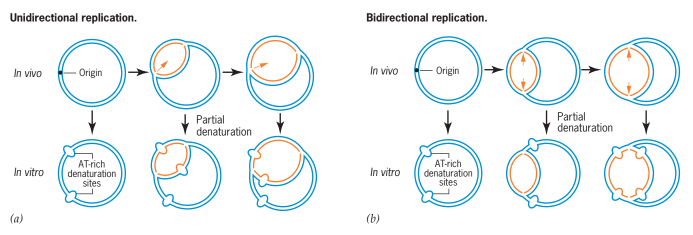
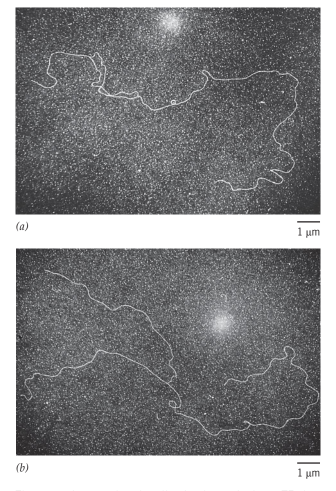


FIGURE 10.10 Rationale of the denaturation mapping procedure used by Schnös and Inman to distinguish between (a) unidirectional and (b) bidirectional modes of chromosome replication.



■ FIGURE 10.11 Electron micrographs of replicating bacteriophage T7 chromosomes. The T7 chromosomes, unlike the *E. coli* and lambda chromosomes, replicates as a linear structure. Its origin of replication is located 17 percent from the left end of the chromosome. The chromosome in (a) illustrates the "eye" (->-) form characteristic of the early stages of replication. Parental strand separation and DNA synthesis proceed bidirectionally outward from the origin. When the leftward moving fork reaches the left end of the chromosome, a Y-shaped structure results, such as the one shown in (b). Replication continues with the remaining rightward fork until two linear chromosomes are produced. For chromosomes much larger than T7, such as eukaryotic chromosomes, replication occurs from multiple origins, giving rise to numerous simultaneously growing 'eyes'. *Original micrographs courtesy of David Dressler, Harvard University.*

structure (**Figure 10.11***b*) continues until the second fork reaches the other end of the molecule, producing two progeny chromosomes.

Replication of chromosomal DNA in eukaryotes is also bidirectional in those cases where it has been investigated. However, bidirectional replication is not universal. The chromosome of coliphage P2, which replicates as a θ -shaped structure like the lambda chromosome, replicates unidirectionally from a unique origin.

KEY POINTS

• DNA replicates by a semiconservative mechanism: as the two complementary strands of a parental double helix unwind and separate, each serves as a template for the synthesis of a new complementary strand.

- The hydrogen-bonding potentials of the bases in the template strands specify complementary base sequences in the nascent DNA strands.
- Replication is initiated at unique origins and usually proceeds bidirectionally from each origin.

DNA Replication in Prokaryotes

The results of studies of DNA replication by autoradiography and electron microscopy indicate that the two progeny strands being synthesized at each replicating fork are being extended in the same over-

all direction at the macromolecular level. Because the complementary strands of a double helix have opposite polarity, synthesis is occurring at the 5' end of one strand $(3' \rightarrow 5' \text{ extension})$ and the 3' end of the other strand $(5' \rightarrow 3' \text{ extension})$. However, the enzymes that catalyze the replication of DNA, **DNA polymerases** (see Focus on DNA Synthesis *In Vitro*), have an absolute requirement for a free 3'-hydroxyl; they only carry out $5' \rightarrow 3'$ synthesis (**True Figure 10.12**). These apparently contradictory results created an interesting paradox. For many years biochemists searched for new polymerases that could catalyze $3' \rightarrow 5'$ synthesis. No such polymerase was ever found. Instead, experimental evidence has shown that all DNA synthesis occurs in the $5' \rightarrow 3'$ direction.

Clearly, the mechanism of DNA replication must be more complex than researchers originally thought (see Figure 10.1). Given the absolute requirement of

FOCUS ON

DNA SYNTHESIS IN VITRO

Involved in biological processes by disrupting cells, separating the various organelles, macromolecules, and other components, and then reconstituting systems in the test tube, so-called *in vitro* systems that are capable of carrying out particular metabolic events. Such *in vitro* systems can be dissected biochemically much more easily than *in vivo* systems, and, therefore, they have contributed immensely to our understanding of biological processes. However, we should never assume that a phenomenon demonstrated *in vitro* occurs *in vivo*. Such an extrapolation should be made only when independent evidence from *in vivo* studies validates the results of the *in vitro* studies.

DNA replication is one area where *in vitro* studies have proven, and continue to prove, invaluable. Much of our knowledge about the process of DNA replication was initially deduced from such studies. In 1957, Arthur Kornberg and his coworkers were the first to demonstrate that DNA synthesis could occur *in vitro*. Kornberg received a Nobel Prize for this work just two years later (1959), which demonstrated just how important other scientists considered this breakthrough. Kornberg and colleagues isolated an enzyme from *E. coli* that catalyzes the covalent addition of nucleotides to preexisting DNA chains. Initially called DNA polymerase or "Kornberg's enzyme," this enzyme is now known as **DNA polymerase I** because of the subsequent discovery of several other DNA polymerases in *E. coli*.

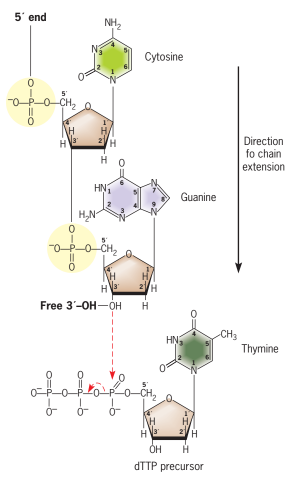
Over many years Kornberg and colleagues carried out extensive *in vitro* studies on the mechanism by which this enzyme catalyzed the synthesis of DNA, and much of what we know about DNA polymerases is based on their results. DNA polymerase I requires the 5'-triphosphates of each of the four deoxyribonucleosides deoxyadenosine triphosphate (dATP), deoxythymidine triphosphate (dTTP), deoxyguanosine triphosphate (dGTP), and deoxycytidine triphosphate (dCTP)—and is active only in the presence of Mg²⁺ ions and preexisting DNA. This DNA must be at least partially double-stranded and partially single-stranded, and must contain a free 3'-hydroxyl (3'-OH) group. The enzyme catalyzes the addition of nucleotides to the 3'-OH group of preexisting DNA strands. Therefore, it catalyzes the covalent extension of DNA chains in only the 5' \rightarrow 3' direction.

DNA replication is a complex process, requiring the

concerted action of a large number of proteins.

DNA polymerase I is a single polypeptide with a molecular weight of 103,000 encoded by a gene called *polA*. However, DNA polymerase I is not the true "DNA replicase" in *E. coli*. In 1969, Paula DeLucia and John Cairns reported that DNA replication occurred in an *E. coli* strain lacking the polymerase activity of DNA polymerase I due to a mutation in the *polA* gene. However, DeLucia and Cairns also discovered that this *polA1* mutant was extremely sensitive to ultraviolet light (UV). We now know that a major function of DNA polymerase I in *E. coli* is to repair defects in DNA, such as those induced by UV (Chapter 13). However, as we will see later in this chapter (Initiation of DNA Chains with RNA Primers), DNA polymerase I also plays an important role in chromosome replication.

Today, *in vitro* studies are being used to characterize DNA polymerases in many different organisms. Several recently discovered polymerases, called **translesion polymerases**, because they can replicate past lesions or defects in DNA that block replication by most polymerases, are proving especially interesting. In humans and other mammals, **DNA polymerase eta** (η) plays an important role in replicating damaged DNA. Individuals who are homozygous for loss-of-function mutations in the *POLH* gene (also called the *XPV* gene) that encodes polymerase η have one form of an inherited disorder called xeroderma pigmentosum (XP). Individuals with XP are extremely sensitive to sunlight; they develop multiple skin cancers after exposure to UV in sunlight (see Chapter 13).



■ FIGURE 10.12 Mechanism of action of DNA polymerases: covalent extension of a DNA primer strand in the 5' \rightarrow 3' direction. The existing chain terminates at the 3' end with the nucleotide deoxyguanylate (deoxyguanosine-5'-phosphate). The diagram shows the DNA polymerase-catalyzed addition of deoxythymidine monophosphate (from the precursor deoxythymidine triphosphate, dTTP) to the 3' end of the chain with the release of pyrophosphate (P₂O₇). DNA polymerases for a free 3'-OH on the primer strand, these enzymes cannot begin the synthesis of a new strand *de novo*. How is the synthesis of a new DNA strand initiated? Since the two parental strands of DNA must be unwound, we have to deal with the need for a swivel or an axis of rotation, especially for circular DNA molecules like the one present in the *E. coli* chromosome. Finally, how does the localized zone of strand separation or replication bubble form at the origin? These considerations and others indicate that DNA replication is more complicated than scientists thought when Watson and Crick proposed the semiconservative mechanism of replication in 1953.

CONTINUOUS SYNTHESIS OF ONE STRAND; DISCONTINUOUS SYNTHESIS OF THE OTHER STRAND

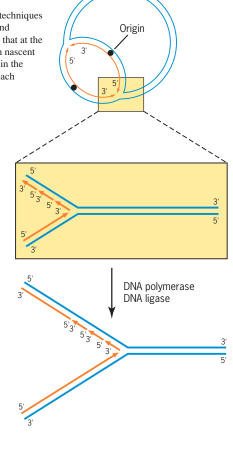
As previously discussed, the two nascent DNA strands being synthesized at each replicating fork are being extended in the same direction at the macromolecular level. Because the complementary strands of a DNA double helix have opposite chemical polarity, one strand is being extended in an overall $5' \rightarrow 3'$ direction and the other strand is being extended in an overall $3' \rightarrow 5'$ direction (**Figure 10.13a**). But DNA polymerases can only catalyze synthesis in the $5' \rightarrow 3'$ direction. This paradox was resolved with the demonstration that the synthesis of one strand of DNA is continuous, whereas synthesis of the other strand is discontinuous. At the molecular level, synthesis of the complementary strands of DNA is occurring in opposite physical directions (**Figure 10.13b**), but both new strands are extended in the same $5' \rightarrow 3'$ chemical direction. The synthesis of the strand being extended in the overall $5' \rightarrow 3'$ direction, called the leading strand, is continuous. The strand being extended in the overall $3' \rightarrow 5'$ direction, called the lagging strand, grows by the synthesis of short fragments (synthesized $5' \rightarrow 3'$) and the subsequent covalent joining of these short fragments. Thus, the synthesis of the lagging strand occurs by a discontinuous mechanism.

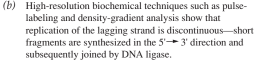
The first evidence for this discontinuous mode of DNA replication came from studies in which intermediates in DNA synthesis were radioactively labeled by growing E. coli cells and bacteriophage T4-infected E. coli cells for very short periods of time in medium containing ³H-thymidine (pulse-labeling experiments). The labeled DNAs were then isolated, denatured, and characterized by measuring their velocity of sedimentation through sucrose gradients during high-speed centrifugation. When E. coli cells were pulse-labeled for 5, 10, or 30 seconds, for example, much of the label was found in small fragments of DNA, 1000 to 2000 nucleotides long (**Figure 10.13c**). These small fragments of DNA have been named Okazaki fragments after Reiji Okazaki and Tuneko Okazaki, the scientists who discovered them in the late 1960s. In eukaryotes, the Okazaki fragments are only 100 to 200 nucleotides in length. When longer pulse-labeling periods are used, more of the label is recovered in large DNA molecules, presumably the size of E. coli or phage T4 chromosomes. If cells are pulse-labeled with ³H-thymidine for a short period and then are transferred to nonradioactive medium for an extended period of growth (pulse-chase experiments), the labeled thymidine is present in chromosome-size DNA molecules. The results of these pulse-chase experiments are important because they indicate that the Okazaki fragments are true intermediates in DNA replication and not some type of metabolic by-product.

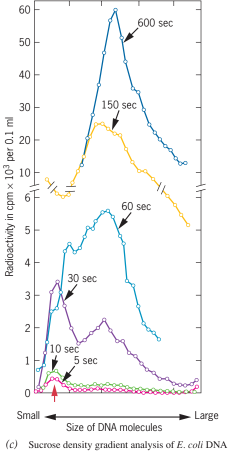
COVALENT CLOSURE OF NICKS IN DNA BY DNA LIGASE

If the lagging strand of DNA is synthesized discontinuously as described in the preceding section, a mechanism is needed to link the Okazaki fragments together to produce the large DNA strands present in mature chromosomes. This mechanism is provided by the enzyme **DNA ligase**. DNA ligase catalyzes the covalent closure of nicks (missing phosphodiester linkages; no missing bases) in DNA molecules by using energy from Relatively low-resolution techniques such as autoradiography and electron microscopy show that at the macromolecular level both nascent DNA chains are extended in the same overall direction at each replication fork.

(a)





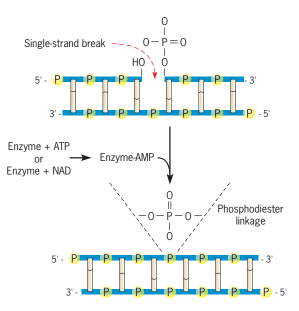


Sucrose density gradient analysis of E. coli DNA pulse-labeled with ³H-thymidine, extracted, and denatured during centrifugation. The gradients used separate DNA molecules based on size.

FIGURE 10.13 Evidence for discontinuous synthesis of the lagging strand. (a) Although both strands of nascent DNA synthesized at a replication fork appear to be extended in the same direction, (b) at the molecular level, they are actually being synthesized in opposite directions. (c) The results of pulse-labeling experiments of Reiji and Tuneko Okazaki and colleagues showing that nascent DNA in *E. coli* exists in short fragments 1000 to 2000 nucleotides long. The red arrow shows the position of the "Okazaki fragments" in the gradient.

nicotinamide adenine dinucleotide (NAD) or adenosine triphosphate (ATP). The *E. coli* DNA ligase uses NAD as a cofactor, but some DNA ligases use ATP. The reaction catalyzed by DNA ligase is shown in **Figure 10.14**. First, AMP of the ligase-AMP intermediate forms a phosphoester linkage with the 5'-phosphate at the nick, and then a nucleophilic attack by the 3'-OH at the nick on the DNA-proximal phosphorus atom produces a phosphodiester linkage between the adjacent nucleotides at the site of the nick. DNA ligase alone has no activity at breaks in DNA where one or more nucleotides are missing—so-called gaps. Gaps can be filled in and sealed only by the combined action of a DNA polymerase and DNA ligase. DNA ligase plays an essential role not only in DNA replication, but also in DNA repair and recombination (Chapter 13).

FIGURE 10.14 DNA ligase catalyzes the covalent closure of nicks in DNA. The energy required to form the ester linkage is provided by either adenosine triphosphate (ATP) or nicotinamide-adenine dinucleotide (NAD), depending on the species.



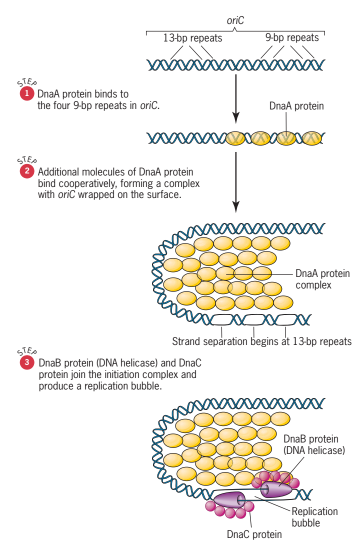
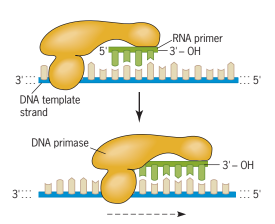


FIGURE 10.15 Prepriming of DNA replication at *oriC* in the *E. coli* chromosome.



■ FIGURE 10.16 The initiation of DNA strands with RNA primers. The enzyme DNA primase catalyzes the synthesis of short (10 to 60 nucleotides long) RNA strands that are complementary to the template strands.

INITIATION OF DNA REPLICATION

The replication of the E. coli chromosome begins at oriC, the unique sequence at which replication is initiated, with the formation of a localized region of strand separation called the *replication bubble*. This replication bubble is formed by the interaction of *prepriming proteins* with oriC (**\blacksquare Figure 10.15**). The first step in prepriming appears to be the binding of four molecules of the *dnaA* gene product—DnaA protein-to the four 9-base-pair (bp) repeats in oriC. Next, DnaA proteins bind cooperatively to form a core of 20 to 40 polypeptides with *oriC* DNA wound on the surface of the protein complex. Strand separation begins within the three tandem 13-bp repeats in *oriC* and spreads until the replication bubble is created. A complex of DnaB protein (the hexameric DNA helicase) and DnaC protein (six molecules) joins the initiation complex and contributes to the formation of two bidirectional replication forks. The DnaT protein also is present in the prepriming protein complex, but its function is unknown. Other proteins associated with the initiation complex at oriC are DnaJ protein, DnaK protein, PriA protein, PriB protein, PriC protein, DNA-binding protein HU, DNA gyrase, and single-strand DNAbinding (SSB) protein. In some cases, however, their functional involvement in the prepriming process has not been established; in other cases, they are known to be involved, but their roles are unknown. The DnaA protein appears to be largely responsible for the localized strand separation at *oriC* during the initiation process.

INITIATION OF DNA CHAINS WITH RNA PRIMERS

All known DNA polymerases have an absolute requirement for a free 3'-OH on the end of the DNA strand being extended and an appropriate DNA template strand (specifying the complementary nascent strand) for activity. No known DNA polymerase can initiate the synthesis of a new strand of DNA. Thus, some special mechanism must

exist to initiate or prime the synthesis of new DNA chains once a replication bubble has formed.

RNA polymerase, a complex enzyme that catalyzes the synthesis of RNA molecules from DNA templates, has long been known to be capable of initiating the synthesis of new RNA chains at specific sites on the DNA. When this occurs, an RNA-DNA hybrid is formed in which the nascent RNA is hydrogen bonded to the DNA template. Because DNA polymerases are capable of extending either DNA or RNA chains containing a free 3'-OH, scientists began testing the idea that DNA synthesis might be initiated by using RNA primers. Their results proved that this idea is correct.

Subsequent research demonstrated that each new DNA chain is initiated by a short **RNA primer** synthesized by **DNA primase** (**■** Figure 10.16). The *E. coli* DNA primase is the product of the *dnaG* gene. In prokaryotes, these RNA primers are 10 to 60 nucleotides long, whereas in eukaryotes they are shorter, only about 10 nucleotides long. The RNA primers provide the free 3'-OHs required for covalent extension of polynucleotide chains by DNA polymerases. In *E. coli*, the enzyme that catalyzes the semiconservative replication of the chromosome is a polymerase called **DNA polymerase III** (see the section Multiple DNA Polymerases and Proofreading). DNA polymerase III catalyzes the addition of deoxyribonucleotides to RNA primers, either continuously on the leading strand or discontinuously by the synthesis of Okazaki fragments on the lagging strand. DNA polymerase III terminates an Okazaki fragment when it bumps into the RNA primer of the preceding Okazaki fragment.

The RNA primers subsequently are excised and replaced with DNA chains. This step is accomplished by DNA polymerase I in *E. coli*. In addition to the $5' \rightarrow 3'$

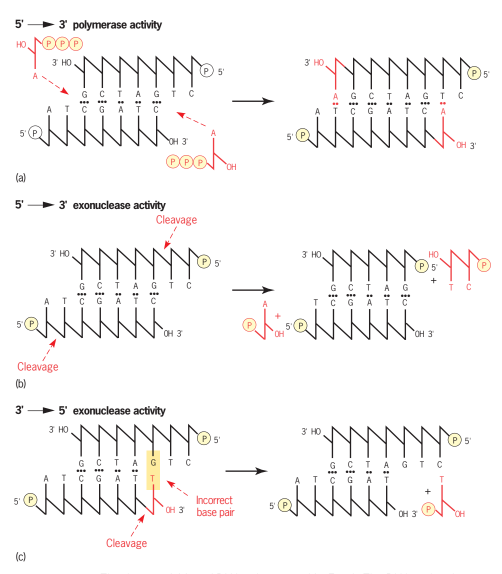
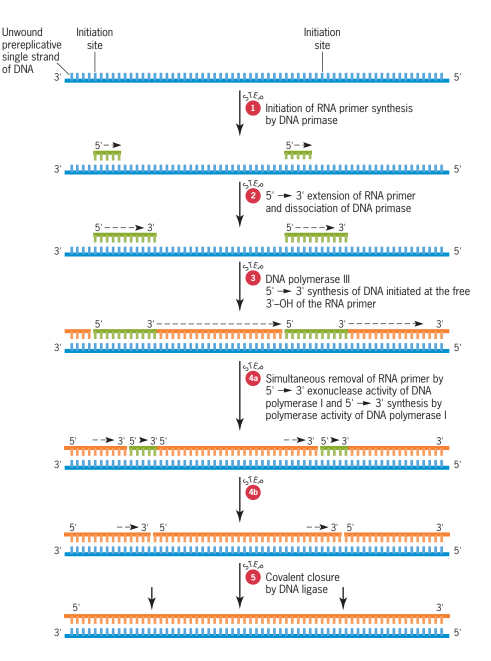


FIGURE 10.17 The three activities of DNA polymerase I in *E. coli*. The DNA molecules are shown here using flattened "stick" diagrams with one complementary strand on the top and the other on the bottom. "Stick" diagrams nicely emphasize the opposite chemical polarity $(5' \rightarrow 3' \text{ and } 3' \rightarrow 5')$ of the complementary strands. As is discussed in the text, all three activities— $(a) 5' \rightarrow 3'$ polymerase activity, $(b) 5' \rightarrow 3'$ exonuclease activity, and $(c) 3' \rightarrow 5'$ exonuclease activity—play important roles in *E. coli* cells.

polymerase activity illustrated in Figure 10.12, DNA polymerase I possesses two exonuclease activities: a $5' \rightarrow 3'$ exonuclease activity, which cuts back DNA strands starting at 5' termini, and a $3' \rightarrow 5'$ exonuclease activity, which cleaves off nucleotides from the 3' termini of DNA strands. Therefore, DNA polymerase I contains three distinct enzyme activities (**■** Figure 10.17), and all three activities play important roles in the replication of the *E. coli* chromosome.

The $5' \rightarrow 3'$ exonuclease activity of DNA polymerase I excises the RNA primer, and, at the same time, the $5' \rightarrow 3'$ polymerase activity of the enzyme replaces the RNA with a DNA chain by using the adjacent Okazaki fragment with its free 3'-OH as a primer. As we might expect based on this mechanism of primer replacement, *E. coli polA* mutants that lack the $5' \rightarrow 3'$ exonuclease activity of DNA polymerase I are defective in the excision of RNA primers and the joining of Okazaki fragments. After DNA polymerase I has replaced the RNA primer with a DNA chain, the 3'-OH of one Okazaki fragment is next to the 5'-phosphate group of the preceding Okazaki fragment. This product is an appropriate substrate for DNA ligase, which catalyzes the formation of a phosphodiester linkage between the adjacent Okazaki fragments. ■ FIGURE 10.18 Synthesis and replacement of RNA primers during replication of the lagging strand of DNA. A short RNA strand is synthesized to provide a 3'-OH primer for DNA synthesis (see Figure 10.16). The RNA primer is subsequently removed and replaced with DNA by the dual 5' \rightarrow 3' exonuclease and 5' \rightarrow 3' polymerase activities built into DNA polymerase I. DNA ligase then covalently closes the nascent DNA chain, catalyzing the formation of phosphodiester linkages between adjacent 3'-hydroxyls and 5'-phosphates (see Figure 10.14).

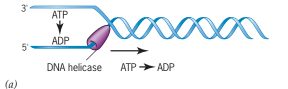


The steps involved in the synthesis and replacement of RNA primers during the discontinuous replication of the lagging strand are illustrated in **Figure 10.18**.

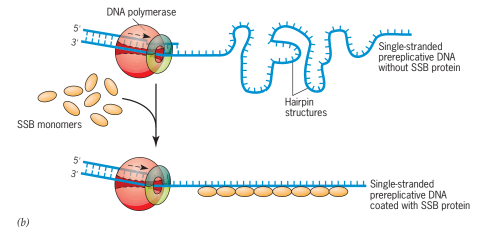
UNWINDING DNA WITH HELICASES, DNA-BINDING PROTEINS, AND TOPOISOMERASES

Semiconservative replication requires that the two strands of a parental DNA molecule be separated during the synthesis of new complementary strands. Since a DNA double helix contains two strands that cannot be separated without untwisting them turn by turn, DNA replication requires an unwinding mechanism. Given that each gyre, or turn, is about 10 nucleotide pairs long, a DNA molecule must be rotated 360° once for each 10 replicated base pairs. In *E. coli*, DNA replicates at a rate of about 30,000 nucleotides per minute. Thus, a replicating DNA molecule must spin at 3000 revolutions per minute to facilitate the unwinding of the parental DNA strands. The unwinding process (**Tigure 10.19a**) involves enzymes called **DNA helicases.** The major replicative DNA helicase in *E. coli* is the product of the *dnaB* gene. DNA helicases unwind DNA molecules using energy derived from ATP.

DNA helicase catalyzes the unwinding of the parental double helix.



Single-strand DNA-binding (SSB) protein keeps the unwound strands in an extended form for replication.



Once the DNA strands are unwound by DNA helicase, they must be kept in an extended single-stranded form for replication. They are maintained in this state by a coating of single-strand DNA-binding protein (SSB protein) (**Figure 10.19b**). The binding of SSB protein to single-stranded DNA is cooperative; that is, the binding of the first SSB monomer stimulates the binding of additional monomers at contiguous sites on the DNA chain. Because of the cooperativity of SSB protein binding, an entire single-stranded region of DNA is rapidly coated with SSB protein. Without the SSB protein coating, the complementary strands could renature or form intrastrand hairpin structures by hydrogen bonding between short segments of complementary or partially complementary nucleotide sequences. Such hairpin structures are known to impede the activity of DNA polymerases. In *E. coli*, the SSB protein is encoded by the *ssb* gene.

Recall that the *E. coli* chromosome contains a circular molecule of DNA. With the *E. coli* DNA spinning at 3000 revolutions per minute to allow the unwinding of the parental strands during replication (**True Figure 10.20**), what provides the swivel or axis of rotation that prevents the DNA from becoming tangled (positively supercoiled) ahead of the replication fork? The required axes of rotation during the replication of circular DNA molecules are provided by enzymes called **DNA topoisomerases**. The topoisomerases catalyze transient breaks in DNA molecules but use covalent linkages to themselves to hold on to the cleaved molecules. The topoisomerases are of two types: (1) DNA topoisomerase I enzymes produce temporary single-strand breaks or nicks in DNA, and (2) DNA topoisomerase II enzymes produce transient double-strand breaks in DNA. An important result of this difference is that topoisomerase I activities remove supercoils from DNA one at a time, whereas topoisomerase II enzymes remove and introduce supercoils two at a time.

The transient single-strand break produced by the activity of topoisomerase I provides an axis of rotation that allows the segments of DNA on opposite sides of the break to spin independently, with the phosphodiester bond in the intact strand serving as a swivel (**Figure 10.21**). Topoisomerase I enzymes are energy-efficient. They conserve the energy of the cleaved phosphodiester linkages by storing it in covalent linkages between themselves and the phosphate groups at the cleavage sites; they then reuse this energy to reseal the breaks.

DNA topoisomerase II enzymes induce transient double-strand breaks and add negative supercoils or remove positive supercoils two at a time by an energy ■ FIGURE 10.19 The formation of functional template DNA requires (a) DNA helicase, which unwinds the parental double helix, and (b) single-strand DNA-binding (SSB) protein, which keeps the unwound DNA strands in an extended form. In the absence of SSB protein, DNA single strands can form hairpin structures by intrastrand base-pairing (b, top), and the hairpin structures will retard or arrest DNA synthesis.

(a)

To unwind the template strands in *E. coli*, the DNA helix in front of the replication fork must spin at 3000 rpm.

Without a swivel or axis of rotation, the unwinding process would produce positive supercoils in front of the replication forks.

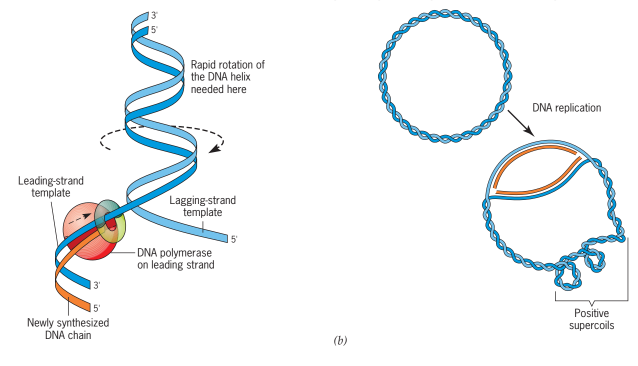


FIGURE 10.20 A swivel or axis of rotation is required during the replication of circular molecules of DNA like those in the *E. coli* or phage λ chromosomes. (*a*) During replication, the DNA in front of a replication fork must spin to allow the strands to be unwound by the helicase. (*b*) In the absence of an axis of rotation, unwinding will result in the production of positive supercoils in the DNA in front of a replication fork.

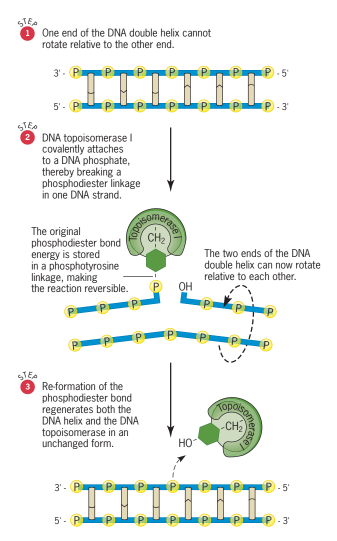
(ATP)-requiring mechanism. They carry out this process by cutting both strands of DNA, holding on to the ends at the cleavage site via covalent bonds, passing the intact double helix through the cut, and resealing the break (**Figure 10.22**). In addition to relaxing supercoiled DNA and introducing negative supercoils into DNA, topoisomerase II enzymes can separate interlocking circular molecules of DNA.

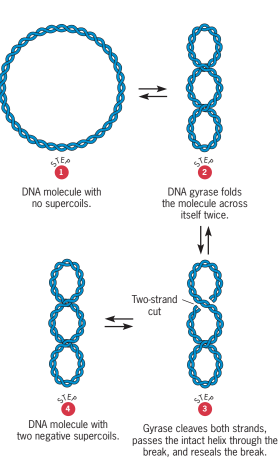
The best-characterized type II topoisomerase is an enzyme named **DNA gyrase** in *E. coli*. DNA gyrase is a tetramer with two α subunits encoded by the *gyrA* gene (originally *nalA*, for nalidixic acid) and two β subunits specified by the *gyrB* gene (formerly *cou*, for coumermycin). Nalidixic acid and coumermycin are antibiotics that block DNA replication in *E. coli* by inhibiting the activity of DNA gyrase. Nalidixic acid and coumermycin inhibit DNA synthesis by binding to the α and β subunits, respectively, of DNA gyrase. Thus, DNA gyrase activity is required for DNA replication to occur in *E. coli*.

Recall that chromosomal DNA is negatively supercoiled in *E. coli* (Chapter 9). The negative supercoils in bacterial chromosomes are introduced by DNA gyrase, with energy supplied by ATP. This activity of DNA gyrase provides another solution to the unwinding problem. Instead of creating positive supercoils ahead of the replication fork by unwinding the complementary strands of relaxed DNA, replication may produce relaxed DNA ahead of the fork by unwinding negatively supercoiled DNA. Because superhelical tension is reduced during unwinding—that is, strand separation is energetically favored—the negative supercoiling behind the fork may drive the unwinding process. If so, this mechanism nicely explains why DNA gyrase activity is required for DNA replication in bacteria. Alternatively, gyrase may simply remove positive supercoils that form ahead of the replication fork.

MULTIPLE DNA POLYMERASES AND PROOFREADING

DNA polymerases are processive enzymes that catalyze the covalent extension at the 3' termini of growing polynucleotide chains. All polymerases require preexisting DNA





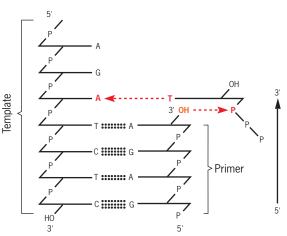
■ **FIGURE 10.22** Mechanism of action of DNA gyrase, an *E. coli* DNA topoisomerase II required for DNA replication.

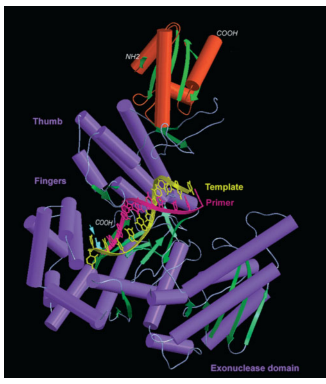
■ FIGURE 10.21 DNA topoisomerase I produces transient single-strand breaks in DNA that act as axes of rotation or swivels during DNA replication.

with two essential components, one providing a primer function and the other a template function (\blacksquare Figure 10.23).

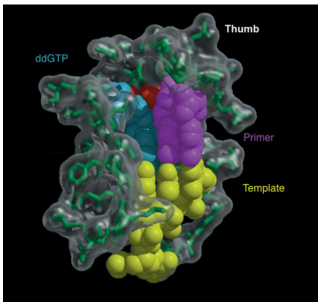
- The *primer DNA* provides a terminus with a free 3'-OH to which nucleotides are added during DNA synthesis. No DNA polymerase can initiate the synthesis of DNA chains *de novo*. All DNA polymerases have an absolute requirement for a free 3'-hydroxyl on a preexisting polynucleotide chain. They catalyze the formation of a phosphodiester bridge between the 3'-OH at the end of the primer DNA chain and the 5'-phosphate of the incoming deoxyribonucleotide.
- **2.** The *template DNA* provides the nucleotide sequence that specifies the complementary sequence of the growing DNA chain. DNA polymerases require a DNA template whose base sequence dictates, by its base-pairing potential, the synthesis of a complementary base sequence in the strand being synthesized.

■ FIGURE 10.23 Template and primer requirements of DNA polymerases. The DNA molecule is shown here as a flattened "stick" diagram, like the ones shown in Figure 10.17. All DNA polymerases require a primer strand (shown on the right) with a free 3'-hydroxyl. The primer strand is covalently extended by the addition of nucleo-tides (such as dTMP, derived from the incoming precursor dTTP shown). In addition, DNA polymerases require a template strand (shown on the left), which determines the base sequence of the strand being synthesized. The new strand will be complementary to the template strand.









(b)

■ FIGURE 10.24 Schematic diagram (a) and space-filling model (b) of the structure of the complex between the phage T7 DNA polymerase, template-primer DNA, and a nucleoside triphosphate (ddGTP) precursor molecule. The template strand, primer strand, and nucleoside triphosphate are shown in yellow, magenta, and cyan, respectively. Protein components are shown in purple, green, orange, and gray. Note the tight juxtaposition between the nucleoside triphosphate, the primer terminus, and the template strand in (b).

The reaction catalyzed by DNA polymerases is a nucleophilic attack by the 3'-OH at the terminus of the primer strand on the nucleotidyl or interior phosphorus atom of the nucleoside triphosphate precursor with the elimination of pyrophosphate. This reaction mechanism explains the absolute requirement of DNA polymerases for a free 3'-OH group on the primer DNA strand that is being covalently extended and dictates that *the direction of synthesis is always* $5' \rightarrow 3'$ (see Figure 10.12).

E. coli contains at least five DNA polymerases: DNA polymerase I, DNA polymerase II, DNA polymerase III, DNA polymerase IV, and DNA polymerase V. DNA polymerases I and II are DNA repair enzymes. Unlike DNA polymerases I and II, DNA polymerase III is a complex enzyme composed of many different subunits. Like DNA polymerase I, DNA polymerase III has $5' \rightarrow 3'$ polymerase and $3' \rightarrow 5'$ exonuclease activities; however, it has a $5' \rightarrow 3'$ exonuclease that is active only on single-stranded DNA. The more recently characterized DNA polymerases IV and V, along with polymerase II, play important roles in the replication of damaged DNA, with the polymerase involved depending on the type of damage (see Chapter 13).

Eukaryotic organisms encode even more polymerases—with at least 15 different DNA polymerases having been identified so far. The eukaryotic DNA polymerases have been named α , β , γ , δ , ε , κ , ζ , η , θ , κ , λ , μ , σ , ϕ , and Rev1. Two or more of the DNA polymerases (α , δ , and/or ε) work together to carry out the semiconservative replication of nuclear DNA. DNA polymerase γ is responsible for the replication of DNA in mitochondria, and DNA polymerases β , ε , κ , ζ , η , θ , κ , λ , μ , σ , ϕ , and Rev1 are DNA repair enzymes or perform other metabolic functions. Some of the eukaryotic DNA polymerases lack the $3' \rightarrow 5'$ exonuclease activity that is present in most prokaryotic DNA polymerases.

All of the DNA polymerases studied to date, prokaryotic and eukaryotic, catalyze the same basic reaction: a nucleophilic attack by the free 3'-OH at the primer strand terminus on the nucleotidyl phosphorus of the nucleoside triphosphate precursor. Thus, all DNA polymerases have an absolute requirement for a free 3'-hydroxyl group on a preexisting primer strand. None of these DNA polymerases can initiate new DNA chains *de novo*, and all DNA synthesis occurs in the 5' \rightarrow 3' direction.

The major replicative DNA polymerases are amazingly accurate, incorporating incorrect nucleotides with an initial frequency of 10^{-5} to 10⁻⁶. (Some repair polymerases are error-prone—see Chapter 13.) Studies of the crystal structure of the complex formed by a monomeric DNA polymerase, a nucleoside triphosphate precursor, and a templateprimer DNA have contributed to our understanding of the high fidelity of DNA synthesis. In these studies, published in 1998, Sylvie Doublié and colleagues determined the structure of the phage T7 polymerase, which is similar to DNA polymerase I of E. coli, with resolution to 0.22 nm. The results show that the polymerase is shaped like a little hand, with the incoming nucleoside triphosphate, the template, and the primer terminus all tightly grasped between the thumb, the fingers, and the palm (**Figure 10.24**). The enzyme positions the incoming nucleoside triphosphate in juxtaposition with the terminus of the primer strand in a position to form hydrogen bonds with the first unpaired base in the template strand. Thus, the structure of this polymerase complex provides a simple explanation for the template-directed selection of incoming nucleotides during DNA synthesis.

DNA polymerase III, the "replicase" in *E. coli*, is a multimeric enzyme (an enzyme with many subunits) with a molecular mass of about 900,000 daltons in its complete or **holoenzyme** form. The minimal core that has catalytic activity *in vitro* contains

three subunits: α (the *dnaE* gene product), ε (the *dnaQ* product), and θ (the *holE* product). Addition of the τ subunit (the *dnaX* product) results in dimerization of the catalytic core and increased activity. The catalytic core synthesizes rather short DNA strands because of its tendency to fall off the DNA template. In order to synthesize the long DNA molecules present in chromosomes, this frequent dissociation of the polymerase from the template must be eliminated. The β subunit (the *dnaN* gene product) of DNA polymerase III forms a dimeric clamp that keeps the polymerase from falling off the template DNA (**Figure 10.25**). The β -dimer forms a ring that encircles the replicating DNA molecule and allows DNA polymerase III to slide along the DNA while remaining tethered to it. The DNA polymerase III holoenzyme, which is responsible for the synthesis of both nascent DNA strands at a replication fork, contains at least 20 polypeptides. The structural complexity of the DNA polymerase III holoenzyme is illustrated in **Figure 10.26**; the diagram shows 16 of the best-characterized polypeptides encoded by seven different genes.

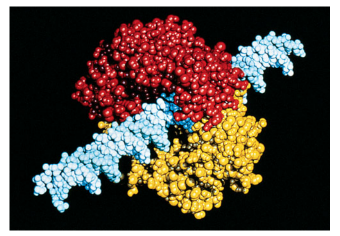
As we discussed earlier, the fidelity of DNA duplication is amazing—with only about one error present in every billion base pairs shortly after synthesis. This high fidelity is necessary to keep the mutation load at a tolerable level, especially in large genomes such as those of mammals, which contain $3 \times 10^{\circ}$ nucleotide pairs. Without the high fidelity of DNA replication, the monozygotic twins discussed at the beginning of this chapter would be less similar in phenotype. Indeed, based on the dynamic structures of the four nucleotides in DNA, the observed fidelity of DNA replication is much higher than expected. The thermodynamic changes in nucleotides that allow the formation of hydrogen-bonded base pairs other than A:T and G:C predict error rates of 10^{-5} to 10^{-4} , or one error per 10,000 to 100,000 incorporated nucleotides. The predicted error rate of 10,000 times the observed error rate raises the question of how this high fidelity of DNA replication can be achieved.

Living organisms have solved the potential problem of insufficient fidelity during DNA replication by evolving a mechanism for **proofread**ing the nascent DNA chain as it is being synthesized. The proofreading process involves scanning the termini of nascent DNA chains for errors and correcting them. This process is carried out by the $3' \rightarrow 5'$ exonucle-

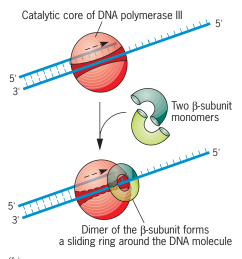
as activities of DNA polymerases (see Figure 10.17). When a template-primer DNA has a terminal mismatch (an unpaired or incorrectly paired base or sequence of bases at the 3' end of the primer), the $3' \rightarrow 5'$ exonuclease activity of the DNA polymerase clips off the unpaired base or bases (**■** Figure 10.27). When an appropriately base-paired terminus is produced, the $5' \rightarrow 3'$ polymerase activity of the enzyme begins resynthesis by adding nucleotides to the 3' end of the primer strand.

In monomeric enzymes like DNA polymerase I of *E. coli*, the $3' \rightarrow 5'$ exonuclease activity is built in. In multimeric enzymes, the $3' \rightarrow 5'$ proofreading exonuclease activity is often present on a separate subunit. In the case of DNA polymerase III of *E. coli*, this proofreading function is carried out by the ε subunit. DNA polymerase IV of *E. coli* contains no exonuclease activity. In eukaryotes, DNA polymerases γ , δ , and ε contain $3' \rightarrow 5'$ proofreading exonuclease activities, but polymerases α and β lack this activity.

Without proofreading during DNA replication, Merry and Sherry, the twins discussed at the beginning of this chapter, would be less similar in appearance. Without proofreading, changes would have accumulated in their genes during the billions of cell divisions that occurred during their growth from small embryos to adults. Indeed, the identity of the genotypes of identical twins depends both on DNA proofreading during replication and on the activity of an army of DNA repair enzymes (Chapter 13). These enzymes continually scan DNA for various types of damage and make repairs before the alterations cause inherited genetic changes.



(a)



(b)

FIGURE 10.25 Space-filling model (*a*) and diagram (*b*) showing how two β -subunits (light and dark green) of DNA polymerase III clamp the enzyme to the DNA molecule (blue).

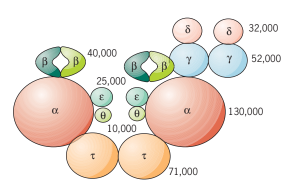
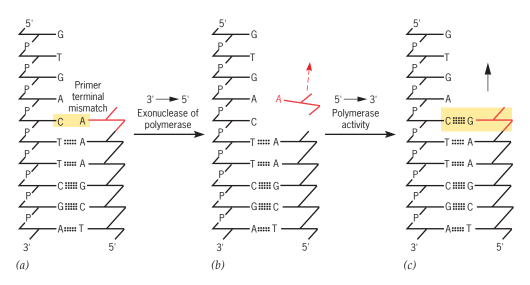
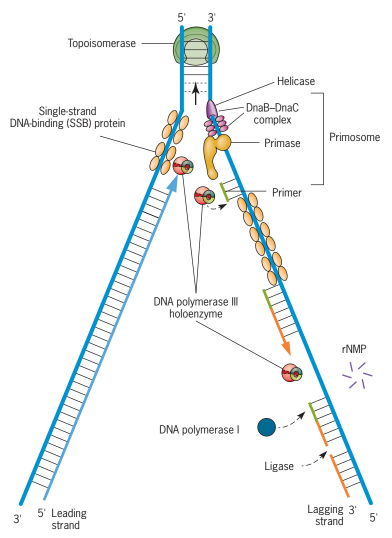


FIGURE 10.26 Structure of the *E. coli* DNA polymerase III holoenzyme. The numbers give the masses of the subunits in daltons.



■ FIGURE 10.27 Proofreading by the $3' \rightarrow 5'$ exonuclease activity of DNA polymerases during DNA replication. As introduced in Figure 10.17, the DNA molecules are shown as "stick" diagrams. If DNA polymerase is presented with a template and primer containing a 3' primer terminal mismatch (a), it will not catalyze covalent extension (polymerization). Instead, the $3' \rightarrow 5'$ exonuclease activity, an integral part of many DNA polymerases, will cleave off the mismatched terminal nucleotide (b). Then, presented with a correctly base-paired primer terminus, DNA polymerase will catalyze $5' \rightarrow 3'$ covalent extension of the primer strand (c).

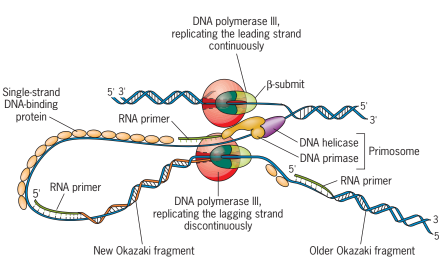


THE PRIMOSOME AND THE REPLISOME

The initiation of Okazaki fragments on the lagging strand is carried out by the primosome, a protein complex containing DNA primase and DNA helicase. The primosome moves along a DNA molecule, powered by the energy of ATP. As it proceeds, DNA helicase unwinds the parental double helix, and DNA primase synthesizes the RNA primers needed to initiate successive Okazaki fragments. The RNA primers are covalently extended with the addition of deoxyribonucleotides by DNA polymerase III. DNA topoisomerases provide transient breaks in the DNA that serve as swivels for DNA unwinding and keep the DNA untangled. Single-strand DNA binding protein coats the unwound prereplicative DNA and keeps it in an extended state for DNA polymerase III. The RNA primers are replaced with DNA by DNA polymerase I, and the single-strand nicks left by polymerase I are sealed by DNA ligase. This sequence of events occurring at each replication fork during the semiconservative replication of the E. coli chromosome is illustrated in ■ Figure 10.28.

As a replication fork moves along a parental double helix, two DNA strands (the leading strand and the lagging strand) are replicated in the highly coordinated series of reactions described above. The complete replication apparatus moving along the DNA molecule at a replication fork is called the **replisome** (Figure 10.29). The replisome contains the DNA polymerase III holoenzyme; one catalytic core replicates the leading strand, the second catalytic core replicates the lagging strand, and

■ **FIGURE 10.28** Diagram of a replication fork in *E. coli* showing the major components of the replication apparatus. rNMP = ribonucleoside monophosphates.



■ FIGURE 10.29 Diagram of the *E. coli* replisome, showing the two catalytic cores of DNA polymerase III replicating the leading and lagging strands and the primosome unwinding the parental double helix and initiating the synthesis of new chains with RNA primers. The entire replisome moves along the parental double helix, with each component performing its respective function in a concerted manner. Actually, the replication complex probably does not move. Instead, the DNA is pulled through the replisome. Replication is proceeding from left to right. *Original micrograph courtesy of David Dressler, Harvard University.*

the primosome unwinds the parental DNA molecule and synthesizes the RNA primers needed for the discontinuous synthesis of the lagging strand. In order for the two catalytic cores of the polymerase III holoenzyme to synthesize both the nascent leading and lagging strands, the lagging strand is thought to form a loop from the primosome to the second catalytic core of DNA polymerase III (**■** Figure 10.29).

In *E. coli*, the termination of replication occurs at variable sites within regions called *terA* and *terB*, which block the movement of replication forks advancing in the counterclockwise and clockwise directions, respectively. DNA topoisomerases or special recombination enzymes then facilitate the separation of the nascent DNA molecules. The DNA is condensed into the nucleoid, or folded genome, of *E. coli*, in part through the negative supercoiling introduced by DNA gyrase.

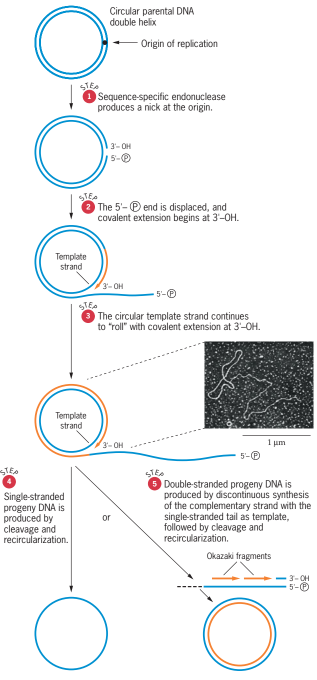
At the beginning of this chapter, we noted the striking fidelity of DNA replication. Now that we have examined the cellular machinery responsible for DNA replication in living organisms, this fidelity seems less amazing. A very sophisticated apparatus, with built-in safeguards against malfunctions, has evolved to assure that the genetic information of *E. coli* is transmitted accurately from generation to generation.

ROLLING-CIRCLE REPLICATION

In the preceding sections of this chapter, we have considered θ -shaped, eye-shaped, and Y-shaped replicating DNAs. We will now examine another important type of DNA replication called **rolling-circle replication**. Rolling-circle replication is used (1) by many viruses to duplicate their genomes, (2) in bacteria to transfer DNA from donor cells to recipient cells during one type of genetic exchange (Chapter 8), and (3) in amphibians to amplify extrachromosomal DNAs carrying clusters of ribosomal RNA genes during oogenesis.

As the name implies, rolling-circle replication is a mechanism for replicating circular DNA molecules. The unique aspect of rolling-circle replication is that one parental circular DNA strand remains intact and rolls (thus the name rolling circle) or spins while serving as a template for the synthesis of a new complementary strand (**■** Figure 10.30). Replication is initiated when a sequence-specific endonuclease cleaves one strand at

FIGURE 10.30 The rolling-circle mechanism of DNA replication. Material for progeny chromosomes (in this case, single stranded DNA for the virus Φ X174) is produced by continuous copying around a nicked, double-stranded DNA circle, with the intact strand serving as a template. Electron micrograph courtesy of David Dressler, Harvard University.



the origin, producing 3'-OH and 5'-phosphate termini. The 5' terminus is displaced from the circle as the intact template strand turns about its axis. Covalent extension occurs at the 3'-OH of the cleaved strand. Since the circular template DNA may turn 360° many times, with the synthesis of one complete or unit-length DNA strand during each turn, rolling-circle replication generates single-stranded tails longer than the contour length of the circular chromosome (Figure 10.30). Rolling-circle replication can produce either single-stranded or double-stranded progeny DNAs. Circular single-stranded progeny molecules are produced by site-specific cleavage of the single-stranded tails at the origins of replication and recircularization of the resulting unit-length molecules. To produce double-stranded progeny molecules, the singlestranded tails are used as templates for the discontinuous synthesis of complementary strands prior to cleavage and circularization. The enzymes involved in rolling-circle replication and the reactions catalyzed by these enzymes are basically the same as those responsible for DNA replication involving θ -type intermediates.

KEY POINTS • DNA replication is complex, requiring the participation of a large number of proteins.

- DNA synthesis is continuous on the progeny strand that is being extended in the overall $5' \rightarrow 3'$ direction, but is discontinuous on the strand growing in the overall $3' \rightarrow 5'$ direction.
- New DNA chains are initiated by short RNA primers synthesized by DNA primase.
- DNA synthesis is catalyzed by enzymes called DNA polymerases.
- All DNA polymerases require a primer strand, which is extended, and a template strand, which is copied.
- All DNA polymerases have an absolute requirement for a free 3'-OH on the primer strand, and all DNA synthesis occurs in the 5' to 3' direction.
- The $3' \rightarrow 5'$ exonuclease activities of DNA polymerases proofread nascent strands as they are synthesized, removing any mispaired nucleotides at the 3' termini of primer strands.
- The enzymes and DNA-binding proteins involved in replication assemble into a replisome at each replication fork and act in concert as the fork moves along the parental DNA molecule.

Unique Aspects of Eukaryotic Chromosome Replication

Although the main features of DNA replication are the same in all organisms, some processes occur only in eukaryotes.

Most of the information about DNA replication has resulted from studies of *E. coli* and some of its viruses. Less information is available about DNA replication in eukaryotic organisms. However, enough information is available to conclude that most aspects of DNA replication are simi-

lar in prokaryotes and eukaryotes, including humans. RNA primers and Okazaki fragments are shorter in eukaryotes than in prokaryotes, but the leading and lagging strands replicate by continuous and discontinuous mechanisms, respectively, in eukaryotes just as in prokaryotes. Nevertheless, a few aspects of eukaryotic DNA replication are unique to these structurally more complex species. For example, DNA synthesis takes place within a small portion of the cell cycle in eukaryotes, not continuously as in prokaryotes. The giant DNA molecules present in eukaryotic chromosomes would take much too long to replicate if each chromosome contained a single origin. Thus, eukaryotic chromosomes contain multiple origins of replication. Rather than using two catalytic complexes of one DNA polymerase to replicate the leading and lagging strands at each replication fork, eukaryotic organisms utilize two or more different polymerases.

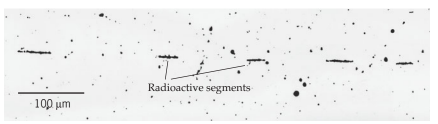
As we discussed in Chapter 9, eukaryotic DNA is packaged in histone-containing structures called nucleosomes. Do these nucleosomes impede the movement of replication forks? If not, how does a replisome move past a nucleosome? Is the nucleosome completely or partially disassembled, or does the fork somehow slide past the

nucleosome as the replisome duplicates the DNA molecule while it is still present on the surface of the nucleosome? Lastly, eukaryotic chromosomes contain linear DNA

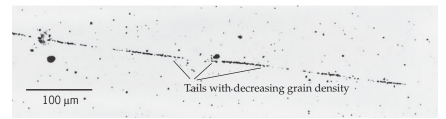
molecules, and the discontinuous replication of the ends of linear DNA molecules creates a special problem. We will address these aspects of chromatin replication in eukaryotes in the final sections of this chapter.

THE CELL CYCLE

When bacteria are growing on rich media, DNA replication occurs nonstop throughout the cell cycle. However, in eukaryotes, DNA replication is restricted to the S phase (for synthesis; Chapter 2). Recall that a normal eukaryotic cell cycle consists of G_1 phase (immediately following the completion of mitosis; G for gap), S phase, G_2 phase (preparation for mitosis), and M phase (mitosis) (see Chapter 2 for details). In rapidly dividing embryonic cells, G_1 and G_2 are very short or nonexistent. In all cells, decisions to continue on through the cell cycle occur at two points: (1) entry into S phase and (2) entry into mitosis. These checkpoints help to ensure that the DNA replicates once and only once during each cell division.



(*a*) Autoradiograph of a portion of a DNA molecule from a Chinese hamster cell that had been pulse-labeled with ³H-thymidine.



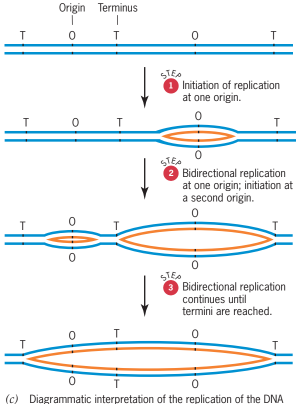
(b) Autoradiograph of a segment of a DNA molecule from a Chinese hamster cell that was pulse-labeled with ³H-thymidine and then transferred to nonradioactive medium for an additional growth period.

MULTIPLE REPLICONS PER CHROMOSOME

The giant DNA molecules in the largest chromosomes of *Drosophila melano*gaster contain about 6.5×10^7 nucleotide pairs. The rate of DNA replication in *Drosophila* is about 2600 nucleotide pairs per minute at 25°C. A single replication fork would therefore take about 17.5 days to replicate one of these giant DNA molecules. With two replication forks moving bidirectionally from a central origin, such a DNA molecule could be replicated in just over 8.5 days. Given that the chromosomes of *Drosophila* embryos replicate within 3 to 4 minutes and the nuclei divide once every 9 to 10 minutes during the early cleavage divisions, it is clear that each giant DNA molecule must contain many origins of replication. Indeed, the complete replication of the DNA of the largest *Drosophila* chromosome within 3.5 minutes would require over 7000 replication forks distributed at equal intervals along the molecule. Thus, multiple origins of replication are required to allow the very large DNA molecules in eukaryotic chromosomes to replicate within the observed cell division times.

The first evidence for multiple origins in eukaryotic chromosomes resulted from pulse-labeling experiments with Chinese hamster cells growing in culture. In 1968, when Joel Huberman and Arthur Riggs pulse-labeled cells with ³H-thymidine for a few minutes, extracted the DNA, and performed autoradiographic analysis of the labeled DNA, they observed tandem arrays of exposed silver grains (**■** Figure 10.31*a*). The simplest interpretation of their results is that individual macromolecules of DNA contain multiple origins of replication. When the pulse-labeling period was followed by a short interval of growth in nonradioactive medium (pulse-chase experiments), the tandem arrays contained central regions of high-grain density with tails of decreasing grain density at *both* ends (**■** Figure 10.31*b*). This result indicates that replication in eukaryotes is bidirectional just as it is in most prokaryotes. The tails of decreasing grain density result from the gradual dilution of the intracellular pools of ³H-thymidine by ¹H-thymidine as the replication forks move bidirectionally from central origins toward replication termini (**■** Figure 10.31*c*).

A segment of DNA whose replication is under the control of one origin and two termini is called a **replicon**. In prokaryotes, the entire chromosome is usually one



c) Diagrammatic interpretation of the replication of the DNA molecules visualized in (*a*) and (*b*).

■ FIGURE 10.31 Evidence for bidirectional replication of the multiple replicons in the giant DNA molecules of eukaryotes. The tandem arrays of radioactivity in (a) indicate that replication occurs at multiple origins; tails with decreasing grain density observed in (b) indicate that replication occurs bidirectionally from each origin (c).

Solve It!

Understanding Replication of the Human X Chromosome

According to the Entrez Genome Database of the National Center for Biotechnology Information, the first human X chromosome to be sequenced contained 154,913,754 nucleotide pairs. If this X chromosome is present in a somatic cell with an S phase of the cell cycle of 10 hours and a replication rate of 3000 nucleotides per minute, what is the minimum number of origins of replication required for its replication? If the average size of the Okazaki fragments formed during the replication of this chromosome is 150 nucleotides, how many Okazaki fragments are produced during its replication? How many RNA primers? In answering these questions, assume that the reported sequence does not include the TTAGGG telomere repeat sequences at the ends of the chromosome.

• To see the solution to this problem, visit the Student Companion site.

replicon. The existence of multiple replicons per eukaryotic chromosome has been verified directly by autoradiography and electron microscopy in several different species. The genomes of humans and other mammals contain about 10,000 origins of replication distributed throughout the chromosomes at 30,000- to 300,000-base-pair intervals. Clearly, the number of replicons per chromosome is not fixed throughout the growth and development of a multicellular eukaryote. Replication is initiated at more sites during the very rapid cell divisions of embryogenesis than during later stages of development. Unfortunately, geneticists don't know what factors determine which origins are operational at any given time or in a particular type of cell. Go to Solve It: Understanding Replication of the Human X Chromosome to test your comprehension of the concepts discussed here.

TWO OR MORE DNA POLYMERASES AT A SINGLE REPLICATION FORK

Given the complexity of the replisome in the simple bacterium *E. coli* (see Figures 10.28 and 10.29), it seems likely that the replication apparatus is even more complex in eukaryotes. Although knowledge of the structure of the replicative machinery in eukaryotes is still limited, many features of DNA replication are similar in eukaryotes and prokaryotes.

As in the case of prokaryotes, much of the information about DNA synthesis in eukaryotes has come from the development and dissection of *in vitro* DNA replication systems. Studies of the replication of DNA viruses of eukaryotes have proven informative, and of these viruses, Simian virus 40 (SV40) has proven particularly useful. The replication of SV40 is carried out almost entirely by the host cell's replication apparatus. Only one viral protein, the so-called T antigen, is required for replication of the SV40 chromosome.

As in prokaryotes, the unwinding of the parental DNA strands requires a DNA topoisomerase and a DNA helicase. The unwound strands are kept in the extended

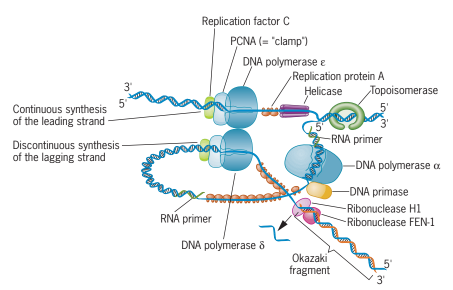


FIGURE 10.32 Some of the important components of a replisome in eukaryotes. Each replisome contains three different polymerases, α , δ , and ϵ . The DNA polymerase α -DNA primase complex synthesizes the RNA primers and adds short segments of DNA. DNA polymerase δ then completes the synthesis of the Okazaki fragments in the lagging strand, and polymerase ϵ catalyzes the continuous synthesis of the leading strand. PCNA (proliferating cell nuclear antigen) is equivalent to the β subunit of *E. coli* DNA polymerase III; it clamps polymerases δ and ϵ to the DNA molecule facilitating the synthesis of long DNA chains. Ribonucleases H1 and FEN-1 (F1 nuclease 1) remove the RNA primers, polymerase δ fills in the gap, and DNA ligase (not shown) seals the nicks, just as in *E. coli* (see Figure 10.18).

state by a single-strand DNA-binding protein called replication protein A (Rp-A). However, unlike the process in prokaryotes, the replication of chromosomal DNA in eukaryotes requires the activity of three different DNA polymerases—polymerase α (Pol α), polymerase δ (Pol δ), and polymerase ϵ (Pol ϵ). At least two polymerases, perhaps all three, are present in each replication fork (replisome), and each polymerase contains multiple subunits. Also, whereas the *E. coli* replisome contains 13 known proteins, the replisomes of yeast and mammals contain at least 27 different polypeptides.

In eukaryotes, Pol α is required for the initiation of replication at origins and for the priming of Okazaki fragments during the discontinuous synthesis of the lagging strand. Pol α exists in a stable complex with DNA primase; indeed, they copurify during isolation. The primase synthesizes the RNA primers, which are then extended with deoxyribonucleotides by Pol a to produce an RNA-DNA chain about 30 nucleotides in total length. These RNA-DNA primer chains are then extended by Pol δ . Pol δ completes the replication of the lagging strand, while polymerase ε catalyzes the replication of the leading strand. Pol 8 must interact with proteins PCNA (proliferating cell nuclear antigen) and replication factor C (Rf-C) to be active (■ Figure 10.32). PCNA is the sliding clamp that tethers Pol δ to the DNA to

allow processive replication (to prevent the polymerase from falling off the template); PCNA is equivalent to the β subunit of DNA polymerase III in *E. coli* (see Figure 10.25). Rf-C is required for PCNA to load onto DNA. PCNA is a trimeric protein that forms a closed ring; Rf-C induces a change in the conformation of PCNA that allows it to encircle DNA, providing the essential sliding clamp.

Polymerases δ and ε both contain the 3' \rightarrow 5' exonuclease activity required for proofreading (see Figure 10.27). However, they do not have 5' \rightarrow 3' exonuclease activity; thus, they cannot remove RNA primers like DNA polymerase I of *E. coli* does. Instead, the RNA primers are excised by two nucleases, ribonuclease H1 (which degrades RNA present in RNA–DNA duplexes) and ribonuclease FEN-1 (F1 nuclease 1). Pol δ then fills in the gaps, and DNA ligase seals the nicks, producing covalently closed progeny strands.

As mentioned earlier, there are at least 15 different DNA polymerases— α , β , γ , δ , ε , κ , ζ , η , θ , κ , λ , μ , σ , ϕ , and Rev1—in eukaryotes. DNA polymerase γ is responsible for the replication of DNA in mitochondria, and the other DNA polymerases have important roles in DNA repair and other pathways (Chapter 13).

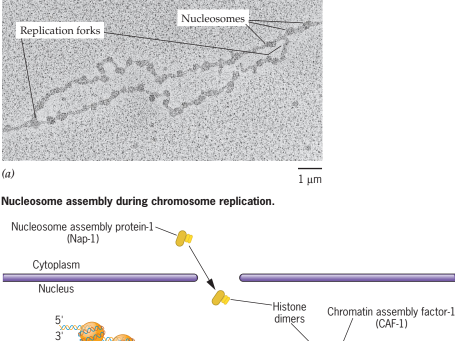
DUPLICATION OF NUCLEOSOMES AT REPLICATION FORKS

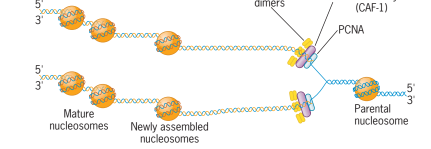
As we discussed in Chapter 9, the DNA in eukaryotic interphase chromosomes is packaged in approximately 11-nm beads called nucleosomes. Each nucleosome contains 166 nucleotide pairs of DNA wound in two turns around an octamer of histone molecules. Given the size of nucleosomes and the large size of DNA replisomes, it seems unlikely that a replication fork can move past an intact nucleosome. Yet, electron micrographs of replicating chromatin in Drosophila clearly show nucleosomes with approximately normal structure and spacing on both sides of replication forks (**Figure 10.33***a*); that is, nucleosomes appear to have the same structure and spacing immediately behind a replication fork (postreplicative DNA) as they do in front of a replication fork (prereplicative DNA). This observation suggests that nucleosomes must be disassembled to let the replisome duplicate the DNA packaged in them and then be quickly reassembled; that is, DNA replication and nucleosome assembly must be tightly coupled.

Since the mass of the histones in nucleosomes is equivalent to that of the DNA, large quantities of histones must be synthesized during each cell generation in order for the nucleosomes to duplicate. Although histone synthesis occurs throughout the cell cycle, there is a burst of histone biosynthesis during S phase that generates enough histones for chromatin duplication. When density-transfer experiments were performed to examine the mode of nucleosome duplication, the nucleosomes on both progeny DNA molecules were found to contain both old (prereplicative) histone complexes and new (postreplicative)

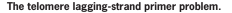
(b)

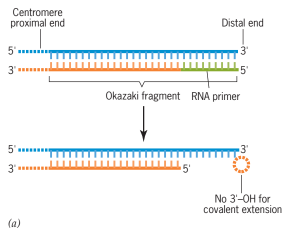
Nucleosome spacing in replicating chromatin.

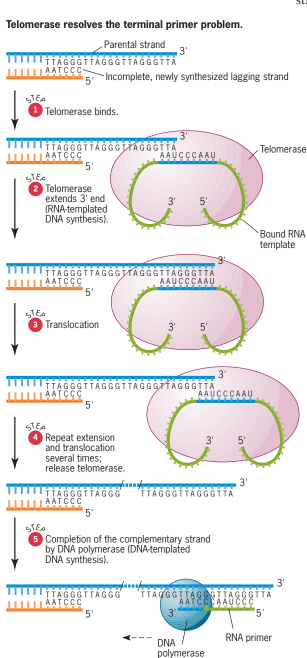




■ FIGURE 10.33 The disassembly and assembly of nucleosomes during the replication of chromosomes in eukaryotes. (a) An electron micrograph showing nucleosomes on both sides of two replication forks in *Drosophila*. Recall that DNA replication is bidirectional; thus, each branch point is a replication fork. (b) The assembly of new nucleosomes during chromosome replication requires proteins that transport histones from the cytoplasm to the nucleus and that concentrate them at the site of nucleosome assembly. PCNA = proliferating cell nuclear antigen (see Figure 10.32).







complexes. Thus, at the protein level, nucleosome duplication appears to occur by a dispersive mechanism.

A number of proteins are involved in the disassembly and assembly of nucleosomes during chromosome replication in eukaryotes. Two of the most important are *n*ucleosome *assembly protein-1* (Nap-1) and *c*hromatin *assembly factor-1* (CAF-1). Nap-1 transports histones from their site of synthesis in the cytoplasm to the nucleus, and CAF-1 carries them to the chromosomal sites of nucleosome assembly (**Figure 10.33b**). CAF-1 delivers histones to the sites of DNA replication by binding to PNCA (*proliferating cell nuclear antigen*)—the clamp that tethers DNA polymerase δ to the DNA template (see Figure 10.32). CAF-1 is an essential protein in *Drosophila*, but not in yeast where other proteins can perform some of its functions.

Many other proteins affect nucleosome structure. Some are involved in chromatin remodeling—changing nucleosome structure in ways that activate or silence the expression of the genes packaged therein. Others modify nucleosome structure by adding methyl or acetyl groups to specific histones. In addition,

eukaryotes contain several minor histones with structures slightly different from the major histones, and the incorporation of these minor histones into nucleosomes can change their structure. In *Drosophila*, for example, the incorporation of histone H3.3 into nucleosomes results in high levels of transcription of the genes therein. Thus, nucleosome structure is not invariant; to the contrary, it plays an important role in modulating gene expression (see On the Cutting Edge: Chromatin Remodeling and Gene Expression in Chapter 11 and the section Chromatin Remodeling in Chapter 19).

TELOMERASE: REPLICATION OF CHROMOSOME TERMINI

We discussed the unique structures of telomeres, or chromosome ends, in Chapter 9. An early reason for thinking that telomeres must have special structures was that DNA polymerases cannot replicate the terminal DNA segment of the lagging strand of a linear chromosome. At the end of the DNA molecule being replicated discontinuously, there would be no DNA strand to provide a free 3'-OH (primer) for polymerization of deoxyribonucleotides after the RNA primer of the terminal Okazaki fragment has been excised (**Figure 10.34***a*). Either (1) the telomere must have a unique structure that facilitates its replication or (2) there must be a special enzyme that resolves this enigma of replicating the terminus of the lagging strand. Indeed, evidence has shown that both are correct. The special structure of telomeres provides a neat mechanism for the addition of telomeres by an RNA-containing enzyme called telomerase. This unique enzyme was discovered in 1985 by Elizabeth Blackburn and Carol Greider. They shared the 2009 Nobel Prize in Physiology or Medicine with Jack Szostak, who, along with Blackburn, determined how the unique structures of telomeres protected them from degradation.

■ FIGURE 10.34 Replication of chromosome telomeres. (a) Because of the requirement for a free 3'-OH at the end of the primer strand, DNA polymerases cannot replace an RNA primer that initiates DNA synthesis close to or at the terminus of the lagging strand. (b) These termini of chromosomes are replicated by a special enzyme called telomerase, which prevents the ends of chromosomes from becoming shorter during each replication. The nucleotide sequence at the terminus of the lagging strand is specified by a short RNA molecule present as an essential component of telomerase. The telomere sequence shown is that of humans.

The telomeres of humans, which contain the tandemly repeated sequence TTAGGG, will be used to illustrate how telomerase adds ends to chromosomes (**Televice 10.34b**). Telomerase recognizes the G-rich telomere sequence on the 3' overhang and extends it $5' \rightarrow 3'$ one repeat unit at a time. Telomerase does not fill in the gap opposite the 3' end of the template strand; it simply extends the 3' end of the template strand; it simply extends the 3' end of the template strand. The unique feature of telomerase is that it contains a built-in RNA template. After several telomere repeat units are added by telomerase, DNA polymerase catalyzes the synthesis of the complementary strand. Without telomerase activity, linear chromosomes would become progressively shorter. If the resulting terminal deletions extended into an essential gene or genes, this chromosome shortening would be lethal.

One change observed in many cancer cells is that the genes encoding telomerase are expressed, whereas they are not expressed in most somatic cells. Thus, one approach to cancer treatments has been to try to develop telomerase inhibitors, so that the chromosomes in cancer cells will lose their telomeres and the cells will die. However, other cancer cells do not contain active telomerase, making this approach problematic.

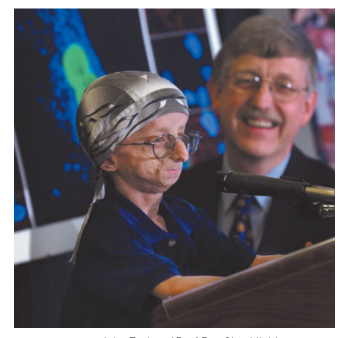
TELOMERE LENGTH AND AGING IN HUMANS

Unlike germ-line cells, most human somatic cells lack, or have very low levels of, telomerase activity. When human somatic cells are grown in culture, they divide only a limited number of times (usually only 20 to 70 cell generations) before senescence and death occur. When telomere lengths are measured in various somatic cell cultures, a correlation is observed

between telomere length and the number of cell divisions preceding senescence and death. Cells with longer telomeres survive longer—go through more cell divisions—than cells with shorter telomeres. As would be expected in the absence of telomerase activity, telomere length decreases as the age of the cell culture increases. Occasionally, somatic cells are observed to acquire the ability to proliferate in culture indefinitely, and these immortal cells have been shown to contain telomerase activity, unlike their progenitors. Since the one common feature of all cancers is uncontrolled cell division or immortality, scientists have proposed that one way to combat human cancers would be to inhibit the telomerase activity in cancer cells.

Further evidence of a relationship between telomere length and aging in humans has come from studies of individuals with disorders called progerias, inherited diseases characterized by premature aging. In the most severe form of progeria, Hutchinson–Gilford syndrome (**Figure 10.35**), senescence—wrinkles, baldness, and other symptoms of aging-begins immediately after birth, and death usually occurs in the teens. This syndrome is caused by a dominant mutation in the gene encoding lamin A, a protein involved in the control of the shape of nuclei in cells. Why this mutation leads to premature aging is unknown. In a less severe form of progeria, Werner syndrome, senescence begins in the teenage years, with death usually occurring in the 40s. Werner syndrome is caused by a recessive mutation in the WRN gene, which encodes a protein involved in DNA repair processes. Again, we still do not understand how the loss of this protein leads to premature aging. However, the somatic cells of individuals with both forms of progeria have short telomeres and exhibit decreased proliferative capacity when grown in culture, which is consistent with the hypothesis that decreasing telomere length contributes to the aging process.

At present, the relationship between telomere length and cell senescence is entirely correlative. There is no direct evidence indicating that telomere shortening causes aging. Nevertheless, the correlation is striking, and the hypothesis that telomere shortening contributes to the aging process in humans warrants further study.



■ FIGURE 10.35 John Tacket, 15, of Bay City, Michigan, speaks about his illness, progeria, during a news conference called in Washington, April 16, 2003, to announce the discovery of the gene that causes this rare, fatal genetic condition, characterized by the appearance of accelerated aging. To Tacket's right is Dr. Francis S. Collins, director of the National Institutes of Health.

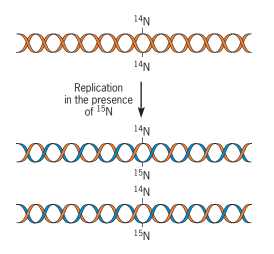
KEY POINTS

- **NTS** The large DNA molecules in eukaryotic chromosomes replicate bidirectionally from multiple origins.
 - Three DNA polymerases (α , δ , and ε) are present at each replication fork in eukaryotes.
 - Telomeres, the unique sequences at the ends of chromosomes, are added to chromosomes by a unique enzyme called telomerase.

Basic Exercises

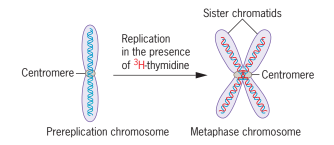
Illustrate Basic Genetic Analysis

- 1. *E. coli* cells that have been growing on normal medium containing ¹⁴N are transferred to medium containing only the heavy isotope of nitrogen, ¹⁵N, for one generation of growth. How will the ¹⁴N and ¹⁵N be distributed in the DNA of these bacteria after one generation?
- **Answer:** Because DNA replicates semiconservatively, the parental strands of DNA containing ¹⁴N will be conserved and used as templates to synthesize new complementary strands containing ¹⁵N. Thus, each DNA double helix will contain one light strand and one heavy strand, as shown in the following diagram.

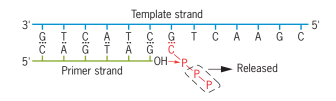


- 2. Radioactive (³H) thymidine is added to the culture medium in which a mouse cell is growing. This cell had not previously been exposed to any radioactivity. If the cell is entering S phase at the time the ³H-thymidine is added, what distribution of radioactivity will be present in the chromosomal DNA at the subsequent metaphase (the first metaphase after the addition of ³H-thymidine)?
- Answer: Remember that each prereplication chromosome contains a single giant DNA molecule extending from one end of the chromosome through the centromere all the way to the other end. This DNA molecule will replicate semiconservatively just like the DNA molecules in *E. coli* discussed above. However, at metaphase, the two progeny double

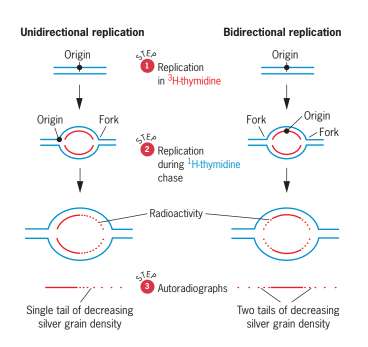
helices will be present in sister chromatids still joined at the centromere, as shown in the following diagram.



- **3.** DNA polymerases are only able to synthesize DNA in the presence of both a template strand and a primer strand. Why? What are the functions of these two strands?
- **Answer:** DNA polymerases can only extend DNA chains with a free 3'-OH because the mechanism of extension involves a nucleophilic attack by the 3'-OH on the interior phosphorus of the deoxyribonucleoside triphosphate precursor with the elimination of pyrophosphate. The strand with the 3'-OH is the primer strand; it is extended during synthesis. The template strand specifies the nucleotide sequence of the strand being synthesized; the new strand will be complementary to the template strand. These functions are illustrated as follows:



- 4. How can autoradiography be used to distinguish between uni- and bidirectional replication of DNA?
- **Answer:** If cells are grown in medium containing ³H-thymidine for a short period of time and are then transferred to nonradioactive medium for further growth (a pulse-chase experiment), uni- and bidirectional replication predict different labeling patterns, and these patterns can be distinguished by autoradiography, as shown here:

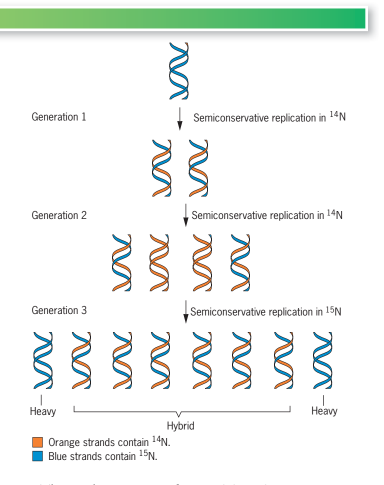


- **5.** Why do most somatic cells generally stop dividing after a limited number of cell divisions? What would happen if they kept dividing? How do cancer cells overcome this obstacle?
- **Answer:** Most somatic cells possess little or no telomerase activity. As a result, the telomeres of chromosomes become shorter during each cell division. If somatic cells kept dividing in the absence of telomerase, chromosomes would lose their telomeres, and, eventually, essential genes near the ends chromosomes would be lost, causing cell death. One of the essential steps in the conversion of a normal somatic cell to a cancer cell is turning on or increasing the synthesis of telomerase so that telomeres are not lost during the uncontrolled cell divisions of cancer cells.

Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. Escherichia coli cells were grown for many generations in a medium in which the only available nitrogen was the heavy isotope ¹⁵N. The cells were then collected by centrifugation, washed with a buffer, and transferred to a medium containing ¹⁴N (the normal light nitrogen isotope). After two generations of growth in the ¹⁴Ncontaining medium, the cells were transferred back to ¹⁵N-containing medium for one final generation of growth. After this final generation of growth in the presence of ¹⁵N, the cells were collected by centrifugation. The DNA was then extracted from these cells and analyzed by CsCl equilibrium density-gradient centrifugation. How would you expect the DNA from these cells to be distributed in the gradient?
- Answer: Meselson and Stahl demonstrated that DNA replication in E. coli is semiconservative. Their control experiments showed that DNA double helices with (1) ¹⁴N in both strands, (2) ¹⁴N in one strand and ¹⁵N in the other strand, and (3) ¹⁵N in both strands separated into three distinct bands in the gradient, called (1) the light band, (2) the hybrid band, and (3) the heavy band, respectively. If you start with a DNA double helix with ¹⁵N in both strands, and replicate it semiconservatively for two generations in the presence of ¹⁴N and then for one generation in the presence of ¹⁵N, you will end up with eight DNA molecules, two with ¹⁵N in both strands and six with ¹⁴N in one strand and ¹⁵N in the other strand, as shown in the following diagram. Therefore, 75 percent (6/8) of the DNA will appear in the hybrid band, and 25 percent (2/8) will appear in the heavy band.



2. The X chromosome of *Drosophila melanogaster* contains a giant DNA molecule 22,422,827 nucleotide pairs long. During the early cleavage stages of embryonic development, nuclear division takes only 10 minutes. If each replication fork travels at the rate of 2600 nucleotide pairs per minute, how many replication forks would be required to replicate the entire X chromosome in 10 minutes? Assume that these replication forks are evenly spaced along the DNA molecule.

Cell division occurs more slowly in the somatic cells of the adult fruit fly. If you are studying somatic cells with a generation time of 20 hours and an S phase of 8 hours, how many replication forks would be needed to complete the replication of the X chromosome during the S phase of mitosis?

If the average size of Okazaki fragments in *Drosophila* is 250 nucleotides, how many Okazaki fragments are synthesized during the replication of the X chromosome? How many RNA primers are needed?

Answer: If a replication fork moves at the rate of 2600 nucleotide pairs per minute, it will traverse 26,000 nucleotide pairs during 10 minutes and catalyze the synthesis of DNA chains 26,000 nucleotides long in each of the two daughter double helices. Given the presence of 22,422,827 nucleotide pairs in the X chromosome and the replication of 26,000 nucleotide pairs by each replication

fork in 10 minutes, the complete replication of the DNA in this chromosome during the cleavage stages of embryonic development would require 862 replication forks (22,422,827 nucleotide pairs/26,000 nucleotide pairs replicated per fork in 10 minutes) evenly spaced along the DNA molecule.

Similarly, in the case of the somatic cells of the adult fly with an S phase of 8 hours, 18 replication forks would need to be evenly spaced along the DNA in the X chromosome to complete replication in 8 hours. One replication fork would replicate 1,248,000 nucleotide pairs in 8 hours (2600 nucleotide pairs per minute \times 480 minutes). Therefore, if replication forks were evenly spaced, 18 of them could replicate the DNA molecule in the X chromosome in 8 hours (22,422,827 nucleotide pairs/1,248,000 nucleotide pairs per fork per 8 hours).

The replication of the giant DNA molecule in the X chromosome of *Drosophila* will require the synthesis and subsequent joining of 89,691 Okazaki fragments (22,422,827 nucleotide pairs/250 nucleotides per Okazaki fragment). It will also require the synthesis of 89,691 RNA primers because the synthesis of each Okazaki fragment is initiated with an RNA primer.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

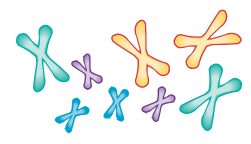
- **10.1** DNA polymerase I of *E. coli* is a single polypeptide of molecular weight 103,000.
 - (a) What enzymatic activities other than polymerase activity does this polypeptide possess?
 - (b) What are the *in vivo* functions of these activities?
 - (c) Are these activities of major importance to an *E. coli* cell? Why?
- **10.2 (c)** *Escherichia coli* cells are grown for many generations in a medium in which the only available nitrogen is the heavy isotope ¹⁵N. They are then transferred to a medium containing ¹⁴N as the only source of nitrogen.
 - (a) What distribution of ¹⁵N and ¹⁴N would be expected in the DNA molecules of cells that had grown for one generation in the ¹⁴N-containing medium assuming that DNA replication was (i) conservative, (ii) semiconservative, or (iii) dispersive?
 - (b) What distribution would be expected after two generations of growth in the ¹⁴N-containing medium assuming
 (i) conservative, (ii) semiconservative, or (iii) dispersive replication?
- **10.3** Why do DNA molecules containing ¹⁵N band at a different position than DNA molecules containing ¹⁴N when centrifuged to equilibrium in *6M* CsCl?
- **10.4** A DNA template plus primer with the structure

3' P —TGCGAATTAGCGACAT— P 5' 5' P —ATCGGTACGACGCTTAAC—OH 3'

(where P = a phosphate group) is placed in an *in vitro* DNA synthesis system (Mg²⁺, an excess of the four deoxyribonucleoside triphosphates, etc.) containing a mutant form of *E. coli* DNA polymerase I that lacks $5' \rightarrow 3'$ exonuclease activity. The $5' \rightarrow 3'$ polymerase and $3' \rightarrow 5'$ exonuclease activities of this aberrant enzyme are identical to those of normal *E. coli* DNA polymerase I. It simply has no $5' \rightarrow 3'$ exonuclease activity.

- (a) What will be the structure of the final product?
- (b) What will be the first step in the reaction sequence?
- **10.5** How might continuous and discontinuous modes of DNA replication be distinguished experimentally?
- **10.6** *E. coli* cells contain five different DNA polymerases— I, II, III, IV, and V. Which of these enzymes catalyzes the semiconservative replication of the bacterial chromosome during cell division? What are the functions of the other four DNA polymerases in *E. coli*?
- **10.7** The Boston barberry is an imaginary plant with a diploid chromosome number of 4, and Boston barberry cells are easily grown in suspended cell cultures. ³H-Thymidine was added to the culture medium in which a G₁-stage cell of this

plant was growing. After one cell generation of growth in ³H-thymidine-containing medium, colchicine was added to the culture medium. The medium now contained both ³H-thymidine and colchicine. After two "generations" of growth in ³H-thymidine-containing medium (the second "generation" occurring in the presence of colchicine as well), the two progeny cells (each now containing eight chromosomes) were transferred to culture medium containing non-radioactive thymidine (¹H-thymidine) plus colchicine. Note that a "generation" in the presence of colchicine consists of a normal cell cycle's chromosomal duplication but no cell division. The two progeny cells were allowed to continue to grow, proceeding through the "cell cycle," until each cell contained a set of metaphase chromosomes that looked like the following.



If autoradiography were carried out on these metaphase chromosomes (four large plus four small), what pattern of radioactivity (as indicated by silver grains on the autoradiograph) would be expected? (Assume no recombination between DNA molecules.)

- **10.8** Suppose that the experiment described in Problem 10.7 was carried out again, except this time replacing the ³H-thymidine with nonradioactive thymidine at the same time that the colchicine was added (after one cell generation of growth in ³H-thymidine-containing medium). The cells were then maintained in colchicine plus nonradioactive thymidine until the metaphase shown in Problem 10.7 occurred. What would the autoradiographs of these chromosomes look like?
- **10.9** Suppose that the DNA of cells (growing in a cell culture) in a eukaryotic species was labeled for a short period of time by the addition of ³H-thymidine to the medium. Next assume that the label was removed and the cells were resuspended in nonradioactive medium. After a short period of growth in nonradioactive medium, the DNA was extracted from these cells, diluted, gently layered on filters, and autoradiographed. If autoradiographs of the type

were observed, what would this indicate about the nature of DNA replication in these cells? Why?

_ _

10.10 Arrange the following enzymes in the order of their action during DNA replication in *E. coli*: (1) DNA polymerase I, (2) DNA polymerase III, (3) DNA primase, (4) DNA gyrase, and (5) DNA helicase.

- **10.11** Fifteen distinct DNA polymerases—α, β, γ, δ, ε, κ, ζ, η, θ, κ, λ, μ, σ, φ, and Rev1—have been characterized in mammals. What are the intracellular locations and functions of these polymerases?
- 10.12 a The E. coli chromosome contains approximately 4×10^6 nucleotide pairs and replicates as a single bidirectional replicon in approximately 40 minutes under a wide variety of growth conditions. The largest chromosome of D. melanogaster contains about 6×10^7 nucleotide pairs. (a) If this chromosome contains one giant molecule of DNA that replicates bidirectionally from a single origin located precisely in the middle of the DNA molecule, how long would it take to replicate the entire chromosome if replication in Drosophila occurred at the same rate as replication in E. coli? (b) Actually, replication rates are slower in eukaryotes than in prokaryotes. If each replication bubble grows at a rate of 5000 nucleotide pairs per minute in Drosophila and 100,000 nucleotide pairs per minute in E. coli, how long will it take to replicate the largest Drosophila chromosome if it contains a single bidirectional replicon as described in (a) above? (c) During the early cleavage divisions in Drosophila embryos, the nuclei divide every 9 to 10 minutes. Based on your calculations in (a) and (b) above, what do these rapid nuclear divisions indicate about the number of replicons per chromosome in Drosophila?
- **10.13** *E. coli* cells that have been growing in ¹⁴N for many generations are transferred to medium containing only ¹⁵N and allowed to grow in this medium for four generations. Their DNA is then extracted and analyzed by equilibrium CsCl density-gradient centrifugation. What proportion of this DNA will band at the "light," "hybrid," and "heavy" positions in the gradient?
- 10.14 The bacteriophage lambda chromosome has several AT-rich segments that denature when exposed to pH 11.05 for 10 minutes. After such partial denaturation, the linear packaged form of the lambda DNA molecule has the structure shown in Figure 10.9a. Following its injection into an E. coli cell, the lambda DNA molecule is converted to a covalently closed circular molecule by hydrogen bonding between its complementary singlestranded termini and the action of DNA ligase. It then replicates as a θ -shaped structure. The entire lambda chromosome is 17.5 µm long. It has a unique origin of replication located 14.3 µm from the left end of the linear form shown in Figure 10.9a. Draw the structure that would be observed by electron microscopy after both (1) replication of an approximately 6-µm-long segment of the lambda chromosomal DNA molecule (in vivo) and (2) exposure of this partially replicated DNA molecule to pH 11.05 for 10 minutes (in vitro), (a) if replication had proceeded bidirectionally from the origin and (b) if replication had proceeded unidirectionally from the origin.
- **10.15** What enzyme activity catalyzes each of the following steps in the semiconservative replication of DNA in pro-karyotes?

- (a) The formation of negative supercoils in progeny DNA molecules.
- (b) The synthesis of RNA primers.
- (c) The removal of RNA primers.
- (d) The covalent extension of DNA chains at the 3'-OH termini of primer strands.
- (e) Proofreading of the nucleotides at the 3'-OH termini of DNA primer strands.
- **10.16** One species of tree has a very large genome consisting of 2.0×10^{10} base pairs of DNA.
 - (a) If this DNA was organized into a single linear molecule, how long (meters) would this molecule be?
 - (b) If the DNA is evenly distributed among 10 chromosomes and each chromosome has one origin of DNA replication, how long would it take to complete the S phase of the cell cycle, assuming that DNA polymerase can synthesize 2 × 10⁴ bp of DNA per minute?
 - (c) An actively growing cell can complete the S phase of the cell cycle in approximately 300 minutes. Assuming that the origins of replication are evenly distributed, how many origins of replication are present on each chromosome?
 - (d) What is the average number of base pairs between adjacent origins of replication?
- **10.17** Why must each of the giant DNA molecules in eukaryotic chromosomes contain multiple origins of replication?
- **10.18** In *E. coli*, viable *polA* mutants have been isolated that produce a defective gene product with little or no $5' \rightarrow 3'$ polymerase activity, but normal $5' \rightarrow 3'$ exonuclease activity. However, no *polA* mutant has been identified that is completely deficient in the $5' \rightarrow 3'$ exonuclease activity, while retaining $5' \rightarrow 3'$ polymerase activity, of DNA polymerase I. How can these results be explained?
- **10.19** Other *polA* mutants of *E. coli* lack the $3' \rightarrow 5'$ exonuclease activity of DNA polymerase I. Will the rate of DNA synthesis be altered in these mutants? What effect(s) will these *polA* mutations have on the phenotype of the organism?
- **10.20** Many of the origins of replication that have been characterized contain AT-rich core sequences. Are these AT-rich cores of any functional significance? If so, what?
- 10.21 (a) Why isn't DNA primase activity required to initiate rolling-circle replication? (b) DNA primase is required for the discontinuous synthesis of the lagging strand, which occurs on the single-stranded tail of the rolling circle. Why?
- **10.22** DNA polymerase I is needed to remove RNA primers during chromosome replication in *E. coli*. However, DNA polymerase III is the true replicase in *E. coli*. Why doesn't DNA polymerase III remove the RNA primers?
- **10.23** In *E. coli*, three different proteins are required to unwind the parental double helix and keep the unwound strands in an extended template form. What are these proteins, and what are their respective functions?
- **10.24** How similar are the structures of DNA polymerase I and DNA polymerase III in *E. coli*? What is the structure of

the DNA polymerase III holoenzyme? What is the function of the *dnaN* gene product in *E. coli*?

- **10.25** The *dnaA* gene product of *E. coli* is required for the initiation of DNA synthesis at *oriC*. What is its function? How do we know that the DnaA protein is essential to the initiation process?
- **10.26** What is a primosome, and what are its functions? What essential enzymes are present in the primosome? What are the major components of the *E. coli* replisome? How can geneticists determine whether these components are required for DNA replication?
- **10.27** The chromosomal DNA of eukaryotes is packaged into nucleosomes during the S phase of the cell cycle. What obstacles do the size and complexity of both the replisome and the nucleosome present during the semiconservative replication of eukaryotic DNA? How might these obstacles be overcome?
- 10.28 Two mutant strains of *E. coli* each have a temperaturesensitive mutation in a gene that encodes a product required for chromosome duplication. Both strains replicate their DNA and divide normally at 25°C, but are unable to replicate their DNA or divide at 42°C. When cells of one strain are shifted from growth at 25°C to growth at 42°C, DNA synthesis stops immediately. When cells of the other strain are subjected to the same temperature shift, DNA synthesis continues, albeit at a decreasing rate, for about a half hour. What can you conclude about the functions of the products of these two genes?
- **10.29** In what ways does chromosomal DNA replication in eukaryotes differ from DNA replication in prokaryotes?
- 10.30 (a) The chromosome of the bacterium Salmonella typhimurium contains about 4 × 10⁶ nucleotide pairs. Approximately how many Okazaki fragments are produced during one complete replication of the *S. typhimurium* chromosome? (b) The largest chromosome of *D. melanogaster* contains approximately 6 × 10⁷ nucleotide pairs. About how many Okazaki fragments are produced during the replication of this chromosome?
- **10.31** In the yeast *S. cerevisiae*, haploid cells carrying a mutation called *est1* (for *ever-shorter telomeres*) lose distal telomere sequences during each cell division. Predict the ultimate phenotypic effect of this mutation on the progeny of these cells.
- **10.32** Assume that the sequence of a double-stranded DNA shown in the following diagram is present at one end of a large DNA molecule in a eukaryotic chromosome.

5'-(centromere sequence)-GATTCCCCGGGAAGCTTGGGGGGGCCCATCTTCGTACGTCTTTGCA-3' 3'-(centromere sequence)-CTAAGGGGCCCTTCGAACCCCCCGGGTAGAAGCATGCAGAAACGT-5'

You have reconstituted a eukaryotic replisome that is active *in vitro*. However, it lacks telomerase activity. If you isolate the DNA molecule shown above and replicate it in your *in vitro* system, what products would you expect?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

- 1. DNA polymerase III catalyzes the semiconservative replication of the chromosome in *E. coli*. How many genes encode structural proteins of DNA polymerase III in *E. coli* strain K12? Which genes encode which subunits? Are these genes clustered to a specific region of the *E. coli* chromosome, or are they distributed throughout the chromosome? How large is the gene encoding the alpha subunit of DNA polymerase III in *E. coli* K12?
- 2. A single gene encodes DNA polymerase I in *E. coli*. What is the name of this gene? How large is the gene? Where is it located on the *E. coli* chromosome? What is the molecular weight of DNA polymerase I? How many amino acids does it contain?

Hint: At the NCBI web site, click Entrez Home \rightarrow Entrez Gene \rightarrow Search using protein name and organism, namely, DNA polymerase III AND *Escherichia K12*[orgn]. If you do not limit your search to strain K12, your search results will include the same genes for all of the other *E. coli* strains that have been sequenced. So, for the sake of simplicity, it is best to include K12 in the organism designator. In the search results, click Primary Source "Ecogene" for more information, including nucleotide coordinates and map position of the gene, protein size, and similar information.

Transcription and RNA Processing

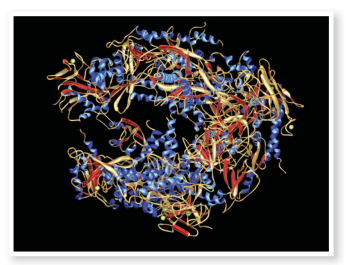
CHAPTER OUTLINE

11

- Transfer of Genetic Information: The Central Dogma
- The Process of Gene Expression
- Transcription in Prokaryotes
- Transcription and RNA Processing in Eukaryotes
- Interrupted Genes in Eukaryotes: Exons and Introns
- Removal of Intron Sequences by RNA Splicing

Storage and Transmission of Information with Simple Codes

We live in the age of the computer. It has an impact on virtually all aspects of our lives, from driving to work to watching spaceships land on the moon. These electronic wizards can store, retrieve, and analyze data with lightning-like speed. The "brain" of the computer is a small chip of silicon, the microprocessor, which contains a sophisticated and integrated array of electronic circuits capable of responding almost instantaneously to coded bursts of electrical energy. In carrying out its amazing feats, the computer uses a binary code, a language based on 0's and 1's. Thus, the alphabet used by computers is like that of the Morse code (dots and dashes) used in telegraphy. Both consist of only two symbols—in marked contrast to the 26 letters of the English alphabet. Obviously, if the computer can perform its wizardry with a binary alphabet, vast amounts of information can be stored and retrieved without using complex codes or lengthy alphabets. In this and the following chapter, we examine (1) how the genetic information of living creatures is written in an alphabet with just four letters, the four base pairs in DNA, and (2) how this genetic



Computer model of the structure of RNA polymerase II, which catalyzes transcription of nuclear genes in eukaryotes.

information is expressed during the growth and development of an organism. We will see that RNA plays a key role in the process of gene expression.

Transfer of Genetic Information: The Central Dogma

Flow of genetic information:

According to the central dogma of molecular biology, genetic information usually flows (1) from DNA to DNA during its transmission from generation to generation and (2) from DNA to protein during its phenotypic expression in an organism

(**Figure 11.1**). During the replication of RNA viruses, information is also transmitted from RNA to RNA. The transfer of genetic information from DNA to protein involves two steps: (1) **transcription**, the transfer of the genetic information from DNA to RNA, and (2) **translation**, the transfer of information from RNA to protein. In addition, genetic information flows from RNA to DNA during the conversion of the genomes of RNA tumor viruses to their DNA proviral forms (Chapter 21). Thus, the transfer of genetic information from RNA to R

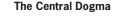
TRANSCRIPTION AND TRANSLATION

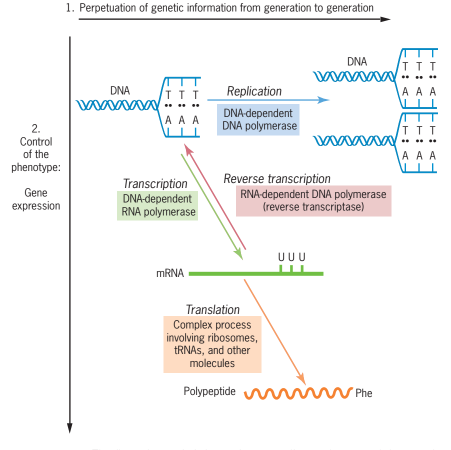
As we discussed earlier, the expression of genetic information occurs in two steps: transcription and translation (Figure 11.1). During transcription, one strand of DNA of a gene is used as a template to synthesize a complementary strand of RNA, called

the gene transcript. For example, in Figure 11.1, the DNA strand containing the nucleotide sequence AAA is used as a template to produce the complementary sequence UUU in the RNA transcript. During translation, the sequence of nucleotides in the RNA transcript is converted into the sequence of amino acids in the polypeptide gene product. This conversion is governed by the genetic code, the specification of amino acids by nucleotide triplets called codons in the gene transcript. For example, the UUU triplet in the RNA transcript shown in Figure 11.1 specifies the amino acid phenylalanine (Phe) in the polypeptide gene product. Translation takes place on intricate macromolecular machines called ribosomes, which are composed of three to five RNA molecules and 50 to 90 different proteins. However, the process of translation also requires the participation of many other macromolecules. This chapter focuses on transcription; translation is the subject of Chapter 12.

The RNA molecules that are translated on ribosomes are called messenger RNAs (mRNAs). In prokaryotes, the product of transcription, the primary transcript, usually is equivalent to the mRNA molecule (**Figure 11.2a**). In eukaryotes, primary transcripts often must be processed by the excision of specific sequences and the modification of both termini before they can be translated (**Figure 11.2***b*). Thus, in eukaryotes, primary transcripts usually are precursors to mRNAs and, as such, are called **pre-mRNAs**. Most of the nuclear genes in higher eukaryotes and some in lower eukaryotes contain noncoding sequences called *introns* that separate the expressed sequences or exons of these genes. The entire sequences of these split genes are transcribed into pre-mRNAs, and the noncoding intron sequences are subsequently removed by splicing reactions carried out on macromolecular structures called *spliceosomes*.

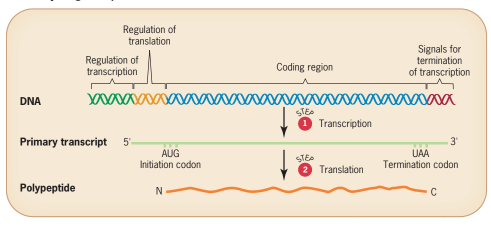
The central dogma of biology is that information stored in DNA is transferred to RNA molecules during transcription and to proteins during translation.



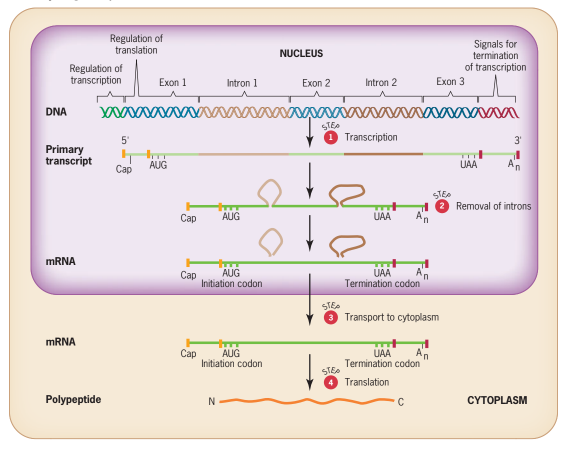


■ FIGURE 11.1 The flow of genetic information according to the central dogma of molecular biology. Replication, transcription, and translation occur in all organisms; reverse transcription occurs in cells infected with certain RNA viruses. Not shown is the transfer of information from RNA to RNA during the replication of RNA viruses.

(a) Prokaryotic gene expression



(b) Eukaryotic gene expression



■ FIGURE 11.2 Gene expression involves two steps: transcription and translation, in both prokaryotes (*a*) and eukaryotes (*b*). In eukaryotes, the primary transcripts or pre-mRNAs often must be processed by the excision of introns and the addition of 5' 7-methyl guanosine caps (CAP) and 3' poly(A) tails [(A)_n]. In addition, eukaryotic mRNAs must be transported from the nucleus to the cytoplasm where they are translated.

FIVE TYPES OF RNA MOLECULES

Five different classes of RNA molecules play essential roles in gene expression. We have already discussed messenger RNAs, the intermediaries that carry genetic information from DNA to the ribosomes where proteins are synthesized. Transfer RNAs (tRNAs) are small RNA molecules that function as adaptors between amino acids and the codons in mRNA during translation. Ribosomal RNAs (rRNAs) are structural and

catalytic components of the ribosomes, the intricate machines that translate nucleotide sequences of mRNAs into amino acid sequences of polypeptides. **Small nuclear RNAs** (snRNAs) are structural components of spliceosomes, the nuclear organelles that excise introns from gene transcripts. Micro RNAs (miRNAs) are short 20- to 22-nucleotide single-stranded RNAs that are cleaved from small hairpin-shaped precursors and block the expression of complementary or partially complementary mRNAs by either causing their degradation or repressing their translation. The roles of mRNAs and snRNAs are discussed in this chapter. The structures and functions of tRNAs and rRNAs will be discussed in detail in Chapter 12. The mechanisms by which miRNAs regulate gene expression are discussed in Chapter 19.

All five types of RNA—mRNA, tRNA, rRNA, snRNA, and miRNA—are produced by transcription. Unlike mRNAs, which specify polypeptides, the final products of tRNA, rRNA, snRNA, and miRNA genes are RNA molecules. Transfer RNA, ribosomal RNA, snRNA, and miRNA molecules are not translated. ■ Figure 11.3 shows an overview of gene expression in eukaryotes, emphasizing the transcriptional origin and functions of the five types of RNA molecules. The process is similar in prokaryotes. However, in prokaryotes, the DNA is not separated from the ribosomes by a nuclear envelope. In addition, prokaryotic genes seldom contain noncoding sequences that are removed during RNA transcript processing.

- The central dogma of molecular biology is that genetic information flows from DNA to DNA during chromosome replication, from DNA to RNA during transcription, and from RNA to protein during translation.
- Transcription involves the synthesis of an RNA transcript complementary to one strand of DNA of a gene.
- Translation is the conversion of information stored in the sequence of nucleotides in the RNA transcript into the sequence of amino acids in the polypeptide gene product, according to the specifications of the genetic code.

KEY POINTS

The Process of Gene Expression

How do genes control the phenotype of an organism? How do the nucleotide sequences of genes direct the growth and development of a cell, a tissue, an organ, or an entire living creature? Geneticists know that the phenotype of an organism is produced by the combined effects of all its genes acting within the constraints imposed by the environment. They also know that the Information stored in the nucleotide sequences of genes is translated into the amino acid sequences of proteins through unstable intermediaries called messenger RNAs.

number of genes in an organism varies over an enormous range, with gene number increasing with the developmental complexity of the species. The RNA genomes of the smallest viruses such as phage MS2 contain only four genes, whereas large viruses such as phage T4 have about 200 genes. Bacteria such as *E. coli* have approximately 4000 genes, and mammals, including humans, have about 20,500 genes. In this and the following chapter, we focus on the mechanisms by which genes direct the synthesis of their products, namely, RNAs and proteins. The mechanisms by which these gene products collectively control the phenotypes of mature organisms are discussed in subsequent chapters, especially Chapter 20.

AN mRNA INTERMEDIARY

If most of the genes of a eukaryote are located in the nucleus, and if proteins are synthesized in the cytoplasm, how do these genes control the amino acid sequences of their protein products? The genetic information stored in the sequences of nucleotide

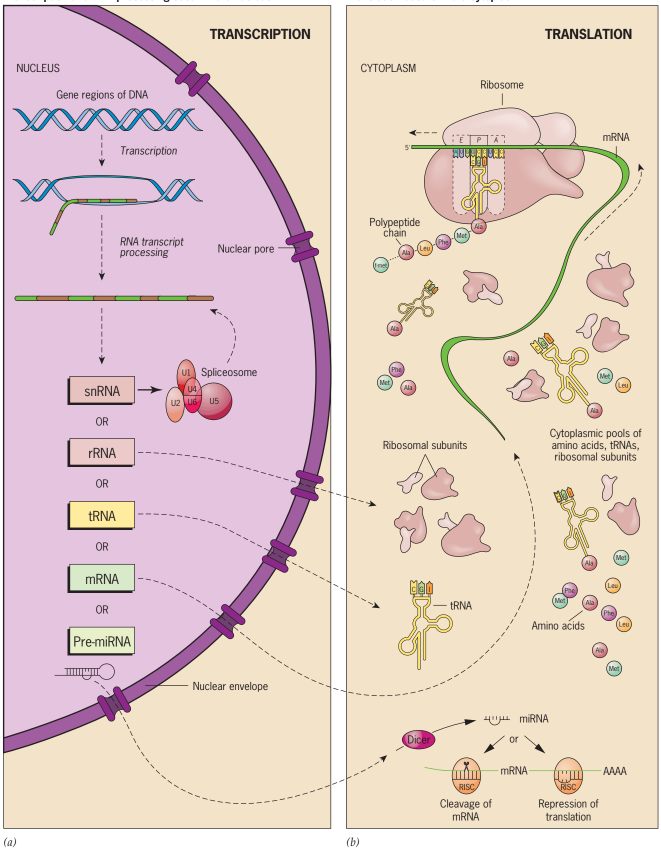


FIGURE 11.3 An overview of gene expression, emphasizing the transcriptional origin of miRNA, snRNA, tRNA, rRNA, and mRNA, the splicing function of snRNA, the regulation of gene expression by miRNA, and the translational roles of tRNA, rRNA, mRNA, and ribosomes. Dicer is a nuclease that processes the miRNA precursor into miRNA, and RISC is the *R*NA-*i*nduced silencing complex.

Transcription and RNA processing occur in the nucleus.

Translation occurs in the cytoplasm.

pairs in genes must somehow be transferred to the sites of protein synthesis in the cytoplasm. Messengers are needed to transfer genetic information from the nucleus to the cytoplasm. Although the need for such messengers is most obvious in eukaryotes, the first evidence for their existence came from studies of prokaryotes. Some of the early evidence for the existence of short-lived messenger RNAs is discussed in Appendix D: Evidence for an Unstable Messenger RNA.

GENERAL FEATURES OF RNA SYNTHESIS

RNA synthesis occurs by a mechanism that is similar to that of DNA synthesis (Chapter 10) except that (1) the precursors are *ribo-nucleoside triphosphates* rather than deoxyribonucleoside triphosphates, (2) only one strand of DNA is used as a template for the synthesis of a complementary RNA chain in any given region, and (3) RNA chains can be initiated *de novo*, without any requirement for a preexisting primer strand. The RNA molecule produced will be complementary and antiparallel to the DNA **template strand** and identical, except that uridine residues replace thymidines, to the DNA **nontemplate strand**

(**Figure 11.4**). If the RNA molecule is an mRNA, it will specify amino acids in the protein gene product. Therefore, mRNA molecules are coding strands of RNA. They are also called **sense strands** of RNA because their nucleotide sequences "make sense" in that they specify sequences of amino acids in the protein gene products.

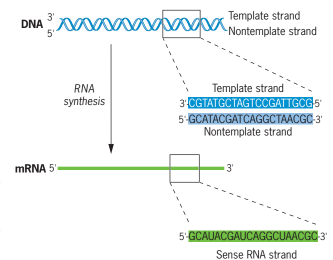
An RNA molecule that is complementary to an mRNA is referred to as **antisense RNA**. This terminology is sometimes extended to the two strands of DNA. However, usage of the terms *sense* and *antisense* to denote DNA strands has been inconsistent. Thus, we will use *template strand* and *nontemplate strand* to refer to the transcribed and nontranscribed strands, respectively, of a gene.

The synthesis of RNA chains, like DNA chains, occurs in the 5' \rightarrow 3' direction, with the addition of ribonucleotides to the 3'-hydroxyl group at the end of the chain (**■** Figure 11.5). The reaction involves a nucleophilic attack by the 3'-OH on the nucleotidyl (interior) phosphorus atom of the ribonucleoside triphosphate precursor with the elimination of pyrophosphate, just as in DNA synthesis. This reaction is catalyzed by enzymes called **RNA polymerases.** The overall reaction is as follows:

$$n(\text{RTP}) \xrightarrow[\text{RNA polymerase}]{\text{RNA polymerase}} (\text{RMP})_n + n(\text{PP})$$

where n is the number of moles of ribonucleotide triphosphate (RTP) consumed, ribonucleotide monophosphate (RMP) incorporated into RNA, and pyrophosphate (PP) produced.

RNA polymerases bind to specific nucleotide sequences called **promoters**, and with the help of proteins called transcription factors, initiate the synthesis of RNA molecules at transcription start sites near the promoters. The promoters in eukaryotes are typically more complex than those of prokaryotes. A single RNA polymerase carries out all transcription in most prokaryotes, whereas five different RNA polymerases are present in eukaryotes, with each polymerase responsible for the synthesis of a distinct class of RNAs. RNA synthesis takes place within a locally unwound



■ **FIGURE 11.4** RNA synthesis utilizes only one DNA strand of a gene as template.

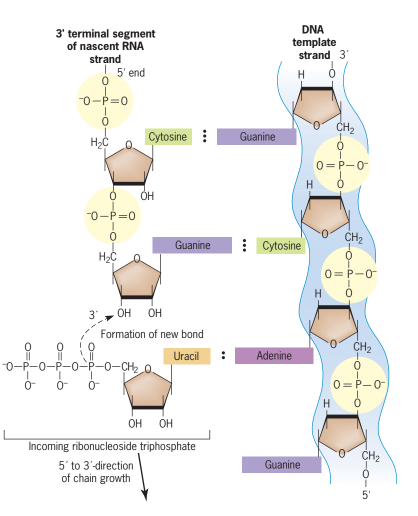
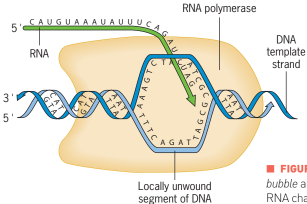


FIGURE 11.5 The RNA chain elongation reaction catalyzed by RNA polymerase.



segment of DNA, sometimes called a **transcription bubble**, which is produced by RNA polymerase (**Figure 11.6**). The nucleotide sequence of an RNA molecule is complementary to that of its DNA template strand, and RNA synthesis is governed by the same base-pairing rules as DNA synthesis, but uracil replaces thymine. As a result, the origin of RNA transcripts can be determined by studying their hybridization to DNAs from different sources such as the chromosome(s) of the cell, viruses, and other infectious organisms (see Problem-Solving Skills: Distinguishing RNAs Transcribed from Viral and Host DNAs).

FIGURE 11.6 RNA synthesis occurs within a locally unwound segment of DNA. This *transcription bubble* allows a few nucleotides in the template strand to base-pair with the growing end of the RNA chain. The unwinding and rewinding of the DNA molecule are catalyzed by RNA polymerase.

PROBLEM-SOLVING SKILLS

Distinguishing RNAs Transcribed from Viral and Host DNAs

THE PROBLEM

E. coli cells that have been infected with a virus present the opportunity for the cells to make two types of RNA transcripts: bacterial and viral. If the virus is a lytic bacteriophage such as T4, only viral transcripts are made; if it is a nonlytic bacteriophage such as M13, both viral and bacterial transcripts are made; and if it is a quiescent prophage such as lambda, only bacterial transcripts are made. Suppose that you have just identified a new DNA virus. How could you determine which types of RNA transcripts are made in cells infected with this virus?

FACTS AND CONCEPTS

- 1. During the first step in gene expression (transcription), one strand of DNA is used as a template for the synthesis of a complementary strand of RNA.
- **2.** RNA can be labeled with ³H by growing cells in medium containing ³H-uridine.
- **3.** DNA can be denatured—separated into its constituent single strands—by exposing it to high temperature or high pH.
- 4. Viral DNAs and host cell DNAs can both be purified, denatured, and bound to membranes for use in subsequent hybridization experiments (see Figure 1 in Appendix D: Evidence for an Unstable Messenger RNA).
- **5.** Under the appropriate conditions, complementary single-stranded RNA and DNA molecules will form stable double helices *in vitro*.

ANALYSIS AND SOLUTION

The source of the RNA transcripts being synthesized in virusinfected cells can be determined by incubating the infected cells for a short period of time in medium containing ³H-uridine, purifying the RNA from these cells, and then hybridizing it to single-stranded viral and bacterial DNAs.

a. You should prepare one membrane with denatured viral DNA bound to it, a second membrane with denatured host DNA bound to it, and a third membrane with no DNA to serve as a control to measure nonspecific binding of ³H-labeled RNA.

- **b.** You should then prepare an appropriate hybridization solution and place the three membranes—one with viral DNA, one with host DNA, and one with no DNA—in this solution.
- c. You next add a sample of the purified ³H-labeled RNA and allow it to hybridize with the DNA on the membranes. Then you wash the membranes thoroughly to remove any nonhybridized RNA. The RNA that remains has either bound specifically to DNA on the membrane or it has bound nonspecifically to the membrane itself. The extent of the RNA binding can be determined by measuring how radioactive each membrane is.
- **d.** Radioactivity on the membrane that had no DNA represents nonspecific "background" binding of RNA to the membrane. This radioactivity can be subtracted from the levels of radioactivity on the other two membranes to measure the specific binding of RNA to viral or bacterial DNA. The results will tell you whether the labeled transcripts were synthesized from viral DNA templates, bacterial DNA templates, or both. With phage T4-infected cells, phage M13-infected cells, and cells containing lambda prophages the results might be summarized as follows. (The plus signs indicate the presence of RNA transcripts that hybridize specifically.)

	RNA Hybridized to Membrane Containing	
	E. coli DNA	Phage DNA
Phage T4-infected <i>E. coli</i> cells	_	+
Phage M13-infected <i>E. coli</i> cells	+	+
<i>E. coli</i> cells carrying lambda prophages	+	_

Which pattern do you observe in cells infected with the newly discovered virus?

For further discussion visit the Student Companion site.

- In eukaryotes, genes are present in the nucleus, whereas polypeptides are synthesized in the cytoplasm.
- Messenger RNA molecules function as intermediaries that carry genetic information from DNA to the ribosomes, where proteins are synthesized.
- RNA synthesis, catalyzed by RNA polymerases, is similar to DNA synthesis in many respects.
- RNA synthesis occurs within a localized region of strand separation, and only one strand of DNA functions as a template for RNA synthesis.

KEY POINTS

Transcription in Prokaryotes

The basic features of transcription are the same in both prokaryotes and eukaryotes, but many of the details—such as the promoter sequences—are different. The RNA polymerase of *E. coli* has been studied in great detail and will be discussed here. It catalyzes all RNA synthesis in this species. The RNA polymerases of archaea have quite different structures; they will not be discussed here. Transcription—the first step in gene expression transfers the genetic information stored in DNA (genes) into messenger RNA molecules that carry the information to the ribosomes—the sites of protein synthesis—in the cytoplasm.

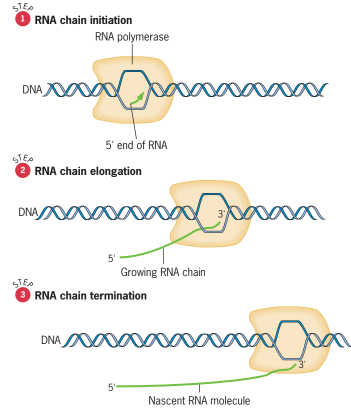
A segment of DNA that is transcribed to produce one RNA molecule is called a **transcription unit**. Transcription units may be equivalent to individual genes, or they may include several contiguous genes. Large transcripts that carry the coding sequences of several genes are common in bacteria. The process of transcription can be divided into three stages: (1) **initiation** of a new RNA chain, (2) **elongation** of the chain, and (3) **termination** of transcription and release of the nascent RNA molecule (**■ Figure 11.7**).

When discussing transcription, biologists often use the terms *upstream* and *down-stream* to refer to regions located toward the 5' end and the 3' end, respectively, of the transcript from some site in the mRNA molecule. These terms are based on the fact that RNA synthesis always occurs in the 5' to 3' direction. Upstream and downstream regions of genes are the DNA sequences specifying the corresponding 5' and 3' segments of their transcripts relative to a specific reference point.

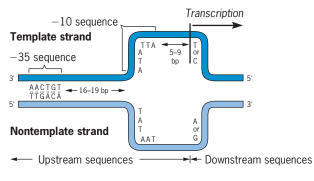
RNA POLYMERASES: COMPLEX ENZYMES

The RNA polymerases that catalyze transcription are complex, multimeric proteins. The *E. coli* RNA polymerase has a molecular weight of about 480,000 and consists of five polypeptides. Two of these are identical; thus, the enzyme contains four distinct polypeptides. The complete RNA polymerase molecule, the **holoenzyme**, has the composition $\alpha_2\beta\beta'\sigma$. The α subunits are involved in the assembly of the *tetrameric core* ($\alpha_2\beta\beta'$) of RNA polymerase. The β subunit contains the ribonucleoside triphosphate binding site, and the β' subunit harbors the DNA template-binding region.

One subunit, the **sigma** (σ) **factor**, is involved only in the initiation of transcription; it plays no role in chain elongation. After RNA chain initiation has occurred, the σ factor is released, and chain elongation (see Figure 11.5) is catalyzed by the core enzyme ($\alpha_2\beta\beta'$). The function of sigma is to recognize and bind RNA polymerase to the transcription initiation or promoter sites in DNA. The core enzyme (with no σ) will catalyze RNA synthesis from DNA templates *in vitro*, but, in so doing, it will initiate RNA chains at random sites on both strands of DNA. In contrast, the holoenzyme (σ present) initiates RNA chains *in vitro* only at sites used *in vivo*.

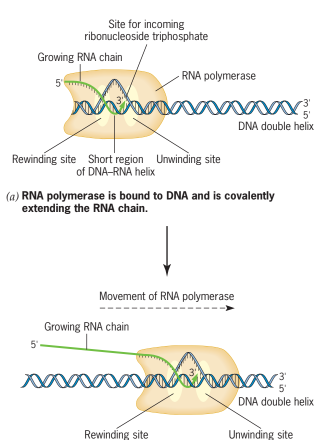






Localized unwinding

■ FIGURE 11.8 Structure of a typical promoter in *E. coli*. RNA polymerase binds to the −35 sequence of the promoter and initiates unwinding of the DNA strands at the AT-rich −10 sequence. Transcription begins within the transcription bubble at a site five to nine base pairs beyond the −10 sequence.



(b) RNA polymerase has moved downstream from its position in (a), processively extending the nascent RNA chain.

FIGURE 11.9 Elongation of an RNA chain catalyzed by RNA polymerase in *E. coli*.

INITIATION OF RNA CHAINS

Initiation of RNA chains involves three steps: (1) binding of the RNA polymerase holoenzyme to a promoter region in DNA; (2) the localized unwinding of the two strands of DNA by RNA polymerase, providing a template strand free to base-pair with incoming ribonucleotides; and (3) the formation of phosphodiester bonds between the first few ribonucleotides in the nascent RNA chain. The holoenzyme remains bound at the promoter region during the synthesis of the first eight or nine bonds; then the sigma factor is released, and the core enzyme begins the elongation phase of RNA synthesis. During initiation, short chains of two to nine ribonucleotides are synthesized and released. This abortive synthesis stops once chains of 10 or more ribonucleotides have been synthesized and RNA polymerase has begun to move downstream from the promoter.

By convention, the nucleotide pairs or nucleotides within and adjacent to transcription units are numbered relative to the transcript initiation site (designated +1)—the nucleotide pair corresponding to the first (5') nucleotide of the RNA transcript. Base pairs preceding the initiation site are given minus (-) prefixes; those following (relative to the direction of transcription) the initiation site are given plus (+) prefixes. Nucleotide sequences preceding the initiation site are referred to as **upstream sequences**; those following the initiation site

are called **downstream sequences**.

As mentioned earlier, the sigma subunit of RNA polymerase mediates its binding to promoters in DNA. Hundreds of *E. coli* promoters have been sequenced and found to have surprisingly little in common. Two short sequences within these promoters are

sufficiently conserved to be recognized, but even these are seldom identical in two different promoters. The midpoints of the two conserved sequences occur at about 10 and 35 nucleotide pairs, respectively, before the transcription-initiation site (■ Figure 11.8). Thus they are called the -10 sequence and the -35 sequence, respectively. Although these sequences vary slightly from gene to gene, some nucleotides are highly conserved. The nucleotide sequences that are present in such conserved genetic elements most often are called **consensus sequences.** The -10 consensus sequence in the nontemplate strand is TATAAT; the -35 consensus sequence is TTGACA. The sigma subunit initially recognizes and binds to the -35 sequence; thus, this sequence is sometimes called the recognition sequence. The AT-rich -10 sequence facilitates the localized unwinding of DNA, which is an essential prerequisite to the synthesis of a new RNA chain. The distance between the -35 and -10 sequences is highly conserved in *E. coli* promoters, never being less than 15 or more than 20 nucleotide pairs in length. In addition, the first or 5' base in E. coli RNAs is usually (>90 percent) a purine.

ELONGATION OF RNA CHAINS

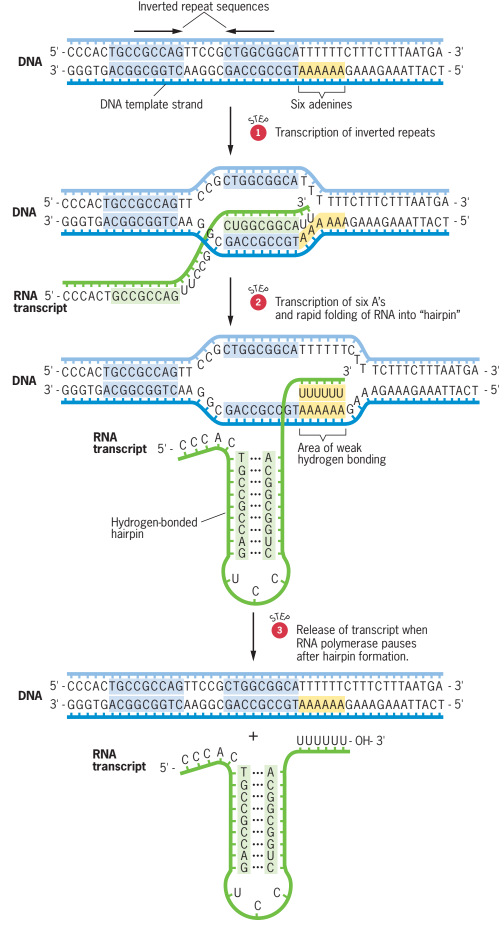
Elongation of RNA chains is catalyzed by the RNA polymerase core enzyme, after the release of the σ subunit. The covalent extension of RNA chains (see Figure 11.5) takes place within the transcription bubble, a locally unwound segment of DNA. The RNA polymerase molecule contains both DNA unwinding and DNA rewinding activities. RNA polymerase continuously unwinds the DNA double helix ahead of the polymerization site and rewinds the complementary DNA strands behind the polymerization site as it moves along the double helix (**■** Figure 11.9). In *E. coli*, the average length of a transcription bubble is 18 nucleotide pairs, and about 40 ribonucleotides are incorporated into the growing RNA chain per second. The nascent RNA chain is displaced from the DNA template strand as RNA polymerase moves along the DNA molecule. The region of transient base-pairing between the growing chain and the DNA template strand is very short, perhaps only three base pairs in length. The stability of the transcription complex is maintained primarily by the binding of the DNA and the growing RNA chain to RNA polymerase, rather than by the basepairing between the template strand of DNA and the nascent RNA.

TERMINATION OF RNA CHAINS

Termination of RNA chains occurs when RNA polymerase encounters a **termination signal**. When it does, the transcription complex dissociates, releasing the nascent RNA molecule. There are two types of transcription terminators in *E. coli*. One type results in termination only in the presence of a protein called *rho* (ρ); therefore, such termination sequences are called *rhodependent terminators*. The other type results in the termination of transcription without the involvement of rho; such sequences are called *rho-independent terminators*.

Rho-independent terminators contain a GC-rich region followed by six or more AT base pairs, with the A's present in the template strand (**Figure 11.10**, top). The nucleotide sequence of the GC-rich region contains inverted repeats-sequences of nucleotides in each DNA strand that are inverted and complementary. When transcribed, these inverted repeat regions produce single-stranded RNA sequences that can base-pair and form hairpin structures (Figure 11.10, bottom). The RNA hairpin structures form immediately after the synthesis of the participating regions of the RNA chain and retard the movement of RNA polymerase molecules along the DNA, causing pauses in chain extension. Since AU base-pairing is weak, requiring less energy

■ FIGURE 11.10 Mechanism of rho-independent termination of transcription. As transcription proceeds along a DNA template, a region of DNA is encountered that contains inverted repeat sequences (shaded). When these repeat sequences are transcribed, the RNA transcript will contain sequences that are complementary to each other. As a result, they will hydrogen bond and form a hairpin structure. When RNA polymerase encounters this hairpin, it will pause, and the weak hydrogen bonds between the A's that follow in the template strand and the U's in the newly synthesized transcript will break, releasing the transcript from the DNA.

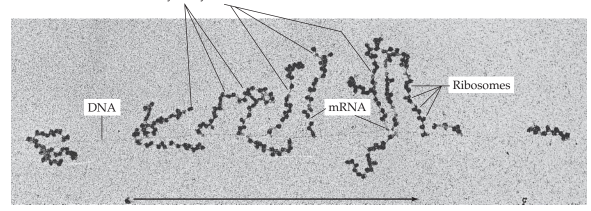


to separate the bases than any of the other standard base pairs, the run of U's after the hairpin region facilitates the release of the newly synthesized RNA chains from the DNA template when the hairpin structure causes RNA polymerase to pause at this site.

The mechanism by which rho-dependent termination of transcription occurs is similar to that of rho-independent termination in that both involve the formation of a hydrogen-bonded hairpin structure upstream from the site of termination. In both cases, these hairpins impede the movement of RNA polymerase, causing it to pause. However, rho-dependent terminators contain two additional sequences: a 50–90 nucleotide-pair sequence upstream from the inverted repeat sequences that produces an RNA strand with many C's but few G's, which therefore forms no hairpins or other secondary structures, and a sequence specifying a rho protein binding site called *rut* (for <u>rho ut</u>ilization) near the 3' end of the transcript. Rho protein binds to the *rut* sequence in the transcript and moves 5' to 3' following RNA polymerase. When polymerase encounters the hairpin, it pauses, allowing rho to catch up, pass through the hairpin, and use its helicase activity to unwind the DNA/RNA base-pairing at the terminus and release the RNA transcript.

CONCURRENT TRANSCRIPTION, TRANSLATION, AND mRNA DEGRADATION

In prokaryotes, the translation and degradation of an mRNA molecule often begin before its synthesis (transcription) is complete. Since mRNA molecules are synthesized, translated, and degraded in the 5' to 3' direction, all three processes can occur simultaneously on the same RNA molecule. In prokaryotes, the polypeptide-synthesizing machinery is not separated by a nuclear envelope from the site of mRNA synthesis. Therefore, once the 5' end of an mRNA has been synthesized, it can immediately be used as a template for polypeptide synthesis. Indeed, transcription and translation often are tightly coupled in prokaryotes. Oscar Miller, Barbara Hamkalo, and colleagues developed techniques that allowed them to visualize this coupling between transcription and translation in bacteria by electron microscopy. One of their photographs showing the coupled transcription of a gene and translation of its mRNA product in *E. coli* is reproduced in **Figure 11.11**.



Gene transcripts (RNA) being simultaneously translated by many ribosomes

Direction of transcription

0.5 μm

FIGURE 11.11 Electron micrograph prepared by Oscar Miller and Barbara Hamkalo showing the coupled transcription and translation of a gene in *E. coli*. DNA, mRNAs, and the ribosomes translating individual mRNA molecules are visible. The nascent polypeptide chains being synthesized on the ribosomes are not visible as they fold into their three-dimensional configuration during synthesis.

- RNA synthesis occurs in three stages: (1) initiation, (2) elongation, and (3) termination.
- RNA polymerases—the enzymes that catalyze transcription—are complex multimeric proteins.
- The covalent extension of RNA chains occurs within locally unwound segments of DNA.
- Chain elongation stops when RNA polymerase encounters a transcription-termination signal.
- Transcription, translation, and degradation of mRNA molecules often occur simultaneously in prokaryotes.

Transcription and RNA Processing in Eukaryotes

Although the overall process of RNA synthesis is similar in prokaryotes and eukaryotes, the process is considerably more complex in eukaryotes. In eukaryotes, RNA is synthesized in the nucleus, and most RNAs that encode proteins must be transported to the cytoplasm for translation on ribosomes. There is evidence suggesting that some translation occurs in the nucleus; however, the vast majority clearly occurs in the cytoplasm.

Prokaryotic mRNAs often contain the coding regions of two or more genes; such mRNAs are said to be multigenic. In contrast, many of the eukaryotic transcripts that have been characterized contain the coding region of a single gene (are

monogenic). Nevertheless, up to one-fourth of the transcription units in the small worm *Caenorhabditis elegans* may be multigenic. Clearly, eukaryotic mRNAs may be either monogenic or multigenic.

Five different RNA polymerases are present in eukaryotes, and each enzyme catalyzes the transcription of a specific class of genes. Moreover, in eukaryotes, the majority of the primary transcripts of genes that encode polypeptides undergo three major modifications prior to their transport to the cytoplasm for translation (**■** Figure 11.12).

- 1. 7-Methyl guanosine caps are added to the 5' ends of the primary transcripts.
- **2.** Poly(A) tails are added to the 3' ends of the transcripts, which are generated by cleavage rather than by termination of chain extension.
- **3.** When present, intron sequences are spliced out of transcripts.

The 5' cap on most eukaryotic mRNAs is a 7-methyl guanosine residue joined to the initial nucleoside of the transcript by a 5'-5' phosphate linkage. The 3' poly(A) tail is a polyadenosine tract 20 to 200 nucleotides long.

In eukaryotes, the population of primary transcripts in a nucleus is called **heterogeneous nuclear RNA (hnRNA)** because of the large variation in the sizes of the RNA molecules present. Major portions of these hnRNAs are noncoding intron sequences, which are excised from the primary transcripts and degraded in the nucleus. Thus, much of the hnRNA actually consists of pre-mRNA molecules undergoing various processing events before leaving the nucleus. Also, in eukaryotes, RNA transcripts are coated with RNAbinding proteins during or immediately after their synthesis. These proteins protect gene transcripts from degradation by ribonucleases, enzymes that degrade RNA molecules, during processing and transport to the cytoplasm. The average half-life of a gene transcript in eukaryotes is about five hours, in contrast to an average half-life of less than five minutes in *E. coli*. This enhanced stability of gene transcripts in eukaryotes is provided, at least in part, by their interactions with RNA-binding proteins.

FIVE RNA POLYMERASES/FIVE SETS OF GENES

Whereas a single RNA polymerase catalyzes all transcription in *E. coli*, eukaryotes ranging in complexity from the single-celled yeasts to humans contain from three to five different RNA polymerases. Three enzymes, designated **RNA polymerases I**, **II**, and **III**, are known to be present in most, if not all, eukaryotes. All three are more

Five different enzymes catalyze transcription in eukaryotes, and the resulting RNA transcripts undergo three important modifications, including the excision of noncoding sequences called introns. The nucleotide sequences of some RNA transcripts are modified posttranscriptionally by RNA editing.



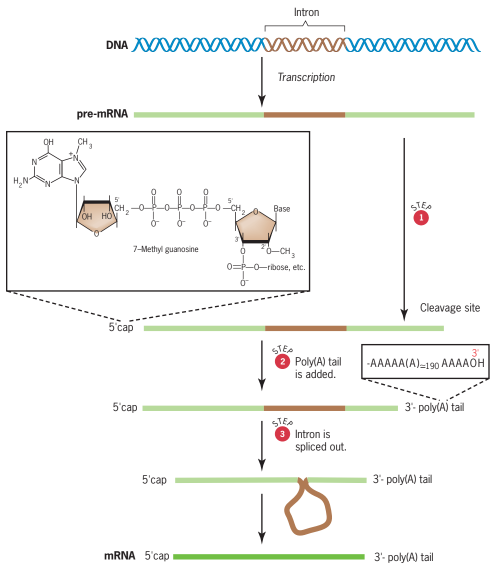


FIGURE 11.12 In eukaryotes, most gene transcripts undergo three different types of posttranscriptional processing.

complex, with 10 or more subunits, than the *E. coli* RNA polymerase. Moreover, unlike the *E. coli* enzyme, all eukaryotic RNA polymerases require the assistance of other proteins called **transcription factors** in order to initiate the synthesis of RNA chains.

The key features of the five eukaryotic RNA polymerases are summarized in Table 11.1. RNA polymerase I is located in the nucleolus, a distinct region of the nucleus where rRNAs are synthesized and combined with ribosomal proteins. RNA polymerase I catalyzes the synthesis of all ribosomal RNAs except the small 5S rRNA. RNA polymerase II transcribes nuclear genes that encode proteins and perhaps other genes specifying hnRNAs. RNA polymerase III catalyzes the synthesis of the transfer RNA molecules, the 5S rRNA molecules, and small nuclear RNAs. To date, RNA polymerases IV and V have been identified only in plants; however, there are hints that they may exist in other eukaryotes, especially fungi.

RNA polymerases IV and V play important roles in turning off the transcription of genes by modifying the structure of chromosomes, a process called *chromatin remodeling* (see On the Cutting Edge: Chromatin Remodeling and Gene Expression and Chapter 19). Chromatin remodeling occurs when the histone tails in nucleosomes (see Figure 9.18) are chemically modified and proteins interact with these modified groups, causing the chromatin to become more or less condensed. RNA polymerase IV synthesizes transcripts

that are processed into short RNAs called *small interfering RNAs* (siRNAs) that are important regulators of gene expression (see Chapter 19). One mechanism of action involves interacting with other proteins to modify (condense or relax) chromatin structure. RNA polymerase V synthesizes a subset of siRNAs and noncoding (antisense) transcripts of genes that are regulated by siRNAs. Although the details of the process are still being worked out, it seems likely that the siRNAs interact with these noncoding transcripts and nucleosome-associated proteins—some well characterized, others unknown—to condense chromatin into structures that cannot be transcribed.

TABLE 11.1 Characteristics of the Five RNA Polymerases of Eukaryotes Enzyme Location Products RNA polymerase I Ribosomal RNAs, excluding 5S rRNA Nucleolus RNA polymerase II Nucleus Nuclear pre-mRNAs RNA polymerase III tRNAs, 5S rRNA, and other small nuclear Nucleus **RNAs** Nucleus (plant) RNA polymerase IV Small interfering RNAs (siRNAs) RNA polymerase V Nucleus (plant) Some siRNAs plus noncoding (antisense) transcripts of siRNA target genes.

ON THE CUTTING EDGE

CHROMATIN REMODELING AND GENE EXPRESSION

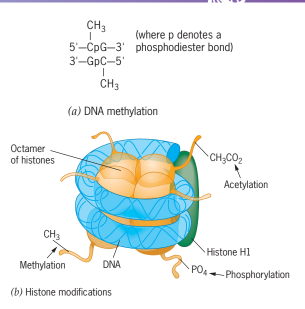
he DNA of eukaryotes is packaged into roughly 11-nm spheres called nucleosomes, which consist of DNA wound on the surface of histone octamers (see Figure 9.18). Within these nucleosomes, the charged amino-terminal tails of the histones bind tightly to DNA, keeping the structures quite compact. How, then, can the transcription factors and the large RNA polymerase complexes gain access to the promoters and transcribe the genes in nucleosomes? The answer is that the structures of nucleosomes containing genes that need to be expressed must be modified to make the promoters available to the proteins required for transcription; that is, **chromatin remodeling** must occur before transcription can begin.

There are several types of chromatin remodeling, some of which are discussed in more detail in Chapter 19. All involve chromatinremodeling proteins, usually multimeric protein complexes. Some require the input of energy from ATP. Chromatin remodeling can occur (1) by sliding nucleosomes along DNA so that specific DNA sequences are located between nucleosomes, (2) by changing the spacing between nucleosomes, or (3) by displacing histone octamers to create nucleosome-free gaps. But what controls these chromatin-remodeling processes? What signals are required to initiate a specific pathway of chromatin remodeling?

The signals that control chromatin remodeling are still being worked out. However, chemical modifications of nucleotides in DNA and of amino acids in the protruding tails of histones in nucleosomes play key roles (**Figure 1**). Many of the genes of mammals contain sequences rich in the dinucleotide sequence

5'-CpG-3' 3'-GpC-5'

upstream from their transcription start sites. These CpG-rich regions are called CpG islands and are important regulatory sequences. The cytosines in the CpG islands are subject to methylation, the addition of methyl (CH₃) groups, and methylated CpG islands, in turn, are binding sites for proteins that regulate transcription. In many cases, the methylation of CpG islands results in the repression of transcription of nearby genes. However, in some cases, recent studies indicate that chromatin remodeling can lead to global changes in gene expression, including both repression and activation.

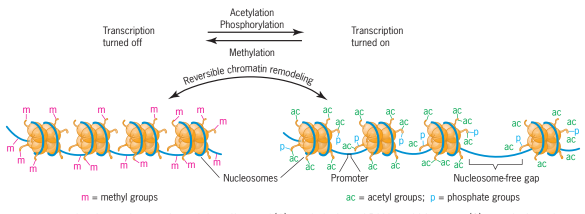


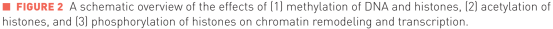


Amino acids in the protruding histone tails of nucleosomes also undergo methylation, and these methyl groups on DNA and histones work together to compact chromatin and repress gene expression. However, methylation is not the whole story! The protruding histone tails undergo two additional modifications: acetylation, the addition of acetyl (CH_3CO_2) groups; and phosphorylation, the addition of phosphate (PO_4) groups.

Acetylation is the more important of these modifications. Acetyl groups are added to specific lysine residues in the histone tails by enzymes called acetylases, neutralizing the positive charges of these lysines (see Figure 12.1). As a result, acetylation decreases the interaction between the negatively charged DNA and the histone tails and sets the stage for the chromatin remodeling required for the initiation of transcription.

In mammals, a protein complex called the *enhanceosome* initiates the activation process by binding to DNA upstream from the promoter and recruiting acetylase, which then adds acetyl groups to histone tails protruding from nucleosomes. Chromatin-remodeling proteins then modify the structure of the complex and make the promoter accessible to transcription factors and RNA polymerase. **Figure 2** shows a





ON THE CUTTING EDGE (continued)

schematic overview of the effects of methylation, acetylation, and phosphorylation on chromatin remodeling and transcription.

Recent evidence indicates that global changes in gene expression—with some genes upregulated and other genes downregulated can be caused by chromatin remodeling. Indeed, several human disorders are now known to result from genetic defects in chromatin remodeling. One form of acute lymphoblastic leukemia and Rett syndrome, a severe neurological disorder, both result from defects in chromatin remodeling. Rett syndrome, which results in loss of motor skills and mental retardation within four years of birth, is caused by mutations in the *MECP2* gene, encoding *methyl-CpG-binding protein 2*, on the X chromosome. Recent evidence suggests that MECP2 is made in large quantities and binds directly to nucleosomes, actually competing with histone H1 for common binding sites. Indeed, when MECP2 binds to nucleosomes, it changes their architecture and alters the expression of genes packaged therein. Exactly how do mutations in the *MECP2* gene alter chromatin remodeling and change gene expression in neurons? The details still must be worked out. However, given the ongoing research in this field, new information will undoubtedly become available in the near future.

MANY

INITIATION OF RNA CHAINS

Unlike their prokaryotic counterparts, eukaryotic RNA polymerases cannot initiate transcription by themselves. All five eukaryotic RNA polymerases require the assistance of protein transcription factors to start the synthesis of an RNA chain. Indeed, these transcription factors must bind to a promoter region in DNA and form an appropriate initiation complex before RNA polymerase will bind and initiate transcription. Different promoters and transcription factors are utilized by RNA polymerases. In this section, we focus on the initiation of pre-mRNA synthesis by RNA polymerase II, which transcribes the vast majority of eukaryotic genes.

In all cases, the initiation of transcription involves the formation of a locally unwound segment of DNA, providing a DNA strand that is free to function as a template for the synthesis of a complementary strand of RNA (see Figure 11.6). The formation of the locally unwound segment of DNA required to initiate transcription involves the interaction of several transcription factors with specific sequences in the promoter for the transcription unit. The promoters recognized by RNA polymerase II consist of short conserved elements, or modules, located upstream from the transcription startpoint. The components of the promoter of the mouse thymidine kinase gene are shown in **Figure 11.13**. Other promoters that are recognized by RNA polymerase II contain some, but not all, of these components. The conserved element closest to the transcription start site (position +1) is called the **TATA box**; it has the consensus sequence TATAAAA (reading 5' to 3' on the nontemplate strand) and is centered at about position -30. The TATA box plays an important role in positioning the transcription startpoint. The second conserved element is called the **CAAT box**; it usually occurs near position -80 and has the consensus sequence GGCCAATCT. Two other

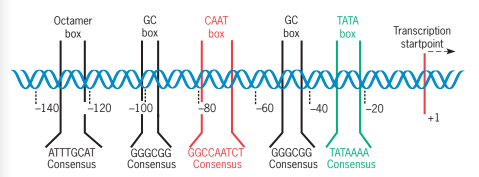


FIGURE 11.13 Structure of a promoter recognized by RNA polymerase II. The TATA and CAAT boxes are located at about the same positions in the promoters of most nuclear genes encoding proteins. The GC and octamer boxes may be present or absent; when present, they occur at many different locations, either singly or in multiple copies. The sequences shown here are the consensus sequences for each of the promoter elements. The conserved promoter elements are shown at their locations in the mouse thymidine kinase gene.

Solve It!

Initiation of Transcription by RNA Polymerase II in Eukaryotes

The nucleotide sequence of the nontemplate strand of a portion of the human *HBB* (β -globin) gene and the amino-terminus of its product, human β -globin (using the single-letter amino acid code), are given as follows. Remember that the nontemplate strand will have the same sequence as the transcript of the gene, but with T's in place of U's.

5'-CCTGTGGAGC	CACACCCTAG	GGTTGGCCAA	TCTACTCCCA	
GGAGCAGGGA	GGGCAGGAGC	CAGGGCTGGG	CATAAAAGTC	
AGGGCAGAGC	CATCTATTGC	TTACATTTGC	TTCTGACACA	
ACTGTGTTCA	CTAGCAACCT	CAAACAGACA	CCATGGTGCA	
β-globin amino terminus: M V H				
TCTGACTCCT C	GAGGAGAAGT (CTGCCGTTAC T	GCCCTGTGG-3	
LTP	EEK	SAVT	A L W —	

Note: Every other codon is underlined in the coding region of the gene.

Does the TATA box in this gene have the consensus sequence? If not, what is its sequence? Does this gene contain a CAAT box? Does it have the consensus sequence? Given that transcription of eukaryotic genes by RNA polymerase II almost always starts (+1 site) at an A preceded by two pyrimidines, predict the sequence of the 5'-terminus of the primary transcript of this gene.

• To see the solution to this problem, visit the Student Companion site.

conserved elements, the *GC box*, consensus GGGCGG, and the *octamer box*, consensus ATTTGCAT, often are present in RNA polymerase II promoters; they influence the efficiency of a promoter in initiating transcription. Try Solve It: Initiation of Transcription by RNA Polymerase II in Eukaryotes to see how these conserved promoter sequences work in the human *HBB* (β -globin) gene.

The initiation of transcription by RNA polymerase II requires the assistance of several **basal transcription factors**. Still other transcription factors and regulatory sequences called *enhancers* and *silencers* modulate the efficiency of initiation (Chapter 19). The basal transcription factors must interact with promoters in the correct sequence to initiate transcription effectively (**Transcription Factor X** for RNA polymerase **II**, where **X** is a letter identifying the individual factor).

TFIID is the first basal transcription factor to interact with the promoter; it contains a TATA-binding protein (TBP) and several small TBPassociated proteins (Figure 11.14). Next, TFIIA joins the complex, followed by TFIIB. TFIIF first associates with RNA polymerase II, and then TFIIF and RNA polymerase II join the transcription initiation complex together. TFIIF contains two subunits, one of which has DNA-unwinding activity. Thus, TFIIF probably catalyzes the localized unwinding of the DNA double helix required to initiate transcription. TFIIE then joins the initiation complex, binding to the DNA downstream from the transcription startpoint. Two other factors, TFIIH and TFIIJ, join the complex after TFIIE, but their locations in the complex are unknown. TFIIH has helicase activity and travels with RNA polymerase II during elongation, unwinding the strands in the region of transcription (the "transcription bubble").

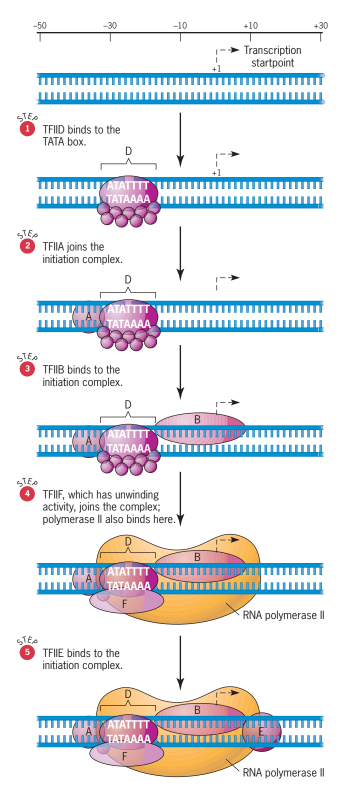
RNA polymerases I and III initiate transcription by processes that are similar, but somewhat simpler, than the one used by polymerase II, whereas the processes used by RNA polymerases IV and V are currently under investigation. The promoters of genes transcribed by polymerases I and III are quite different from those utilized by polymerase II, even though they sometimes contain some of the same regulatory elements. RNA polymerase I promoters are bipartite, with a core sequence extending from about -45 to +20, and an upstream control element extending from -180 to about -105. The two regions have similar sequences, and both are GC-rich. The core sequence is sufficient for initiation; however, the efficiency of initiation is strongly enhanced by the presence of the upstream control element.

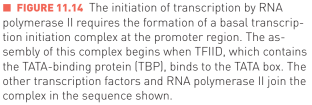
Interestingly, the promoters of most of the genes transcribed by RNA polymerase III are located within the transcription units, downstream from the transcription startpoints, rather than upstream as in units transcribed by RNA polymerases I and II. The promoters of other genes transcribed by polymerase III are located upstream of the transcription start site, just as for polymerases I and II. Actually, polymerase III promoters can be divided into three classes, two of which have promoters located within the transcription unit.

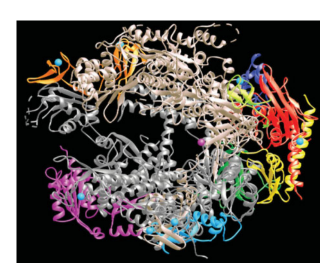
RNA CHAIN ELONGATION AND THE ADDITION OF 5' METHYL GUANOSINE CAPS

Once eukaryotic RNA polymerases have been released from their initiation complexes, they catalyze RNA chain elongation by the same mechanism as the RNA polymerases of prokaryotes (see Figures 11.5 and 11.6). Studies on the crystal structures of various RNA polymerases have

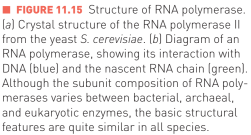
provided a good picture of key features of this important enzyme. Although the RNA polymerases of bacteria, archaea, and eukaryotes have different substructures, their



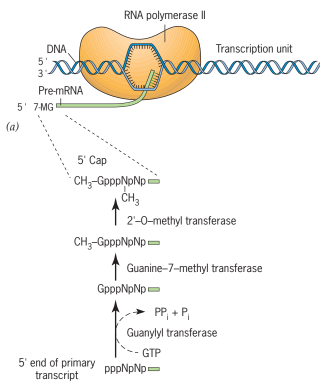




(a) Crystal structure of yeast RNA polymerase II.

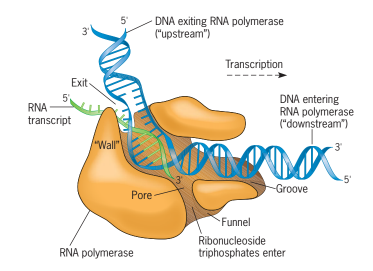


Early stage in the transcription of a gene by RNA polymerase II.



(b) Pathway of biosynthesis of the 7-MG cap.

■ FIGURE 11.16 7-Methyl guanosine (7-MG) caps are added to the 5' ends of pre-mRNAs shortly after the elongation process begins.



(b) Diagram of the interaction between DNA and RNA polymerase based on crystal structures and other structural analyses.

key features and mechanisms of action are quite similar. The crystal structure of RNA polymerase II (resolution = .28 nm) of S. cerevisiae is shown in ■ Figure 11.15a. A schematic diagram showing structural features of an RNA polymerase and its interaction with DNA and the growing RNA transcript is shown in **Figure 11.15***b*.

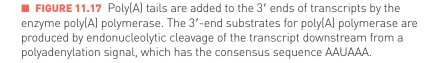
Early in the elongation process, the 5' ends of eukaryotic pre-mRNAs are modified by the addition of 7-methyl guanosine (7-MG) caps. These 7-MG caps are added

when the growing RNA chains are only about 30 nucleotides long (**Figure 11.16**). The 7-MG cap contains an unusual 5'-5' triphosphate linkage (see Figure 11.12) and two or more methyl groups. These 5' caps are added co-transcriptionally by the biosynthetic pathway shown in Figure 11.16. The 7-MG caps are recognized by protein factors involved in the initiation of translation (Chapter 12) and also help protect the growing RNA chains from degradation by nucleases.

Recall that eukaryotic genes are present in chromatin organized into nucleosomes (Chapter 9). How does RNA polymerase transcribe DNA packaged in nucleosomes? Does the nucleosome have to be disassembled before the DNA within it can be transcribed? Surprisingly, RNA polymerase II is able to move past nucleosomes with the help of a protein complex called FACT (facilitates chromatin transcription), which removes histone H2A/H2B dimers from the nucleosomes leaving histone "hexasomes." After polymerase II moves past the nucleosome, FACT and other accessory proteins help redeposit the histone dimers, restoring nucleosome structure. Also, we should note that chromatin that contains genes actively being transcribed has a less compact structure than chromatin that contains inactive genes. Chromatin in which active genes are packaged tends to contain histones with lots of acetyl groups (Chapter 9), whereas chromatin with inactive genes contains histones with fewer acetyl groups. These differences are discussed further in Chapter 19.

TERMINATION BY CHAIN CLEAVAGE AND THE ADDITION OF 3' POLY(A) TAILS

The 3' ends of RNA transcripts synthesized by RNA polymerase II are produced by endonucleolytic cleavage of the primary transcripts rather than by the termination of transcription (**Figure 11.17**). The actual transcription termination events often occur at multiple sites that



are located 1000 to 2000 nucleotides downstream from the site that will become the 3' end of the mature transcript. That is, transcription proceeds beyond the site that will become the 3' terminus, and the distal segment is removed by endonucleolytic cleavage. The cleavage event that produces the 3' end of a transcript usually occurs at a site 11 to 30 nucleotides downstream from a conserved polyadenylation signal, consensus AAUAAA, and upstream from a GU-rich sequence located near the end of the transcript. After cleavage, the enzyme **poly(A) polymerase** adds poly(A) tails, tracts of adenosine monophosphate residues about 200 nucleotides long, to the 3' ends of the transcripts (Figure 11.17). The addition of poly(A) tails to eukaryotic mRNAs is called **polyadenylation**. To examine the polyadenylation signal of the human *HBB* (β -globin) gene, check out Solve It: Formation of the 3'-Terminus of an RNA Polymerase II Transcript.

The formation of poly(A) tails on transcripts requires a specificity component that recognizes and binds to the AAUAAA sequence, a

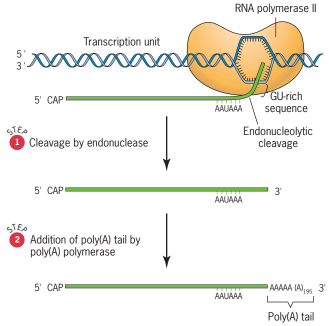
stimulatory factor that binds to the GU-rich sequence, an endonuclease, and the poly(A) polymerase. These proteins form a multimeric complex that carries out both the cleavage and the polyadenylation in tightly coupled reactions. The poly(A) tails of eukaryotic mRNAs enhance their stability and play an important role in their transport from the nucleus to the cytoplasm.

In contrast to RNA polymerase II, both RNA polymerase I and III respond to discrete termination signals. RNA polymerase I terminates transcription in response to an 18-nucleotide-long sequence that is recognized by an associated terminator protein. RNA polymerase III responds to a termination signal that is similar to the rho-independent terminator in *E. coli* (see Figure 11.10).

RNA EDITING: ALTERING THE INFORMATION CONTENT OF mRNA MOLECULES

According to the central dogma of molecular biology, genetic information flows from DNA to RNA to protein during gene expression. Normally, the genetic information is not altered in the mRNA intermediary. However, the discovery of **RNA editing** has shown that exceptions do occur. RNA editing processes alter the information content of gene transcripts in two ways: (1) by changing the structures of individual bases and (2) by inserting or deleting uridine monophosphate residues.

The first type of RNA editing, which results in the substitution of one base for another base, is rare. This type of editing was discovered in studies of the apolipoprotein-B (apo-B) genes and mRNAs in rabbits and humans. Apolipoproteins are blood proteins that transport certain types of fat molecules in the circulatory system. In the liver, the *apo-B* mRNA encodes a large protein 4563 amino acids long. In the intestine, the *apo-B* mRNA directs the synthesis of a protein only 2153 amino acids long. Here, a C residue in the pre-mRNA is converted to a U, generating an internal UAA translationtermination codon, which results in the truncated apolipoprotein (**Figure 11.18**). UAA is one of three codons that terminates polypeptide chains during translation. If a UAA codon is produced within the coding region of an mRNA, it will prematurely terminate the polypeptide during translation, yielding an incomplete gene product. The $C \rightarrow U$ conversion is catalyzed by a sequence-specific RNA-binding protein with an activity that removes amino groups from cytosine residues. A similar example of RNA editing has been documented for an mRNA specifying a protein (the glutamate receptor) present in rat brain cells. More extensive mRNA editing of the $C \rightarrow U$ type occurs in the mitochondria of plants, where most of the gene transcripts are edited to some degree. Mitochondria have their own DNA genomes and protein-synthesizing



Solve It!

Formation of the 3'-Terminus of an RNA Polymerase II Transcript

The nucleotide sequence of the nontemplate strand of a portion of the human HBB (β -globin) gene and the carboxyl-terminus of its product, human β -globin (using the single-letter amino acid code; see Figure 12.1), are given as follows. Remember that the nontemplate strand will have the same sequence as the transcript of the gene, but with T's in place of U's.

5-GGTGTGGCTA ATGCCCTGGC CCACAGTAT CACTAAGCTC GCTTTCTTGC G V A N A L A H K Y H COOHterminus of pglobin TGTCCAATT CTATTAAAGG TTCCTTGTT CCCTAAGTCC AACTACTAAAA CACGGGATA TTATGAAAGGG CCTTGAGCAT CTGGATTCTG ACTATAAAA AACATTTATT TTCATTGCAA TGATATTAT AAATTATTTC TGAATATTT-3

Note that every other codon is underlined in the coding region of the gene. Also, note that the GT-rich sequence involved in cleavage is located far downstream, near the end of the transcription unit, and is not shown. Can you predict the exact endonucleolytic cleavage site that produces the 3' end of the transcript? Can you predict the approximate cleavage site? Will the 3' end of the transcript produced by this cleavage event undergo any subsequent modification(s)? If so, what?

To see the solution to this problem, visit the Student Companion site.

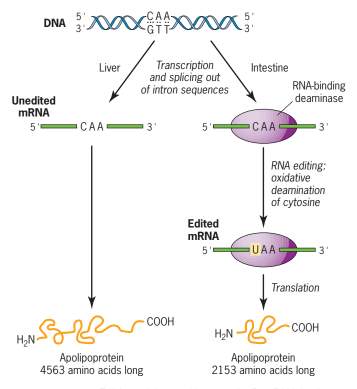


FIGURE 11.18 Editing of the apolipoprotein-B mRNA in the intestines of mammals.

machinery (Chapter 15). In some transcripts present in plant mitochondria, most of the C's are converted to U residues.

A second, more complex type of RNA editing occurs in the mitochondria of trypanosomes (a group of flagellated protozoa that causes sleeping sickness in humans). In this case, uridine monophosphate residues are inserted into (occasionally deleted from) gene transcripts, causing major changes in the polypeptides specified by the mRNA molecules. This RNA editing process is mediated by **guide RNAs** transcribed from distinct mitochondrial genes. The guide RNAs contain sequences that are partially complementary to the pre-mRNAs to be edited. Pairing between the guide RNAs and the pre-mRNAs results in gaps with unpaired A residues in the guide RNAs. The guide RNAs serve as templates for editing, as U's are inserted in the gaps in premRNA molecules opposite the A's in the guide RNAs.

Why do these RNA editing processes occur? Why are the final nucleotide sequences of these mRNAs not specified by the sequences of the mitochondrial genes as they are in most nuclear genes? As yet, answers to these interesting questions are purely speculative. Trypanosomes are primitive single-celled eukaryotes that diverged from other eukaryotes early in evolution. Some evolutionists have speculated that RNA editing was common in ancient cells, where many reactions are thought to have been catalyzed by RNA molecules instead of proteins. Another view is that RNA editing is a primitive mechanism for altering patterns of gene expression. For whatever reason, RNA editing plays a major role in the expression of genes in the mitochondria of trypanosomes and plants.

- **KEY POINTS** *Three to five different RNA polymerases are present in eukaryotes, and each polymerase transcribes a distinct set of genes.*
 - Eukaryotic gene transcripts usually undergo three major modifications: (1) the addition of 7-methyl guanosine caps to 5' termini, (2) the addition of poly(A) tails to 3' ends, and (3) the excision of noncoding intron sequences.
 - The information content of some eukaryotic transcripts is altered by RNA editing, which changes the nucleotide sequences of transcripts prior to their translation.

Interrupted Genes in Eukaryotes: Exons and Introns

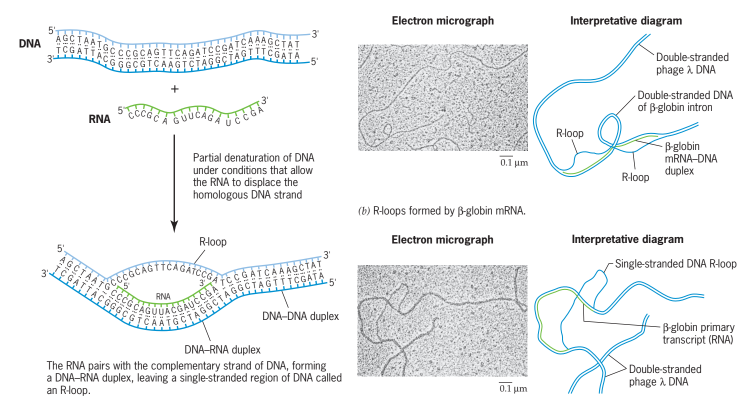
Most eukaryotic genes contain noncoding sequences called introns that interrupt the coding sequences, or exons. The introns are excised from RNA transcripts prior to their transport to the cytoplasm. Most of the well-characterized genes of prokaryotes consist of continuous sequences of nucleotide pairs, which specify colinear sequences of amino acids in the polypeptide gene products. However, in 1977, molecular analyses of three eukaryotic genes yielded a major surprise. Studies of mouse and rabbit β -globin (one of two different proteins in hemoglobin) genes and the chicken

ovalbumin (an egg storage protein) gene revealed that they contain noncoding sequences intervening between coding sequences. They were subsequently found in the nontranslated regions of some genes. They are called **introns** (for **int**ervening sequences.) The sequences that remain present in mature mRNA molecules (both coding and noncoding sequences) are called **exons** (for **ex**pressed sequences).

Some of the earliest evidence for introns in mammalian β -globin genes resulted from the visualization of genomic DNA–mRNA hybrids by electron microscopy. Because DNA–RNA duplexes are more stable than DNA double helices, when partially denatured DNA double helices are incubated with homologous RNA molecules under the appropriate conditions, the RNA strands will hybridize with the complementary DNA strands, displacing the equivalent DNA strands (**Figure 11.19a**). The resulting DNA–RNA hybrid structures will contain single-stranded regions of DNA called **R-loops**, where RNA molecules have displaced DNA strands to form DNA–RNA duplex regions. These R-loops can be visualized directly by electron microscopy.

When Shirley Tilghman, Philip Leder, and colleagues hybridized purified mouse β -globin mRNA to a DNA molecule that contained the mouse β -globin gene, they observed two R-loops separated by a loop of double-stranded DNA (**Figure 11.19b**). Their results demonstrated the presence of a sequence of nucleotide pairs in the middle of the β -globin gene that is not present in β -globin mRNA and, therefore, does not encode amino acids in the β -globin polypeptide. When Tilghman and coworkers repeated the R-loop experiments using purified β -globin gene transcripts isolated from nuclei and believed to be primary gene transcripts or pre-mRNA molecules, in place of cytoplasmic β -globin mRNA, they observed only one R-loop (**Figure 11.19c**). This result indicated that the primary transcript contains the complete structural gene sequence, including both exons and introns. Together, the R-loop results obtained with cytoplasmic mRNA and nuclear pre-mRNA demonstrate that the intron sequence is excised and the exon sequences are spliced together during processing events that convert the primary transcript to the mature mRNA.

Tilghman and coworkers confirmed their interpretation of the R-loop results by comparing the sequence of the mouse β -globin gene with the predicted amino acid sequence of the β -globin polypeptide. Their results showed that the gene contained a noncoding intron at this position in the gene. Subsequent research showed that the mouse β -globin gene actually contains two introns. For details of these studies and additional information on the discovery of introns, see A Milestone in Genetics: Introns on the Student Companion site.



(a) The technique of R-loop hybridization.

(c) R-loop formed by β -globin primary transcript (pre-mRNA).

FIGURE 11.19 R-loop evidence for an intron in the mouse β -globin gene. (a) R-loop hybridization. (b) When mouse β -globin genes and mRNAs were hybridized under R-loop conditions, two R-loops were observed in the resulting DNA-RNA hybrids. (c) When primary transcripts or pre-mRNAs of mouse β -globin genes were used in the R-loop experiments, a single R-loop was observed. These results demonstrate that the intron sequence is present in the primary transcript but is removed during the processing of the primary transcript to produce the mature mRNA.

SOME VERY LARGE EUKARYOTIC GENES

Subsequent to the pioneering studies on the mammalian globin genes and the chicken ovalbumin gene (see Milestone on the Student Companion site), noncoding introns have been demonstrated in a large number of eukaryotic genes. In fact, interrupted genes are much more common than uninterrupted genes in higher animals and plants. For example, the *Xenopus laevis* gene that encodes vitellogenin A2 (which ends up as egg yolk protein) contains 33 introns, and the chicken 1 α 2 collagen gene contains at least 50 introns. The collagen gene spans 37,000 nucleotide pairs but gives rise to an mRNA molecule only about 4600 nucleotides long. Other genes contain relatively few introns, but some of the introns are very large. For example, the *Ultrabithorax* (*Ubx*) gene of *Drosophila* contains an intron that is approximately 70,000 nucleotide pairs in length. The largest gene characterized to date is the human *DMD* gene, which causes Duchenne muscular dystrophy when rendered nonfunctional by mutation. The *DMD* gene spans 2.5 million nucleotide pairs and contains 78 introns.

Although introns are present in most genes of higher animals and plants, they are not essential because not all such genes contain introns. The sea urchin histone genes and four *Drosophila* heat-shock genes were among the first animal genes shown to lack introns. We now know that many genes of higher animals and plants lack introns.

INTRONS: BIOLOGICAL SIGNIFICANCE?

At present, scientists know relatively little about the biological significance of the exon–intron structure of eukaryotic genes. Introns are highly variable in size, ranging from about 50 nucleotide pairs to thousands of nucleotide pairs in length. This fact has led to speculation that introns may play a role in regulating gene expression. Although it is unclear how introns regulate gene expression, new research has shown that some introns contain sequences that can regulate gene expression in either a positive or negative fashion. Other introns contain alternative tissue-specific promoters; still others contain sequences that enhance the accumulation of transcripts. The fact that introns accumulate new mutations much more rapidly than exons indicates that many of the specific nucleotide-pair sequences of introns, excluding the ends, are not very important.

In some cases, the different exons of genes encode different functional domains of the protein gene products. This is most apparent in the case of the genes encoding heavy and light antibody chains (see Figure 20.17). In the case of the mammalian globin genes, the middle exon encodes the heme-binding domain of the protein. There has been considerable speculation that the exon–intron structure of eukaryotic genes has resulted from the evolution of new genes by the fusion of uninterrupted (single exon) ancestral genes. If this hypothesis is correct, introns may merely be relics of the evolutionary process.

Alternatively, introns may provide a selective advantage by increasing the rate at which coding sequences in different exons of a gene can reassort by recombination, thus speeding up the process of evolution. In some cases, alternate ways of splicing a transcript produce a family of related proteins. In these cases, introns result in multiple products from a single gene. The alternate splicing of the rat troponin T transcript is illustrated in Figure 19.2. In the case of the mitochondrial gene of yeast encoding cytochrome b, the introns contain exons of genes encoding enzymes involved in processing the primary transcript of the gene. Thus, different introns may indeed play different roles, and many introns may have no biological significance. Since many eukaryotic genes contain no introns, these noncoding regions are not required for normal gene expression.

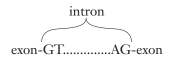
KEY POINTS

- Most, but not all, eukaryotic genes are split into coding sequences called exons and noncoding sequences called introns.
- Some genes contain very large introns; others harbor large numbers of small introns.
- The biological significance of introns is still open to debate.

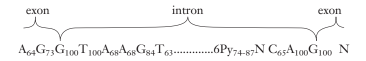
Removal of Intron Sequences by RNA Splicing

Most nuclear genes that encode proteins in multicellular eukaryotes contain introns. Fewer, but still many, of the genes of unicellular eukaryotes such as the yeasts contain introns. Rare genes of archaea and of a few viruses of prokaryotes also contain introns. In the case of these "split" genes, the primary transcript contains the entire sequence of the gene, and the intron sequences are excised during RNA processing (see Figure 11.12).

For genes that encode proteins, the splicing mechanism must be precise; it must join exon sequences with accuracy to the single nucleotide to assure that codons in exons distal to introns are read correctly (**■** Figure 11.20). Accuracy to this degree would seem to require precise splicing signals, presumably nucleotide sequences within introns and at the exon–intron junctions. However, in the primary transcripts of nuclear genes, the only completely conserved sequences of different introns are the dinucleotide sequences at the ends of introns, namely,



The sequences shown here are for the DNA nontemplate strand (equivalent to the RNA transcript, but with T rather than U). In addition, there are short consensus sequences at the exon–intron junctions. For nuclear genes, the consensus junctions are



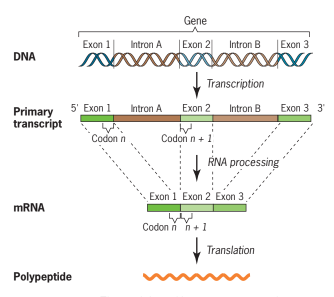
The numerical subscripts indicate the percentage frequencies of the consensus bases at each position; thus, a 100 subscript indicates that a base is always present at that position. N and Py indicate that any of the four standard nucleotides or either

pyrimidine, respectively, may be present at the indicated position. The exon-intron junctions are different for tRNA genes and structural genes in mitochondria and chloroplasts, which utilize different RNA splicing mechanisms. However, different species do show some sequence conservation at exon-intron junctions.

Recent research has shown that splicing and intron sequences can influence gene expression. Direct evidence for their importance has been provided by mutations at these sites that cause mutant phenotypes in many different eukaryotes. Indeed, such mutations are sometimes responsible for inherited diseases in humans, such as hemoglobin disorders.

The discovery of noncoding introns in genes stimulated intense interest in the mechanism(s) by which intron sequences are removed during gene expression. The early demonstration that the intron sequences in eukaryotic genes were transcribed along with the exon sequences focused research on the processing of primary gene transcripts. Just as *in vitro* systems provided important information about the mechanisms of transcription and translation, the key to understanding RNA splicing events was the development of *in vitro* splicing systems. By using these systems, researchers have shown that there are three distinct types of intron excision from RNA transcripts.

1. The introns of tRNA precursors are excised by precise endonucleolytic cleavage and ligation reactions catalyzed by special splicing endonuclease and ligase activities.



■ FIGURE 11.20 The excision of intron sequences from primary transcripts by RNA splicing. The splicing mechanism must be accurate to the single nucleotide to assure that codons in downstream exons are translated correctly to produce the right amino acid sequence in the polypeptide product.

The noncoding introns are excised from gene transcripts by several different mechanisms.

- **2.** The introns of some rRNA precursors are removed autocatalytically in a unique reaction mediated by the RNA molecule itself. (No protein enzymatic activity is involved.)
- **3.** The introns of nuclear pre-mRNA (hnRNA) transcripts are spliced out in two-step reactions carried out by complex ribonucleoprotein particles called spliceosomes.

These three mechanisms of intron excision are discussed in the following three sections. There are other mechanisms of intron excision, but for the sake of brevity they are not discussed here.

tRNA PRECURSOR SPLICING: UNIQUE NUCLEASE AND LIGASE ACTIVITIES

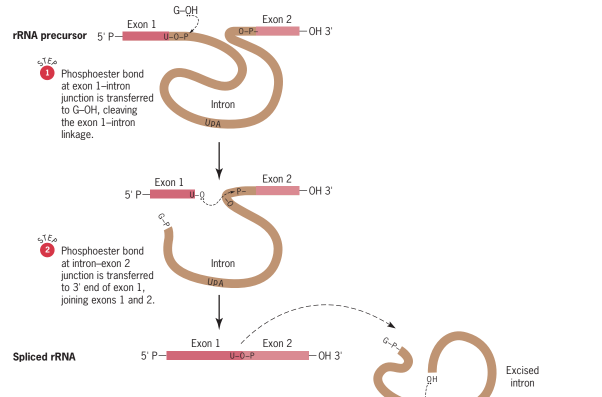
The tRNA precursor splicing reaction has been worked out in detail in the yeast *Saccharomyces cerevisiae*. Both *in vitro* splicing systems and temperature-sensitive splicing mutants have been used in dissecting the tRNA splicing mechanism in *S. cerevisiae*. The excision of introns from yeast tRNA precursors occurs in two stages. In stage I, a nuclear membrane-bound *splicing endonuclease* makes two cuts precisely at the ends of the intron. Then, in stage II, a *splicing ligase* joins the two halves of the tRNA to produce the mature form of the tRNA molecule. The specificity for these reactions resides in conserved three-dimensional features of the tRNA precursors, not in the nucleotide sequences per se.

AUTOCATALYTIC SPLICING

A general theme in biology is that metabolism occurs via sequences of enzymecatalyzed reactions. These all-important enzymes are generally proteins, sometimes single polypeptides and sometimes complex heteromultimers. Occasionally, enzymes require nonprotein cofactors to perform their functions. When covalent bonds are being altered, it is usually assumed that the reaction is being catalyzed by an enzyme. Thus, the 1982 discovery by Thomas Cech and his coworkers that the intron in the rRNA precursor of Tetrahymena thermophila was excised without the involvement of any protein catalytic activity was quite surprising. However, it is now clearly established that the splicing activity that excises the intron from this rRNA precursor is intrinsic to the RNA molecule itself. Indeed, Cech and Sidney Altman shared the 1989 Nobel Prize in Chemistry for their discovery of catalytic RNAs. Moreover, such self-splicing or autocatalytic activity has been shown to occur in rRNA precursors of several lower eukaryotes and in a large number of rRNA, tRNA, and mRNA precursors in mitochondria and chloroplasts of many different species. In the case of many of these introns, the self-splicing mechanism is the same as or very similar to that utilized by the *Tetrahymena* rRNA precursors (see Figure 11.21). For others, the self-splicing mechanism is similar to the splicing mechanism observed with nuclear mRNA precursors, but without the involvement of the spliceosome (see the next section).

The autocatalytic excision of the intron in the *Tetrahymena* rRNA precursor and certain other introns requires no external energy source and no protein catalytic activity. Instead, the splicing mechanism involves a series of phosphoester bond transfers, with no bonds lost or gained in the process. The reaction requires a guanine nucleoside or nucleotide with a free 3'-OH group (GTP, GDP, GMP, or guanosine all work) as a cofactor plus a monovalent cation and a divalent cation. The requirement for the G-3'-OH is absolute; no other base can be substituted in the nucleoside or nucleotide cofactor. The intron is excised by means of two phosphoester bond transfers, and the excised intron can subsequently circularize by means of another phosphoester bond transfer. These reactions are diagrammed in Figure 11.21.

The autocatalytic circularization of the excised intron suggests that the selfsplicing of these rRNA precursors resides primarily, if not entirely, within the intron structure itself. Presumably, the autocatalytic activity is dependent on the secondary



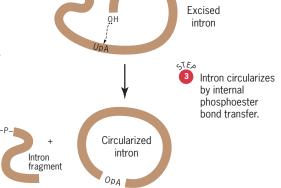
structure of the intron or at least the secondary structure of the RNA precursor molecule. The secondary structures of these self-splicing RNAs must bring the reactive groups into close juxtaposition to allow the phosphoester bond transfers to occur. Since the self-splicing phosphoester bond transfers are potentially reversible reactions, rapid degradation of the excised introns or export of the spliced rRNAs to the cytoplasm may drive splicing in the forward direction.

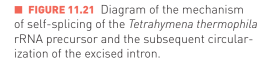
Note that the autocatalytic splicing reactions are intramolecular in nature and thus are not dependent on concentration. Moreover, the RNA precursors are capable of forming an active center in which the guanosine-3'-OH cofactor binds. The autocatalytic splicing of these rRNA precursors demonstrates that catalytic sites are not restricted to proteins; however, there is no *trans* catalytic activity as for enzymes, only *cis* catalytic activity. Some scientists believe that autocatalytic RNA splicing may be a relic of an early RNA-based world.

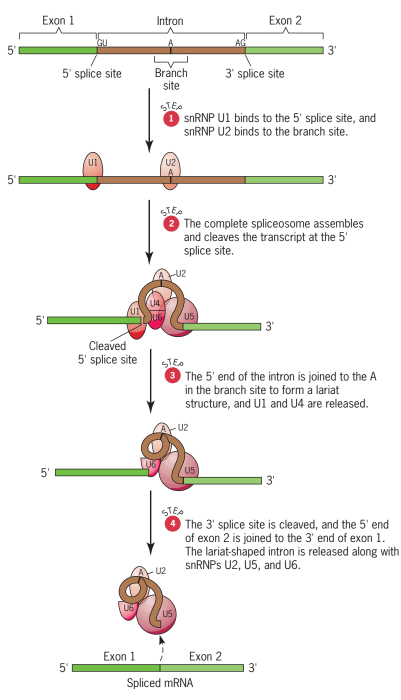
PRE-mRNA SPLICING: snRNAs, snRNPs, AND THE SPLICEOSOME

The introns in nuclear pre-mRNAs are excised in two steps like the introns in yeast tRNA precursors and *Tetrahymena* rRNA precursors that were discussed in the preceding two sections. However, the introns are not excised by simple splicing nucleases and ligases or autocatalytically, and no guanosine cofactor is required. Instead, nuclear pre-mRNA splicing is carried out by complex RNA-protein structures called **spliceosomes**. These structures are in many ways like small ribosomes. They contain a set of small RNA molecules called snRNAs (small nuclear RNAs) and about 40 different proteins. The two stages in nuclear pre-mRNA splicing are known (**Figure 11.22**); however, some of the details of the splicing process are still uncertain.

Five snRNAs, called U1, U2, U4, U5, and U6, are involved in nuclear pre-mRNA splicing as components of the spliceosome. (snRNA U3 is localized in the nucleolus and probably is involved in the formation of ribosomes.) In mammals, these snRNAs range in size from 100 nucleotides (U6) to 215 nucleotides (U3). Some of the snRNAs







■ FIGURE 11.22 The postulated roles of the snRNA-containing snRNPs in nuclear pre-mRNA splicing.

in the yeast *S. cerevisiae* are much larger. These snRNAs do not exist as free RNA molecules. Instead, they are present in small nuclear RNA–protein complexes called **snRNPs** (small nuclear ribonucleoproteins). Spliceosomes are assembled from four different snRNPs and protein splicing factors during the splicing process.

Each of the snRNAs U1, U2, and U5 is present by itself in a specific snRNP particle. snRNAs U4 and U6 are present together in a fourth snRNP; U4 and U6 snRNAs contain two regions of intermolecular complementarity that are basepaired in the U4/U6 snRNP. Each of the four types of snRNP particles contains a subset of seven well-characterized snRNP proteins plus one or more proteins unique to the particular type of snRNP particle.

The first stage in nuclear pre-mRNA splicing involves cleavage at the 5' intron splice site (\downarrow GU-intron) and the formation of an intramolecular phosphodiester linkage between the 5' carbon of the G at the cleavage site and the 2' carbon of a conserved A residue near the 3' end of the intron. This stage occurs on complete spliceosomes (Figure 11.22) and requires the hydrolysis of ATP. Evidence indicates that the U1 snRNP must bind at the 5' splice site prior to the initial cleavage reaction. Recognition of the cleavage site at the 5' end of the intron probably involves base-pairing between the consensus sequence at this site and a complementary sequence near the 5' terminus of snRNA U1. However, the specificity of the binding of at least some of the snRNPs to intron consensus sequences involves both the snRNAs and specific snRNP proteins.

The second snRNP to be added to the splicing complex appears to be the U2 snRNP; it binds at the consensus sequence that contains the conserved A residue that forms the branch point in the lariat structure of the spliced intron. Thereafter, the U5 snRNP binds at the 3' splice site, and the U4/U6 snRNP is added to the complex to yield the complete spliceosome (Figure 11.22). When the 5' intron splice site is cleaved in step 1, the U4 snRNA is released from the spliceosome. In step 2 of the splicing reaction, the 3' splice site of the intron is cleaved, and the two exons are joined by a normal 5' to 3' phosphodiester linkage (Figure 11.22). The spliced mRNA is now ready for export to the cytoplasm and translation on ribosomes.

- **KEY POINTS** Noncoding intron sequences are excised from RNA transcripts in the nucleus prior to their transport to the cytoplasm.
 - Introns in tRNA precursors are removed by the concerted action of a splicing endonuclease and ligase, whereas introns in some rRNA precursors are spliced out autocatalytically—with no catalytic protein involved.
 - The introns in nuclear pre-mRNAs are excised on complex ribonucleoprotein structures called spliceosomes.
 - The intron excision process must be precise, with accuracy to the nucleotide level, to ensure that codons in exons distal to introns are read correctly during translation.

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. If the template strand of a segment of a gene has the nucleotide sequence 3'-GCTAAGC-5', what nucleotide sequence will be present in the RNA transcript specified by this gene segment?
- **Answer:** The RNA transcript will be complementary to the template strand and will have the opposite chemical polarity, as in the following illustration:

DNA template strand:	3'-GCTAAGC-5'
RNA transcript:	5'-CGAUUCG-3'

2. If the nontemplate strand of a gene in *E. coli* had the sequence:

5'-TTGACA-(18 bases)-TATAAT-(8 bases)-GCCTTCCAGTG-3'

what nucleotide sequence would be present in the RNA transcript of this gene?

Answer: The gene contains perfect -35 and -10 promoter sequences. Transcription should be initiated at a site five to nine bases downstream from the -10 TATAAT sequence, and the 5'-terminus of the transcript should contain a purine. The template strand and the 5'-end of the transcript should have the following structure:

DNA template strand:

(-35 sequence) (-10 sequence) 3'-AACTGT-(18 bases)-ATATTA-(8 bases)-CGGAAGGTCAC-5' RNA transcript: 5'-GCCUUCCAGUG-3'

- **3.** If the nontemplate strand shown in Exercise 2 were part of a gene in *Drosophila* rather than *E. coli*, would the same transcript be produced?
- **Answer:** No, because the promoter sequences that control transcription in eukaryotes such as *Drosophila* are different from the promoters in prokaryotes such as *E. coli*. Therefore, the *E. coli* gene would probably not be transcribed if present in *Drosophila*.
- 4. The primary transcript or pre-mRNA of a nuclear gene in a chimpanzee has the sequence:

5'-G-exon 1-AGGUAAGC-intron-CAGUC-exon 2-A-3'

After the intron has been excised, what is the most likely sequence of the mRNA?

Answer: Introns contain highly conserved dinucleotide termini: 5'-GT—AG-3' in the DNA nontemplate strand or 5'-GU—AG-3' in the RNA transcript. Thus, the intron sequence is almost certain to be 5'-GUUAAGC—intron— CAG-3'. With precise excision of the intron, the sequence of the mRNA will be:

5'-G-exon 1-AGUC-exon 2-A-3'

Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. Certain medically important human proteins such as insulin and growth hormone are now being produced in bacteria. By using the tools of genetic engineering, DNA sequences encoding these proteins have been introduced into bacteria. You wish to introduce a human gene into *E. coli* and have that gene produce large amounts of the human gene product in the bacterial cells. Assuming that the human gene of interest can be isolated and introduced into *E. coli*, what problems might you encounter in attempting to achieve your goal?
- **Answer:** The promoter sequences that are required to initiate transcription are very different in mammals and bacteria. Therefore, your gene will not be expressed in *E. coli* unless you first fuse its coding region to a bacterial promoter. In addition, your human gene probably will contain introns. Since *E. coli* cells do not contain spliceosomes or equivalent machinery with which to excise introns from RNA transcripts, your human gene will not be expressed correctly if

it contains introns. As you can see, expressing eukaryotic genes in prokaryotic cells is not a trivial task.

2. A human β -globin gene has been purified and inserted into a linear bacteriophage lambda chromosome, producing the following DNA molecule:

 λ DNA
 Exon 1
 Intron 1
 Exon 2
 Intron 2
 Exon 3
 λ DNA

If this DNA molecule is hybridized to human β -globin mRNA using conditions that favor DNA–RNA duplexes over DNA–DNA duplexes (R-loop mapping conditions) and the product is visualized by electron microscopy, what nucleic acid structure would you expect to see?

Answer: The primary transcript of this human β -globin gene will contain both introns and all three exons. However, prior to its export to the cytoplasm, the intron sequences will be spliced out of the transcript. Thus, the mature mRNA molecule will contain the three exon sequences spliced together with no intron sequences present. Under R-loop conditions, the mRNA will hybridize with the complementary strand of DNA, displacing the equivalent DNA strand. However, since the mRNA contains no intron sequences, the introns will remain as regions of double-stranded DNA as shown in the following diagram.

^{"R-loops"} λ DNA Exon 1 Exon 2 Exon 3 λ DNA Intron 1 Intron 2

Displaced single-stranded exon DNA

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- 11.1 Distinguish between DNA and RNA (a) chemically, (b) functionally, and (c) by location in the cell.
- **11.2** What bases in the mRNA transcript would represent the following DNA template sequence: 5'-TGCAGACA-3'?
- **11.3** What bases in the transcribed strand of DNA would give rise to the following mRNA base sequence: 5'-CUGAU-3'?
- **11.4** On the basis of what evidence was the messenger RNA hypothesis established?
- **11.5** At what locations in a eukaryotic cell does protein synthesis occur?
- **11.6** List three ways in which the mRNAs of eukaryotes differ from the mRNAs of prokaryotes.
- **11.7** What different types of RNA molecules are present in prokaryotic cells? in eukaryotic cells? What roles do these different classes of RNA molecules play in the cell?
- **11.8** Many eukaryotic genes contain noncoding introns that separate the coding sequences or exons of these genes. At what stage during the expression of these split genes are the noncoding intron sequences removed?
- **11.9** For several decades, the dogma in biology has been that molecular reactions in living cells are catalyzed by enzymes composed of polypeptides. We now know that the introns of some precursor RNA molecules such as the rRNA precursors in *Tetrahymena* are removed autocatalytically ("self-spliced") with no involvement of any catalytic protein. What does the demonstration of autocatalytic splicing indicate about the dogma that biological reactions are always catalyzed by proteinaceous enzymes?
- **11.10** What role(s) do spliceosomes play in pathways of gene expression? What is their macromolecular structure?
- **11.11** What components of the introns of nuclear genes that encode proteins in higher eukaryotes are conserved and required for the correct excision of intron sequences from primary transcripts by spliceosomes?
- 11.12 Match one of the following terms with each of the descriptions given. *Terms:* (1) sigma (σ) factor; (2) poly(A) tail; (3) TATAAT; (4) exons; (5) TATAAAA; (6) RNA polymerase III; (7) intron; (8) RNA polymerase II; (9) heterogeneous nuclear RNA (hnRNA);

(10) snRNA; (11) RNA polymerase I; (12) TTGACA;(13) GGCCAATCT (CAAT box).

Descriptions:

- (a) Intervening sequence found in many eukaryotic genes.
- (b) A conserved nucleotide sequence (-30) in eukaryotic promoters involved in the initiation of transcription.
- (c) Small RNA molecules that are located in the nuclei of eukaryotic cells, most as components of the spliceosome, that participate in the excision of introns from nuclear gene transcripts.
- (d) A sequence (-10) in the nontemplate strand of the promoters of *E. coli* that facilitates the localized unwinding of DNA when complexed with RNA polymerase.
- (e) The RNA polymerase in the nucleus that catalyzes the synthesis of all rRNAs except for the small 5S rRNA.
- (f) The subunit of prokaryotic RNA polymerase that is responsible for the initiation of transcription at promoters.
- (g) An *E. coli* promoter sequence located 35 nucleotides upstream from the transcription–initiation site; it serves as a recognition site for the sigma factor.
- (h) The RNA polymerase in the nucleus that catalyzes the synthesis of the transfer RNA molecules and small nuclear RNAs.
- (i) A polyadenosine tract 20 to 200 nucleotides long that is added to the 3' end of most eukaryotic messenger RNAs.
- (j) The RNA polymerase that transcribes nuclear genes that encode proteins.
- (k) A conserved sequence in the nontemplate strand of eukaryotic promoters that is located about 80 nucleotides upstream from the transcription start site.
- (l) Segments of a eukaryotic gene that correspond to the sequences in the final processed RNA transcript of the gene.
- (m) The population of primary transcripts in the nucleus of a eukaryotic cell.
- **11.13** (a) Which of the following nuclear pre-mRNA nucleotide sequences potentially contains an intron?
 - (1) 5'-UGACCAUGGCGCUAACACUGCCAAUUG-GCAAUACUGACCUGAUAGCAUCAGCCAA-3'
 - (2) 5'- U A G U C U C A U C U G U C C A U U G A C U U C -GAAACUGAAUCGUAACUCCUACGUCUAUGGA-3'
 - (3) 5'-UAGCUGUUUGUCAUGACUGACUGGUCACU-AUCGUACUAACCUGUCAUGCAAUGUC-3'

- (4) 5'-UAGCAGUUCUGUCGCCUCGUGGUGCUGCUG GCCCUUCGUCGCUCGGGCUUAGCUA-3'
- (5) 5'-UAGGUUCGCAUUGACGUACUUCUGAAAC-UACUAACUACUAACGCAUCGAGUCUCAA-3'

(b) One of the five pre-mRNAs shown in (a) may undergo RNA splicing to excise an intron sequence. What mRNA nucleotide sequence would be expected to result from this splicing event?

- 11.14 What is the function of the introns in eukaryotic genes?
- 11.15 A particular gene is inserted into the phage lambda chromosome and is shown to contain three introns. (a) The primary transcript of this gene is purified from isolated nuclei. When this primary transcript is hybridized under R-loop conditions with the recombinant lambda chromosome carrying the gene, what will the R-loop structure(s) look like? Label your diagram. (b) The mRNA produced from the primary transcript of this gene is then isolated from cytoplasmic polyribosomes and similarly examined by the R-loop hybridization procedure using the recombinant lambda chromosome carrying the gene. Diagram what the R-loop structure(s) will look like when the cytoplasmic mRNA is used. Again, label the components of your diagram.
- **11.16** A segment of DNA in *E. coli* has the following sequence of nucleotide pairs:

When this segment of DNA is transcribed by RNA polymerase, what will be the sequence of nucleotides in the RNA transcript if the promoter is located to the left of the sequence shown?

11.17 A segment of DNA in *E. coli* has the following sequence of nucleotide pairs:

ATGCTACTGCTATTCGCTGTATCG-5'

When this segment of DNA is transcribed by RNA polymerase, what will be the sequence of nucleotides in the RNA transcript?

11.18 A segment of DNA in *E. coli* has the following sequence of nucleotide pairs:

GCAATGGGCTGTATCGATGCTACTGCTAT-5'

When this segment of DNA is transcribed by RNA polymerase, what will be the sequence of nucleotides in the RNA transcript?

11.19 A segment of human DNA has the following sequence of nucleotide pairs:

CGTTACCCGACATAGCTACGATGACGATA-3

When this segment of DNA is transcribed by RNA polymerase, what will be the sequence of nucleotides in the RNA transcript?

- **11.20** The genome of a human must store a tremendous amount of information using the four nucleotide pairs present in DNA. What do the Morse code and the language of computers tell us about the feasibility of storing large amounts of information using an alphabet composed of just four letters?
- **11.21** What is the central dogma of molecular genetics? What impact did the discovery of RNA tumor viruses have on the central dogma?
- 11.22 The biosynthesis of metabolite X occurs via six steps catalyzed by six different enzymes. What is the minimal number of genes required for the genetic control of this metabolic pathway? Might more genes be involved? Why?
- **11.23** What do the processes of DNA synthesis, RNA synthesis, and polypeptide synthesis have in common?
- **11.24** What are the two stages of gene expression? Where do they occur in a eukaryotic cell? a prokaryotic cell?
- **11.25** Compare the structures of primary transcripts with those of mRNAs in prokaryotes and eukaryotes. On average, in which group of organisms do they differ the most?
- **11.26** What five types of RNA molecules participate in the process of gene expression? What are the functions of each type of RNA? Which types of RNA perform their function(s) in (a) the nucleus and (b) the cytoplasm?
- **11.27** Why was the need for an RNA intermediary in protein synthesis most obvious in eukaryotes? How did researchers first demonstrate that RNA synthesis occurred in the nucleus and that protein synthesis occurred in the cytoplasm?
- **11.28** Two eukaryotic genes encode two different polypeptides, each of which is 335 amino acids long. One gene contains a single exon; the other gene contains an intron 41,324 nucleotide pairs long. Which gene would you expect to be transcribed in the least amount of time? Why? When the mRNAs specified by these genes are translated, which mRNA would you expect to be translated in the least time? Why?

- **11.29** Design an experiment to demonstrate that RNA transcripts are synthesized in the nucleus of eukaryotes and are subsequently transported to the cytoplasm.
- 11.30 Total RNA was isolated from human cells growing in culture. This RNA was mixed with nontemplate strands (single strands) of the human gene encoding the enzyme thymidine kinase, and the RNA–DNA mixture was incubated for 12 hours under renaturation conditions. Would you expect any RNA–DNA duplexes to be formed during the incubation? If so, why? If not, why not? The same experiment was then performed using the template strand of the thymidine kinase gene. Would you expect any RNA–DNA duplexes to be formed in this second experiment? If so, why? If not, why not?
- **11.31** Two preparations of RNA polymerase from *E. coli* are used in separate experiments to catalyze RNA synthesis *in vitro* using a purified fragment of DNA carrying the *argH* gene as template DNA. One preparation catalyzes the synthesis of RNA chains that are highly heterogeneous in size. The other preparation catalyzes the synthesis of RNA chains that are all the same length. What is the most likely difference in the composition of the RNA polymerases in the two preparations?
- **11.32** Transcription and translation are coupled in prokaryotes. Why is this not the case in eukaryotes?
- **11.33** What two elements are almost always present in the promoters of eukaryotic genes that are transcribed by RNA

polymerase II? Where are these elements located relative to the transcription start site? What are their functions?

- **11.34** In what ways are most eukaryotic gene transcripts modified? What are the functions of these posttranscriptional modifications?
- **11.35** How does RNA editing contribute to protein diversity in eukaryotes?
- **11.36** How do the mechanisms by which the introns of tRNA precursors, *Tetrahymena* rRNA precursors, and nuclear pre-mRNAs are excised differ? In which process are snRNAs involved? What role(s) do these snRNAs play?
- **11.37** A mutation in an essential human gene changes the 5'-splice site of a large intron from GT to CC. Predict the phenotype of an individual homozygous for this mutation.
- 11.38 Total RNA was isolated from nuclei of human cells growing in culture. This RNA was mixed with a purified, denatured DNA fragment that carried a large intron of a housekeeping gene (a gene expressed in essentially all cells), and the RNA–DNA mixture was incubated for 12 hours under renaturation conditions. Would you expect any RNA–DNA duplexes to be formed during the incubation? If so, why? If not, why not? The same experiment was then performed using total cytoplasmic RNA from these cells. Would you expect any RNA–DNA duplexes to be formed in this second experiment? If so, why? If not, why not?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

Duchenne muscular dystrophy (DMD) is an X-linked recessive disease in humans that affects about one in 3300 newborn males. Individuals with DMD undergo progressive muscle degeneration starting early in life. They are usually confined to wheel chairs by their teens and commonly die in their late teens or early twenties. The disorder is caused by mutations in the human *DMD* gene, which encodes a protein called dystrophin. This protein is associated with the intracellular membranes of muscle cells. The *DMD* gene is one of the largest genes known and is composed of many exons and introns. Because of its medical importance, the NCBI web site contains a large amount of information on the *DMD* gene and its product dystrophin.

- 1. How large is the human *DMD* gene? How many exons and introns does it contain? How large is the *DMD* mRNA? the *DMD* protein coding sequence?
- 2. What is the largest exon in the human *DMD* gene? the smallest exon? Where are the mutations located that cause

Duchenne muscular dystrophy? Some of the mutations in this gene cause a less severe form of muscular dystrophy called Becker muscular dystrophy. Where are these mutations located?

3. Do other species contain genes that are closely related to the human *DMD* gene and encode similar dystrophins? What species? How similar are these genes to each other and to the human *DMD* gene?

Hint: At the NCBI home page (click on each of the following) \rightarrow Human genome resources \rightarrow Gene Database \rightarrow Search with query DMD AND human[orgn] \rightarrow 1. DMD \rightarrow Primary Source: HGNC:2928 \rightarrow under Gene Symbol Links, click GENATLAS \rightarrow DMD \rightarrow See the exons. To view homologous genes in other organisms, go back to the results of your DMD gene search, and click on HomoloGene. Also, search the OMIM (the Online Medical Inheritance in Man) database for more information about Duchenne and Becker muscular dystrophies.

Translation and the Genetic Code



CHAPTER OUTLINE

- Protein Structure
- One Gene—One Colinear Polypeptide
- Protein Synthesis: Translation
- The Genetic Code
- Codon-tRNA Interactions

before publishing his observations in 1910. In his paper, Herrick emphasized the chronic nature of the anemia and the presence of



Scanning electron micrograph of normal and crescent-shaped red blood cells in a patient with sickle-cell anemia.

the sickle-shaped red cells. In 1916, at age 32, the patient died from severe anemia and kidney damage.

James Herrick was the first to publish a description of sickle-cell anemia, the first inherited human disease to be understood at the molecular level. Hemoglobin contains four polypeptidestwo α -globin chains and two β -globin chains—and an iron-containing heme group. In 1957, Vernon Ingram and colleagues demonstrated that the sixth amino acid of the β-chain of sickle-cell hemoglobin was valine, whereas glutamic acid was present at this position in normal adult human hemoglobin. This single amino acid change in a single polypeptide chain is responsible for all the symptoms of sickle-cell anemia.

Sickle-Cell Anemia: Devastating Effects of a Single Base-Pair Change In 1904 James Herrick, a Chicago physician, and Ernest Irons, a medical intern working under Herrick's supervision, examined

medical intern working under Herrick's supervision, examined the blood cells of one of their patients. They noticed that many of the red blood cells of the young man were thin and elongated, in striking contrast to the round, donutlike red cells of their other patients. They obtained fresh

blood samples and repeated their microscopic examinations several times, always with the same result. The blood of this patient always contained cells shaped like the sickles that farmers used to harvest grain at that time.

The patient was a 20-yearold college student who was experiencing periods of weakness and dizziness. In many respects, the patient seemed normal, both physically and mentally. His major problem was fatigue. However, a physical exam showed an enlarged heart and enlarged lymph nodes. His heart always seemed to be working too hard, even when he was resting. Blood tests showed that the patient was anemic; the hemoglobin content of his blood was about half the normal level. Hemoglobin is the complex protein that carries oxygen from the lungs to other tissues. Herrick charted this patient's symptoms for six years

How does the genetic information of an organism, stored in the sequence of nucleotide pairs in DNA, control the phenotype of the organism? How does a nucleotide-pair change in a gene—like the mutation that causes sickle-cell anemia—alter the structure of a protein, the emissary through which the gene acts? In Chapter 11, we discussed the transfer of genetic information stored in the sequences of nucleotide pairs in DNA to the sequences of nucleotides in mRNA molecules, which, in eukaryotes, carry that information from the nucleus to the sites of protein synthesis in the cytoplasm. The transfer of information from DNA to RNA (transcription) and RNA processing occur in the nucleus. In this chapter, we examine the process by which genetic information stored in sequences of nucleotides in mRNAs is used to specify the sequences of amino acids in polypeptide gene products. This process, **translation**, takes place in the cytoplasm on complex workbenches called ribosomes and requires the participation of many macromolecules.

Protein Structure

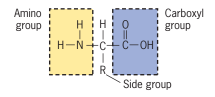
Proteins are complex macromolecules composed of 20 different amino acids.

Collectively, the proteins constitute about 15 percent of the wet weight of cells. Water molecules account for 70 percent of the total weight of living cells. With the exception of water, proteins are by far the most prevalent component of living organisms in terms of

total mass. Not only are proteins major components in terms of cell mass, but they also play many roles vital to the lives of all cells. Before discussing the synthesis of proteins, we need to become more familiar with their structure.

POLYPEPTIDES: TWENTY DIFFERENT AMINO ACID SUBUNITS

Proteins are composed of polypeptides, and every polypeptide is encoded by a gene. Each polypeptide consists of a long sequence of amino acids linked together by covalent bonds. Twenty different amino acids are present in most proteins. Occasionally, one or more of the amino acids are chemically modified after a polypeptide is synthesized, yielding a novel amino acid in the mature protein. The structures of the 20 common amino acids are shown in **Figure 12.1**. All the amino acids except proline contain a *free amino group* and a *free carboxyl group*.



The amino acids differ from each other by the *side groups* (designated **R** for *R*adical) that are present. The highly varied side groups provide the structural diversity of proteins. These side chains are of four types: (1) hydrophobic or nonpolar groups, (2) hydrophilic or polar groups, (3) acidic or negatively charged groups, and (4) basic or positively charged groups (Figure 12.1). The chemical diversity of the side groups of the amino acids is responsible for the enormous structural and functional versatility of proteins.

A **peptide** is a compound composed of two or more amino acids. Polypeptides are long sequences of amino acids, ranging in length from 51 amino acids in insulin to over 1000 amino acids in the silk protein fibroin. Given the 20 different amino acids commonly found in polypeptides, the number of different polypeptides that are possible is truly enormous. For example, the number of different amino acid sequences that can occur in a polypeptide containing 100 amino acids is 20¹⁰⁰. Since 20¹⁰⁰ is too large to comprehend, let's consider a short peptide. There are 1.28 billion (207) different amino acid sequences possible in a peptide seven amino acids long. The amino acids in polypeptides are covalently joined by linkages called **peptide bonds**. Each

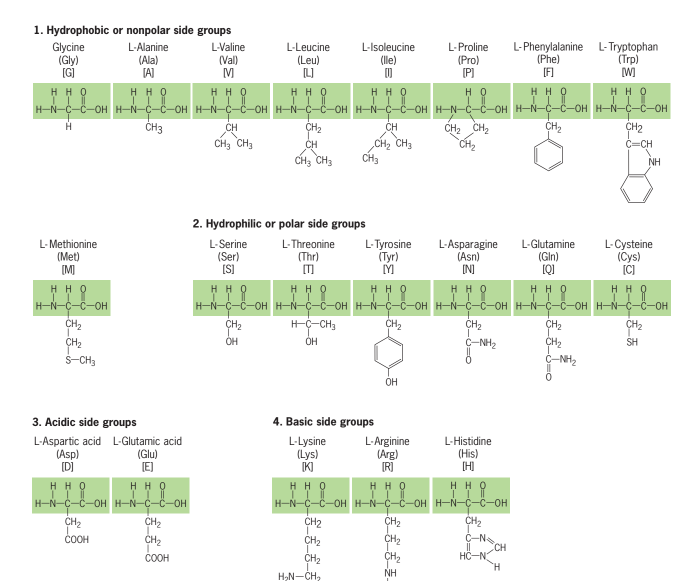


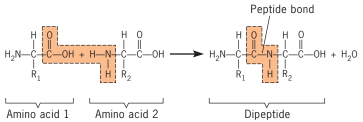
FIGURE 12.1 Structures of the 20 amino acids commonly found in proteins. The amino and carboxyl groups, which participate in peptide bond formation during protein synthesis, are shown in the shaded areas. The side groups, which are different for each amino acid, are shown below the shaded areas. The standard three-letter abbreviations are shown in parentheses. The one-letter symbol for each amino acid is given in brackets.

Ċ=NH | NH₂

peptide bond is formed by a reaction between the amino group of one amino acid and the carboxyl group of a second amino acid with the elimination of a water molecule (**Figure 12.2**).

PROTEINS: COMPLEX THREE-DIMENSIONAL STRUCTURES

Four different levels of organization—primary, secondary, tertiary, and quaternary—are distinguished in the complex three-dimensional structures of proteins. The *primary structure* of a polypeptide is its amino acid sequence, which is specified by the nucleotide sequence of a gene. The *secondary structure* of a polypeptide refers to the spatial interrelationships of the amino acids in segments of the polypeptide. The *tertiary structure* of a polypeptide refers to its overall



■ FIGURE 12.2 The formation of a peptide bond between two amino acids by the removal of water. Each peptide bond connects the amino group of one amino acid and the carboxyl group of the adjacent amino acid.

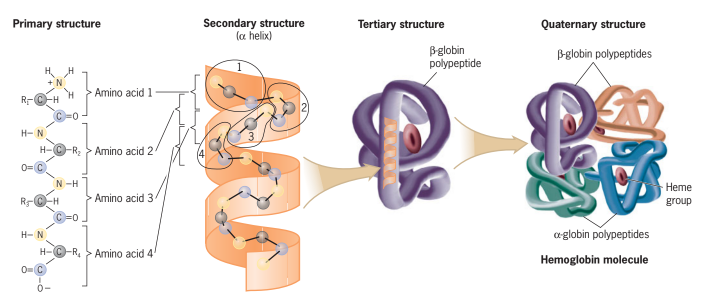


FIGURE 12.3 The four levels of organization in proteins—(1) primary, (2) secondary, (3) tertiary, and (4) quaternary structures—are illustrated using human hemoglobin as an example.

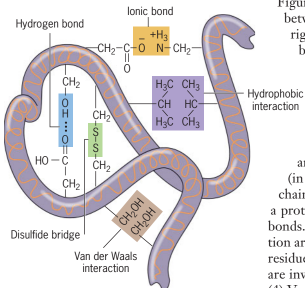
organization (**Figure 12.3**). Most polypeptides will fold spontaneously into specific conformations dictated by their primary structures. If denatured (unfolded) by treatment with appropriate solvents, most proteins will re-form their original conformations when the denaturing agent is removed. Thus, in most cases, all of the information required for shape determination resides in the primary structure of the protein. In some cases, protein folding involves interactions with proteins called **chaperones** that help nascent polypeptides form the proper three-dimensional structure.

folding in three-dimensional space, and the *quaternary structure* refers to the association of two or more polypeptides in a multimeric protein. Hemoglobin provides an excellent example of the complexity of proteins, exhibiting all four levels of structural

The two most common types of secondary structure in proteins are α *helices* (see Figure 12.3) and β *sheets*. Both structures are maintained by hydrogen bonding between peptide bonds located in close proximity to one another. The α helix is a rigid cylinder in which each peptide bond is hydrogen bonded to the peptide bond between amino acids three and four residues away. Because of its rigid structure, proline cannot be present within an α helix. A β sheet occurs when a polypeptide folds back upon itself, sometimes repeatedly, and the parallel segments are held in place by hydrogen bonding between neighboring peptide bonds.

Whereas the spatial organization of adjacent amino acids and segments of a polypeptide determine its secondary structure, the overall folding of the complete polypeptide defines its tertiary structure, or *conformation*. In general, amino acids with hydrophilic side chains are located on the surfaces of proteins (in contact with the aqueous cytoplasm), whereas those with hydrophobic side chains interact with each other in the interior regions. The tertiary structure of a protein is maintained primarily by a large number of relatively weak noncovalent bonds. The only covalent bonds that play a significant role in protein conformation are disulfide (S—S) bridges that form between appropriately positioned cysteine residues (**■ Figure 12.4**). However, four different types of noncovalent interactions are involved: (1) ionic bonds, (2) hydrogen bonds, (3) hydrophobic interactions, and (4) Van der Waals interactions (Figure 12.4).

lonic bonds occur between amino acid side chains with opposite charges—for example, the side groups of lysine and glutamic acid (see Figure 12.1). Ionic bonds are strong forces under some conditions, but they are relatively weak interactions in the aqueous interiors of living cells because the polar water molecules partially neutralize or shield the charged groups. **Hydrogen bonds** are weak interactions between electronegative atoms



■ **FIGURE 12.4** The five types of molecular interactions that determine the tertiary structure, or three-dimensional conformation, of a polypeptide. The disulfide bridge is a covalent bond; all other interactions are noncovalent.

(which have a partial negative charge) and hydrogen atoms (which are electropositive) that are linked to other electronegative atoms. **Hydrophobic interactions** are associations of nonpolar groups with each other when present in aqueous solutions because of their insolubility in water. Hydrogen bonds and hydrophobic interactions play important roles in DNA structure; thus, we have discussed them in some detail in Chapter 9 (see Table 9.2). **Van der Waals interactions** are weak attractions that occur between atoms when they are placed in close proximity to one another. Van der Waals forces are very weak, with about one one-thousandth of the strength of a covalent bond, but they play an important role in maintaining the conformations of closely aligned regions of macromolecules.

Quaternary structure exists only in proteins that contain more than one polypeptide. Hemoglobin provides a good illustration of quaternary structure, being a tetrameric molecule composed of two α -globin chains and two β -globin chains, plus four iron-containing heme groups (see Figure 12.3).

In a few cases, the primary translation products contain short amino acid sequences, called **inteins**, which excise themselves from the nascent polypeptides. Inteins occur in both eukaryotes and prokaryotes. For example, one of the first inteins discovered is in the RecA protein, which is involved in recombination and DNA repair in *Mycobacterium tuberculosis*, the bacterium that causes tuberculosis.

Since the secondary, tertiary, and quaternary structures of proteins and intein excision usually are determined by the primary structure(s) of the polypeptide(s) involved, in the rest of this chapter we will focus on the mechanisms by which genes control the primary structures of polypeptides.

- Most genes exert their effect(s) on the phenotype of an organism through proteins, which are large macromolecules composed of polypeptides.
- Each polypeptide is a chainlike polymer assembled from different amino acids.
- The amino acid sequence of each polypeptide is specified by the nucleotide sequence of a gene.
- The vast functional diversity of proteins results in part from their complex three-dimensional structures.

One Gene-One Colinear Polypeptide

Most genes encode polypeptides. Before exploring how they do this—that is, how a gene's nucleotide sequence specifies a polypeptide's amino acid sequence—let's consider two classic genetic studies that enhanced our understanding of the connection between genes and their polypeptide products. The sequence of nucleotide pairs in a gene specifies a colinear sequence of amino acids in its polypeptide product.

BEADLE AND TATUM: ONE GENE-ONE ENZYME

During the late 1930s, George Beadle and Boris Ephrussi performed pioneering experiments on *Drosophila* eye color mutants. They identified genes that are required for the synthesis of specific eye pigments, indicating that enzyme-catalyzed metabolic pathways are under genetic control. Their results motivated Beadle to search for the ideal organism to use in extending this work. He chose the salmoncolored bread mold *Neurospora crassa* because it can grow on medium containing only (1) inorganic salts, (2) a simple sugar, and (3) one vitamin, biotin. *Neurospora* growth medium containing only these components is called "minimal medium." Beadle and his new collaborator, Edward Tatum, reasoned that *Neurospora* must be capable of synthesizing all the other essential metabolites, such as the purines, pyrimidines, amino acids, and other vitamins, *de novo*. Furthermore, they reasoned that the biosynthesis of these growth factors must be under genetic control. If so,

KEY POINTS

mutations in genes whose products are involved in the biosynthesis of essential metabolites would be expected to produce mutant strains with additional growth-factor requirements.

Beadle and Tatum tested this prediction by irradiating asexual spores (conidia) of wild-type *Neurospora* with X rays or ultraviolet light, and screening the clones produced by the mutagenized spores for new growth-factor requirements (**Figure 12.5**). In order

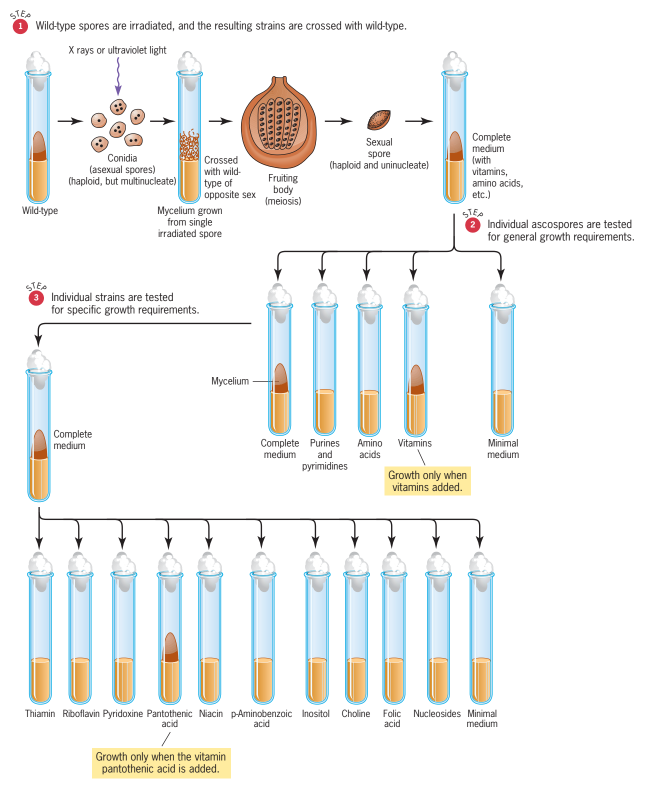


FIGURE 12.5 Diagram of Beadle and Tatum's experiment with *Neurospora* that led to the one gene-one enzyme hypothesis.

to select strains with a mutation in only one gene, they studied only mutant strains that yielded a 1:1 mutant to wild-type progeny ratio when crossed with wild type. They identified mutants that grew on medium supplemented with all the amino acids, purines, pyrimidines, and vitamins (called "complete medium") but could not grow on minimal medium. They analyzed the ability of these mutants to grow on medium supplemented with just amino acids, or just vitamins, and so on (Figure 12.5, step 2). For example, Beadle and Tatum identified mutant strains that grew in the presence of vitamins but could not grow in medium supplemented with amino acids or other growth factors. They next investigated the ability of these vitamin-requiring strains to grow on media supplemented with each of the vitamins separately (Figure 12.5, step 3).

In this way, Beadle and Tatum demonstrated that each mutation resulted in a requirement for one growth factor. By correlating their genetic analyses with biochemical studies of the mutant strains, they demonstrated in several cases that one mutation resulted in the loss of one enzyme activity. This work, for which Beadle and Tatum received a Nobel Prize in 1958, was soon verified by similar studies of many other organisms in many laboratories. The *one gene–one enzyme* concept thus became a central tenet of molecular genetics.

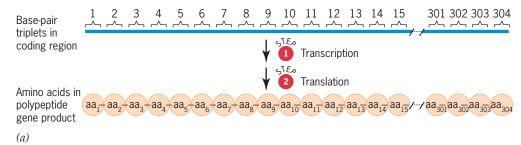
Subsequent to the work of Beadle and Tatum, many enzymes and structural proteins were shown to be heteromultimeric—that is, to contain two or more different polypeptide chains, with each polypeptide encoded by a separate gene. For example, in *E. coli*, the enzyme tryptophan synthetase is a heterotetramer composed of two α polypeptides encoded by the *trpA* gene and two β polypeptides encoded by the *trpB* gene. Similarly, the hemoglobins, which transport oxygen from our lungs to all other tissues of our bodies, are tetrameric proteins that contain two α -globin chains and two β -globin chains, as well as four oxygen-binding heme groups (see Figure 12.3). Other enzymes, for example, *E. coli* DNA polymerase III (Chapter 10) and RNA polymerase II (Chapter 11), contain many different polypeptide subunits, each encoded by a separate gene. Thus, the one gene–one enzyme concept was modified to *one gene–one polypeptide*.

COLINEARITY BETWEEN THE CODING SEQUENCE OF A GENE AND ITS POLYPEPTIDE PRODUCT

Now that we have established the one gene—one polypeptide relationship, we can ask whether the nucleotide pair sequences in genes are colinear with the amino acid sequences of the polypeptides that they encode. That is, do the first base pairs of the coding sequence of a gene specify the first amino acid of the polypeptide, and so on, in a systematic way? The answer is that genes and their polypeptide products are, indeed, colinear structures; this relationship is illustrated in **Figure 12.6a**. As was discussed in Chapter 11, most of the genes of multicellular eukaryotes are interrupted by noncoding introns. However, the presence of introns in genes simply means that there is no direct correlation in physical distances between the positions of base pairs in a gene and the positions of amino acids in the polypeptide specified by that gene (**Figure 12.6b**).

The first strong evidence for colinearity between a gene and its polypeptide product resulted from studies by Charles Yanofsky and colleagues on the *E. coli* gene that encodes the α subunit of the enzyme tryptophan synthetase. As mentioned earlier, this enzyme contains two α polypeptides encoded by the *trpA* gene and two β polypeptides encoded by the *trpB* gene. Yanofsky and coworkers performed a detailed genetic analysis of mutations in the *trpA* gene and correlated the genetic data with biochemical data on the sequences of the wild-type and mutant tryptophan synthetase α polypeptides. They demonstrated that there was a direct correlation between the map positions of mutations in the *trpA* gene and the positions of the resultant amino acid substitutions in the tryptophan synthetase α

Coding region of typical uninterrupted prokaryotic gene.



Coding region of typical intron-interrupted eukaryotic gene.

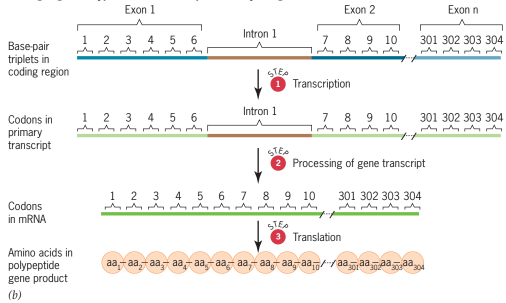


FIGURE 12.6 Colinearity between the coding regions of genes and their polypeptide products.

polypeptide (**Figure 12.7**). Definitive evidence for colinearity has been provided by direct comparisons of the nucleotide sequences of genes and the amino acid sequences of their polypeptide products.

	trpA gene		
Genetic map —	A3 A33 A446 A487 A223 A23 A46 A187 A78 A58 A169 A96		
Map distances (not to scale)	0 1.4 0.04 0.3 0.4 0.001 0.6 0.5 0.001 0.02 0.3		
Amino acid in wild-type polypeptide	Glu Glu Tyr Leu Thr Gly Gly Gly Gly Gly Ser Gln		
Amino acid in mutant polypeptide	Val Met Cys Arg IIe Arg Glu Val Cys Asp Leu (term.)		
Position of amino acid change in polypeptide H_2	N—1—49—49—175—177—183—211—211—213—234—234—235—243—268—COOH		

peptide product, the α polypeptide of tryptophan synthetase. The map positions of mutations in the *trpA* gene are shown at the top, and the locations of the amino acid substitutions produced by these mutations are shown below the map.

KEY POINTS

■ **FIGURE 12.7** Colinearity between the *E. coli trpA* gene and its poly-

• Beadle and Tatum's experiments with Neurospora led to the one gene-one enzyme hypothesis, which was subsequently modified to the one gene-one polypeptide concept.

The sequences of nucleotide pairs in a gene and amino acids in its polypeptide product are colinear.

Protein Synthesis: Translation

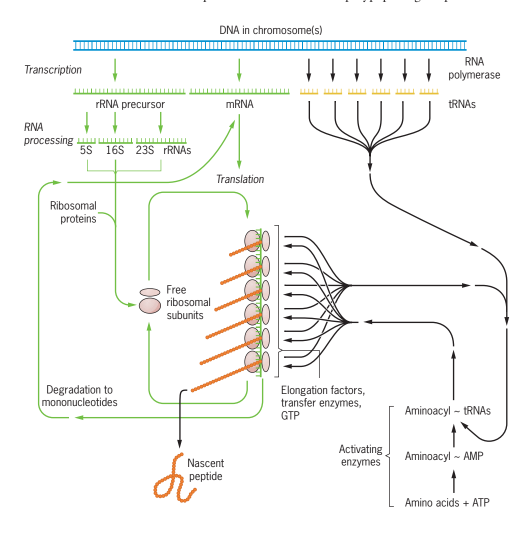
The process by which the genetic information stored in the sequence of nucleotides in an mRNA is translated, according to the specifications of the genetic code, into the sequence of amino acids in the polypeptide gene product is complex, requiring the functions of a large

The genetic information in mRNA molecules is translated into the amino acid sequences of polypeptides according to the specifications of the genetic code.

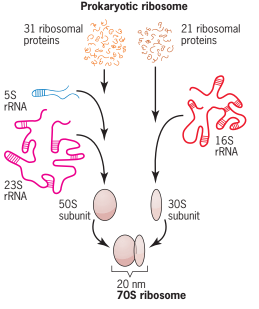
number of macromolecules. These include (1) over 50 polypeptides and three to five RNA molecules present in each ribosome (the exact composition varies from species to species), (2) at least 20 amino acid-activating enzymes, (3) 40 to 60 different tRNA molecules, and (4) numerous soluble proteins involved in polypeptide chain initiation, elongation, and termination. Because many of these macromolecules, particularly the components of the ribosome, are present in large quantities in each cell, the translation system makes up a major portion of the metabolic machinery of each cell.

OVERVIEW OF PROTEIN SYNTHESIS

Before focusing on the details of the translation process, we should preview the process of protein synthesis in its entirety. An overview of protein synthesis, illustrating its complexity and the major macromolecules involved, is presented in **Figure 12.8**. The first step in gene expression, transcription, involves the transfer of information stored in genes to messenger RNA (mRNA) intermediaries, which carry that information to the sites of polypeptide synthesis in the cytoplasm. Transcription is discussed in detail in Chapter 11. The second step, translation, involves the transfer of the information in mRNA molecules into the sequences of amino acids in polypeptide gene products.



■ FIGURE 12.8 Overview of protein synthesis. The sizes of the rRNA molecules shown are correct for bacteria; larger rRNAs are present in eukaryotes. For simplicity, all RNA species have been transcribed from contiguous segments of a single DNA molecule. In reality, the various RNAs are transcripts of genes located at different positions on from one to many chromosomes. Details of the various stages of protein synthesis are discussed in subsequent sections of this chapter.



Eukaryotic (mammalian) ribosome

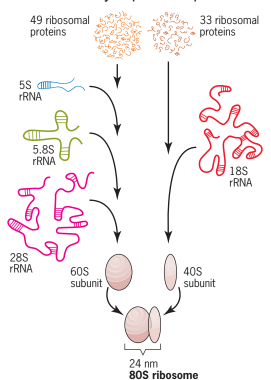


FIGURE 12.9 Macromolecular composition of prokaryotic and eukaryotic ribosomes.

Translation occurs on ribosomes, which are complex macromolecular structures located in the cytoplasm. Translation involves three types of RNA, all of which are transcribed from DNA templates (chromosomal genes). In addition to mRNAs, three to five RNA molecules (rRNA molecules) are present as part of the structure of each ribosome, and 40 to 60 small RNA molecules (tRNA molecules) function as adaptors by mediating the incorporation of the proper amino acids into polypeptides in response to specific nucleotide sequences in mRNAs. The amino acids are attached to the correct tRNA molecules by a set of activating enzymes called **aminoacyl-tRNA synthetases**.

The nucleotide sequence of an mRNA molecule is translated into the appropriate amino acid sequence according to the dictations of the genetic code. Some nascent polypeptides contain short amino acid sequences at the amino or carboxyl termini that function as signals for their transport into specific cellular compartments such as the endoplasmic reticulum, mitochondria, chloroplasts, or nuclei. Nascent secretory proteins, for example, contain a short *signal sequence* at the amino terminus that directs the emerging polypeptide to the membranes of the endoplasmic reticulum. Similar targeting sequences are present at the amino termini of proteins destined for import into mitochondria and chloroplasts. Some nuclear proteins contain targeting extensions at the carboxyl termini. In many cases, the targeting peptides are removed enzymatically by specific peptidases after transport of the protein into the appropriate cellular compartment.

The ribosomes may be thought of as workbenches, complete with machines and tools needed to make a polypeptide. They are nonspecific in the sense that they can synthesize any polypeptide (any amino acid sequence) encoded by a particular mRNA molecule, even an mRNA from a different species. Each mRNA molecule is simultaneously translated by several ribosomes, resulting in the formation of a polyribosome, or polysome. Given this brief overview of protein synthesis, we will now examine some of the more important components of the translation machinery more closely.

COMPONENTS REQUIRED FOR PROTEIN SYNTHESIS: RIBOSOMES

Living cells devote more energy to the synthesis of proteins than to any other aspect of metabolism. About one-third of the total dry mass of most cells consists of molecules that participate directly in the biosynthesis of proteins. In *E. coli*, the approximately 200,000 ribosomes account for 25 percent of the dry weight of each cell. This commitment of a major proportion of the metabolic machinery of cells to the process of protein synthesis documents its importance in the life forms that exist on our planet.

When the sites of protein synthesis were labeled in cells grown for short intervals in the presence of radioactive amino acids and were visualized by autoradiography, the results showed that proteins are synthesized on the ribosomes. In prokaryotes, ribosomes are distributed throughout cells; in eukaryotes, they are located in the cytoplasm, frequently on the extensive intracellular membrane network of the endoplasmic reticulum.

Ribosomes are approximately half protein and half RNA (**■** Figure 12.9). They are composed of two subunits, one large and one small, which dissociate when the translation of an mRNA molecule is completed and reassociate during the initiation of translation. Each subunit contains a large, folded RNA molecule on which the ribosomal proteins assemble. Ribosome sizes are most frequently expressed in terms of their rates of sedimentation during centrifugation, in Svedberg (S) units. [One Svedberg unit is equal to a sedimentation coefficient (velocity/centrifugal force) of 10^{-13} seconds.] The *E. coli* ribosome, like the ribosomes of other prokaryotes, has a molecular weight of 2.5×10^6 , a size of 70S, and dimensions of about 20 nm $\times 25$ nm. The ribosomes of eukaryotes are larger (usually about 80S); however, size varies from species to species. The ribosomes present in the mitochondria and chloroplasts of eukaryotic cells are smaller (usually about 60S).

Although the size and macromolecular composition of ribosomes vary, the overall three-dimensional structure of the ribosome is basically the same in all organisms. In

E. coli, the small (30S) ribosomal subunit contains a 16S (molecular weight about 6×10^5) RNA molecule plus 21 different polypeptides, and the large (50S) subunit contains two RNA molecules (5S, molecular weight about 4×10^4 , and 23S, molecular weight about 1.2×10^6) plus 31 polypeptides. In mammalian ribosomes, the small subunit contains an 18S RNA molecule plus 33 polypeptides, and the large subunit contains three RNA molecules of sizes 5S, 5.8S, and 28S plus 49 polypeptides. In organelles, the corresponding rRNA sizes are 5S, 13S, and 21S.

Masayasu Nomura and his colleagues were able to disassemble the 30S ribosomal subunit of E. coli into the individual macromolecules and then reconstitute functional 30S subunits from the components. In this way, they studied the functions of individual rRNA and ribosomal protein molecules.

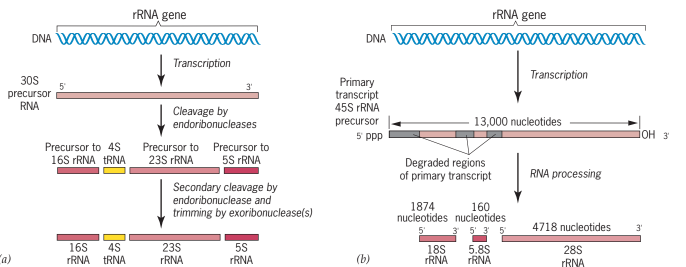
The ribosomal RNA molecules, like mRNA molecules, are transcribed from a DNA template. In eukaryotes, rRNA synthesis occurs in

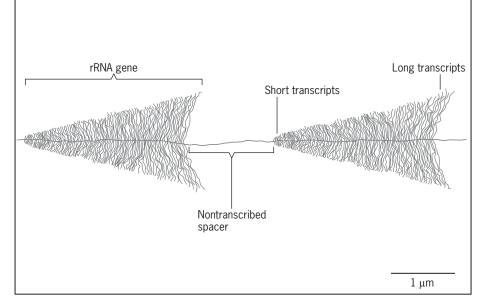
the nucleolus (see Figure 2.1) and is catalyzed by RNA polymerase I. The nucleolus is a highly specialized component of the nucleus devoted exclusively to the synthesis of rRNAs and their assembly into ribosomes. The ribosomal RNA genes are present in tandemly duplicated arrays separated by intergenic spacer regions. The transcription of these tandem sets of rRNA genes can be visualized directly by electron microscopy. (**Figure 12.10**) shows a schematic diagram of the observed transcription.

The transcription of the rRNA genes produces RNA precursors that are much larger than the RNA molecules found in ribosomes. These rRNA precursors undergo posttranscriptional processing to produce the mature rRNA molecules. In E. coli, the rRNA gene transcript is a 30S precursor, which undergoes endonucleolytic cleavages to produce the 5S, 16S, and 23S rRNAs plus one 4S transfer RNA molecule (**Figure 12.11a**). In mammals, the 5.8S, 18S, and 28S rRNAs are cleaved from a 45S precursor (**Figure 12.11b**), whereas the 5S rRNA is produced by posttranscriptional processing of a separate gene transcript. In addition to the posttranscriptional cleavages of rRNA precursors, many of the nucleotides in rRNAs are posttranscriptionally methylated. The methylation is thought to protect rRNA molecules from degradation by ribonucleases.

Multiple copies of the genes for rRNA are present in the genomes of all organisms that have been studied to date. This redundancy of rRNA genes is not surprising considering the large number of ribosomes present per cell. In E. coli, seven rRNA genes ■ FIGURE 12.10 Schematic diagram of an electron micrograph showing the transcription of tandemly repeated rRNA genes in the nucleolus of the new Triturus viridescens. A gradient of fibrils of increasing length is observed for each rRNA gene, and nontranscribed spacer regions separate the genes.

FIGURE 12.11 Synthesis and processing of (a) the 30S rRNA precursor in E. coli and (b) the 45S rRNA precursor in mammals.





(a)

(*rrnA—rrnE*, *rrnG*, *rrnH*) are distributed among three distinct sites on the chromosome. In eukaryotes, the rRNA genes are present in hundreds to thousands of copies. The 5.8S-18S-28S rRNA genes of eukaryotes are present in tandem arrays in the **nucleolar organizer regions** of the chromosomes. In some eukaryotes, such as maize, there is a single pair of nucleolar organizers (on chromosome 6 in maize). In *Drosophila* and the South African clawed toad, *Xenopus laevis*, the sex chromosomes carry the nucleolar organizers. Humans have five pairs of nucleolar organizers located on the short arms of chromosomes 13, 14, 15, 21, and 22. The 5S rRNA genes in eukaryotes are not located in the nucleolar organizer regions. Instead, they are distributed over several chromosomes. However, the 5S rRNA genes are highly redundant, just as are the 5.8S-18S-28S rRNA genes.

COMPONENTS REQUIRED FOR PROTEIN SYNTHESIS: TRANSFER RNAs

Although the ribosomes provide many of the components required for protein synthesis, and the specifications for each polypeptide are encoded in an mRNA molecule, the translation of a coded mRNA message into a sequence of amino acids in a polypeptide requires one additional class of RNA molecules, the transfer RNA (tRNA) molecules. Chemical considerations suggested that direct interactions between the amino acids and the nucleotide triplets or codons in mRNA were unlikely. Thus, in 1958, Francis Crick proposed that some kind of an adaptor molecule must mediate the specification of amino acids by codons in mRNAs during protein synthesis. The adaptor molecules were soon identified by other researchers and shown to be small (4S, 70–95 nucleotides long) RNA molecules. These molecules, first called soluble RNA (sRNA) molecules and subsequently transfer RNA (tRNA) molecules, contain a triplet nucleotide sequence, the anticodon, which is complementary to and base-pairs with the codon sequence in mRNA during translation. There are one to four tRNAs for each of the 20 amino acids.

The amino acids are attached to the tRNAs by high-energy (very reactive) bonds (symbolized ~) between the carboxyl groups of the amino acids and the 3'-hydroxyl termini of the tRNAs. The tRNAs are activated or charged with amino acids in a two-step process, with both reactions catalyzed by the same enzyme, aminoacyl-tRNA synthetase. There is at least one aminoacyl-tRNA synthetase for each of the 20 amino acids. The first step in aminoacyl-tRNA synthesis involves the activation of the amino acid using energy from adenosine triphosphate (ATP):

amino acid + ATP
aminoacyl-tRNA
synthetase
amino acid ~ AMP +
$$(\mathbb{P}) \sim (\mathbb{P})$$

The amino acid~AMP intermediate is not normally released from the enzyme before undergoing the second step in aminoacyl-tRNA synthesis, namely, the reaction with the appropriate tRNA:

The aminoacyl~tRNAs are the substrates for polypeptide synthesis on ribosomes, with each activated tRNA recognizing the correct mRNA codon and presenting the amino acid in a steric configuration (three-dimensional structure) that facilitates peptide bond formation.

The tRNAs are transcribed from genes. As in the case of rRNAs, the tRNAs are transcribed in the form of larger precursor molecules that undergo posttranscriptional processing (cleavage, trimming, methylation, and so forth). The mature tRNA molecules contain several nucleosides that are not present in the primary tRNA gene transcripts. These unusual nucleosides, such as inosine, pseudouridine, dihydrouridine, 1-methyl guanosine, and several others, are produced by posttranscriptional, enzyme-catalyzed modifications of the four nucleosides incorporated into RNA during transcription.

Because of their small size (most are 70 to 95 nucleotides long), tRNAs have been more amenable to structural analysis than the other, larger molecules of RNA involved in protein synthesis. The complete nucleotide sequence and proposed cloverleaf structure of the alanine tRNA of yeast (**Figure 12.12**) were published by Robert W. Holley and colleagues in 1965; Holley shared the 1968 Nobel Prize in Physiology or Medicine for this work. The three-dimensional structure of the phenylalanine tRNA of yeast was determined by X-ray diffraction studies in 1974 (**Figure 12.13**). The anticodon of each tRNA US occurs within a loop (nonhydrogen-bonded region) near the middle of 1 the molecule. Di

It should be apparent that tRNA molecules must contain a great deal of specificity despite their small size. Not only must they (1) have the correct anticodon sequences, so as to respond to the right codons, but they also must (2) be recognized by the correct aminoacyl-tRNA synthetases, so that they are activated with the correct amino acids, and (3) bind to the appropriate sites on the ribosomes to carry out their adaptor functions.

There are three tRNA binding sites on each ribosome (■ Figure 12.14a-b). The A or aminoacyl site binds the incoming aminoacyl-tRNA, the tRNA carrying the next amino acid to be added to the growing polypeptide chain. The **P** or **peptidyl site** binds the tRNA to which the growing polypeptide is attached. The *E* or exit site binds the departing uncharged tRNA.

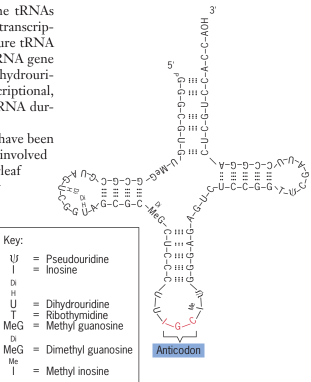
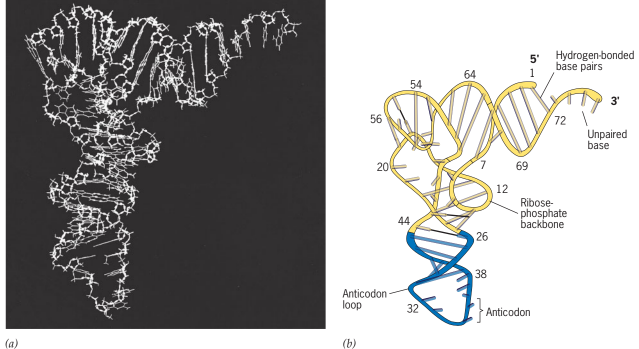


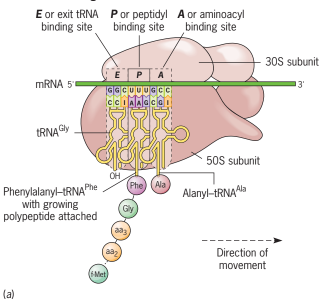
FIGURE 12.12 Nucleotide sequence and cloverleaf configuration of the alanine tRNA of S. cerevisiae. The names of the modified nucleosides present in the tRNA are shown in the inset.



(a)

FIGURE 12.13 Photograph (a) and interpretative drawing (b) of a molecular model of the yeast phenylalanine tRNA based on X-ray diffraction data.

70S ribosome diagram



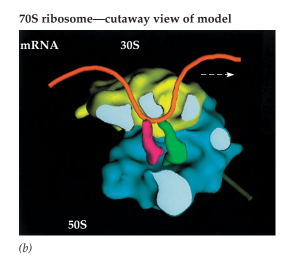


FIGURE 12.14 Ribosome structure in E. coli. (a) Each ribosome/mRNA complex contains three aminoacyl-tRNA binding sites. The A or aminoacyl-tRNA site is occupied by alanyl-tRNA^{Ala}. The P or peptidyl site is occupied by phenylalanyl-tRNA^{Phe}, with the growing polypeptide chain covalently linked to the phenylalanine tRNA. The E or exit site is occupied by tRNA^{Gly} prior to its release from the ribosome. (b) An mRNA molecule (orange), which is attached to the 30S subunit (light green) of the ribosome, contributes specificity to the tRNA-binding sites, which are located largely on the 50S subunit (blue) of the ribosome. The aminoacyl-tRNAs located in the P and A sites are shown in red and dark green, respectively. The E site is unoccupied.

The three-dimensional structure of the 70S ribosome of the bacterium *Thermus thermophilus* has been solved with resolution to 0.55 nm by X-ray crystallography (**■** Figure 12.15*a*-*c*). The crystal structure shows the positions of the three tRNA binding sites at the 50S–30S interface and the relative positions of the rRNAs and ribosomal proteins.

Although the aminoacyl-tRNA binding sites are located largely on the 50S subunit and the mRNA molecule is bound by the 30S subunit, the specificity for aminoacyltRNA binding in each site is provided by the mRNA codon that makes up part of the binding site (see Figure 12.14*b*). As the ribosome moves along an mRNA (or as the mRNA is shuttled across the ribosome), the specificity for the aminoacyl-tRNA binding in the *A*, *P*, and *E* sites changes as different mRNA codons move into register in the binding sites. The ribosomal binding sites by themselves (minus mRNA) are thus capable of binding any aminoacyl-tRNA.

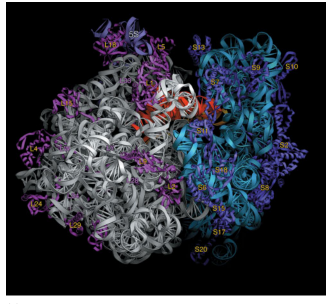
TRANSLATION: THE SYNTHESIS OF POLYPEPTIDES USING mRNA TEMPLATES

We now have reviewed all the major components of the protein-synthesizing system. The mRNA molecules provide the specifications for the amino acid sequences of the polypeptide gene products. The ribosomes provide many of the macromolecular components required for the translation process. The tRNAs provide the adaptor molecules needed to incorporate amino acids into polypeptides in response to codons in mRNAs. In addition, several soluble proteins participate in the process. The translation of the sequence of nucleotides in an mRNA molecule into the sequence of amino acids in its polypeptide product can be divided into three stages: (1) polypeptide chain initiation, (2) chain elongation, and (3) chain termination.

Translation: Polypeptide Chain Initiation

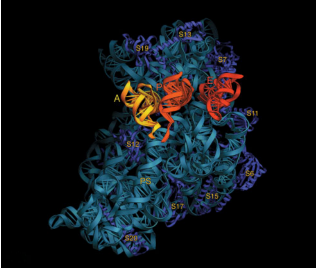
The initiation of translation includes all events that precede the formation of a peptide bond between the first two amino acids of the new polypeptide chain. Although several aspects of the initiation process are the same in prokaryotes and eukaryotes, some are different. Accordingly, we will first examine the initiation of polypeptide chains in *E. coli*, and we will then look at the unique aspects of translational initiation in eukaryotes.

70S ribosome—crystal structure



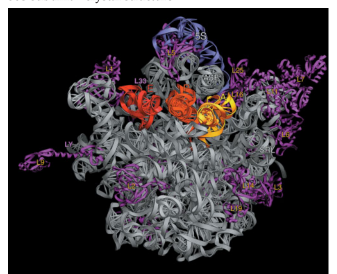
(a)

30S subunit—crystal structure





50S subunit—crystal structure



(b)

■ FIGURE 12.15 Ribosome structure in *Thermus thermophilus*. Crystal structure of the 70S ribosome with 0.55 nm resolution, showing the complete ribosome (*a*) and the interfaces of the 50S (*b*) and 30S (*c*) subunits. (*a*) 50S subunit on the left; 30S subunit on the right. (*b*, *c*) Interfaces of the 50S subunit and the 30S subunit obtained by rotating the structures shown in (*a*) 90° to the left (*b*) or to the right (*c*), respectively. The tRNAs in the *A*, *P*, and *E* sites are shown in gold, orange, and red, respectively. Components: 16S rRNA (cyan); 23S rRNA (gray); 5S rRNA (light blue); 30S subunit proteins (dark blue); and 50S subunit proteins (magenta). L1, large subunit protein 1; S7, small subunit protein 7.

In *E. coli*, the initiation process involves the 30S subunit of the ribosome, a special initiator tRNA, an mRNA molecule, three soluble protein initiation factors: IF-1, IF-2, and IF-3, and one molecule of GTP (**Figure 12.16**). Translation occurs on 70S ribosomes, but the ribosomes dissociate into their 30S and 50S subunits each time they complete the synthesis of a polypeptide chain. In the first stage of the initiation of translation, a free 30S subunit interacts with an mRNA molecule and the initiation factors. The 50S subunit joins the complex to form the 70S ribosome in the final step of the initiation process.

The synthesis of polypeptides is initiated by a special tRNA, designated $tRNA_{f}^{Met}$, in response to a translation initiation codon (usually AUG, sometimes GUG). Therefore, all polypeptides begin with methionine during synthesis. The amino-terminal methionine is subsequently cleaved from many polypeptides. Thus, functional proteins need not have an amino-terminal methionine. The methionine on the initiator $tRNA_{f}^{Met}$ has the

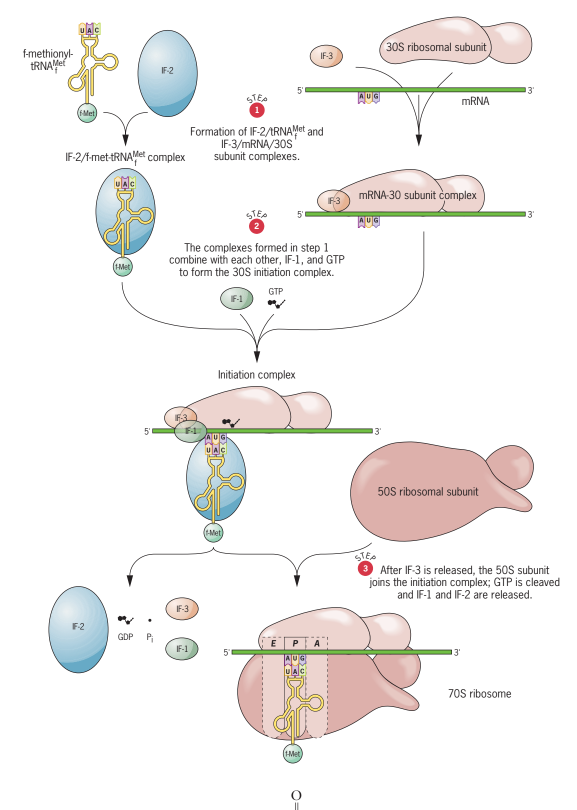


FIGURE 12.16 The initiation of translation in *E. coli*.

(a valine codon when present at internal positions), that occurs in some mRNA molecules.

Polypeptide chain initiation begins with the formation of two complexes: (1) one contains initiation factor IF-2 and methionyl-tRNA_f^{Met}, and (2) the other contains an mRNA molecule, a 30S ribosomal subunit and initiation factor IF-3 (Figure 12.16). The 30S subunit/mRNA complex will form only in the presence of IF-3; thus, IF-3 controls the ability of the 30S subunit to begin the initiation process. The formation of the 30S subunit/mRNA complex depends in part on base-pairing between a

nucleotide sequence near the 3' end of the 16S rRNA and a sequence near the 5' end of the mRNA molecule (**Figure 12.17**). Prokaryotic mRNAs contain a conserved polypurine tract, consensus AGGAGG, located about seven nucleotides upstream from the AUG initiation codon. This conserved hexamer, called the **Shine-Dalgarno sequence** after the scientists who discovered it, is complementary to a sequence near the 3' terminus of the 16S ribosomal RNA. When the Shine-Dalgarno sequences of mRNAs are experimentally modified so that they can no longer base-pair with the 16S rRNA, the modified mRNAs either are not translated or are translated very inefficiently, indicating that this base-pairing plays an important role in translation.

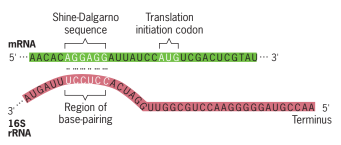
The IF-2/methionyl-tRNA_f^{Met} complex and the mRNA/30S subunit/IF-3 complex subsequently combine with each other and with initiation factor IF-1 and one molecule of GTP to form the complete 30S initiation complex. The final step in the initiation of translation is the addition of the 50S subunit to the 30S initiation complex to produce the complete 70S ribosome. Initiation factor IF-3 must be released from the complex before the 50S subunit can join the complex; IF-3 and the 50S subunit are never found to be associated with the 30S subunit at the same time. The addition of the 50S subunit requires energy from GTP and the release of initiation factors IF-1 and IF-2.

The addition of the 50S ribosomal subunit to the complex positions the initiator tRNA, methionyl-tRNA_f^{Met}, in the peptidyl (*P*) site with the anticodon of the tRNA aligned with the AUG initiation codon of the mRNA. Methionyl-tRNA_f^{Met} is the only aminoacyl-tRNA that can enter the *P* site directly, without first passing through the aminoacyl (*A*) site. With the initiator AUG positioned in the *P* site, the second codon of the mRNA is in register with the *A* site, dictating the aminoacyl-tRNA binding specificity at that site and setting the stage for the second phase in polypeptide synthesis, chain elongation.

The initiation of translation is more complex in eukaryotes, involving several soluble initiation factors. Nevertheless, the overall process is similar except for two features. (1) The amino group of the methionine on the initiator tRNA is not formylated as in prokaryotes. (2) The initiation complex forms at the 5' terminus of the mRNA, not at the Shine-Dalgarno/AUG translation start site as in *E. coli*. In eukaryotes, the initiation complex scans the mRNA, starting at the 5' end, searching for an AUG translation-initiation codon. Thus, in eukaryotes, translation frequently begins at the AUG closest to the 5' terminus of the mRNA molecule, although the efficiency with which a given AUG is used to initiate translation depends on the contiguous nucleotide sequence. The optimal initiation sequence is 5'-GCC(A or G)CCAUGG-3'. The purine (A or G) three bases upstream from the AUG initiator codon and the G immediately following it are the most important—influencing initiation efficiency by tenfold or more. Changes of other bases in the sequence cause smaller decreases in initiation efficiency. These sequence requirements for optimal translation initiation in eukaryotes are called Kozak's rules, after Marilyn Kozak, who first proposed them.

Like prokaryotes, eukaryotes contain a special initiator tRNA, tRNA,^{Met} ("i" for initiator), but the amino group of the methionyl-tRNA,^{Met} is not formylated. The initiator methionyl-tRNA,^{Met} interacts with a soluble initiation factor and enters the *P* site directly during the initiation process, just as in *E. coli*.

In eukaryotes, a cap-binding protein (CBP) binds to the 7-methyl guanosine cap at the 5' terminus of the mRNA. Then, other initiation factors bind to the CBP-mRNA complex, followed by the small (40S) subunit of the ribosome. The entire initiation complex moves $5' \rightarrow 3'$ along the mRNA molecule, searching for an AUG codon.



■ FIGURE 12.17 Base-pairing between the Shine-Dalgarno sequence in a prokaryotic mRNA and a complementary sequence near the 3' terminus of the 16S rRNA is involved in the formation of the mRNA/30S ribosomal subunit initiation complex. When an AUG triplet is found, the initiation factors dissociate from the complex, and the large (60S) subunit binds to the methionyl-tRNA/mRNA/40S subunit complex, forming the complete (80S) ribosome. The 80S ribosome/mRNA/tRNA complex is ready to begin the second phase of translation, chain elongation. Try Solve It: Control of Translation in Eukaryotes to explore this process further.

Translation: Polypeptide Chain Elongation

The process of polypeptide chain **elongation** is basically the same in both prokaryotes and eukaryotes. The addition of each amino acid to the growing polypeptide occurs in three steps: (1) binding of an aminoacyl-tRNA to the A site of the ribosome, (2) transfer of the growing polypeptide chain from the tRNA in the P site to the tRNA in the A site by the formation of a new peptide bond, and (3) translocation of the ribosome along the mRNA to position the next codon in the A site (**TRNA**).

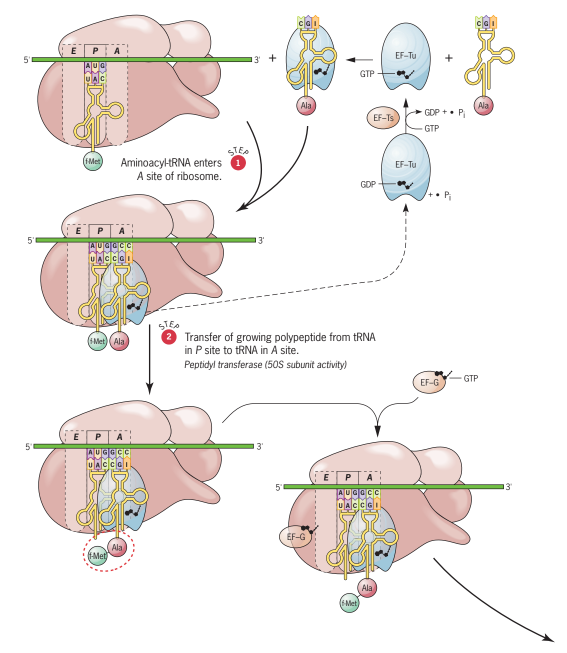


FIGURE 12.18 Polypeptide chain elongation in *E. coli*.

During step 3, the nascent polypeptide-tRNA and the uncharged tRNA are translocated from the *A* and *P* sites to the *P* and *E* sites, respectively. These three steps are repeated in a cyclic manner throughout the elongation process. The soluble factors involved in chain elongation in *E. coli* are described here. Similar factors participate in chain elongation in eukaryotes.

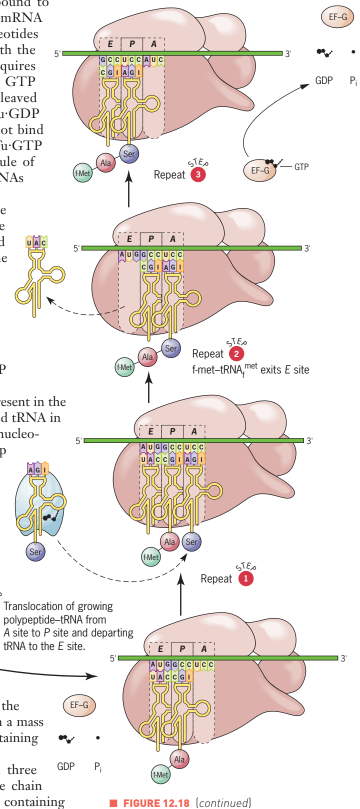
In the first step, an aminoacyl-tRNA enters and becomes bound to the *A* site of the ribosome, with the specificity provided by the mRNA codon in register with the *A* site (Figure 12.18). The three nucleotides in the anticodon of the incoming aminoacyl-tRNA must pair with the nucleotides of the mRNA codon present at the *A* site. This step requires **elongation factor Tu** carrying a molecule of GTP (**EF-Tu-GTP**). The GTP is required for aminoacyl-tRNA binding at the *A* site but is not cleaved until the peptide bond is formed. After the cleavage of GTP, EF-Tu-GDP is released from the ribosome. EF-Tu-GDP is inactive and will not bind to aminoacyl-tRNAs. EF-Tu-GDP is converted to the active EF-Tu-GTP form by **elongation factor Ts (EF-Ts)**, which hydrolyzes one molecule of GTP in the process. EF-Tu interacts with all of the aminoacyl-tRNAs except methionyl-tRNA.

The second step in chain elongation is the formation of a peptide bond between the amino group of the aminoacyl-tRNA in the A site and the carboxyl terminus of the growing polypeptide chain attached to the tRNA in the P site. This uncouples the growing chain from the tRNA in the P site and covalently joins the chain to the tRNA in the A site (Figure 12.18). This key reaction is catalyzed by **peptidyl transferase**, an enzymatic activity built into the 50S subunit of the ribosome. We should note that the peptidyl transferase activity resides in the 23S rRNA molecule rather than in a ribosomal protein, perhaps another relic of an early RNA-based world. Peptide bond formation requires the hydrolysis of the molecule of GTP brought to the ribosome by EF-Tu in step 1.

During the third step in chain elongation, the peptidyl-tRNA present in the A site of the ribosome is translocated to the P site, and the uncharged tRNA in the P site is translocated to the E site, as the ribosome moves three nucleotides toward the 3' end of the mRNA molecule. The translocation step requires GTP and **elongation factor G (EF-G)**. The ribosome undergoes changes in conformation during the translocation process, suggesting that it may shuttle along the mRNA molecule. The energy for the movement of the ribosome is provided by the hydrolysis of GTP. The translocation of the peptidyl-tRNA from the A site to the P site leaves the A site unoccupied and the ribosome ready to begin the next cycle of chain elongation.

The elongation of one eukaryotic polypeptide, the silk protein fibroin, can be visualized with the electron microscope by using techniques developed by Oscar Miller, Barbara Hamkalo, and colleagues. Most proteins fold up on the surface of the ribosome during their synthesis. However, fibroin remains extended from the surface of the ribosome under the conditions used by Miller and coworkers. As a result, nascent polypeptide chains of increasing length can be seen attached to the ribosomes as they are scanned from the 5' end of the mRNA to the 3' end (**■** Figure 12.19). Fibroin is a large protein with a mass of over 200,000 daltons; it is synthesized on large polyribosomes containing 50 to 80 ribosomes.

Polypeptide chain elongation proceeds rapidly. In *E. coli*, all three ^{GI} steps required to add one amino acid to the growing polypeptide chain occur in about 0.05 second. Thus, the synthesis of a polypeptide containing



Solve It!

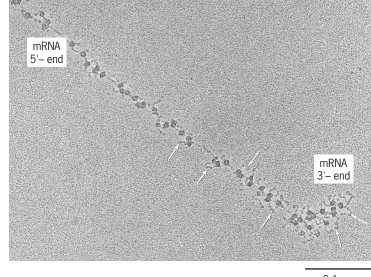
Control of Translation in Eukaryotes

The nucleotide sequence of the nontemplate strand of a portion of the human *HBB* (β -globin) gene specifying the 5'-terminus of the *HBB* mRNA is given below. Remember that the nontemplate strand will have the same sequence as the transcript of the gene, but with Ts in place of Us. Position 1 is the nucleotide corresponding to the 5'-end of the mRNA.

1 ACATTTGCT	T CTGACACAAC
TGTGTTCACT	AGCAACCTCA
AACAGACACC	ATGGTGCATC
TGACTCCTGA	GGAGAAGTCT
GCCGTTACTG	CCCTGTGGGG

Based on this sequence, the genetic code (see Table 12.1), and your knowledge of the initiation of translation in eukaryotes, predict the amino-terminal amino acid sequence of human β -globin.

To see the solution to this problem, visit the Student Companion site.



0.1 μm

FIGURE 12.19 Visualization of the elongation of fibroin polypeptides in the posterior silk gland of the silkworm *Bombyx mori*. The arrows point to growing fibroin polypeptides. Note their increasing length as one approaches the 3' end of the mRNA molecule.

300 amino acids takes only about 15 seconds. Given its complexity, the accuracy and efficiency of the translational apparatus are indeed amazing.

Translation: Polypeptide Chain Termination

Polypeptide chain elongation undergoes **termination** when any of three **chain-termination codons** (UAA, UAG, or UGA) enters the *A* site on the ribosome (**Telease factors (RFs)**. In three stop codons are recognized by soluble proteins called **release factors (RFs)**. In *E. coli*, there are two release factors, RF-1 and RF-2. RF-1 recognizes termination codons UAA and UAG; RF-2 recognizes UAA and UGA. In eukaryotes, a single release factor (**eRF**) recognizes all three termination codons. The presence of a release factor in the *A* site alters the activity of peptidyl transferase such that it adds a water molecule to the carboxyl terminus of the nascent polypeptide. This reaction releases the polypeptide from the tRNA molecule in the *P* site and triggers the translocation of the free tRNA to the *E* site. Termination is completed by the release of the mRNA molecule from the ribosome and the dissociation of the ribosome into its subunits. The ribosomal subunits are then ready to initiate another round of protein synthesis, as previously described.

KEY POINTS

- Genetic information carried in the sequences of nucleotides in mRNA molecules is translated into sequences of amino acids in polypeptide gene products by intricate macromolecular machines called ribosomes.
 - The translation process is complex, requiring the participation of many different RNA and protein molecules.
 - Transfer RNA molecules serve as adaptors, mediating the interaction between amino acids and codons in mRNA.
 - The process of translation involves the initiation, elongation, and termination of polypeptide chains and is governed by the specifications of the genetic code.

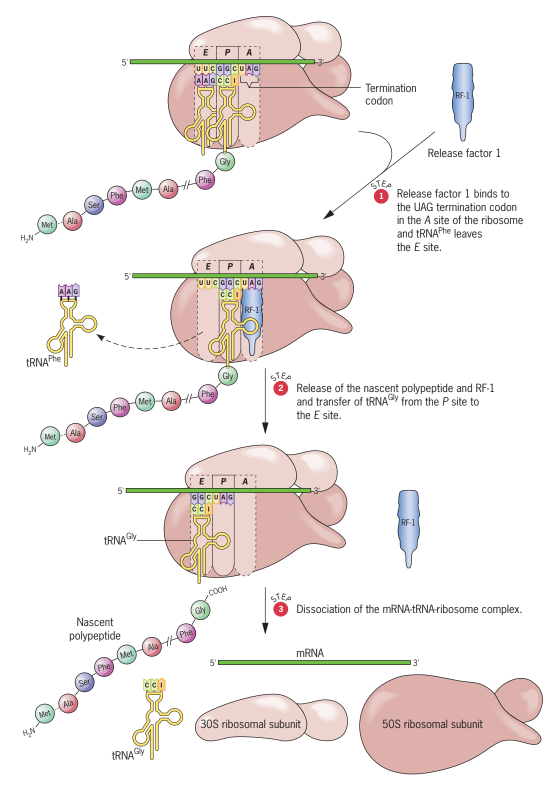


FIGURE 12.20 Polypeptide chain termination in *E. coli*. The formyl group of formylmethionine is removed during translation.

The Genetic Code

The genetic code is a nonoverlapping code, with each amino acid plus polypeptide initiation and termination specified by RNA codons composed of three nucleotides. As it became evident that genes controlled the structure of polypeptides, attention focused on how the sequence of the four different nucleotides in DNA could control the sequence of the 20 amino acids present in proteins. With the discovery of the mRNA

intermediary (Chapter 11), the question became one of how the sequence of the four bases present in mRNA molecules could specify the amino acid sequence of a polypeptide. What is the nature of the genetic code relating mRNA base sequences to amino acid sequences? Clearly, the symbols or letters used in the code must be the bases; but what comprises a codon, the unit or word specifying one amino acid or, actually, one aminoacyl-tRNA?

PROPERTIES OF THE GENETIC CODE: AN OVERVIEW

The main features of the genetic code were worked out during the 1960s. Cracking the code was one of the most exciting events in the history of science, with new information reported almost daily. By the mid-1960s, the genetic code was largely solved. Before focusing on specific features of the code, let us consider its most important properties.

- **1.** *The genetic code is composed of nucleotide triplets.* Three nucleotides in mRNA specify one amino acid in the polypeptide product; thus, each codon contains three nucleotides.
- **2.** *The genetic code is nonoverlapping*. Each nucleotide in mRNA belongs to just one codon except in rare cases where genes overlap and a nucleotide sequence is read in two different reading frames.
- **3.** *The genetic code is comma-free*. There are no commas or other forms of punctuation within the coding regions of mRNA molecules. During translation, the codons are read consecutively.
- **4.** *The genetic code is degenerate.* All but two of the amino acids are specified by more than one codon.
- **5.** *The genetic code is ordered.* Multiple codons for a given amino acid and codons for amino acids with similar chemical properties are closely related, usually differing by a single nucleotide.
- **6.** *The genetic code contains start and stop codons.* Specific codons are used to initiate and to terminate polypeptide chains.
- **7.** *The genetic code is nearly universal.* With minor exceptions, the codons have the same meaning in all living organisms, from viruses to humans.

THREE NUCLEOTIDES PER CODON

Twenty different amino acids are incorporated into polypeptides during translation. Thus, at least 20 different codons must be formed with the four bases available in mRNA. Two bases per codon would result in only 4² or 16 possible codons—clearly not enough. Three bases per codon yields 4³ or 64 possible codons—an apparent excess.

In 1961, Francis Crick and colleagues published the first strong evidence in support of a *triplet code* (three nucleotides per codon). Crick and coworkers carried out a genetic analysis of mutations induced at the *r*II locus of bacteriophage T4 by the chemical proflavin. Proflavin is a mutagenic agent that causes single base-pair additions and deletions (Chapter 13). Phage T4 *r*II mutants are unable to grow in cells of *E. coli* strain K12, but grow like wild-type phage in cells of *E. coli* strain B. Wild-type T4 grows equally well on either strain. Crick and coworkers isolated proflavin-induced

revertants of a proflavin-induced *r*II mutation. These revertants were shown to result from the occurrence of additional mutations at nearby sites rather than reversion of the original mutation. Second-site mutations that restore the wild-type phenotype in a mutant organism are called **suppressor mutations** because they cancel, or suppress, the effect(s) of the original mutation.

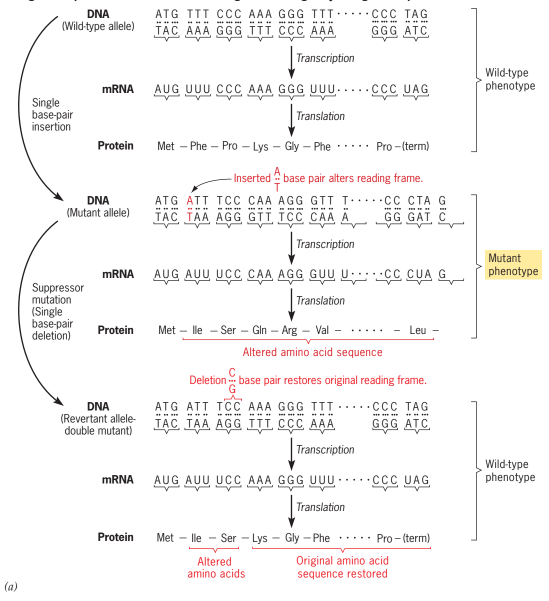
Crick and colleagues reasoned that if the original mutation was a single base-pair addition or deletion, then the suppressor mutations must be single base-pair deletions or additions, respectively, occurring at a site or sites near the original mutation. If sequential nucleotide triplets in an mRNA specify amino acids, then every nucleotide sequence can be recognized or read during translation in three different ways. For example, the sequence AAAGGGCCCTTT can be read (1) AAA, GGG, CCC, TTT, (2) A, AAG, GGC, CCT, TT, or (3) AA, AGG, GCC, CTT, T. The **reading frame** of an mRNA is the series of nucleotide triplets that are read (positioned in the A site of the ribosome) during translation. A single base-pair addition or deletion will alter the reading frame of the gene and mRNA for that portion of the gene distal to the mutation. This effect is illustrated in **Figure 12.21a**. The suppressor mutations were then isolated as single mutants by screening progeny of backcrosses to wild-type. Like the original mutation, the suppressor mutations were found to produce *r*II mutant phenotypes. Crick and colleagues next isolated proflavin-induced suppressor mutations, and so on.

Crick and colleagues then classified all the isolated mutations into two groups, plus (+) and minus (-) (for additions and deletions, although they had no idea which group was which), based on the reasoning that a (+) mutation would suppress a (-) mutation but not another (+) mutation, and vice versa (Figure 12.21). Then, Crick and coworkers constructed recombinants that carried various combinations of the (+) and the (-) mutations. Like the single mutants, recombinants with two (+) mutations or two (-) mutations always had the mutant phenotype. The critical result was that recombinants with three (+) mutations (\blacksquare Figure 12.21b) or three (-) mutations often exhibited the wild-type phenotype. This indicated that the addition of three base pairs or the deletion of three base pairs left the distal portion of the gene with the wild-type reading frame. This result would be expected only if each codon contained three nucleotides.

Evidence from *in vitro* translation studies soon supported the results of Crick and colleagues and firmly established the triplet nature of the code. Some of the more important results follow: (1) Trinucleotides were sufficient to stimulate specific binding of aminoacyl-tRNAs to ribosomes. For example, 5'-UUU-3' stimulated the binding of phenylalanyl-tRNA^{Phe} to ribosomes. (2) Chemically synthesized mRNA molecules that contained repeating dinucleotide sequences directed the synthesis of copolymers (large chainlike molecules composed of two different subunits) with alternating amino acid sequences. For example, when poly(UG), was used as an artificial mRNA in an *in vitro* translation system, the repeating copolymer (cys-val), was synthesized. (The subscripts n and m refer to the number of nucleotides and amino acids in the respective polymers.) (3) In contrast, mRNAs with repeating trinucleotide sequences directed the synthesis of a mixture of three homopolymers (initiation being at random on such mRNAs in the *in vitro* systems). For example, poly(UUG), directed the synthesis of a mixture of polyleucine, polycysteine, and polyvaline. These results are consistent only with a triplet code, with its three different reading frames. When poly(UUG), is translated in reading frame 1, UUG, UUG, polyleucine is produced, whereas translation in reading frame 2, UGU, UGU, yields polycysteine, and translation in reading frame 3, GUU, GUU, produces polyvaline. Ultimately, the triplet nature of the code was definitively established by comparing the nucleotide sequences of genes and mRNAs with the amino acid sequences of their polypeptide products.

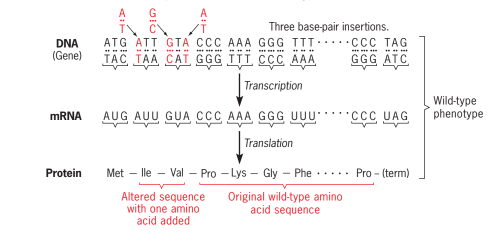
DECIPHERING THE CODE

The cracking of the genetic code in the 1960s took several years and involved intense competition between many different research laboratories. New information



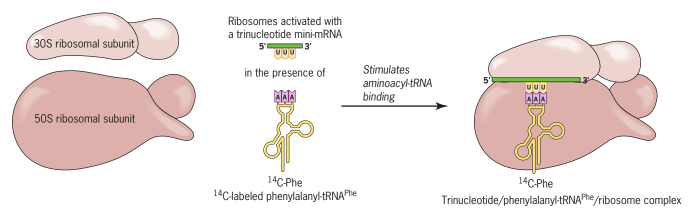
A single base-pair deletion restores the reading frame changed by a single base-pair addition.

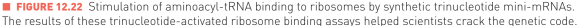
Recombinant containing three single base-pair additions has the wild-type reading frame.



(b)

FIGURE 12.21 Early evidence that the genetic code is a triplet code. See the text for details.





accumulated rapidly but sometimes was inconsistent with earlier data. Indeed, cracking the code proved to be a major challenge.

Deciphering the genetic code required scientists to obtain answers to several questions. (1) Which codons specify each of the 20 amino acids? (2) How many of the 64 possible triplet codons are utilized? (3) How is the code punctuated? (4) Do the codons have the same meaning in viruses, bacteria, plants, and animals? The answers to these questions were obtained primarily from the results of two types of experiments, both of which were performed with cell-free systems. The first type of experiment involved translating artificial mRNA molecules *in vitro* and determining which of the 20 amino acids were incorporated into proteins. In the second type of experiment, ribosomes were activated with mini-mRNAs just three nucleotides long. Then, researchers determined which aminoacyl-tRNAs were stimulated to bind to ribosomes activated with each of the trinucleotide messages (**■ Figure 12.22**).

The decade of the 1960s—the era of the cracking of the genetic code—was one of the most exciting times in the history of biology. Deciphering the genetic code was a difficult and laborious task, and progress came in a series of breakthroughs. We discuss these important developments in A Milestone in Genetics: Cracking the Genetic Code on the Student Companion web site. By combining the results of *in vitro* translation experiments performed with synthetic mRNAs and trinucleotide binding assays, Marshall Nirenberg, Severo Ochoa, H. Ghobind Khorana, Philip Leder, and their colleagues worked out the meaning of all 64 triplet codons (Table 12.1). Nirenberg and Khorana shared the 1968 Nobel Prize in Physiology or Medicine for their work on the code with Robert Holley, who determined the complete nucleotide sequence of the yeast alanine tRNA. Ochoa had already received the 1959 Nobel Prize for his discovery of RNA polymerase.

INITIATION AND TERMINATION CODONS

The genetic code also provides for punctuation of genetic information at the level of translation. In both prokaryotes and eukaryotes, the codon AUG is used to initiate polypeptide chains (Table 12.1). In rare instances, GUG is used as an initiation codon. In both cases, the initiation codon is recognized by an initiator tRNA, $tRNA_f^{Met}$ in prokaryotes and $tRNA_i^{Met}$ in eukaryotes. In prokaryotes, an AUG codon must follow an appropriate nucleotide sequence, the Shine-Delgarno sequence, in the 5' nontranslated segment of the mRNA molecule in order to serve as translation initiation codon. In eukaryotes, the codon must be the first AUG encountered by the ribosome as it scans from the 5' end of the mRNA molecule. At internal positions, AUG is recognized by tRNA^{Met}, and GUG is recognized by a valine tRNA.

TABLE 12.1								
The Genetic Code ^a								
	Second letter							
		U	С	А	G			
First (5 ') letter	U	UUU > Phe (F)	Γυου	UAU > Tyr (Y)	UGU > Cys (C)	U		
			UCC VCA UCG			С		
		UUA > Leu (L)		UAA Stop (terminator)	UGA Stop (terminator)	A		
				UAG Stop (terminator)	UGG Trp (W)	G		
	С	CUU	CCU	CAU > His (H)	CGU	U		
		CUC > Leu (L)			CGC	С		
		CUA	> Pro (P)	CAA S GIn (O)	> Arg (R) CGA	А	'n	
		CUG	CCG		CGG_	G) let	
	A	AUU	ACU	AAU > Asn (N)	AGU > Ser (S)	U	Third (3') letter	
ιĒ		AUC > lle (l)	ACC	AAC	AGC	С	Ē	
		AUA	ACA ACG	AAA > Lys (K)	AGA Arg (R)	А		
		AUG Met (M) (initiator)		AAG	AGG	G		
	G	GUU	GCU	GAU > Asp (D)	GGU	U		
		GUC > Val (V)	GCC > Ala (A)	GAC	GGC > Gly (G)	С	= Polypeptide chain	
		GUA	GCA	GAA Glu (E)	GGA	А	initiation codon	
		gug_	GCG	GAG	GGG_	G	= Polypeptide chain termination codon	
			ce or codon refers to he indicated amino aci					
specifies the incorporation of the indicated amino acid or polypeptide chain termination. The one-letter symbols for the amino acids are given in parentheses after the standard three-letter abbreviations.								

Three codons—UAG, UAA, and UGA—specify polypeptide chain termination (Table 12.1). These codons are recognized by protein release factors, rather than by tRNAs. Prokaryotes contain two release factors, RF-1 and RF-2. RF-1 terminates polypeptides in response to codons UAA and UAG, whereas RF-2 causes termination at UAA and UGA codons. Eukaryotes contain a single release factor that recognizes all three termination codons.

A DEGENERATE AND ORDERED CODE

All the amino acids except methionine and tryptophan are specified by more than one codon (Table 12.1). Three amino acids—leucine, serine, and arginine—are each specified by six different codons. Isoleucine has three codons. The other amino acids each have either two or four codons. The occurrence of more than one codon per amino acid is called **degeneracy** (although the usual connotations of the term are hardly appropriate). The degeneracy in the genetic code is not at random; instead, it is highly ordered. In most cases, the multiple codons specifying a given amino acid differ by only one base, the third or 3' base of the codon. The degeneracy is primarily of two types. (1) Partial degeneracy occurs when the third base may be either of the two pyrimidines (U or C) or, alternatively, either of the two purines (A or G). With partial degeneracy, changing the third base from a purine to a pyrimidine, or vice versa, will change the amino acid specified by the codon. (2) In the case of complete degeneracy, any of the four bases may be present at the third position in the codon, and the codon will still specify the same amino acid. For example, valine is encoded by GUU, GUC, GUA, and GUG (Table 12.1).

Scientists have speculated that the **order** in the genetic code has evolved as a way of minimizing mutational lethality. Many base substitutions at the third position of codons do not change the amino acid specified by the codon. Moreover, amino acids

PROBLEM-SOLVING SKILLS

Predicting Amino Acid Substitutions Induced by Mutagens

THE PROBLEM

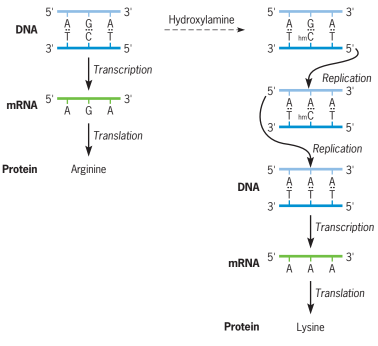
The chemical hydroxylamine (NH₂OH) transfers a hydroxyl (-OH) group to cytosine producing hydroxymethylcytosine (hmC), which, unlike cytosine, pairs with adenine. Therefore, hydroxylamine induces G:C to A:T base-pair substitutions in DNA. If you treat the double-stranded DNA of a virus such as phage T4 with hydroxylamine, what amino acid substitutions will be induced in the proteins encoded by the virus?

FACTS AND CONCEPTS

- **1.** The nature of the genetic code—the meaning of the 64 triplet nucleotide sequences in mRNA—is shown in Table 12.1.
- **2.** Complete degeneracy occurs when the first two nucleotides in an mRNA codon are sufficient to determine the amino acid in the polypeptide specified by the mRNA.
- **3.** Partial degeneracy occurs when the same amino acid is specified if the base in the 3' nucleotide of a codon is either of the two pyrimidines or either of the two purines.
- **4.** Hydroxylamine will only alter codons specified by DNA basepair triplets that contain G:C base pairs.
- 5. If the G:C base pair occupies the third (3') position of the triplet, hydroxylamine will induce amino acid substitutions only in cases where the genetic code is NOT degenerate, that is, where the base present as the 3' nucleotide of the codon determines its meaning. Only two codons are not degenerate at the 3' position; they are 5'-AUG-3' (methionine) and 5'-UGG-3' (tryptophan).
- 6. For codons with complete or partial degeneracy at the 3' position, hydroxylamine will not induce amino acid substitutions by modifying the base pair specifying the 3' base in the codon. It will induce G:C → A:T and C:G → T:A substitutions (where the first base given is in the template strand). However, given the partial or complete degeneracy, the resulting codons will still specify the same amino acids. An AAG lysine codon, for example, could be changed to an AAA lysine codon, or a UUC phenylalanine codon could be changed to a UUU phenylalanine codon. However, no amino acid substitution will occur in either case.

ANALYSIS AND SOLUTION

The answer to the question of which amino acid substitutions will be induced by hydroxylamine requires a careful analysis of the nature of the genetic code (Table 12.1). Potential targets of hydroxylamine mutagenesis are DNA triplets specifying mRNA codons containing C's and G's at the first (5') and second positions in the codons and triplets specifying nondegenerate codons with G's or C's at the third (3') position. Indeed, there are more potential targets in genomes than nontargets; 51 of the 64 DNA triplets contain G:C or C:G base pairs. Consider as an example the arginine codon 5'-AGA-3'; it will be transcribed from a DNA template strand with the sequence 3'-TCT-5' (reversing the polarity to keep the bases in the same order). The C in this sequence can be hydroxymethylated, producing hmC, which will pair with adenine. After two semiconservative replications, the DNA template strand will contain the sequence 3'-TTT-5' at this site, and transcription of this sequence will yield a 5'-AAA-3' mRNA codon. Translation of the mRNA will result in the insertion of lysine in the resulting polypeptide because AAA is a lysine codon. Thus, one example of the effects of hydroxylamine will be the replacement of arginine residues with lysines. This process is diagrammed below.



The only amino acids specified by codons with no targets of hydroxylamine-induced amino acid substitutions are phenylalanine (UUU & UUC), isoleucine (AUU, AUC, & AUA), tyrosine (UAU & UAC), asparagine (AAU & AAC), and lysine (AAA & AAG). The other amino acids are all specified by DNA base-pair triplets that contain one or more G:C's, with the C's being potential targets of hydroxylamine mutagenesis. For further discussion visit the Student Companion site.

with similar chemical properties (such as leucine, isoleucine, and valine) have codons that differ from each other by only one base. Thus, many single base-pair substitutions will result in the substitution of one amino acid for another amino acid with very similar chemical properties (for example, valine for isoleucine). In most cases, conservative substitutions of this type will yield active gene products, which minimizes the effects of mutations. Try Problem-Solving Skills: Predicting Amino Acid Substitutions Induced by Mutagens to test your understanding of the genetic code.

A NEARLY UNIVERSAL CODE

Vast quantities of information are now available from *in vitro* studies, from amino acid replacements due to mutations, and from correlated nucleic acid and polypeptide sequencing, which allow a comparison of the meaning of the 64 codons in different species. These data all indicate that the genetic code is nearly **universal**; that is, the codons have the same meaning, with minor exceptions, in all species.

The most important exceptions to the universality of the code occur in mitochondria of mammals, yeast, and several other species. Mitochondria have their own chromosomes and protein-synthesizing machinery (Chapter 15). Although the mitochondrial and cytoplasmic systems are similar, there are some differences. In the mitochondria of humans and other mammals, (1) UGA specifies tryptophan rather than chain termination, (2) AUA is a methionine codon, not an isoleucine codon, and (3) AGA and AGG are chain-termination codons rather than arginine codons. The other 60 codons have the same meaning in mammalian mitochondria as in nuclear mRNAs (Table 12.1). There are also rare differences in codon meaning in the mitochondria of other species and in nuclear transcripts of some protozoa. However, since these exceptions are rare, the genetic code should be considered nearly universal.

KEY POINTS • 1

• Each of the 20 amino acids in proteins is specified by one or more nucleotide triplets in mRNA.

- Of the 64 possible triplets, given the four bases in mRNA, 61 specify amino acids and 3 signal chain termination.
- The code is nonoverlapping, with each nucleotide part of a single codon, degenerate, with most amino acids specified by two or four codons, and ordered, with similar amino acids specified by related codons.
- The genetic code is nearly universal; with minor exceptions, the 64 triplets have the same meaning in all organisms.

Codon-tRNA Interactions

Codons in mRNA molecules are recognized by aminoacyl-tRNAs during translation.

The translation of a sequence of nucleotides in mRNA into the correct sequence of amino acids in the polypeptide product requires the accurate recognition of codons by aminoacyl-tRNAs. Because of the degeneracy of the genetic code, either several different tRNAs

must recognize the different codons specifying a given amino acid or the anticodon of a given tRNA must be able to base-pair with several different codons. Actually, both of these phenomena occur. Several tRNAs exist for certain amino acids, and some tRNAs recognize more than one codon.

RECOGNITION OF CODONS BY tRNAs: THE WOBBLE HYPOTHESIS

The hydrogen bonding between the bases in the anticodons of tRNAs and the codons of mRNAs follows strict base-pairing rules only for the first two bases of the codon. The base-pairing involving the third base of the codon is less stringent, allowing what Crick has called *wobble* at this site. On the basis of molecular distances and steric (three-dimensional structure) considerations, Crick proposed that wobble would allow

several types, but not all types, of base-pairing at the third codon base during the codon-anticodon interaction. His proposal has since been strongly supported by experimental data. Table 12.2 shows the base-pairing predicted by Crick's wobble hypothesis.

The **wobble hypothesis** predicted the existence of at least two tRNAs for each amino acid with codons that exhibit complete degeneracy, and this has proven to be true. The wobble hypothesis also predicted the occurrence of three tRNAs for the six serine codons. Three serine tRNAs have been characterized: (1) tRNA^{Ser1} (anti-codon AGG) binds to codons UCU and UCC, (2) tRNA^{Ser2} (anticodon AGU) binds to codons UCG, and (3) tRNA^{Ser3} (anticodon UCG) binds to codons AGU and AGC. These specificities were verified by the trinucleotide-stimulated binding of purified aminoacyl-tRNAs to ribosomes *in vitro*.

Finally, several tRNAs contain the base inosine, which is made from the purine hypoxanthine. Inosine is produced by a posttranscriptional modification of adenosine. Crick's wobble hypothesis predicted that when inosine is present at the 5' end of an anticodon (the wobble position), it would base-pair with uracil, cytosine, or adenine in the codon. In fact, purified alanyl-tRNA containing inosine (I) at the 5' position of the anticodon (see Figure 12.12) binds to ribosomes activated with GCU, GCC, or GCA trinucleotides (**Figure 12.23**). The same result has been obtained with other purified tRNAs with inosine at the 5' position of the anticodon. Thus, Crick's wobble hypothesis nicely explains the relationships between tRNAs and codons given the degenerate, but ordered, genetic code.

SUPPRESSOR MUTATIONS THAT PRODUCE tRNAs WITH ALTERED CODON RECOGNITION

Even if we exclude the mitochondria, the genetic code is not absolutely universal. Minor variations in codon recognition and translation are well documented. In *E. coli* and yeast, for example, some mutations in tRNA genes alter the anticodons and thus the codons recognized by the mutant tRNAs. These mutations were initially detected as *suppressor mutations*, nucleotide substitutions that suppressed the effects of other mutations. The suppressor mutations were subsequently shown to occur in tRNA genes. Many of these suppressor mutations changed the anticodons of the altered tRNAs.

The best-known examples of suppressor mutations that alter tRNA specificity are

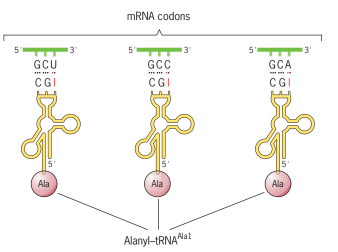
those that suppress UAG chain-termination mutations within the coding sequences of genes. Such mutations, called *amber* mutations (after one of the researchers who discovered them), result in the synthesis of truncated polypeptides. Mutations that produce chain-termination triplets within genes have come to be known as **nonsense mutations**, in contrast to **missense mutations**, which change a triplet so that it specifies a different amino acid. A gene that contains a missense mutation encodes a complete polypeptide, but with an amino acid substitution in the polypeptide gene product. A nonsense mutation results in a truncated polypeptide, with the length of the chain depending on the position of the mutation within the gene. Nonsense mutations frequently result from single base-pair substitutions, as illustrated in **Figure 12.24a**. The polypeptide fragments produced from genes containing nonsense mutations (**Figure 12.24b**) often are completely nonfunctional. See Solve It: Effects of Base-Pair Substitutions in the Coding Region of the *HBB* Gene.

Suppression of nonsense mutations has been shown to result from mutations in tRNA genes that cause the mutant tRNAs to recognize the termination (UAG, UAA, or UGA) codons, albeit with varying efficiencies. These mutant tRNAs are referred to as **suppressor tRNAs**. When the *amber* (UAG) suppressor tRNA produced by the *amber su3* mutation in *E. coli* was sequenced, it was found to have an altered anticodon. This particular *amber* suppressor mutation occurs in the tRNA^{Tyr2} gene (one of two

TABLE 12.2

Base-Pairing between the 5' Base of the Anticodons of tRNAs and the 3' Base of Codons of mRNAs According to the Wobble Hypothesis

Base in Anticodon	Base in Codon
G	U or C
A	G U
U	A or G A. U. or C
I	7, 0, 01 0



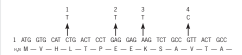
■ FIGURE 12.23 Base-pairing between the anticodon of alanyltRNA^{Ala1} and mRNA codons GCU, GCC, and GCA according to Crick's wobble hypothesis. Trinucleotide-activated ribosome binding assays have shown that alanyl-tRNA^{Ala1} does indeed base-pair with all three codons.

■ FIGURE 12.24 (a) The formation of an amber (UAG) chain-termination mutation. (b) Its effect on the polypeptide gene product in the absence of a suppressor tRNA, and (c) in the presence of a suppressor tRNA. The amber mutation shown here changes a CAG glutamine (Gln) codon to a UAG chain-termination codon. The polypeptide containing the tyrosine inserted by the suppressor tRNA may or may not be functional; however, suppression of the mutant phenotype will occur only when the polypeptide is functional.

Solve It!

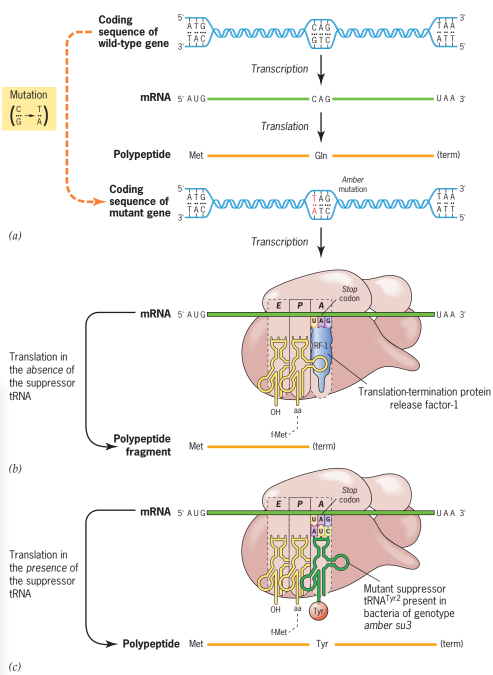
Effects of Base-Pair Substitutions in the Coding Region of the *HBB* Gene

The first 42 nucleotides, shown as triplets corresponding to mRNA codons, in the nontemplate strand of the coding region of the human HBB (β -globin) gene are given below. Recall that the nontemplate strand has the same sequence as the mRNA, but with T's in place of U's. The first (amino-terminal) 14 amino acids of the nascent human β -globin are also given using the single letter code (see Table 12.1). The methionine is subsequently removed to yield mature B-globin. Consider the potential phenotypic effects of the four single nucleotide substitutions numbered 1 through 4 below when present in homozygotes.



Which substitution would you expect to have the largest effect on phenotype? The second largest effect? No effect? No, or a very small, effect?

• To see the solution to this problem, visit the Student Companion site.



tyrosine tRNA genes in *E. coli*). The anticodon of the wild-type (nonsuppressor) tRNA^{Tyr2} was shown to be 5'-G'UA-3' (where G' is a derivative of guanine). The anticodon of the mutant (suppressor) tRNA^{Tyr2} is 5'-CUA-3'. Because of the single-base substitution, the anticodon of the suppressor tRNA^{Tyr2} base-pairs with the 5'-UAG-3' *amber* codon (recall that base-pairing always involves strands of opposite polarity); that is,

mRNA: 5'-UAG-3' (codon) tRNA: 3'-AUC-5' (anticodon)

Thus, suppressor tRNAs allow complete polypeptides to be synthesized from mRNAs containing termination codons within genes (**■** Figure 12.24c). Such polypeptides will be functional if the amino acid inserted by the suppressor tRNA does not significantly alter the protein's chemical properties. In addition, see On the Cutting Edge: Seleno-cysteine, the 21st Amino Acid.

- The wobble hypothesis explains how a single tRNA can respond to two or more codons.
- Some suppressor mutations alter the anticodons of tRNAs so that the mutant tRNAs recognize chain-termination codons and insert amino acids in response to their presence in mRNA molecules.

ON THE CUTTING EDGE

SELENOCYSTEINE, THE 21ST AMINO ACID

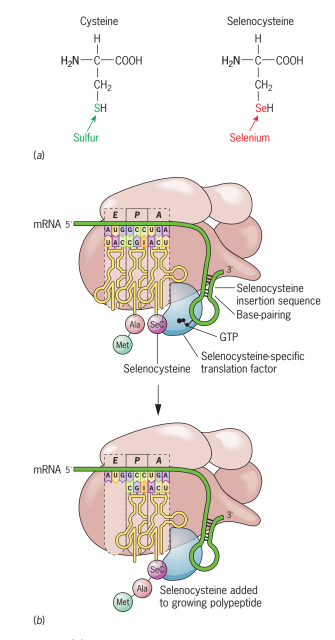
he structures of the 20 amino acids that are found in most proteins are shown in Figure 12.1, and the codons that specify each of these amino acids and the initiation and termination of polypeptide chains are shown in Table 12.1. However, in a few proteins, there is another amino acid—**selenocysteine**—that is specified by the genetic code at the time of translation. Selenocysteine contains the essential trace element selenium (atomic number 34) in place of the sulfur group in cysteine (**Figure 1a**). When present in proteins—called **selenoproteins**—the reactive selenium is usually present at the active site and participates in oxidation/ reduction (hydrogen removal/addition) reactions. Selenoproteins play important metabolic roles in all living organisms—prokaryotes, eukaryotes, and archaea.

Selenocysteine is incorporated into polypeptides during translation in response to the codon UGA, which normally functions as a chain-termination signal. The mRNAs encoding selenoproteins contain special <u>selenocysteine</u> insertion <u>sequences</u> (SECIS elements) that interact with specific translation factors leading to the incorporation of selenocysteine into polypeptides in response to UGA codons (**Figure 1b**). The structures and locations of SECIS elements vary among procaryotes, archaea, and eukaryotes. However, in all cases, the SECIS elements form hairpin-like structures similar to those in tRNAs by intrastrand hydrogen bonding. In eukaryotes, these elements are located in the 3'-untranslated regions of mRNAs.

Selenocysteine has its own tRNA with a 5'-UCA-3' anticodon and a unique hairpin domain. This tRNA is activated by the addition of serine, which is then converted to selenocysteine. During the translation of mRNAs encoding selenoproteins, the selenocysteyl-tRNAs respond to UGA codons with the aid of a selenocysteine-specific translation factor (Figure 1*b*). This selenocysteine-specific translation factor replaces elongation factor Tu during the entry of selenocysteine-tRNA into the A site on the ribosome. In the absence of selenium, translation of mRNAs encoding selenoproteins results in the synthesis of truncated polypeptides, with translation being terminated at the UGA selenocysteine codons. Thus, UGA codons in mRNAs lacking SECIS elements specify polypeptide chain termination, whereas UGA codons in mRNAs with downstream SECIS hairpin structures specify selenocysteine.

Are there other modified amino acids that are specified by codons during translation? So far, there is one other documented example, pyrrolysine—lysine with a pyrroline ring on the end of the side chain. Pyrrolysine is incorporated into polypeptides in some archaea and one bacterium, but not eukaryotes, in response to the codon UAG, which normally signals chain termination. The mechanism(s) by which UAG codons specify pyrrolysine incorporation rather than chain termination are still under investigation.

KEY POINTS



■ FIGURE 1 (a) Comparison of the structures of cysteine and selenocysteine. (b) The incorporation of selenocysteine into a growing polypeptide in response to the codon UGA when a selenocysteine insertion sequence is present in the mRNA being translated.

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. The human β -globin polypeptide is 146 amino acids long. How long is the coding portion of the human β -globin mRNA?
- **Answer:** Each amino acid is specified by a codon containing three nucleotides. Therefore, the 146 amino acids in β -globin will be specified by 438 (146 × 3) nucleotides. However, a termination codon must be present at the end of the coding sequence, bringing the length to 438 + 3 = 441 nucleotides. In the case of β -globin and many other proteins, the aminoterminal methionine (specified by the initiation codon AUG) is removed from the β -globin during synthesis. Adding the initiation codon increases the coding sequence of the β -globin mRNA to 444 nucleotides (441 + 3).
- 2. If the coding segment of an mRNA with the sequence 5'-AUGUUUCCCAAAGGG-3' is translated, what amino acid sequence will be produced?
- **Answer:** (Amino-terminus)-methionine-phenylalanine-prolinelysine-glycine-(carboxyl-terminus). The amino acid sequence is deduced using the genetic code shown in **Table 12.1**. AUG is the methionine initiation codon followed by the phenylalanine codon UUU, the proline codon CCC, the lysine codon AAA, and the glycine codon GGG.
- **3.** If a coding segment of the template strand of a gene (DNA) has the sequence 3'-TACAAAGGGTTTCCC-5', what amino acid sequence will be produced if it is transcribed and translated?
- Answer: The mRNA sequence produced by transcription of this segment of the gene will be 5'-AUGUUUCCCAAAGGG-3'. Note that this mRNA has the same nucleotide sequence as the one discussed in Exercise 2. Thus, it will produce the same peptide when translated: NH,-Met-Phe-Pro-Lys-Gly-COOH.
- 4. What sequence of nucleotide pairs in a gene in *Drosophila* will encode the amino acid sequence methionine-

tryptophan (reading from the amino terminus to the carboxyl terminus)?

Answer: The codons for methionine and tryptophan are AUG and UGG, respectively. Thus, the nucleotide sequence in the mRNA specifying the dipeptide sequence methionine-tryptophan must be 5'-AUGUGG-3'. The template DNA strand must be complementary and antiparallel to the mRNA sequence (3'-TACACC-5'), and the other strand of DNA must be complementary to the template strand. Therefore, the sequence of base pairs in the gene must be:

5'-ATGTGG-3' 3'-TACACC-5'

- 5. A wild-type gene contains the trinucleotide-pair sequence:
 - 5'-GAG-3' 3'-CTC-5'

This triplet specifies the amino acid glutamic acid. If the second base pair in this gene segment were to change from A:T to T:A, yielding the following DNA sequence:

5'-GTG-3' 3'-CAC-5'

would it still encode glutamic acid?

Answer: No, it would now specify the amino acid valine. The codon for glutamic acid is 5'-GAG-3', which tells us that the bottom strand of DNA is the template strand. Transcription of the wild-type gene yields the mRNA sequence 5'-GAG-3', which is a glutamic acid codon. Transcription of the altered gene produces the mRNA sequence 5'-GUG-3', which is a valine codon. Indeed, this is exactly the same nucleotide-pair change that gave rise to the altered hemoglobin in Herrick's sickle-cell anemia patient, discussed at the beginning of this chapter. See Figure 1.9 for further details.

Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. The average mass of the 20 common amino acids is about 137 daltons. Estimate the approximate length of an mRNA molecule that encodes a polypeptide with a mass of 65,760 daltons. Assume that the polypeptide contains equal amounts of all 20 amino acids.
- **Answer:** Based on this assumption, the polypeptide would contain about 480 amino acids (65,760 daltons/137 daltons per amino acid). Since each codon contains three nucleotides, the coding region of the mRNA would have to be 1440 nucleotides long (480 amino acids × 3 nucleotides per amino acid).
- 2. The antibiotic streptomycin kills sensitive *E. coli* by inhibiting the binding of $tRNA_{f}^{Met}$ to the *P* site of the ribosome and by causing misreading of codons in mRNA. In sensitive bacteria, streptomycin is bound by protein S12 in the 30S subunit of the ribosome. Resistance to streptomycin can result from a mutation in the gene-encoding protein S12 so that the altered protein will no longer bind the antibiotic. In 1964, Luigi Gorini and Eva Kataja isolated mutants of *E. coli* that grew on minimal medium supplemented with either the amino acid arginine or streptomycin. That is, in

the absence of streptomycin, the mutants behaved like typical arginine-requiring bacteria. However, in the absence of arginine, they were streptomycin-dependent conditional-lethal mutants. That is, they grew in the presence of streptomycin but not in the absence of streptomycin. Explain the results obtained by Gorini and Kataja.

Answer: The streptomycin-dependent conditional-lethal mutants isolated by Gorini and Kataja contained missense mutations in genes encoding arginine biosynthetic enzymes. If arginine was present in the medium, these enzymes were unessential. However, these enzymes were required for growth in the

absence of arginine (one of the 20 amino acids required for protein synthesis).

Streptomycin causes misreading of mRNA codons in bacteria. This misreading allowed the codons that contained the missense mutations to be translated ambiguously—with the wrong amino acids incorporated—when the antibiotic was present. When streptomycin was present in the mutant bacteria, an amino acid occasionally would be inserted (at the site of the mutation) that resulted in an active enzyme, which, in turn, allowed the cells to grow, albeit slowly. In the absence of streptomycin, no misreading occurred, and all of the mutant polypeptides were inactive.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **12.1** In a general way, describe the molecular organization of proteins and distinguish proteins from DNA, chemically and functionally. Why is the synthesis of proteins of particular interest to geneticists?
- 12.2 At what locations in the cell does protein synthesis occur?
- **12.3** Is the number of potential alleles of a gene directly related to the number of nucleotide pairs in the gene? Is such a relationship more likely to occur in prokaryotes or in eukaryotes? Why?
- 12.4 Why was it necessary to modify Beadle and Tatum's one gene-one enzyme concept of the gene to one gene-one polypeptide?
- **12.5** (a) Why is the genetic code a triplet code instead of a singlet or doublet code? (b) How many different amino acids are specified by the genetic code? (c) How many different amino acid sequences are possible in a polypeptide 146 amino acids long?
- **12.6** What types of experimental evidence were used to decipher the genetic code?
- 12.7 In what sense and to what extent is the genetic code (a) degenerate, (b) ordered, and (c) universal?
- 12.8 The thymine analog 5-bromouracil is a chemical mutagen that induces single base-pair substitutions in DNA called transitions (substitutions of one purine for another purine and one pyrimidine for another pyrimidine). Using the known nature of the genetic code (Table 12.1), which of the following amino acid substitutions should you expect to be induced by 5-bromouracil with the highest frequency:
 - (a) Met \rightarrow Val;
 - (b) Met \rightarrow Leu;
 - (c) Lys \rightarrow Thr;
 - (d) Lys \rightarrow Gln;
 - (e) $Pro \rightarrow Arg; or$
 - (f) $Pro \rightarrow Gln$? Why?

- 12.9 Using the information given in Problem 12.8, would you expect 5-bromouracil to induce a higher frequency of His → Arg or His → Pro substitutions? Why?
- **12.10** What is the minimum number of tRNAs required to recognize the six codons specifying the amino acid leucine?
- **12.11** Characterize ribosomes in general as to size, location, function, and macromolecular composition.
- 12.12 (a) Where in the cells of higher organisms do ribosomes originate? (b) Where in the cells are ribosomes most active in protein synthesis?
- **12.13** Identify three different types of RNA that are involved in translation and list the characteristics and functions of each.
- 12.14 (a) How is messenger RNA related to polysome formation? (b) How does rRNA differ from mRNA and tRNA in specificity? (c) How does the tRNA molecule differ from that of DNA and mRNA in size and helical arrangement?
- 12.15 Outline the process of aminoacyl-tRNA formation.
- 12.16 How is translation (a) initiated and (b) terminated?
- 12.17 Of what significance is the wobble hypothesis?
- **12.18 (co)** If the average molecular mass of an amino acid is assumed to be 100 daltons, about how many nucleotides will be present in an mRNA coding sequence specifying a single polypeptide with a molecular mass of 27,000 daltons?
- **12.19** The bases A, G, U, C, I (inosine) all occur at the 5' positions of anticodons in tRNAs.
 - (a) Which base can pair with three different bases at the 3' positions of codons in mRNA?
 - (b) What is the minimum number of tRNAs required to recognize all codons of amino acids specified by codons with complete degeneracy?

- 12.20 a Assume that in the year 2025, the first expedition of humans to Mars discovers several Martian life forms thriving in hydrothermal vents that exist below the planet's surface. Several teams of molecular biologists extract proteins and nucleic acids from these organisms and make some momentous discoveries. Their first discovery is that the proteins in Martian life forms contain only 14 different amino acids instead of the 20 present in life forms on Earth. Their second discovery is that the DNA and RNA in these organisms have only two different nucleotides instead of the four nucleotides present in living organisms on Earth. (a) Assuming that transcription and translation work similarly in Martians and Earthlings, what is the minimum number of nucleotides that must be present in the Martian codon to specify all the amino acids in Martians? (b) Assuming that the Martian code proposed above has translational start-and-stop signals, would you expect the Martian genetic code to be degenerate like the genetic code used on Earth?
- **12.21** What are the basic differences between translation in prokaryotes and translation in eukaryotes?
- **12.22** What is the function of each of the following components of the protein-synthesizing apparatus:
 - (a) aminoacyl-tRNA synthetase,
 - (b) release factor 1,
 - (c) peptidyl transferase,
 - (d) initiation factors,
 - (e) elongation factor G?
- **12.23** An *E. coli* gene has been isolated and shown to be 68 nm long. What is the maximum number of amino acids that this gene could encode?
- 12.24 (a) What is the difference between a nonsense mutation and a missense mutation? (b) Are nonsense or missense mutations more frequent in living organisms? (c) Why?
- **12.25** The human α-globin chain is 141 amino acids long. How many nucleotides in mRNA are required to encode human α-globin?
- **12.26** What are the functions of the *A*, *P*, and *E* aminoacyl-tRNA binding sites on the ribosome?
- 12.27 (a) In what ways does the order in the genetic code minimize mutational lethality? (b) Why do base-pair changes that cause the substitution of a leucine for a valine in the polypeptide gene product seldom produce a mutant phenotype?
- **12.28** (a) What is the function of the Shine-Dalgarno sequence in prokaryotic mRNAs? (b) What effect does the deletion of the Shine-Dalgarno sequence from an mRNA have on its translation?
- 12.29 (a) In what ways are ribosomes and spliceosomes similar?(b) In what ways are they different?

12.30 The 5' terminus of a human mRNA has the following sequence:

5'cap-GAAGAGACAAGGTCAUGGCCAU-AUGCUUGUUCCAAUCGUUAGCUGCGCAG-GAUCGCCCUGGG......3'

When this mRNA is translated, what amino acid sequence will be specified by this portion of the mRNA?

12.31 A partial (5' subterminal) nucleotide sequence of a prokaryotic mRNA is as follows:

> 5'-.....AGGAGGCUCGAACAUGUCAAUAUGC-UUGUUCCAAUCGUUAGCUGCGCAGGACCGU-CCCGGA......3'

> When this mRNA is translated, what amino acid sequence will be specified by this portion of the mRNA?

12.32 • The following DNA sequence occurs in the nontemplate strand of a structural gene in a bacterium (the promoter sequence is located to the left but is not shown):

5'-GAATGTCAGAACTGCCATGCTTCATATGAA-TAGACCTCTAG-3'

- (a) What is the ribonucleotide sequence of the mRNA molecule that is transcribed from this piece of DNA?
- (b) What is the amino acid sequence of the polypeptide encoded by this mRNA?
- (c) If the nucleotide indicated by the arrow undergoes a mutation that changes T to A, what will be the resulting amino acid sequence following transcription and translation?
- 12.33 Alan Garen extensively studied a particular nonsense (chain-termination) mutation in the alkaline phosphatase gene of E. coli. This mutation resulted in the termination of the alkaline phosphatase polypeptide chain at a position where the amino acid tryptophan occurred in the wild-type polypeptide. Garen induced revertants (in this case, mutations altering the same codon) of this mutant with chemical mutagens that induced single base-pair substitutions and sequenced the polypeptides in the revertants. Seven different types of revertants were found, each with a different amino acid at the tryptophan position of the wild-type polypeptide (termination position of the mutant polypeptide fragment). The amino acids present at this position in the various revertants included tryptophan, serine, tyrosine, leucine, glutamic acid, glutamine, and lysine. Did the nonsense mutation studied by Garen contain a UAG, a UAA, or a UGA nonsense mutation? Explain the basis of your deduction.
- **12.34** The following DNA sequence occurs in a bacterium (the promoter sequence is located to the left but is not shown).

5'-CAATCATGGACTGCCATGCTTCATATGAATAGTTGACAT-3' 3'-GTTAGTACCTGACGGTACGAAGTATACTTATCAACTGTA-5

> (a) What is the ribonucleotide sequence of the mRNA molecule that is transcribed from the template strand of this

piece of DNA? Assume that both translational start and termination codons are present.

- (b) What is the amino acid sequence of the polypetide encoded by this mRNA?
- (c) If the nucleotide indicated by the arrow undergoes a mutation that causes this C:G base pair to be deleted, what will be the polypeptide encoded by the mutant gene?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

The genetic code is degenerate, with two to six codons specifying each of the amino acids except for methionine and tryptophan.

- 1. Are all of the codons specifying a given amino acid used with equal frequency, or are some codons used more frequently than others? For example, the codons UUA, UUG, CUU, CUC, CUA, and CUG all specify leucine. Are these six leucine codons present with equal frequency in the coding regions of mRNAs?
- **2.** Are the six codons specifying leucine used with equal frequency in mRNAs transcribed from human nuclear genes? From human mitochondrial genes? Are these codons used at the same frequency in nuclear and mitochondrial genes?
- **3.** Are the leucine codons used at about the same frequencies in different species, for example, in humans and *E. coli* cells? Is there any bias in codon usage (preferred use of specific

codons) related to the AT/GC content of the genomes of different species?

Hint: A search of the databases at the NCBI web site will yield an overwhelming amount of information. In this case, more accessible information can be obtained at the http://www.kazusa.jp/codon web site, which summarizes data on codon usage in 35,799 organisms (many viruses). These data are compiled from NCBI-GenBank File Release 160.0 (June 15, 2007). In the Query Box, type *Homo sapiens* and click "Submit." Your search will yield two results: (1) mitochondrion *Homo sapiens* and *Homo sapiens*. Clicking the first will give you a table of codon usage in human mitochondria, and clicking the second will give you a table of codon usage data for *E. coli* and other species of interest by simply typing the species name in the Query Box.



Mutation, DNA Repair, and Recombination

CHAPTER OUTLINE

- Mutation: Source of the Genetic Variability Required for Evolution
- The Molecular Basis of Mutation
- Mutation: Basic Features of the Process
- Mutation: Phenotypic Effects
- Assigning Mutations to Genes by the Complementation Test
- Screening Chemicals for Mutagenicity: The Ames Test
- DNA Repair Mechanisms
- Inherited Human Diseases with Defects in DNA Repair
- DNA Recombination Mechanisms

loss-of-function mutations in any of nine different autosomal genes. Moreover, other inherited disorders are known to result from the failure to repair DNA damaged by other physical and chemical agents. The life-threatening consequences of these inherited defects in the DNA repair enzymes dramatically emphasize their importance.

Given the key role that DNA plays in living organisms, the evolution of mechanisms to protect its integrity would seem inevitable. Indeed, as we discuss in this chapter, living cells contain numerous enzymes that constantly scan DNA to search for damaged or incorrectly paired nucleotides. When detected, these defects are corrected by a small army of DNA repair enzymes, each having evolved to combat a particular type of damage. In this chapter, we examine the types of changes that occur in DNA, the processes by which these alterations are corrected, and the related processes of recombination between homologous DNA molecules.

Xeroderma Pigmentosum: Defective Repair of Damaged DNA in Humans

The sun shone brightly on a midsummer day—a perfect day for most children to spend at the beach. All of Nathan's friends were dressed in shorts or swimsuits. As Nathan prepared to join his friends, he pulled on full-length sweatpants and a long-sleeved shirt. Then he put on a wide-brimmed hat and applied a thick layer of sunscreen to his hands and face. Whereas his friends enjoy playing in the sunshine, Nathan lives in constant fear of the effects of sunlight. Nathan was born with the inherited disorder xeroderma pigmentosum, an autosomal recessive trait that affects about one out of 250,000 children. Nathan's skin cells are extremely sensitive to ultraviolet radiation—the high-energy rays of sunlight. Ultraviolet light causes chemical changes in the DNA in Nathan's skin cells, changes that lead not only to intense freckling but also to skin cancer.

Nathan's friends gave little thought to playing in the sun; sunburn was their only major concern. Their skin cells contain enzymes that correct the changes in DNA resulting from exposure to ultraviolet light. However, Nathan's skin cells are lacking one of the enzymes required to repair ultraviolet light-induced alterations in the structure of DNA. Xeroderma pigmentosum results from homozygous



Children playing outdoors. The child in the white coveralls has xeroderma pigmentosum, an autosomal recessive disorder characterized by acute sensitivity to sunlight. He must avoid exposure to sunlight to prevent skin cancer.

Mutation: Source of the Genetic Variability Required for Evolution

We know from preceding chapters that inheritance is based on genes that are transmitted from parents to offspring during reproduction and that the genes store genetic information encoded in the sequences of nucleotide pairs in DNA or nucleotides in RNA. Mutations—inherited changes in the genetic material provide new genetic variation that allows organisms to evolve.

We have examined how this genetic information is accurately duplicated during the semiconservative replication of DNA. This accurate replication was shown to depend in part on proofreading activities built into the DNA polymerases that catalyze DNA synthesis. Thus, mechanisms have evolved to facilitate the faithful transmission of genetic information from cell to cell and ultimately from generation to generation. Nevertheless, mistakes in the genetic material do occur. Such heritable changes in the genetic material are called mutations.

The term **mutation** refers to both (1) the change in the genetic material and (2) the process by which the change occurs. An organism that exhibits a novel phenotype resulting from a mutation is called a **mutant**. Used in its broad historical sense, mutation refers to any sudden, heritable change in the genotype of a cell or an organism. However, changes in the genotype, and thus in the phenotype, of an organism that result from recombination events that produce new combinations of preexisting genetic variation must be carefully distinguished from changes caused by new mutations. Both events sometimes give rise to new phenotypes at very low frequencies. Mutational changes in the genotype of an organism include changes in chromosome number and structure (Chapter 6), as well as changes in the structures of individual genes. Mutations that involve changes at specific sites in a gene are referred to as **point mutations**. They include the substitution of one base pair for another or the insertion or deletion of one or a few nucleotide pairs at a specific site in a gene. Today, the term *mutation* sometimes is used in a narrow sense to refer only to changes in the structures of individual genes. In this chapter, we explore the process of mutation as defined in the narrow sense.

Mutation is the ultimate source of all genetic variation; it provides the raw material for evolution. Recombination mechanisms rearrange genetic variability into new combinations, and natural or artificial selection preserves the combinations best adapted to the existing environmental conditions or desired by the plant or animal breeder. Without mutation, all genes would exist in only one form. Alleles would not exist, and classical genetic analysis would not be possible. Most important, populations of organisms would not be able to evolve and adapt to environmental changes. Some level of mutation is essential to provide new genetic variability and allow organisms to adapt to new environments. At the same time, if mutations occurred too frequently, they would disrupt the faithful transfer of genetic information from generation to generation. Moreover, most mutations with easily detected phenotypic effects are deleterious to the organisms in which they occur. As we would expect, the rate of mutation is influenced by genetic factors, and mechanisms have evolved that regulate the level of mutation that occurs under various environmental conditions.

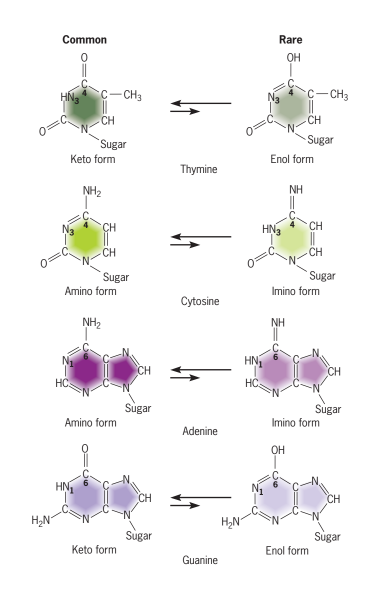
• Mutations are heritable changes in the genetic material that provide the raw material for evolution.

KEY POINT

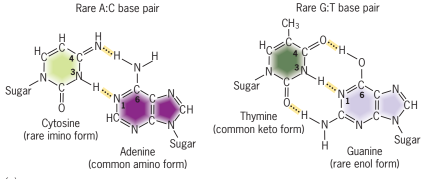
The Molecular Basis of Mutation

When Watson and Crick described the double-helix structure of DNA and proposed its semiconservative replication based on specific base-pairing to account for the accurate transmission of genetic information from generation to generation, they also proposed a mechanism to explain spontaneous Mutations alter the nucleotide sequences of genes in several ways, for example the substitution of one base pair for another or the deletion or addition of one or a few base pairs. mutation. Watson and Crick pointed out that the structures of the bases in DNA are not static. Hydrogen atoms can move from one position in a purine or pyrimidine to another position-for example, from an amino group to a ring nitrogen. Such chemical fluctuations are called tautomeric shifts. Although tautomeric shifts are rare, they may be of considerable importance in DNA metabolism because some alter the pairing potential of the bases. The nucleotide structures that we discussed in Chapter 9 are the common, more stable forms, in which adenine always pairs with thymine and guanine always pairs with cytosine. The more stable keto forms of thymine and guanine and the amino forms of adenine and cytosine may infrequently undergo tautomeric shifts to less stable enol and imino forms, respectively (**Figure 13.1**). The bases would be expected to exist in their less stable tautomeric forms for only short periods of time. However, if a base existed in the rare form at the moment that it was being replicated or being incorporated into a nascent DNA chain, a mutation would result. When the bases are present in their rare imino or enol states, they can form adenine-cytosine and guanine-thymine base pairs (**Figure 13.2a**). The net effect of such an event, and the subsequent replication required to segregate the mismatched base pair, is an A:T to G:C or a G:C to A:T base-pair substitution (**Figure 13.2***b*). See Solve It: Nucleotide-Pair Substitutions in the Human HBB Gene to examine the effects of such changes in the nucleotide sequence of an important gene.

Mutations resulting from tautomeric shifts in the bases of DNA involve the replacement of a purine in one strand of DNA with the other purine and the



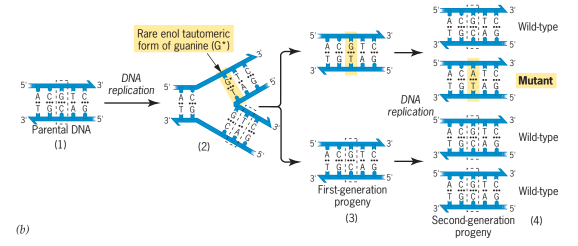
■ **FIGURE 13.1** Tautomeric forms of the four common bases in DNA. The shifts of hydrogen atoms between the number 3 and number 4 positions of the pyrimidines and between the number 1 and number 6 positions of the purines change their base-pairing potential.



Hydrogen-bonded A:C and G:T base pairs that form when cytosine and guanine are in their rare imino and enol tautomeric forms.

(a)

Mechanism by which tautomeric shifts in the bases in DNA cause mutations.



■ FIGURE 13.2 The effects of tautomeric shifts in the nucleotides in DNA on (a) basepairing and (b) mutation. Rare A:C and G:T base pairs like those shown in (a) also form when thymine and adenine are in their rare enol and imino forms, respectively. (b) A guanine (1) undergoes a tautomeric shift to its rare enol form (G*) at the time of replication (2). In its enol form, guanine pairs with thymine (2). During the subsequent replication (3 to 4), the guanine shifts back to its more stable keto form. The thymine incorporated opposite the enol form of guanine (2) directs the incorporation of adenine during the next replication (3 to 4). The net result is a G:C to A:T base-pair substitution.

replacement of a pyrimidine in the complementary strand with the other pyrimidine. Such base-pair substitutions are called **transitions**. Base-pair substitutions involving the replacement of a purine with a pyrimidine and vice versa are called **transversions**. There are three substitutions—one transition and two transversions—possible for every base pair. A total of four different transitions and eight different transversions are possible (**Transversions**. Another type of point mutation involves the addition or deletion of one or a few base pairs. Base-pair additions and deletions within the coding regions of genes are collectively referred to as **frameshift mutations** because they alter the reading frame of all base-pair triplets (DNA triplets that specify codons in mRNA and amino acids in the polypeptide gene product) in the gene that are distal to the site at which the mutation occurs (**Transver 13.3***b*).

All three types of point mutations—transitions, transversions, and frameshift mutations—are present among spontaneously occurring mutations. A surprisingly large proportion of the spontaneous mutations that have been studied in prokaryotes are single base-pair additions and deletions rather than base-pair substitutions. These frameshift mutations almost always result in the synthesis of nonfunctional protein gene products.

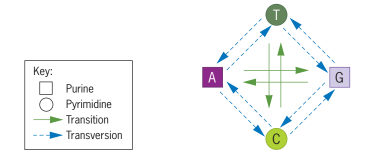
Solve It!

Nucleotide-Pair Substitutions in the Human *HBB* Gene

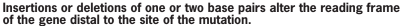
The second amino acid in mature human β -globin is histidine, which is specified by the codon CAU in the *HBB* mRNA. If you consider only single nucleotide-pair substitutions in the portion of the *HBB* gene specifying the histidine at position 2, how many different amino acid substitutions are possible? Which nucleotide-pair substitutions will give rise to each of the amino acid substitutions? Have any of these amino acid substitutions been detected in human β -globins. Have any of these variants been named? If so, what are their names? You will need to perform a web search to answer the last two questions.

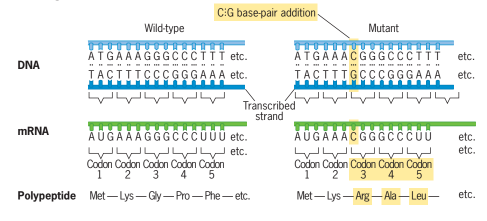
To see the solution to this problem, visit the Student Companion site.

Twelve different base substitutions can occur in DNA.









(b)

■ FIGURE 13.3 Types of point mutations that occur in DNA: (a) base substitutions and (b) frameshift mutations. (a) The base substitutions include four transitions (purine for purine and pyrimidine for pyrimidine; green arrows) and eight transversions (purine for pyrimidine and pyrimidine for purine; blue arrows). (b) A mutant gene (top, right) was produced by the insertion of a C:G base pair between the sixth and seventh base pairs of the wild-type gene (top, left). This insertion alters the reading frame of that portion of the gene distal to the mutation, relative to the direction of transcription and translation (left to right, as diagrammed). The shift in reading frame, in turn, changes all of the codons in the mRNA and all of the amino acids in the polypeptide specified by base-pair triplets distal to the mutation.

Although much remains to be learned about the causes, molecular mechanisms, and frequency of spontaneously occurring mutations, three major factors are (1) the accuracy of the DNA replication machinery, (2) the efficiency of the mechanisms that have evolved for the repair of damaged DNA, and (3) the degree of exposure to mutagenic agents present in the environment. Perturbations of the DNA replication apparatus or DNA repair systems, both of which are under genetic control, have been shown to cause large increases in mutation rates.

INDUCED MUTATIONS

Many naturally occurring mutations were identified and studied by the early geneticists. However, the science of genetics changed dramatically in 1927 when Hermann J. Muller discovered that X rays induced mutations in *Drosophila*. The ability to induce mutations opened the door to a completely new approach to genetic analysis. Geneticists could now induce mutations in genes of interest and then study the effects of the missing gene products.

Muller demonstrated that the treatment of *Drosophila* sperm with X rays sharply increased the frequency of X-linked recessive lethals. (X rays are a form of electromagnetic radiation with shorter wavelengths and higher energy than visible light;

see Figure 13.10.) Muller's study was the first demonstration that mutation could be induced by an external factor. In 1946, he received the Nobel Prize in Physiology or Medicine for this important discovery.

Muller's unambiguous demonstration of the mutagenicity of X rays became possible because he developed a simple and accurate technique that could be used to identify lethal mutations on the X chromosome of *Drosophila*. This technique, called the *CIB* method, is performed with females heterozygous for a normal X chromosome and an altered X chromosome—the *CIB* chromosome—that Muller constructed specifically for use in his experiment.

The *ClB* chromosome has three essential components. (1) *C*, for *c*rossover suppressor, refers to a long inversion that suppresses recombination between the *ClB* chromosome and the structurally normal X chromosome in heterozygous females. The inversion does not prevent crossing over between the two chromosomes, but causes progeny carrying recombinant X chromosomes produced by crossing over between the two chromosomes to abort because of duplications and deficiencies (see Chapter 7). (2) *l* refers to a recessive *l*ethal mutation on the *ClB* chromosome. Homozygous females and hemizygous males carrying this X-linked lethal mutation are not viable. (3) *B* refers to a mutation that causes the *b*ar-eye phenotype, a condition in which the large compound eyes of wild-type flies are reduced in size to narrow, bar-shaped eyes. Because *B* is partially dominant, females heterozygous for the *ClB* chromosome can be identified readily. Both the recessive lethal (*l*) and the bar-eye mutation (*B*) are located within the inverted segment of the *ClB* chromosome.

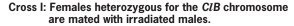
Muller irradiated male flies and mated them with *ClB/+* females **Figure 13.4**). All the bar-eyed daughters of this mating carried the *ClB* chromosome of the female parent and the irradiated X chromosome of the male parent. Because the entire population of reproductive cells of the males was irradiated, each bar-eyed daughter carried a potentially mutated X chromosome. These bar-eyed daughters were then mated individually

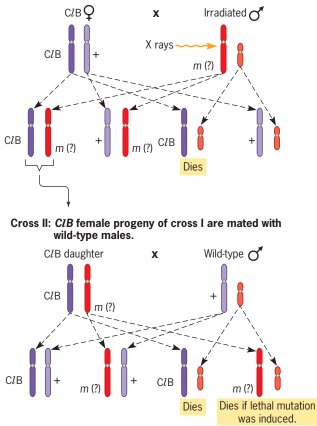
(in separate cultures) with wild-type males. If the irradiated X chromosome carried by a bar-eyed daughter had acquired an X-linked lethal, all the progeny of the mating would be female. Males hemizygous for the *ClB* chromosome would die because of the recessive lethal (*l*) this chromosome carries; in addition, males hemizygous for the irradiated X chromosome would die if a recessive lethal had been induced on it. Matings of bar-eyed daughters carrying an irradiated X chromosome in which no lethal mutation had been induced would produce female and male progeny in a ratio of 2:1 (only the males with the *ClB* chromosome will die). With the *ClB* technique, detecting newly induced recessive, X-linked lethals is unambiguous and error free; it involves nothing more complex than scoring for the presence or absence of male progeny. By this procedure, Muller was able to demonstrate a 150-fold increase in the frequency of X-linked lethals after treating male flies with X rays.

We discuss Muller's demonstration of X-ray-induced mutations on the X chromosome of *Drosophila* further in A Milestone in Genetics: Muller Demonstrates That X Rays Are Mutagenic on the Student Companion site.

Other researchers soon demonstrated that X rays are mutagenic in other organisms, including plants, other animals, and microbes. Moreover, other types of high-energy electromagnetic radiation and many chemicals were soon shown to be potent mutagens. The ability to induce mutations in genes contributed immensely to progress in genetics. It allowed researchers to induce mutations in genes of interest and "knock out" their functions. The mutant organisms could then be studied to gain information about the function of the wild-type gene product. This approach—mutational dissection—of biological processes has proven to be a powerful tool in the analysis of many biological processes.

X rays have many effects on living tissues. Therefore, X-ray-induced mutations provide little information about the molecular mechanisms by which mutations are produced. The discovery of chemical mutagens with specific effects on DNA has led to a better understanding of mutation at the molecular level.





■ FIGURE 13.4 The *ClB* technique used by Muller to detect X-linked recessive lethal mutations (*m*) in *Drosophila*. The mating shown in Cross II will produce only female progeny if an X-linked recessive lethal is present on the irradiated X chromosome. One-third of the progeny produced from Cross II will be males if there is no recessive lethal on the irradiated X chromosome. Thus, scoring for lethal mutations simply involves screening the progeny of Cross II for the presence or absence of males.

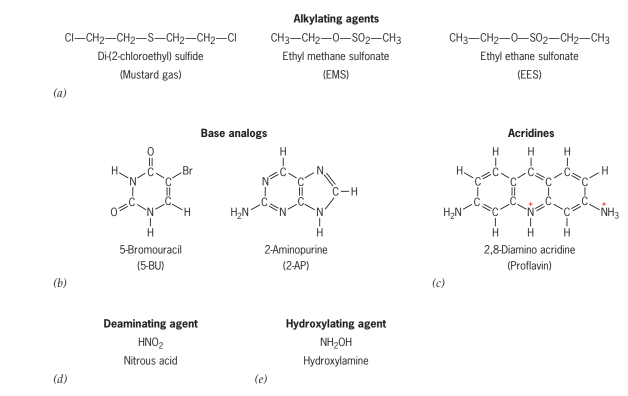
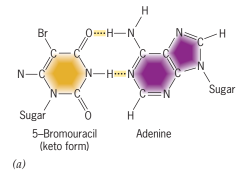
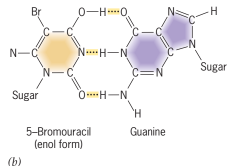


FIGURE 13.5 Some potent chemical mutagens.





5-Bromouracil : guanine base pair.



■ **FIGURE 13.6** Base-pairing between 5-bromouracil and (*a*) adenine or (*b*) guanine.

Mustard gas (sulfur mustard) was the first chemical shown to be mutagenic. Charlotte Auerbach and her associates discovered the mutagenic effects of mustard gas and related compounds during World War II. However, because of the potential use of mustard gas in chemical warfare, the British government placed their results on the classified list. Thus, Auerbach and coworkers could neither publish their results nor discuss them with other geneticists until the war ended. The compounds that they studied are examples of a large class of chemical mutagens that transfer alkyl (CH₃—, CH₃CH₂—, and so forth) groups to the bases in DNA; thus, they are called alkylating agents. Like X rays, mustard gas has many effects on DNA. Later, chemical mutagens that have specific effects on DNA were discovered (**The Figure 13.5**).

MUTATIONS INDUCED BY CHEMICALS

Chemical mutagens can be divided into two groups: (1) those that are mutagenic to both replicating and nonreplicating DNA, such as the alkylating agents and nitrous acid; and (2) those that are mutagenic only to replicating DNA, such as base analogs—purines and pyrimidines with structures similar to the normal bases in DNA. The base analogs must be incorporated into DNA chains in the place of normal bases during replication in order to exert their mutagenic effects. The second group of mutagens also includes the acridine dyes, which intercalate into DNA and increase the probability of mistakes during replication.

The mutagenic **base analogs** have structures similar to the normal bases and are incorporated into DNA during replication. However, their structures are sufficiently different from the normal bases in DNA that they increase the frequency of mispairing, and thus mutation, during replication. The two most commonly used base analogs are 5-bromouracil and 2-aminopurine. The pyrimidine 5-bromouracil is a thymine analog; the bromine at the 5 position is similar in several respects to the methyl ($-CH_3$) group at the 5 position in thymine. However, the bromine at this position changes the charge distribution and increases the frequency of tautomeric shifts (see Figure 13.1). In its more stable keto form, 5-bromouracil pairs with adenine. After a tautomeric shift to its enol form, 5-bromouracil pairs with guanine (**\blacksquare Figure 13.6**).

The mutagenic effect of 5-bromouracil is the same as that predicted for tautomeric shifts in normal bases (see Figure 13.2*b*), namely, transitions.

If 5-bromouracil is present in its less frequent enol form as a nucleoside triphosphate at the time of its incorporation into a nascent strand of DNA, it will be incorporated opposite guanine in the template strand and cause a G:C \rightarrow A:T transition (**Figure 13.7a**). If, however, 5-bromouracil is incorporated in its more frequent keto form opposite adenine (in place of thymine) and undergoes a tautomeric shift to its enol form during a subsequent replication, it will cause an A:T \rightarrow G:C transition (**Figure 13.7b**). Thus, 5-bromouracil induces transitions in both directions, A:T \leftrightarrow G:C. An important consequence of the bidirectionality of 5-bromouracil-induced transitions is that mutations originally induced with this thymine analog can also be induced to mutate back to the wild-type with 5-bromouracil. 2-Aminopurine acts in a similar manner but is incorporated in place of adenine or guanine.

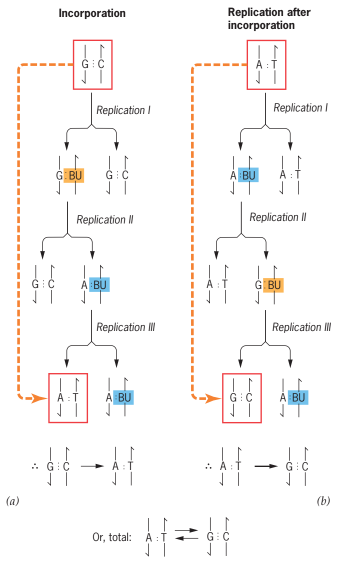
Nitrous acid (HNO₂) is a potent mutagen that acts on either replicating or nonreplicating DNA. Nitrous acid causes oxidative deamination of the amino groups in adenine, guanine, and cytosine. This reaction converts the amino groups to keto groups and changes the hydrogen-bonding potential of the modified bases (**Figure 13.8**). Adenine is deaminated to hypoxanthine, which base-pairs with cytosine rather than thymine. Cytosine is converted to uracil, which base-pairs with adenine instead of guanine. Deamination of guanine produces xanthine, but xanthinejust like guanine-base-pairs with cytosine. Thus, the deamination of guanine is not mutagenic. Because the deamination of adenine results in A:T \rightarrow G:C transitions, and the deamination of cytosine produces $G:C \rightarrow A:T$ transitions, nitrous acid induces transitions in both directions, A:T \leftrightarrow G:C. As a result, nitrous acid-induced mutations also are induced to mutate back to wild-type by nitrous acid. Test your understanding of nitrous acid-induced mutation by working through Problem-Solving Skills: Predicting Amino Acid Changes Induced by Chemical Mutagens.

The **acridine dyes** such as proflavin (see Figure 13.5*c*), acridine orange, and a whole series of related compounds are potent mutagens that induce frameshift mutations (see Figure 13.3*b*). The positively charged acridines intercalate, or sandwich themselves, between the stacked base pairs in DNA (**■** Figure 13.9). In so doing, they increase the rigidity and alter the conformation of the double helix, causing slight bends or kinks in the molecule. When DNA molecules containing intercalated acridines replicate, additional delations of erast as a first here arise acridines replicate,

additions and deletions of one to a few base pairs occur. As we might expect, these small additions and deletions, usually of a single base pair, result in altered reading frames for the portion of the gene distal to the mutation (see Figure 13.3*b*). Thus, acridine-induced mutations in exons of genes usually result in nonfunctional gene products.

Alkylating agents are chemicals that donate alkyl groups to other molecules. They include nitrogen mustard, and methyl and ethyl methane sulfonate (MMS and EMS) (see Figure 13.5*a*)—chemicals that have multiple effects on DNA. Alkylating agents induce all types of mutations, including transitions, transversions, frameshifts, and even chromosome aberrations, with relative frequencies that depend on the reactivity of the agent involved. One mechanism of mutagenesis by alkylating agents involves the transfer of methyl or ethyl groups to the bases, resulting in altered base-pairing potentials. For example, EMS causes ethylation of the bases in DNA at the 7-N and the 6-O positions. When 7-ethylguanine is produced, it base-pairs with thymine to cause G:C \rightarrow A:T transitions. Other base alkylation products activate error-prone DNA repair processes that introduce transitions, transversions, and frameshift mutations during the repair process. Some alkylating agents, particularly difunctional alkylating agents (those with two reactive alkyl groups), cross-link DNA strands or molecules and induce chromosome breaks, which result in various kinds of chromosomal aberrations (Chapter 6). Alkylating agents as a class therefore exhibit less specific mutagenic effects than do base analogs, nitrous acid, or acridines.

Effect of enol form of 5-bromouracil during:



■ FIGURE 13.7 The mutagenic effects of 5-bromouracil. (a) When 5-bromouracil (BU) is present in its less frequent enol form (orange) at the time of incorporation into DNA, it induces G:C → A:T transitions. (b) When 5-bromouracil is incorporated into DNA in its more common keto form (blue) and shifts to its enol form during a subsequent replication, it induces A:T → G:C transitions. Thus, 5-bromouracil can induce transitions in both directions, A:T ↔ G:C.

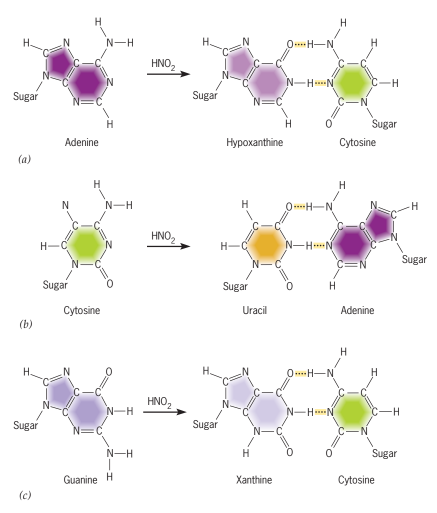
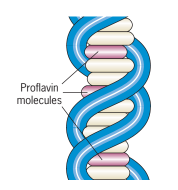


FIGURE 13.8 Nitrous acid induces mutations by oxidative deamination of the bases in DNA. Nitrous acid converts (*a*) adenine to hypoxanthine, causing A:T \rightarrow G:C transitions; (*b*) cytosine to uracil, causing G:C \rightarrow A:T transitions; and (*c*) guanine to xanthine, which is not mutagenic. Together, the effects of nitrous acid on adenine and cytosine explain its ability to induce transitions in both directions, A:T \leftrightarrow G:C.

In contrast to most alkylating agents, the **hydroxylating agent** hydroxylamine (NH_2OH) has a specific mutagenic effect. It induces only $G:C \rightarrow A:T$ transitions. When DNA is treated with hydroxylamine, the amino group of cytosine is hydroxylated. The resulting hydroxylaminocytosine base-pairs with adenine, leading to $G:C \rightarrow A:T$ transitions. Because of its specificity, hydroxylamine has been very useful in classifying transition mutations. Mutations that are induced to revert to wild-type by nitrous acid or base analogs, and therefore were originally caused by transitions, can be divided into two classes on the basis of their revertibility with hydroxylamine. (1) Those with an A:T base pair at the mutant site will not be induced to revert by hydroxylamine. (2) Those with a G:C base pair at the mutant site will be induced to revert by hydroxylamine. Thus, hydroxylamine can be used to determine whether a particular mutation was an A:T \rightarrow G:C or a G:C \rightarrow A:T transition.

MUTATIONS INDUCED BY RADIATION

The portion of the electromagnetic spectrum (**Figure 13.10**) with wavelengths shorter and of higher energy than visible light is subdivided into **ionizing radiation** (X rays, gamma rays, and cosmic rays) and **nonionizing radiation** (ultraviolet light). Ionizing radiations are of high energy and are useful for medical diagnosis because they penetrate living tissues for substantial distances. In the process, these high-energy



■ FIGURE 13.9 Intercalation of proflavin into the DNA double helix. X-ray diffraction studies have shown that these positively charged acridine dyes become sandwiched between the stacked base pairs.

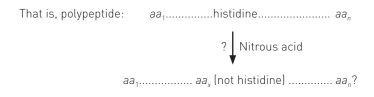
PROBLEM-SOLVING SKILLS



Predicting Amino Acid Changes Induced by Chemical Mutagens

THE PROBLEM

You are given the nature of the genetic code in Table 12.1. As is illustrated in Figure 13.8, the chemical nitrous acid deaminates adenine, cytosine, and guanine (adenine \rightarrow hypoxanthine, which base-pairs with cytosine; cytosine \rightarrow uracil, which base-pairs with adenine; and guanine \rightarrow xanthine, which base-pairs with cytosine). If you treat a population of nonreplicating tobacco mosaic viruses (TMVs) with nitrous acid, would you expect the nitrous acid to induce any mutations that result in the substitution of another amino acid for a histidine (His) residue in a wild-type polypeptide?



If so, what amino acid(s), and by what mechanism(s)? If not, why not?

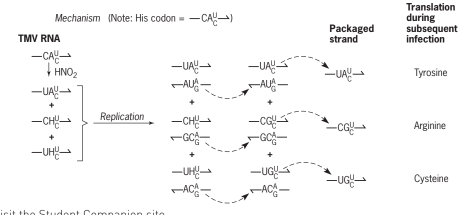
FACTS AND CONCEPTS

- **1.** TMV stores its genetic information in single-stranded RNA that is equivalent to mRNA.
- 2. The TMV genomic RNA replicates like DNA via a complementary (base-paired) double-stranded intermediate.

- **3.** Although the tobacco mosaic viruses are not replicating at the time of treatment with nitrous acid, they will subsequently be allowed to replicate by infecting tobacco leaves in order to determine whether or not any mutations of the indicated type were induced by treatment with nitrous acid.
- The histidine codons are CAU and CAC. Therefore, the TMV genome (RNA) contains one of these sequences at all sites specifying histidine in the polypeptides encoded by TMV.
- **5.** The adenines and cytosines in the TMV genome are potential targets of nitrous acid-induced mutation.

ANALYSIS AND SOLUTION

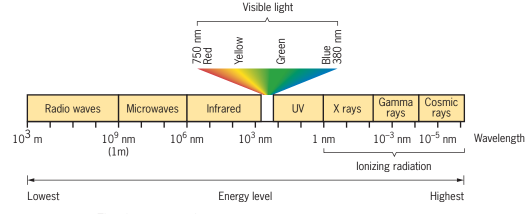
When nitrous acid deaminates adenine and cytosine, it produces hypoxanthine and uracil, respectively. During subsequent replication of the modified TMV RNAs, hypoxanthine pairs with cytosine and uracil pairs with adenine. As a result, some of the A's and C's in TMV RNA will be converted to G's and U's. The deamination of these bases results in tyrosine, arginine, and cysteine codons in the TMV genomes produced by the semiconservative replication of the mutagenized viral RNA. Thus, nitrous acid mutagenesis will lead to the replacement of some histidines in wild-type TMV proteins with tyrosines, arginines, and cysteines in mutant proteins, as shown in the following diagram.



For further discussion visit the Student Companion site.

rays collide with atoms and cause the release of electrons, creating positively charged free radicals or ions. The ions, in turn, collide with other molecules and cause the release of additional electrons. The result is that a cone of ions is formed along the track of each high-energy ray as it passes through living tissues. This process of ionization is induced by machine-produced X rays, protons, and neutrons, as well as by the alpha, beta, and gamma rays released by radioactive isotopes such as ³²P, ³⁵S, and the uranium-238 used in nuclear reactors.

Ultraviolet rays, having lower energy than ionizing radiations, penetrate only the surface layer of cells in higher plants and animals and do not cause ionizations. Ultraviolet rays dissipate their energy to the atoms they encounter, raising the electrons in the outer orbitals to higher energy levels, a state referred to as *excitation*.





Molecules containing atoms in either ionic forms or excited states are chemically more reactive than those containing atoms in their normal stable states. The increased reactivity of atoms present in DNA molecules is responsible for most of the mutagenicity of ionizing radiation and ultraviolet light.

X rays and other forms of ionizing radiation are quantitated in **roentgen** (**r**) units, which are measures of the number of ionizations per unit volume under a standard set of conditions. Specifically, one roentgen unit is a quantity of ionizing radiation that produces $2.083 \times 10^{\circ}$ ion pairs in one cubic centimeter of air at 0°C and a pressure of 760 mm of mercury. Note that the dosage of irradiation in roentgen units does not involve a time scale. The same dosage may be obtained by a low intensity of irradiation over a long period of time or a high intensity of irradiation for a short period of time. This point is important because in most studies the frequency of induced point mutations is directly proportional to the dosage of irradiation (**Tradiation** Figure 13.11). For example, X-irradiation of *Drosophila* sperm causes an approximately 3 percent increase in mutation rate for each 1000-r increase in irradiation dosage. This linear relationship shows that the induction of mutations by X rays exhibits single-hit kinetics, which means that each mutation results from a single ionization event. That is, every ionization has a fixed probability of inducing a mutation under a standard set of conditions.

What is a safe level of irradiation? The development and use of the atomic bomb and the accidents at nuclear power plants have generated concern about exposure to ionizing radiations. The linear relationship between mutation rate and radiation dosage indicates that there is no safe level of irradiation. Rather, the results indicate

> that the higher the dosage of irradiation, the higher the mutation rate, and the lower the dosage, the lower the mutation rate. Even very low levels of irradiation have certain low, but real, probabilities of inducing mutations.

> In *Drosophila* sperm, chronic irradiation (low levels of irradiation over long periods of time) is as effective in inducing mutations as acute irradiation (the same total dosage of irradiation administered at high intensity for short periods of time). However, in mice, chronic irradiation results in fewer mutations than the same dosage of acute irradiation. Moreover, when mice are treated with intermittent doses of irradiation, the mutation frequency is slightly lower than when they are treated with the same total amount of irradiation in a continuous dose. The differential response of fruit flies and mammals to chronic irradiation is thought to result from differences in the efficiency with which these species repair irradiationinduced damage in DNA. Repair mechanisms may exist in the spermatogonia and oocytes of mammals that do not function in *Drosophila* sperm. Nevertheless, we should emphasize that all of these irradiation treatments are mutagenic, albeit to different degrees, in both *Drosophila* and mammals.

Ionizing radiation also induces gross changes in chromosome structure, including deletions, duplications, inversions, and translocations (Chapter 6). These chromosome aberrations result from radiation-induced breaks in chromosomes. Because these

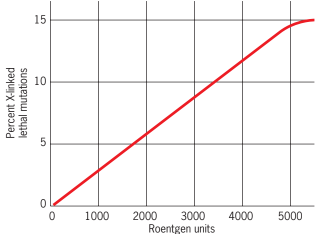


FIGURE 13.11 Relationship between irradiation dosage and mutation frequency in *Drosophila*.

aberrations require two chromosomal breaks, they exhibit two-hit kinetics rather than the single-hit kinetics observed for point mutations.

Ultraviolet (UV) radiation does not possess sufficient energy to induce ionizations. However, it is readily absorbed by many organic molecules such as the purines and pyrimidines in DNA, which then enter a more reactive or excited state. UV rays penetrate tissue only slightly. Thus, in multicellular organisms, only the epidermal layer of cells usually is exposed to the effects of UV. However, ultraviolet light is a potent mutagen for unicellular organisms. The maximum absorption of UV by DNA is at a wavelength of 254 nm. Maximum mutagenicity also occurs at 254 nm, suggesting that the UV-induced mutation process is mediated directly by the absorption of UV by purines and pyrimidines. In vitro studies show that the pyrimidines absorb strongly at 254 nm and, as a result, become very reactive. Two major products of UV absorption by pyrimidines (thymine and cytosine) are pyrimidine hydrates and pyrimidine dimers (**Figure 13.12**). Thymine dimers cause mutations in two ways. (1) Dimers perturb the structure of DNA double

helices and interfere with accurate DNA replication. (2) Errors occur during the cellular processes that repair defects in DNA, such as UV-induced thymine dimers (see the section DNA Repair Mechanisms later in this chapter).

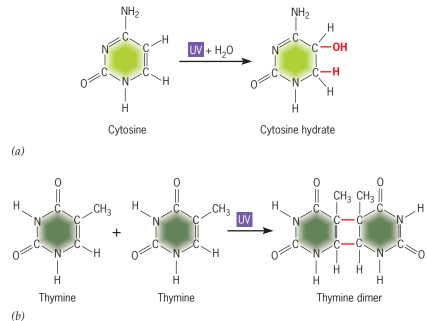
MUTATIONS INDUCED BY TRANSPOSABLE GENETIC ELEMENTS

Living organisms contain remarkable DNA elements that can move from one site in the genome to another site. These **transposons**, or transposable genetic elements, are the subject of Chapter 17. The insertion of a transposon into a gene will often render the gene nonfunctional (**Theorem Figure 13.13**). If the gene encodes an important product, a mutant phenotype is likely to result. Geneticists now know that many of the classical mutants of maize, *Drosophila*, *E. coli*, and other organisms were caused by the insertion of transposable genetic elements into important genes. Indeed, Mendel's *wrinkled* allele in the pea (Chapter 3) and the first mutation (*w*¹) causing white eyes in *Drosophila* (Chapter 5) both resulted from the insertion of transposable elements. See Chapter 17 for additional details about the mechanisms by which transposons move and, in the process, produce mutations.

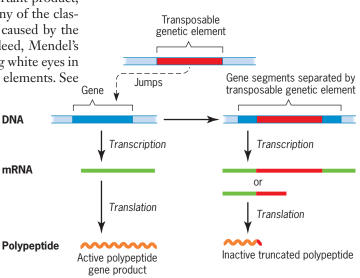
EXPANDING TRINUCLEOTIDE REPEATS AND INHERITED HUMAN DISEASES

All of the types of mutations discussed in the preceding sections of this chapter occur in humans. In addition, another type of mutation occurs that is associated with human diseases. Repeated sequences of one to six nucleotide pairs are known as **simple tandem repeats**. Such repeats are dispersed throughout the human genome. Repeats of three nucleotide pairs, **trinucleotide repeats**, can increase in copy number and cause inherited diseases in

humans. Several trinucleotides have been shown to undergo such increases in copy number. Expanded CGG trinucleotide repeats at the *FRAXA* site on the X chromosome are responsible for fragile X syndrome, the second most common form of inherited mental retardation in humans (Chapter 16). Normal X chromosomes contain from 6 to about 50 copies of the CGG repeat at the *FRAXA* site. Mutant X chromosomes contain up to 1000 copies of the tandem CGG repeat at this site (see Focus on Fragile X Syndrome and Expanded Trinucleotide Repeats in Chapter 16).



■ FIGURE 13.12 Pyrimidine photoproducts of UV irradiation. (a) Hydrolysis of cytosine to a hydrate form that may cause mispairing of bases during replication. (b) Cross-linking of adjacent thymine molecules to form thymine dimers, which block DNA replication.



■ FIGURE 13.13 Mechanism of transposoninduced mutation. The insertion of a transposable genetic element (red) into a wildtype gene (left) will usually render the gene nonfunctional (right). A truncated gene product usually results from transcriptionor translation-termination signals, or both, located within the transposon. CAG and CTG trinucleotide repeats are involved in several inherited neurological diseases, including Huntington disease, myotonic dystrophy, Kennedy disease, dentatorubral pallidoluysian atrophy, Machado-Joseph disease, and spinocerebellar ataxia. In all of these neurological disorders, the severity of the disease is correlated with trinucleotide copy number—the higher the copy number, the more severe the disease symptoms. In addition, the expanded trinucleotides associated with these diseases are unstable in somatic cells and between generations. This instability gives rise to the phenomenon of *anticipation*, which is the increasing severity of the disease or earlier age of onset that occurs in successive generations as the trinucleotide copy number increases. A possible mechanism of trinucleotide expansion is discussed in Chapter 16.

KEY POINTS

- Mutations are induced by chemicals, ionizing irradiation, ultraviolet light, and endogenous transposable genetic elements.
 - Point mutations are of three types: (1) transitions—purine for purine and pyrimidine for pyrimidine substitutions; (2) transversions—purine for pyrimidine and pyrimidine for purine substitutions; and (3) frameshift mutations—additions or deletions of one or two nucleotide pairs, which alter the reading frame of the gene distal to the site of the mutation.
 - Several inherited human diseases are caused by expanded trinucleotide repeats.

Mutation: Basic Features of the Process

Mutations occur in all organisms from viruses to humans. They can occur spontaneously or be induced by mutagenic agents. Mutation is usually a random, nonadaptive process. Mutations occur in all genes of all living organisms. These mutations provide new genetic variability that allows organisms to adapt to environmental changes. Thus, mutations have been, and continue to be, essential to the evolutionary process. Before we discuss the phenotypic effects of mutations, let's consider some of the basic features of this important process.



FIGURE 13.14 The original Delicious apple was the result of a somatic mutation. It has subsequently been modified by the selection of additional somatic mutations.

MUTATION: SOMATIC OR GERMINAL

A mutation may occur in any cell and at any stage in the development of a multicellular organism. The immediate effects of the mutation and its ability to produce a phenotypic change are determined by its dominance, the type of cell in which it occurs, and the time at which it takes place during the life cycle of the organism. In higher animals, the germ-line cells that give rise to the gametes separate from other cell lineages early in development (Chapter 2). All nongerm-line cells are somatic cells. **Germinal mutations** are those that occur in germ-line cells, whereas **somatic mutations** occur in somatic cells.

If a mutation occurs in a somatic cell, the resulting mutant phenotype will occur only in the descendants of that cell. The mutation will not be transmitted through the gametes to the progeny. The Delicious apple (**Figure 13.14**) and the navel orange are examples of mutant phenotypes that resulted from mutations occurring in somatic cells. The Delicious apple was discovered in 1881 by Jessie Hiatt, an Iowa farmer. It has subsequently been modified by the selection of additional somatic mutations. The fruit trees in which the original mutations occurred were somatic mosaics. Fortunately, vegetative propagation was feasible for both the Delicious apple and the navel orange, and today numerous progeny from grafts and buds have perpetuated the original mutations. If dominant mutations occur in germ-line cells, their effects may be expressed immediately in progeny. If the mutations are recessive, their effects are often obscured in diploids. Germinal mutations may occur at any stage in the reproductive cycle of the organism. If the mutation arises in a gamete, only a single member of the progeny is likely to have the mutant gene. If a mutation occurs in a primordial germ-line cell of the testis or ovary, several gametes may receive the mutant gene, enhancing its potential for perpetuation. Thus, the dominance of a mutant allele and the stage in the reproductive cycle at which a mutation occurs are major factors in determining the likelihood that the mutant allele will be manifested in an organism.

The earliest recorded dominant germinal mutation in domestic animals was that observed by Seth Wright in 1791 on his farm by the Charles River in Dover, Massachusetts. Among his flock of sheep, Wright noticed a peculiar male lamb with unusually short legs. It occurred to him that it would be an advantage to have a whole flock of these short-legged sheep, which could not jump over the low stone fences in his New England neighborhood. Wright used the new short-legged ram to breed his ewes in the next season. Two of their lambs had short legs. Short-legged sheep were then bred together, and a line was developed in which the new trait was expressed in all individuals.

MUTATION: SPONTANEOUS OR INDUCED

When a new mutation—such as the one that produced Wright's short-legged sheep occurs, is it caused by some agent in the environment or does it result from an inherent process in living organisms? **Spontaneous mutations** are those that occur without a known cause. They may truly be spontaneous, resulting from a low level of inherent metabolic errors, or they may actually be caused by unknown agents present in the environment. **Induced mutations**, as already discussed, are those resulting from exposure of organisms to physical and chemical agents that cause changes in DNA (or RNA in some viruses). Such agents are called **mutagens**; they include ionizing irradiation, ultraviolet light, and a wide variety of chemicals, as discussed in the preceding section.

Operationally, it is impossible to prove that a particular mutation occurred spontaneously or was induced by a mutagenic agent. Geneticists must restrict such distinctions to the population level. If the mutation rate is increased a hundredfold by treatment of a population with a mutagen, an average of 99 of every 100 mutations present in the population will have been induced by the mutagen. Researchers can thus make valid comparisons between spontaneous and induced mutations statistically by comparing populations exposed to a mutagenic agent with control populations that have not been exposed to the mutagen.

Spontaneous mutations occur infrequently, although the observed frequencies vary from gene to gene and from organism to organism. Measurements of spontaneous mutation frequencies for various genes of phage and bacteria range from about 10^{-8} to 10^{-10} detectable mutations per nucleotide pair per generation. For eukaryotes, estimates of mutation rates range from about 10^{-7} to 10^{-9} detectable mutations per nucleotide pair per generation. For eukaryotes, estimates of mutation rates range from about 10^{-7} to 10^{-9} detectable mutations per nucleotide pair per generation (considering only those genes for which extensive data are available). In comparing mutation rates per nucleotide with mutation rates per gene, the coding region of the average gene is usually assumed to be 1000 nucleotide pairs in length. Thus, the mutation rate per gene varies from about 10^{-4} to 10^{-7} per generation.

Treatment with mutagenic agents can increase mutation frequencies by orders of magnitude. The mutation frequency per gene in bacteria and viruses can be increased to over 1 percent by treatment with potent chemical mutagens. That is, over 1 percent of the genes of the treated organisms will contain a mutation, or, stated differently, over 1 percent of the phage or bacteria in the population will have a mutation in a given gene.

MUTATION: USUALLY A RANDOM, NONADAPTIVE PROCESS

The rats in many cities are no longer affected by the anticoagulants that have traditionally been used as rodent poisons. Many cockroach populations are insensitive to chlordane, the poison used to control them in the 1950s. Housefly populations often exhibit high

levels of resistance to many insecticides. More and more pathogenic microorganisms are becoming resistant to antibiotics developed to control them. The introduction of these pesticides and antibiotics by humans produced new environments for these organisms. Mutations producing resistance to these pesticides and antibiotics occurred; the sensitive organisms were killed; and the mutants multiplied to produce new resistant populations. Many such cases of evolution via mutation and natural selection are well documented.

These examples raise a basic question about the nature of mutation. Is mutation a purely random event in which the environmental stress merely preserves preexisting mutations? Or is mutation directed by the environmental stress? For example, if you cut off the tails of mice for many generations, will you eventually produce a strain of tailless mice? Despite the beliefs of Jean Lamarck and Trofim Lysenko, who believed in the inheritance of "acquired traits"—traits imposed on organisms by environmental factors—the answer is no; the mice will continue to be born with tails.

Today, it is hard to understand how Lysenko could have sold his belief in Lamarckism—the inheritance of acquired traits—to those in power in the Soviet Union from 1937 through 1964. However, disproving Lamarckism was not an easy task, especially in the case of microorganisms, where even small cultures often contain billions of organisms.

As an example, let us consider a population of bacteria such as *E. coli* growing in a streptomycin-free environment. When exposed to streptomycin, most of the bacteria will be killed by the antibiotic. However, if the population is large enough, it will soon give rise to a streptomycin-resistant culture in which all the cells are resistant to the antibiotic. Does streptomycin simply select rare, randomly occurring mutants that preexist in the population, or do all of the cells have some low probability of developing resistance in response to the presence of streptomycin? How can geneticists distinguish between these two possibilities? Resistance to streptomycin can only be detected by treating the culture with the antibiotic. How, then, can a geneticist determine whether resistant bacteria are present prior to exposure to streptomycin, or are induced by the presence of the antibiotic?

In 1952, Joshua and Esther Lederberg developed an important new technique called **replica plating**. This technique allowed them to demonstrate the presence of antibiotic-resistant mutants in bacterial cultures prior to exposure to the antibiotic (**Figure 13.15**). The Lederbergs first diluted the bacterial cultures, spread the bacteria on the surface of semisolid nutrient agar medium in petri dishes, and incubated the plates until each bacterium had produced a visible colony on the surface of the agar. They next inverted each plate and pressed it onto sterile velvet placed over a wood block. Some of the cells from each colony stuck to the velvet. They then gently pressed a sterile plate of nutrient agar medium containing streptomycin onto the velvet. They repeated this replica-plating procedure with many plates, each containing about 200 bacterial colonies. After they incubated the selective plates (those containing streptomycin) overnight, rare streptomycin-resistant colonies had formed.

The Lederbergs subsequently tested the colonies on the nonselective plates (those not containing streptomycin) for their ability to grow on medium containing streptomycin. Their results were definitive. The colonies that grew on the selective replica plates almost always contained streptomycin-resistant cells, whereas colonies that failed to grow on the selective medium seldom contained any resistant cells (Figure 13.15).

If a mutation that makes a bacterium resistant to streptomycin occurs at an early stage in the growth of a colony, the resistant cell will divide and produce two, then four, then eight, and eventually a large number of resistant bacteria. Thus, if mutation is a randomly occurring, nonadaptive process, many of the colonies that form on the nonselective plates will contain more than one antibiotic-resistant bacterium and will give rise to resistant cultures when tested for growth on selective media. However, if mutation is adaptive and the mutations to streptomycin resistance occur only after exposure to the antibiotic, then the colonies on the nonselective plates that gave rise to resistant colonies on the selective plates after replica plating would be no more likely to contain streptomycin-resistant cells than the other colonies on the nonselective plates.

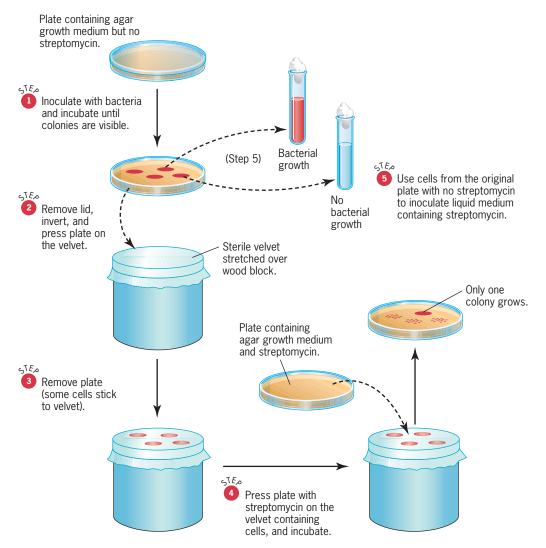


FIGURE 13.15 Joshua and Esther Lederberg's use of replica plating to demonstrate the random or nondirected nature of mutation. For simplicity, only four colonies are shown on each plate, and only two are tested for streptomycin resistance in Step 5. Actually, each plate would contain about 200 colonies, and many plates would be used to find an adequate number of mutant colonies.

Thus, by using their replica-plating technique, the Lederbergs demonstrated the existence of streptomycin-resistant mutants in a population of bacteria prior to their exposure to the antibiotic. Their results, along with those of many other experiments, have shown that environmental stress does not direct or cause genetic changes as Lysenko believed; it simply selects rare preexisting mutations that result in phenotypes better adapted to the new environment.

MUTATION: A REVERSIBLE PROCESS

As we discussed earlier, a mutation in a wild-type gene can produce a mutant allele that results in an abnormal phenotype. However, the mutant allele can also mutate back to a form that restores the wild-type phenotype. That is, mutation is a reversible process.

The mutation of a wild-type gene to a form that results in a mutant phenotype is referred to as *forward mutation*. However, sometimes the designation of the wild-type and mutant phenotypes is quite arbitrary. They may simply represent two different, but normal, phenotypes. For example, geneticists consider the alleles for brown and blue eye color in humans both to be wild-type. However, in a population composed almost entirely of brown-eyed individuals, the allele for blue eyes might be thought of as a mutant allele.

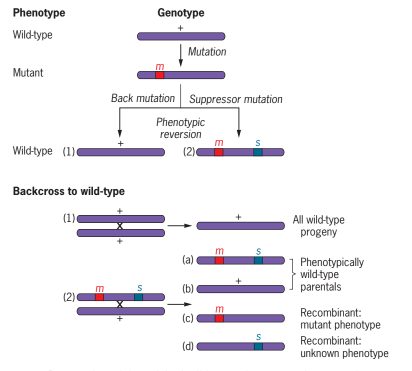


FIGURE 13.16 Restoration of the original wild-type phenotype of an organism may occur by (1) back mutation or (2) suppressor mutation (shown on the same chromosome for simplicity). Some mutants can revert to the wild-type phenotype by both mechanisms. Revertants of the two types can be distinguished by backcrosses to the original wild-type. If back mutation has occurred, all backcross progeny will be wild-type. If a suppressor mutation is responsible, some of the backcross progeny will have the mutant phenotype (2c).

When a second mutation restores the original phenotype lost because of an earlier mutation, the process is called **reversion** or **reverse mutation**. Reversion may occur in two different ways: (1) by **back mutation**, a second mutation at the same site in the gene as the original mutation, restoring the wild-type nucleotide sequence, or (2) by **suppressor mutation**, a second mutation at a different location in the genome, which compensates for the effects of the first mutation (**Figure 13.16**). Back mutation restores the original wild-type nucleotide sequence of the gene, whereas a suppressor mutation does not. Suppressor mutations may occur at distinct sites in the same gene as the original mutation or in different genes, even on different chromosomes.

Some mutations revert primarily by back mutation, whereas others do so almost exclusively through the occurrence of suppressor mutations. Thus, in genetic studies, researchers often must distinguish between these two possibilities by backcrossing the phenotypic revertant with the original wild-type organism. If the wild-type phenotype is restored by a suppressor mutation, the original mutation will still be present and can be separated from the suppressor mutation by recombination (Figure 13.16). If the wild-type phenotype is restored by back mutation, all of the progeny of the backcross will be wild-type.

KEY POINTS

• Mutations occur in both germ-line and somatic cells, but only germ-line mutations are transmitted to progeny.

- Mutations can occur spontaneously or be induced by mutagenic agents in the environment.
- Mutation usually is a nonadaptive process in which an environmental stress simply selects organisms with preexisting, randomly occurring mutations.
- Restoration of the wild-type phenotype in a mutant organism can result from either back mutation or a suppressor mutation.

Mutation: Phenotypic Effects

The effects of mutations on phenotype range from alterations so minor that they can be detected only by special genetic or biochemical techniques, to gross modifications of morphology, to lethals. A gene is a

sequence of nucleotide pairs that usually encodes a specific polypeptide. Any mutation occurring within a given gene will thus produce a new allele of that gene. Genes containing mutations with no effect on phenotype or small effects that can be recognized only by special techniques are called **isoalleles**. Other mutations produce **null alleles** that result in no gene product or totally nonfunctional gene products. If mutations of the latter type occur in genes that are required for the growth of the organism, individuals that are homozygous for the mutation will not survive. Such mutations are called **recessive lethals**.

Mutations can be either recessive or dominant. In monoploid organisms such as viruses and bacteria, both recessive and dominant mutations can be recognized by their effect on the phenotype of the organism in which they occur. In diploid organisms such as fruit flies and humans, recessive mutations will alter the phenotype only when present in the homozygous condition. Thus, in diploids, most recessive mutations will not be recognized at the time of their occurrence because they will be present in the heterozygous state. X-linked recessive mutations are an exception; they will be expressed in the hemizygous state in the heterogametic sex (for example, males in humans and fruit flies; females in birds). X-linked recessive lethal mutations will alter the sex ratio of offspring because hemizygous individuals that carry the lethal will not survive (**Figure 13.17**).

MUTATIONS WITH PHENOTYPIC EFFECTS: USUALLY DELETERIOUS AND RECESSIVE

Most of the mutations that have been identified and studied by geneticists are deleterious and recessive. This result is to be expected if we consider what is known about the genetic control of metabolism and the techniques available for identifying mutations. As we discussed in Chapter 4, metabolism occurs by sequences of chemical reactions, with each step catalyzed by a specific enzyme encoded by one or more genes. Mutations in these genes frequently produce blocks in metabolic pathways (**Figure 13.18**). These blocks occur because alterations in the base-pair sequences of genes often cause changes in the amino acid sequences of polypeptides (**Figure 13.19**), which may result in nonfunctional products (Figure 13.18). Indeed, this is the most commonly observed effect of easily detected mutations. Given a wildtype allele encoding an active enzyme and mutant alleles encoding less active or totally inactive enzymes, it is apparent why most of the observed mutations would be recessive. If a cell contains both active and inactive forms of a given enzyme, the active form usually will catalyze the reaction in question. Therefore, the allele specifying the active product usually will be dominant, and the allele encoding the inactive product will be recessive (Chapter 4).

Because of the degeneracy and order in the genetic code (Chapter 12), many mutations have no effect on the phenotype of the organism; they are called **neutral mutations**. But why should most mutations with phenotypically recognizable effects result in decreased gene-product activity or no gene-product activity? A wild-type allele of a gene encoding a wild-type enzyme or structural protein will have been selected for optimal activity during the course of



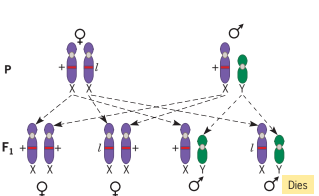


FIGURE 13.17 Alteration of the sex ratio by an X-linked recessive lethal mutation. Females heterozygous for an X-linked recessive lethal will produce female and male progeny in a 2:1 ratio.

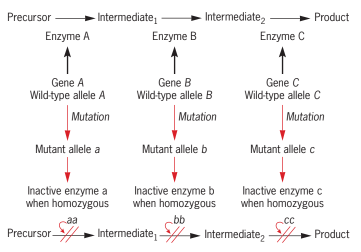


FIGURE 13.18 Recessive mutant alleles often result in blocks in metabolic pathways. The pathways can be only a few steps long, as diagrammed here, or many steps long. The wild-type allele of each gene usually encodes a functional enzyme that catalyzes the appropriate reaction. Most mutations that occur in wild-type genes result in altered forms of the enzyme with reduced or no activity. In the homozygous state, mutant alleles that produce inactive products cause metabolic blocks $(-\lambda \rightarrow)$ owing to the lack of the required enzyme activity.

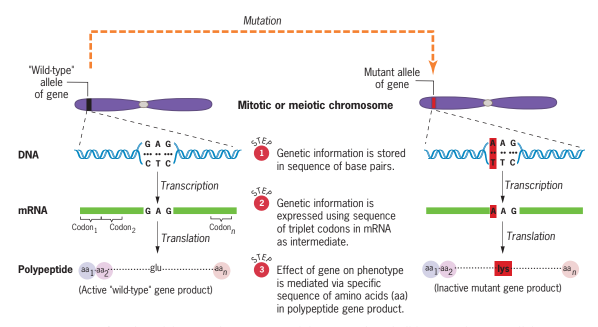


FIGURE 13.19 Overview of the mutation process and the expression of wild-type and mutant alleles. Mutations alter the sequences of nucleotide pairs in genes, which, in turn, cause changes in the amino acid sequences of the polypeptides encoded by these genes. A G:C base pair (top, left) has mutated to an A:T base pair (top, right). This mutation changes one mRNA codon from GAG to AAG and one amino acid in the polypeptide product from glutamic acid (glu) to lysine (lys). Such changes often yield nonfunctional gene products.

evolution. Thus mutations, which cause random changes in the highly adapted amino acid sequences, usually will produce less active or totally inactive products. You can make an analogy with any complex, carefully engineered machine such as a computer or an automobile. If you randomly modify an essential component, the machine is unlikely to perform as well as it did prior to the change. This view of mutation and the interaction between mutant and wild-type alleles fits with the observation that most mutations with recognizable phenotypic effects are recessive and deleterious.

EFFECTS OF MUTATIONS IN HUMAN GLOBIN GENES

Mutant human hemoglobins provide good illustrations of the deleterious effects of mutation. In Chapters 1 and 12, we discussed the structure of hemoglobin and the traumatic effects of one hemoglobin variant, sickle-cell hemoglobin. Recall that the major form of hemoglobin in adults (hemoglobin A) contains two identical *alpha* (α) *chains* and two identical *beta* (β) *chains*. Each α polypeptide consists of a specific sequence of 141 amino acids, whereas each β chain is 146 amino acids long. Because of similarities in their amino acid sequences, all the globin chains (and, thus, their structural genes) are believed to have evolved from a common progenitor.

Many different variants of adult hemoglobin have been identified in human populations, and several of them have severe phenotypic effects. Many of the variants were initially detected by their altered electrophoretic behavior (movement in an electric field due to charge differences—see Chapter 14). The hemoglobin variants provide an excellent illustration of the effects of mutation on the structures and functions of gene products and, ultimately, on the phenotypes of the affected individuals.

When the amino acid sequences of the β chains of hemoglobin A and the hemoglobin in patients with sickle-cell disease (hemoglobin S) were determined and compared, hemoglobin S was found to differ from hemoglobin A at only one position. The sixth amino acid from the amino terminus of the β chain of hemoglobin A is glutamic acid (a negatively charged amino acid). The β chain of hemoglobin S contains valine (no charge at neutral pH) at that position. The α chains of hemoglobin A and hemoglobin S are identical. Thus, the change of a single amino acid in one polypeptide can have severe effects on the phenotype.

In the case of hemoglobin S, the substitution of valine for glutamic acid at the sixth position in the β chain allows a new bond to form, which changes the conformation of the protein and leads to aggregation of hemoglobin molecules. This change results in the grossly abnormal (sickle) shape of the red blood cells. The mutational change in the *HBB*⁴ allele that gave rise to *HBB*^S was a substitution of a T:A base pair for an A:T base pair, with a T in the transcribed strand in the first case and an A in the transcribed strand in the second case (see Figure 1.9). This A:T \rightarrow T:A base-pair substitution was first predicted from protein sequence data and the known codon assignments, and was later verified by sequencing the *HBB*⁴ and *HBB*^S alleles.

Over 100 hemoglobin variants with amino acid changes in the β chain are known (see the Genomics on the Web questions at the end of the chapter). Most of them differ from the normal β chain of hemoglobin A by a single amino acid substitution. Some differ by two amino acids. Numerous variants of the α polypeptide also have been identified.

The hemoglobin examples show that mutation is a process in which changes in gene structure, often changes in one or a few base pairs, can cause changes in the amino acid sequences of the polypeptide gene products. These alterations in protein structure, in turn, cause changes in the phenotype that are recognized as mutant.

MUTATION IN HUMANS: BLOCKS IN METABOLIC PATHWAYS

In Chapter 4, we discussed the genetic control of metabolic pathways, in which each step in a pathway is catalyzed by an enzyme encoded by one or more genes. When mutations occur in such genes, they often cause metabolic blocks (see Figure 13.18) that lead to abnormal phenotypes. This picture of the genetic control of metabolism is valid for all living organisms, including humans (see On the Cutting Edge: Screening Eight-Cell Pre-Embryos for Tay-Sachs Mutations).

We can illustrate the effects of mutations on human metabolism by considering virtually any metabolic pathway. However, the metabolism of the aromatic amino acids phenylalanine and tyrosine provides an especially good example because some of the early studies of mutations in humans revealed blocks in this pathway (see the Chapter 4 Milestone in Genetics: Garrod's Inborn Errors of Metabolism on the Student Companion site). Phenylalanine and tyrosine are essential amino acids required for protein synthesis; they are not synthesized *de novo* in humans as they are in microorganisms. Thus, both amino acids must be obtained from dietary proteins.

The best-known inherited defect in phenylalanine-tyrosine metabolism is phenylketonuria, which is caused by the absence of phenylalanine hydroxylase, the enzyme that converts phenylalanine to tyrosine. Newborns with phenylketonuria, an autosomal recessive disease, develop severe mental retardation if not placed on a diet low in phenylalanine (see the Chapter 4 Milestone on the Student Companion site). The first inherited disorder in the phenylalanine-tyrosine metabolic pathway to be studied in humans was alkaptonuria, which is caused by autosomal recessive mutations that inactivate the enzyme homogentisic acid oxidase. Alkaptonuria played an important role in the evolution of the concept of the gene (see the Chapter 4 Milestone on the Student Companion site).

Two other inherited disorders are caused by mutations in genes encoding enzymes required for the catabolism of tyrosine; both are inherited as autosomal recessives. Tyrosinosis and tyrosinemia result from the lack of the enzymes tyrosine transaminase and *p*-hydroxyphenylpyruvic acid oxidase, respectively. Both enzymes are required to degrade tyrosine to CO_2 and H_2O . Tyrosinosis is very rare; only a few cases have been studied. Individuals with tyrosinosis show pronounced increases in tyrosine levels in their blood and urine and have various congenital abnormalities. Individuals with tyrosinemia have elevated levels of both tyrosine and *p*-hydroxyphenylpyruvic acid in their blood and urine. Most newborns with tyrosinemia die within six months after birth because of liver failure.

Albinism, the absence of pigmentation in the skin, hair, and eyes, results from a mutational block in the conversion of tyrosine to the dark pigment melanin. One type of albinism is caused by the absence of tyrosinase, the enzyme that catalyzes the

ON THE CUTTING EDGE

SCREENING EIGHT-CELL PRE-EMBRYOS FOR TAY-SACHS MUTATIONS

f all the inherited human disorders, Tay-Sachs disease is one of the most tragic. Infants homozygous for the mutant gene that causes Tay-Sachs disease are normal at birth. However, within a few months, they become hypersensitive to loud noises and develop a cherry-red spot on the retina of the eye. These early symptoms of the disease often go undetected by parents and physicians. At six months to one year after birth, Tay-Sachs children begin to undergo progressive neurological degeneration that rapidly leads to mental retardation, blindness, deafness, and general loss of control of body functions. By two years of age, they are usually totally paralyzed and develop chronic respiratory infections. Death commonly occurs at three to four years of age.

Tay-Sachs disease is caused by an autosomal recessive mutation in the *HEXA* gene, which encodes the enzyme hexosaminidase A. This mutation is rare in most populations. However, about 1 of 30 adults in the Ashkenazi Jewish population of Central Europe carries the mutant gene in the heterozygous state, and the disease occurs in about 1 of 3600 of their children. If two individuals from this Jewish population marry, the chance that both will carry the mutant gene is about 1 in 1000 (0.033 \times 0.033). If both parents are carriers, on average, one-fourth of their children will be homozygous for the mutant gene and develop Tay-Sachs disease.

Hexosaminidase acts on a complex lipid called ganglioside G_{M2} , cleaving it into a smaller ganglioside (G_{M3}) and *N*-acetyl-*D*-galactosamine, as shown in **Figure 1**. The function of ganglioside G_{M2} is to coat nerve cells, insulating them from events occurring in neighboring cells and thus speeding up the transmission of nerve impulses. In the absence of the enzyme that breaks it down, ganglioside G_{M2} accumulates and literally smothers nerve cells. This buildup of complex lipids on neurons blocks their action, leading to deterioration of the nervous system and eventually to paralysis.

Although Tay-Sachs disease was described by Warren Tay in 1881 and the biochemical basis has been known for over 25 years, there is still no effective treatment of this disorder. Whereas some

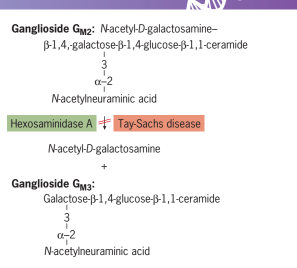


FIGURE 1 The metabolic defect in humans with Tay-Sachs disease.

inherited disorders can be treated by supplying the missing enzyme to patients, this won't work with Tay-Sachs disease because the enzyme cannot penetrate the barrier separating brain cells from the circulatory system. Moreover, somatic-cell gene therapy providing functional copies of the defective gene to somatic cells [Chapter 16]—is not yet possible because there is no established procedure for introducing genes into neurons.

Amniocentesis (Chapter 6) has been used extensively to detect the Tay-Sachs mutation during fetal development. More recently, a DNA test has been developed that permits the detection of the mutant gene using DNA from a single cell. This test can be used to screen eight-cell pre-embryos produced by *in vitro* fertilization for the Tay-Sachs mutation. One cell is used for the DNA test, and the other seven cells retain the capacity to develop into a normal embryo when implanted into the uterus of the mother. Only embryos that test normal—those not homozygous for the deadly Tay-Sachs gene—are implanted. This procedure allows heterozygous parents to have children without worrying about the birth of a child with Tay-Sachs disease.

first step in the synthesis of melanin from tyrosine. Other types of albinism result from blocks in subsequent steps in the conversion of tyrosine to melanin. Albinism is inherited as an autosomal recessive trait; heterozygotes usually have normal levels of pigmentation. Therefore, two albinos who have mutations in different genes will produce normally pigmented children.

Thus, studies of a single metabolic pathway, phenylalanine-tyrosine metabolism, have revealed five different inherited disorders, all caused by mutations in genes that control steps in this pathway. Similar examples of the genetic control of metabolism can be obtained by examining essentially any other metabolic pathway in humans.

CONDITIONAL LETHAL MUTATIONS: POWERFUL TOOLS FOR GENETIC STUDIES

Of all the mutations—from isoalleles to lethals—conditional lethal mutations are the most useful for genetic studies. These are mutations that are (1) lethal in one

environment, the *restrictive condition*, but are (2) viable in a second environment, the *permissive condition*. Conditional lethal mutations allow geneticists to identify and study mutations in essential genes that result in complete loss of gene-product activity even in haploid organisms. Mutants carrying conditional lethals can be propagated under permissive conditions, and information about the functions of the gene products can be inferred by studying the consequences of their absence under the restrictive conditions. Conditional lethal mutations have been used to investigate a vast array of biological processes from development to photosynthesis.

The three major classes of mutants with conditional lethal phenotypes are (1) auxotrophic mutants, (2) temperature-sensitive mutants, and (3) suppressorsensitive mutants. Auxotrophs are mutants that are unable to synthesize an essential metabolite (amino acid, purine, pyrimidine, vitamin, and so forth) that is synthesized by wild-type or *prototrophic* organisms of the same species. The auxotrophs will grow and reproduce when the metabolite is supplied in the medium (the permissive condition); they will not grow when the essential metabolite is absent (the restrictive condition). Temperature-sensitive mutants will grow at one temperature but not at another. Most temperature-sensitive mutants are heat-sensitive; however, some are cold-sensitive. The temperature sensitivity usually results from the increased heat or cold lability of the mutant gene product—for example, an enzyme that is active at low temperature but partially or totally inactive at higher temperatures. Occasionally, only the synthesis of the gene product is sensitive to temperature, and once synthesized, the mutant gene product may be as stable as the wild-type gene product. Suppressor-sensitive mutants are viable when a second genetic factor, a suppressor, is present, but they are nonviable in the absence of the suppressor. The suppressor gene may correct or compensate for the defect in phenotype that is caused by the suppressor-sensitive mutation, or it may cause the gene product altered by the mutation to be nonessential. We have discussed one class of suppressor-sensitive mutations, the *amber* mutations, in Chapter 12.

Now, let's briefly consider how conditional lethal mutations can be used to investigate biological processes—to dissect biological processes into their individual parts or steps. Let's begin with a simple biosynthetic pathway:

Gene A

$$\downarrow$$
 Gene B
 \downarrow \downarrow \downarrow \downarrow \downarrow
Enzyme A
Precursor X \longrightarrow Intermediate Y \longrightarrow Product Z

Intermediate Y is produced from precursor X by the action of enzyme A, the product of gene A, but intermediate Y may be rapidly converted to product Z by enzyme B, the product of gene B. If so, intermediate Y may be present in minute quantities and be difficult to isolate and characterize. However, in a mutant organism that has a mutation in gene B, resulting in the synthesis of either an inactive form of enzyme B or no enzyme B, intermediate Y may accumulate to much higher concentrations, facilitating its isolation and characterization. Similarly, a mutation in gene A may aid in the identification of precursor X. In this way, the sequence of steps in a given metabolic pathway can often be determined.

Morphogenesis in living organisms occurs in part by the sequential addition of proteins to macromolecular structures to produce the final three-dimensional conformations, and the sequence of protein additions can often be determined by isolating and studying mutant organisms with defects in the genes encoding the proteins involved. Because an appropriate mutation will eliminate the activity of a single polypeptide, mutations provide a powerful tool with which to dissect biological processes—to break the processes down into individual steps.

The resolving power of mutational dissection of biological processes has been elegantly documented by the research of Robert Edgar, Jonathan King, William Wood, and colleagues, who worked out the complete pathway of morphogenesis for bacteriophage T4. This complex process involves the products of about 50 of the roughly 200 genes in the T4 genome. Each gene encodes a structural protein of the

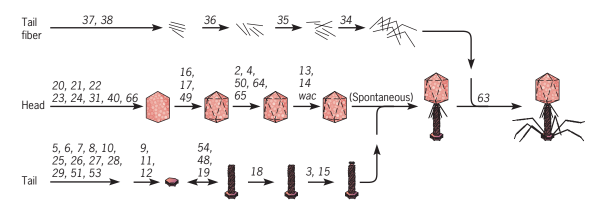


FIGURE 13.20 Abbreviated pathway of morphogenesis in bacteriophage T4. The head, the tail, and the tail fibers are produced via separate branches of the pathway and are then joined in the final stages of morphogenesis. The numbers identify the T4 genes whose products are required at each step in the pathway. The sequences of early steps in head and tail formation are known but are omitted here to keep the diagram concise.

virus or an enzyme that catalyzes one or more steps in the morphogenetic pathway. By (1) isolating mutant strains of phage T4 with temperature-sensitive and suppressorsensitive conditional lethal mutations in each of the approximately 50 genes, and (2) using electron microscopy and biochemical techniques to analyze the structures that accumulate when these mutant strains are grown under the restrictive conditions, Edgar, King, Wood, and coworkers established the complete pathway of phage T4 morphogenesis (**■** Figure 13.20).

Many other biological processes also have been successfully dissected by mutational studies. Examples include the photosynthetic electron transport chains in plants and pathways of nitrogen fixation in bacteria. Currently, mutational dissection is yielding new insights into the processes of differentiation and development in higher plants and animals (Chapter 20). Researchers are also using mutations to dissect behavior and learning in *Drosophila*. In principle, scientists should be able to use mutations to dissect any biological process. Every gene can mutate to a nonfunctional state. Thus, mutational dissection of biological processes is limited only by the ingenuity of researchers in identifying mutations of the desired types.

KEY POINTS

- The effects of mutations on the phenotypes of living organisms range from minor to lethal changes.
 - Most mutations exert their effects on the phenotype by altering the amino acid sequences of polypeptides, the primary gene products.
 - The mutant polypeptides, in turn, cause blocks in metabolic pathways.
 - Conditional lethal mutations provide powerful tools with which to dissect biological processes.

Assigning Mutations to Genes by the Complementation Test

The complementation or *trans* test can be used to determine whether two mutations are located in the same gene or in two different genes.

With the emergence of the one gene–one polypeptide concept (Chapter 12), scientists could define the gene biochemically, but they had no genetic tool to use in determining whether two mutations were in the same or different genes. This deficiency was

resolved in the 1940s when Edward Lewis developed the complementation test for functional allelism. Before we discuss Lewis's work, we need to define some new terms. A double heterozygote, which carries two mutations and their wild-type alleles, that is, m_1 and m_1^+ along with m_2 and m_2^+ , can exist in either of two arrangements (Figure 13.21). When the two mutations are on the same chromosome, the arrangement is called the **coupling** or **cis configuration**, and a heterozygote with this genotype is called a **cis heterozygote** (**I Figure 13.21***a*). When the two mutations are on different chromosomes, the arrangement is called the **repulsion** or **trans configuration**. An organism with this genotype is a **trans heterozygote** (**I Figure 13.21***b*).

In the 1940s and 1950s, Lewis observed that fruit flies carrying certain mutants in the *cis* and *trans* configurations had different phenotypes. We will examine his results with two recessive eye color mutations *white* (*w*) and *apricot* (*apr*). Flies that are homozygous for the X-linked mutations *apr* and *w* have apricot-colored eyes and white eyes, respectively, in contrast to the red eyes of wild-type *Drosophila*. When Lewis produced *cis* heterozygotes with the genotype *apr* $w/apr^+ w^+$, they had red eyes just like wild-type flies (**■** Figure 13.22a). When he constructed *trans* heterozygotes with genotype *apr* $w^+/apr^+ w$, they had light apricot-colored eyes (**■** Figure 13.22b). Both genotypes contained the same mutant and wild-type genetic information but in different arrangements. When organisms that contain the same genetic markers, but in different arrangements, have different phenotypes, the markers are said to exhibit position effects. The type of position effect that Lewis observed is called a *cis-trans* position effect.

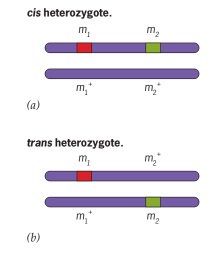
Lewis's discovery of *cis-trans* position effects led to the development of the **complementation test** or **trans test** for functional allelism. This test allows geneticists to determine whether mutations that produce the same or similar phenotypes are in the same gene or in different genes. The mutations must be tested pairwise by determining the phenotypes of *trans* heterozygotes. That is, *trans* heterozygotes must be constructed for each pair of mutations and examined to determine whether they have mutant or wild-type phenotypes.

Ideally, the complementation or *trans* test should be done in conjunction with the *cis* test—a control that is often omitted. *Cis* tests are performed by constructing *cis* heterozygotes with each pair of mutations being studied and determining whether the

heterozygotes have mutant or wild-type phenotypes. Together, the complementation or *trans* test and the *cis* test are referred to as the *cis-trans* test. Each *cis* heterozygote, which contains one wild-type chromosome, should have the wild-type phenotype whether the mutations are in the same gene or in two different genes. Indeed, the *cis* heterozygote must have the wild-type phenotype for the results of the *trans* test to be valid. If the *cis* heterozygote has the mutant phenotype, the *trans* test cannot be used to determine whether the two mutations are in the same gene. Thus, the *trans* test cannot be used to assign dominant mutations to genes.

With diploid organisms, *trans* heterozygotes are produced simply by crossing organisms that are homozygous for each of the mutations of interest. With viruses, *trans* heterozygotes are produced by simultaneously infecting host cells with two different mutants. Regardless of how the *trans* heterozygotes are constructed, the results of the *trans* or complementation tests provide the same information.

- **1.** If a *trans* heterozygote has the mutant phenotype (the phenotype of organisms or cells homozygous for either one of the two mutations), then the two mutations are in the same unit of function, the same gene.
- **2.** If the *trans* heterozygote has the wild-type phenotype, then the two mutations are in two different units of function, two different genes.



■ **FIGURE 13.21** The arrangement of genetic markers in *cis* and *trans* heterozygotes.

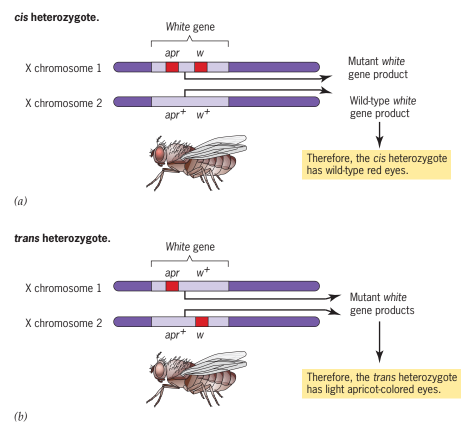


FIGURE 13.22 The *cis-trans* position effect observed by Edward Lewis with the *apr* and *w* mutations of *Drosophila*.

Solve It!

How Can You Assign Mutations to Genes?

Four independently isolated mutants of *E. coli*, all of which are unable to grow in the absence of tryptophan (tryptophan auxotrophs), were examined in all possible *cis* and *trans* heterozygotes (partial diploids). All of the *cis* heterozygotes were able to grow in the absence of tryptophan. The *trans* heterozygotes yielded two different responses: some of them grew in the absence of tryptophan; others did not. The experimental results, using "+" to indicate growth and "0" to indicate no growth, are given in the following table.

Growth of *Trans* Heterozygotes on Medium Lacking Tryptophan

		/ 1		
Mutant:	1	<u>2</u>	<u>3</u>	4
4	+	0	+	0
3	0	+	0	
2	+	0		
1	0			

How many genes are defined by these four mutations? Which mutant strains carry mutations in the same gene(s)?

To see the solution to this problem, visit the Student Companion site. When the two mutations present in a *trans* heterozygote are both in the same gene, both chromosomes will carry defective copies of that gene. As a result, the *trans* heterozygote will contain only nonfunctional products of the gene involved and will have a mutant phenotype.

When a *trans* heterozygote has the wild-type phenotype, the two mutations are said to exhibit complementation or to complement each other and are located in different genes. In this case, the *trans* heterozygote will contain functional products of both genes and, therefore, will exhibit the wild-type phenotype.

Let's illustrate this concept of complementation by examining *trans* tests performed with some well-characterized *amber* mutations of bacteriophage T4. *Amber* mutations in essential genes are conditional lethal mutations (see the section of this chapter entitled Conditional Lethal Mutations: Powerful Tools for Genetic Studies). When present in restrictive host bacteria such as *E. coli* strain B, their phenotype is lethality—that is, no progeny are produced. However, when present in a permissive host cell such as *E. coli* strain CR63, their phenotype is wild-type—that is, about 300 progeny phage are produced per infected cell. With conditional lethal mutations, the distinction between the mutant and wild-type phenotypes is maximal: lethality versus normal growth.

Amber mutations produce translation-termination triplets within the coding regions of genes (see Figure 12.24). As a result, the products of the mutant genes are truncated polypeptides, which are almost always totally nonfunctional. Therefore, complementation tests performed with *amber* mutations are usually unambiguous.

Two of the three *amber* mutations that we will consider (*am*B17 and *am*H32) are located in gene 23, which encodes the major structural protein of the phage head; the other mutation (*am*E18) is in gene 18, which specifies the major structural protein of the phage tail (see Figure 13.20).

In a *trans* heterozygote containing mutations *am*B17 (head gene) and *am*E18 (tail gene), wild-type copies of both genes are present, producing functional head and tail proteins (**■** Figure 13.23*a*). As a result, this *trans* heterozygote exhibits the wild-type phenotype (a normal yield of progeny phage). Mutations *am*B17 and *am*E18 complement one another because they are located in two different genes.

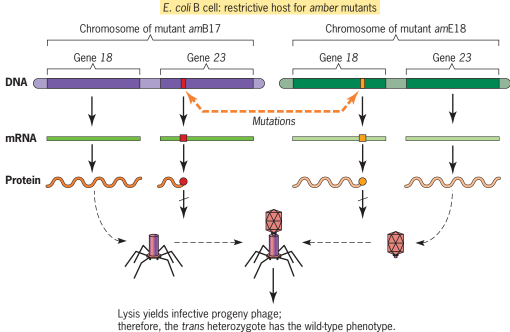
In a *trans* heterozygote containing mutations *am*B17 and *am*H32 (both in head gene 23), on the other hand, no functional gene 23 head protein is made (**\blacksquare Figure 13.23b**). Thus, this *trans* heterozygote has the mutant phenotype (lethality, or no progeny). Mutations *am*B17 and *am*H32 do not complement one another because they are both located in the same gene.

By using the complementation test, a researcher can determine whether independent mutations that result in the same phenotype are in the same gene or different genes. Ten *amber* mutations, for example, could all be in one gene, or one in one gene and nine in a second gene, and so on, with the final possibility being that each mutation could be in a separate gene. In the last case, the 10 mutations would identify 10 different genes. To test your comprehension of this concept, try Solve It: How Can You Assign Mutations to Genes?

Complementation is the result of the functionality of the gene *products* specified by chromosomes carrying two different mutations when they are present in the same protoplasm. Complementation does not depend on recombination of the two chromosomes. *Complementation, or the lack of it, is assessed by the phenotype (wildtype or mutant) of each trans heterozygote. Recombination, in contrast, involves the actual breakage of chromosomes and reunion of parts to produce wild-type and double-mutant chromosomes.*

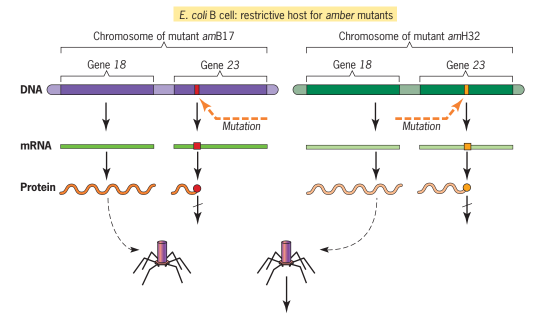
Note that the complementation or *trans* test defines *functional allelism*, that is, whether two mutations are in the same gene or in two different genes. Two mutations that do not complement each other in a *trans* heterozygote are in the same unit of function, the same gene. *Structural allelism*—the occurrence of two or more different mutations at the same site on a chromosome—is determined by the recombination test. Two mutations that do not recombine are structurally allelic; the mutations either occur at the same site or overlap a common site.

Complementation between mutations amB17 and amE18.





Lack of complementation between mutations amB17 and amH32.



No phage heads and, thus, no infective progeny phage are produced in the infected cell; therefore, the *trans* heterozygote has the mutant phenotype.

(b)

■ FIGURE 13.23 Complementation and noncomplementation in *trans* heterozygotes. (a) Complementation between mutation *am*B17 in gene 23, which encodes the major structural protein of the phage T4 head, and mutation *am*E18 in gene 18, which encodes the major structural protein of the phage tail. Phage heads and tails are both synthesized in the cell, with the result that infective progeny phage are produced. (b) When the *trans* heterozygote contains two mutations (*am*B17 and *am*H32) in gene 23, no heads are produced, and no infective progeny phage can be assembled.

Mutations that are both structurally and functionally allelic are called **homoalleles**; they do not complement or recombine with each other. Mutant homoalleles have defects at the same site, or defects that overlap a common site, in the same gene. Mutations that are functionally allelic, but structurally nonallelic, are called **heteroalleles**; they recombine with each other but do not complement one another. Mutant heteroalleles occur at different sites but within the same gene.

KEY POINTS • *The complementation test can be used to determine whether two mutations that produce the same phenotype are in the same gene or in two different genes.*

• Homoalleles are mutations at the same site in a gene; heteroalleles are mutations at different sites in a gene.

Screening Chemicals for Mutagenicity: The Ames Test

The Ames test provides a simple and inexpensive method for detecting the mutagenicity of chemicals.

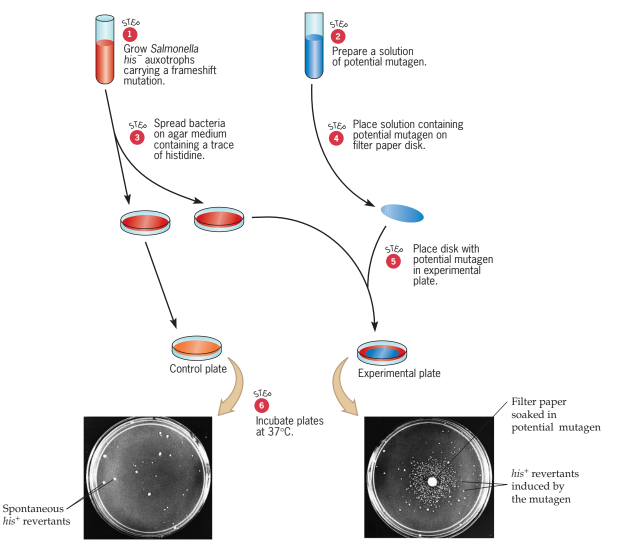
Mutagenic agents are also **carcinogens**; that is, they induce cancers. The one characteristic that the hundreds of types of cancer have in common is that the malignant cells continue to divide after cell division would have

stopped in normal cells. Of course, cell division, like all other biological processes, is under genetic control. Specific genes encode products that regulate cell division in response to intracellular, intercellular, and environmental signals. When these genes mutate to nonfunctional states, uncontrolled cell division sometimes results. Clearly, we wish to avoid being exposed to mutagenic and carcinogenic agents. However, our technological society depends on the extensive use of chemicals in both industry and agriculture. Hundreds of new chemicals are produced each year, and the mutagenicity and carcinogenicity of these chemicals need to be evaluated before their use becomes widespread.

Traditionally, the carcinogenicity of chemicals has been tested on rodents, usually newborn mice. These studies involve feeding or injecting the substance being tested and subsequently examining the animals for tumors. Mutagenicity tests have been done in a similar fashion. However, because mutation is a low-frequency event and because maintaining large populations of mice is an expensive undertaking, the tests have been relatively insensitive; that is, low levels of mutagenicity could not be detected.

Bruce Ames and his associates developed sensitive techniques that allow the mutagenicity of large numbers of chemicals to be tested quickly at relatively low cost. Ames and coworkers constructed auxotrophic strains of the bacterium *Salmonella typhimurium* carrying various types of mutations—transitions, transversions, and frameshifts—in genes required for the biosynthesis of the amino acid histidine. They monitored the reversion of these auxotrophic mutants to prototrophy by placing a known number of mutant bacteria on medium lacking histidine and scoring the number of colonies produced by prototrophic revertants. Because some chemicals are mutagenic only to replicating DNA, they added a small amount of histidine—enough to allow a few cell divisions but not the formation of visible colonies—to the medium. They measured the mutagenicity of a chemical by comparing the frequency of reversion in its presence with the spontaneous reversion frequency (**■** Figure 13.24). They assessed its ability to induce different types of mutations by using a set of tester strains that carry different types of mutations—one strain with a transition, one with a frameshift mutation, and so forth.

Over a period of several years during which they tested thousands of different chemicals, Ames and his colleagues observed a greater than 90 percent correlation



■ FIGURE 13.24 The Ames test for mutagenicity. The medium in each petri dish contains a trace of histidine and a known number of *his*⁻ cells of a specific *Salmonella typhimurium* "tester strain" harboring a frameshift mutation. The control plate shown on the left provides an estimate of the frequency of spontaneous reversion of this particular tester strain. The experimental plate on the right shows the frequency of reversion induced by the potential mutagen, in this case, the carcinogen 2-aminofluorene.

between the mutagenicity and the carcinogenicity of the substances tested. Initially, they found several potent carcinogens to be nonmutagenic to the tester strains. Subsequently, they discovered that many of these carcinogens are metabolized to strongly mutagenic derivatives in eukaryotic cells. Thus, Ames and his associates added a rat liver extract to their assay systems in an attempt to detect the mutagenicity of metabolic derivatives of the substances being tested. Coupling of the rat liver activation system to the microbial mutagenicity tests expanded the utility of the system considerably. For example, nitrates (found in charred meats) are not themselves mutagenic or carcinogenic. However, in eukaryotic cells, nitrates are converted to nitrosamines, which are highly mutagenic and carcinogenic. Ames's mutagenicity tests demonstrated the presence of frameshift mutagens in several components of chemically fractionated cigarette smoke condensates. In some cases, activation by the liver extract preparation was required for mutagenicity; in other cases, activation was not required. The Ames test provides a rapid, inexpensive, and sensitive procedure for testing the mutagenicity of chemicals. Since mutagenic chemicals are also carcinogens, the Ames test can be used to identify chemicals that have a high likelihood of being carcinogenic.

KEY POINT

Bruce Ames and coworkers developed an inexpensive and sensitive method for testing the mutagenicity of chemicals with histidine auxotrophic mutants of Salmonella.

DNA Repair Mechanisms

Living organisms contain many enzymes that scan their DNA for damage and initiate repair processes when damage is detected. The multiplicity of repair mechanisms that have evolved in organisms ranging from bacteria to humans emphatically documents the importance of keeping mutation at a tolerable level. For example, *E. coli* cells possess five well-characterized mechanisms for the repair of defects

in DNA: (1) light-dependent repair or photoreactivation, (2) excision repair, (3) mismatch repair, (4) postreplication repair, and (5) the error-prone repair system (SOS response). Moreover, there are at least two different types of excision repair, and the excision repair pathways can be initiated by several different enzymes, each acting on a specific kind of damage in DNA. Mammals seem to possess all of the repair mechanisms found in *E. coli* except photoreactivation. Because most mammalian cells do not have access to light, photoreactivation would be of relatively little value to them.

The importance of DNA repair pathways to human health is clear. Inherited disorders such as xeroderma pigmentosum, which was discussed at the beginning of this chapter, vividly document the serious consequences of defects in DNA repair. We discuss some of these inherited disorders in a subsequent section of this chapter.

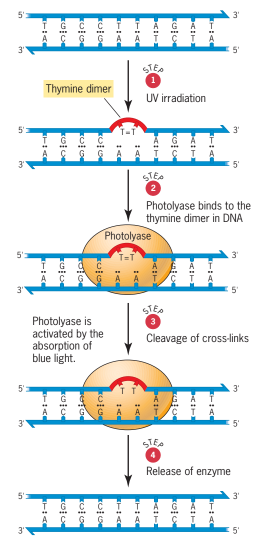
LIGHT-DEPENDENT REPAIR

Light-dependent repair or photoreactivation of DNA in bacteria is carried out by a lightactivated enzyme called DNA photolyase. When DNA is exposed to ultraviolet light, thymine dimers are produced by covalent cross-linkages between adjacent thymine residues (see Figure 13.12b). DNA photolyase recognizes and binds to thymine dimers in DNA, and uses light energy to cleave the covalent cross-links (**Figure 13.25**). Photolyase will bind to thymine dimers in DNA in the dark, but it cannot catalyze cleavage of the bonds joining the thymine moieties without energy derived from visible light, specifically light within the blue region of the spectrum. Photolyase also splits cytosine dimers and cytosine-thymine dimers. Thus, when ultraviolet light is used to induce mutations in bacteria, the irradiated cells are grown in the dark for a few generations to maximize the mutation frequency.

EXCISION REPAIR

Excision repair of damaged DNA involves at least three steps. In step 1, a DNA repair endonuclease or endonuclease-containing enzyme complex recognizes, binds to, and excises the damaged base or bases in DNA. In step 2, a DNA polymerase fills in the gap by using the undamaged complementary strand of DNA as template. In step 3, the enzyme DNA ligase seals the break left by DNA polymerase to complete the repair process. There are two major types of excision repair: **base excision repair** systems remove abnormal or chemically modified bases from DNA, whereas **nucleotide excision repair** pathways remove larger defects like thymine dimers. Both excision pathways are operative in the dark, and both occur by very similar mechanisms in *E. coli* and humans.

Base excision repair (**Figure 13.26**) can be initiated by any of a group of enzymes called DNA glycosylases that recognize abnormal bases in DNA. Each glycosylase recognizes a specific type of altered base, such as deaminated bases, oxidized bases, and so on (step 2). The glycosylases cleave the glycosidic bond between the abnormal base and 2-deoxyribose, creating apurinic or apyrimidinic sites (AP sites) with missing bases (step 3). AP sites are recognized by enzymes called AP endonucleases, which act together with phosphodiesterases to excise the sugar-phosphate groups



■ FIGURE 13.25 Cleavage of thymine dimer cross-links by light-activated photolyase. The arrows indicate the opposite polarity of the complementary strands of DNA. at these sites (step 4). DNA polymerase then replaces the missing nucleotide according to the specifications of the complementary strand (step 5), and DNA ligase seals the nick (step 6).

Nucleotide excision repair removes larger lesions like thymine dimers and bases with bulky side-groups from DNA. In nucleotide excision repair, a unique excision nuclease activity produces cuts on either side of the damaged nucleotide(s) and excises an oligonucleotide containing the damaged base(s). This nuclease is called an **excinuclease** to distinguish it from the endonucleases and exonucleases that play other roles in DNA metabolism.

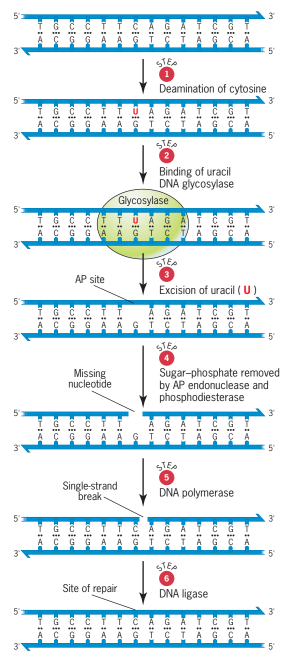
The *E. coli* nucleotide excision repair pathway is shown in Figure 13.27. In *E. coli*, excinuclease activity requires the products of three genes, *uvrA*, *uvrB*, and *uvrC* (designated *uvr* for *UV* repair). A trimeric protein containing two UvrA polypeptides and one UvrB polypeptide recognizes the defect in DNA, binds to it, and uses energy from ATP to bend the DNA at the damaged site. The UvrA dimer is then released, and the UvrC protein binds to the UvrB/DNA complex. The UvrC protein cleaves the fourth or fifth phosphodiester bond from the damaged nucleotide(s) on the 3' side and the eighth phosphodiester linkage from the damage on the 5' side. The *uvrD* gene product, DNA helicase II, releases the excised dodecamer. In the last two steps of the pathway, DNA polymerase I fills in the gap, and DNA ligase seals the remaining nick in the DNA molecule.

Nucleotide excision repair in humans occurs through a pathway similar to the one in *E. coli*, but it involves about four times as many proteins. In humans, the excinuclease activity contains 15 polypeptides. Protein XPA (for *x*eroderma *p*igmentosum protein *A*) recognizes and binds to the damaged nucleotide(s) in DNA. It then recruits the other proteins required for excinuclease activity. In humans, the excised oligomer is 24 to 32 nucleotides long rather than the 12-mer removed in *E. coli*. The gap is filled in by either DNA polymerase δ or ε in humans, and DNA ligase completes the job.

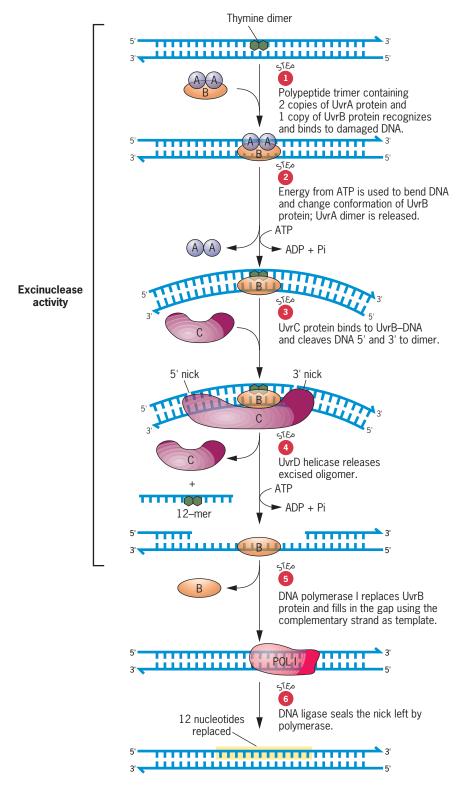
OTHER DNA REPAIR MECHANISMS

During the last few years, research on DNA repair mechanisms has demonstrated the presence of an army of DNA repair enzymes that constantly scan DNA for damage ranging from the presence of thymine dimers induced by ultraviolet light to modifications too diverse and numerous to describe here. New results of this work have shown that several previously unknown DNA polymerases play critical roles in various DNA repair processes. Detailed discussions of these important DNA repair processes are beyond the scope of this text. Nevertheless, the importance of these repair mechanisms cannot be overstated. What is more important to the survival of a species than maintaining the integrity of its genetic blueprint?

In Chapter 10, we examined the mechanism by which the $3' \rightarrow 5'$ exonuclease activity built into DNA polymerases proofreads DNA strands during their synthesis, removing any mismatched nucleotides at the 3' termini of growing strands. Another postreplication DNA repair pathway, mismatch repair, provides a backup to this replicative proofreading by correcting mismatched nucleotides remaining in DNA after replication. Mismatches often involve the normal four bases in DNA. For example, a T may be mispaired with a G. Because both T and G are normal components of DNA, mismatch repair systems need some way to determine whether the T or the G is the correct base at a given site. The repair system makes this distinction by identifying the template strand, which contains the original nucleotide sequence, and the newly synthesized strand, which contains the misincorporated base (the error). In bacteria, this distinction can be made based on the pattern of methylation in newly replicated DNA. In E. coli, the A in GATC sequences is methylated subsequent to its synthesis. Thus, a time interval occurs during which the template strand is methylated, and the newly synthesized strand is unmethylated. The mismatch repair system uses this difference in methylation state to excise the mismatched nucleotide in the nascent



■ FIGURE 13.26 Repair of DNA by the base excision pathway. Base excision repair may be initiated by any one of several different DNA glycosylases. In the example shown, uracil DNA glycosylase starts the repair process.



■ FIGURE 13.27 Repair of DNA by the nucleotide excision pathway in *E. coli*. The excinuclease (excision nuclease) activity requires the products of three genes *uvrA*, *uvrB*, and *uvrC*. Nucleotide excision occurs by a similar pathway in humans, except that many more proteins are involved and a 24-to-32-nucleotide-long oligomer is excised.

strand and replace it with the correct nucleotide by using the methylated parental strand of DNA as template.

In *E. coli*, mismatch repair requires the products of four genes, *mutH*, *mutL*, *mutS*, and *mutU*(=uvrD). The MutS protein recognizes mismatches and binds to them to initiate the repair process. MutH and MutL proteins then join the complex. MutH contains a GATC-specific endonuclease activity that cleaves the unmethylated strand at hemimethylated (that is, half methylated) GATC sites either 5' or 3' to the mismatch. The incision sites may be 1000 nucleotide pairs or more from the mismatch. The subsequent excision process requires MutS, MutL, DNA helicase II (MutU), and an appropriate exonuclease. If the incision occurs at a GATC sequence 5' to the mismatch, a 5' \rightarrow 3' exonuclease like *E. coli* exonuclease VII is required. If the incision occurs 3' to the mismatch, a $3' \rightarrow 5'$ exonuclease activity like that of *E. coli* exonuclease I is needed. After the excision process has removed the mismatched nucleotide from the unmethylated strand, DNA polymerase III fills in the large-up to 1000 bp—gap, and DNA ligase seals the nick.

Homologues of the *E. coli* MutS and MutL proteins have been identified in fungi, plants, and mammals—an indication that similar mismatch repair pathways occur in eukaryotes. In fact, mismatch excision has been demonstrated *in vitro* with nuclear extracts prepared from human cells. Thus, mismatch repair is probably a universal or nearly universal mechanism for safeguarding the integrity of genetic information stored in double-stranded DNA.

In E. coli, light-dependent repair, excision repair, and mismatch repair can be eliminated by mutations in the *phr* (*photoreactivation*), *uvr*; and *mut* genes, respectively. In mutants deficient in more than one of these repair mechanisms, still another DNA repair system, called *postreplication repair*, is operative. When DNA polymerase III encounters a thymine dimer in a template strand, its progress is blocked. DNA polymerase restarts DNA synthesis at some position past the dimer, leaving a gap in the nascent strand opposite the dimer in the template strand. At this point, the original nucleotide sequence has been lost from both strands of the progeny double helix. The damaged DNA molecule is repaired by a recombination-dependent repair process mediated by the E. coli recA gene product. The RecA protein, which is required for homologous recombination, stimulates the exchange of single strands between homologous double helices. During postreplication repair, the RecA protein binds to the single strand of DNA at the gap and mediates pairing with the homologous segment of the sister double helix. The gap opposite

the dimer is filled with the homologous DNA strand from the sister DNA molecule. The resulting gap in the sister double helix is filled in by DNA polymerase, and the nick is sealed by DNA ligase. The thymine dimer remains in the template strand of the original progeny DNA molecule, but the complementary strand is now intact. If the thymine dimer is not removed by the nucleotide excision repair system, this postreplication repair must be repeated after each round of DNA replication.

The DNA repair systems described so far are quite accurate. However, when the DNA of *E. coli* cells is heavily damaged by mutagenic agents such as UV light, the cells take some drastic steps in their attempt to survive. They go through a so-called **SOS response**, during which a whole battery of DNA repair, recombination, and replication proteins are synthesized. Two of these proteins, encoded by the *umuC* and *umuD* (*UV mutable*) genes, are subunits of DNA polymerase V, an enzyme that catalyzes the replication of DNA in damaged regions of the chromosome—regions where replication by DNA polymerase III is blocked. DNA polymerase V allows replication to proceed across damaged segments of template strands, even though the nucleotide sequences in the damaged region cannot be replicated accurately. This *error-prone repair* system eliminates gaps in the newly synthesized strands opposite damaged nucleotides in the template strands but, in so doing, increases the frequency of replication errors.

The mechanism by which the SOS system is induced by DNA damage has been worked out in considerable detail. Two key regulatory proteins—LexA and RecA control the SOS response. Both are synthesized at low background levels in the cell in the absence of damaged DNA. Under this condition, LexA binds to the DNA regions that regulate the transcription of the genes that are induced during the SOS response and keeps their expression levels low. When cells are exposed to ultraviolet light or other agents that cause DNA damage, the RecA protein binds to single-stranded regions of DNA caused by the inability of DNA polymerase III to replicate the damaged regions. The interaction of RecA with DNA activates RecA, which then stimulates LexA to inactivate itself by self-cleavage. With LexA inactive, the level of expression of the SOS genes—including *recA*, *lexA*, *umuC*, *umuD*, and others increases and the error-prone repair system is activated.

The SOS response appears to be a somewhat desperate and risky attempt to escape the lethal effects of heavily damaged DNA. When the error-prone repair system is operative, mutation rates increase sharply.

Recent research on DNA repair mechanisms indicates that many new repair processes remain to be elucidated. During the last few years, several new DNA polymerases that have unique roles in DNA repair have been characterized. The results of these studies suggest that we have much to learn about the mechanisms that safeguard the integrity of our genetic information.

- Multiple DNA repair systems have evolved to safeguard the integrity of genetic information in living organisms.
- **KEY POINTS**

• Each repair pathway corrects a specific type of damage in DNA.

Inherited Human Diseases with Defects in DNA Repair

As we discussed at the beginning of this chapter, individuals with xeroderma pigmentosum (XP) are extremely sensitive to sunlight. Exposure to sunlight results in a high frequency of skin cancer in XP

Several inherited human disorders result from defects in DNA repair pathways.

patients (Figure 13.28). The cells of individuals with XP are deficient in the repair of UV-induced damage to DNA, such as thymine dimers. The XP syndrome can result from defects in any of at least eight different genes. The products of seven of these genes, *XPA*, *XPB*, *XPC*, *XPD*, *XPE*, *XPF*, and *XPG*, are required for nucleotide excision repair (Table 13.1). They have been purified and shown to be essential for excinuclease activity. Since excinuclease activity in humans requires 15 polypeptides,



FIGURE 13.28 Phenotypic effects of the inherited disease xeroderma pigmentosum. Individuals with this malignant disease develop extensive skin tumors after exposure to sunlight.

the list of *XP* genes will probably expand in the future. Two other human disorders, Cockayne syndrome and trichothiodystrophy, also result from defects in nucleotide excision repair. Individuals with Cockayne syndrome exhibit retarded growth and mental skills, but not increased rates of skin cancer. Patients with trichothiodystrophy have short stature, brittle hair, and scaly skin; they also have underdeveloped mental abilities. Individuals with either Cockayne syndrome or trichothiodystrophy are defective in a type of excision repair that is coupled to transcription. However, details of this transcription-coupled repair process are still being worked out.

In addition to the damage to skin cells, some individuals with XP develop neurological abnormalities, which appear to result from the premature death of nerve cells. This effect on the very long-lived nerve cells may have interesting implications with respect to the causes of aging. One theory is that aging results from the accumulation of somatic mutations. If so, a defective repair system would be expected to speed up the aging process, and this appears to be the case with the nerve cells of XP patients. However, at present, there is little evidence linking somatic mutation to senescence.

Hereditary nonpolyposis colon cancer (also called Lynch syndrome) is known to result from inherited defects in the DNA mismatch repair pathway. It can be caused by mutations in at least seven different genes, five of which are listed in **Table 13.1**. Several of these genes are homologues of *E. coli* and *S. cerevisiae* mismatch repair genes. Thus, the mismatch repair pathway of humans is similar to those in bacteria and fungi. This type of colon cancer occurs in about one of every 200 people, so it is a common type of cancer. Once we understand the inherited defects better, perhaps we will be able to develop effective methods of treating these cancers other than surgery, chemotherapy, and radiotherapy.

Ataxia-telangiectasia, Fanconi anemia, Bloom syndrome, Werner syndrome, Rothmund-Thomson syndrome, and Nijmegan breakage syndrome are six other inherited diseases in humans associated with known defects in DNA metabolism. All six disorders exhibit autosomal recessive patterns of inheritance, and all result in a high risk of malignancy, especially leukemia in the case of ataxia-telangiectasia and Fanconi anemia. Cells of patients with ataxia-telangiectasia exhibit an abnormal sensitivity to ionizing radiation, suggesting a defect in the repair of radiation-induced DNA damage.

TABLE 13.1

Inherited Human Diseases Caused by Defects in DNA Repair

inner neu numan Diseases Causeu by Delects in DNA Repair									
Inherited Disorder	Gene	Chromosome	Function of Product	Major Symptoms					
1. Xeroderma pigmentosum	XPA XPB	9 2	DNA-damage-recognition protein $3' \rightarrow 5'$ helicase	UV sensitivity, early onset skin cancers, neurological					
	XPC	3	DNA-damage-recognition protein	disorders					
	XPD	19	$5' \rightarrow 3'$ helicase						
	XPE	11	DNA-damage-recognition protein						
	XPF	16	Nuclease, 3' incision						
	XPG	13	Nuclease, 5' incision						
	XPV	6	Translesion DNA polymerase η						
2. Trichothiodystrophy	TTDA	6	Basal transcription factor IIH	UV sensitivity, neurological					
	XPB	2	3′ → 5′ helicase	disorders, mental retardation					
	XPD	19	$5' \rightarrow 3'$ helicase						
3.Cockayne syndrome	CSA	5	DNA excision repair protein	UV sensitivity, neurological					
	CSB	10	DNA excision repair protein	and developmental disorders, premature aging					
4. Ataxia-telangiectasia	ATM	11	Serine/threonine kinase	Radiation sensitivity, chromosome instability, early onset progressive neurodegeneration, cancer prone					
 Nonpolyposis colon cancer (Lynch syndrome) 	MSH2	2	DNA mismatch recognition protein (like <i>E. coli</i> MutS)	High risk of familial colon cancer					
	MLH1	3	Homolog of <i>E. coli</i> mismatch repair protein MutL						
	MSH6	2	MutS homolog 6						
	PMS2	7	Endonuclease PMS2						
	PMS1	2	Homolog of yeast mismatch repair protein						
6. Fanconi anemia	FA (8 genes, A-H, on 5 different chromosomes)			Sensitivity to DNA-cross-linking agents, chromosome instability, cancer prone					
7. Bloom syndrome	BLM	15	BLM RecQ helicase	Chromosome instability, mental retardation, cancer prone					
8. Werner syndrome	WRN	8	WRN RecQ helicase	Chromosome instability, progressive neurodegeneration, cancer prone					
9. Rothmund-Thomson syndrome	RECQL4	8	RecQ helicase L4	Chromosome instability, mental retardation, cancer prone					
10. Nijmegan breakage syndrome	NBSI	8	DNA-double-strand-break- recognition protein	, Chromosome instability, microcephaly (small cranium), cancer prone					

Cells of individuals with Fanconi anemia are impaired in the removal of DNA interstrand cross-links, such as those formed by the antibiotic mitomycin C. Individuals with Bloom syndrome and Nijmegan breakage syndrome exhibit a high frequency of chromosome breaks that result in chromosome aberrations (Chapter 6) and sister chromatid exchanges. Ataxia-telangiectasia is caused by defects in a kinase involved in the control of the cell cycle, and Bloom syndrome, Werner syndrome, and Rothmund-Thomson syndrome result from alterations in specific DNA helicases (members of the RecQ family of helicases). Table 13.1 lists some of the better known human diseases resulting from inherited defects in DNA repair pathways. KEY POINTS

- The importance of DNA repair pathways is documented convincingly by inherited human disorders that result from defects in DNA repair.
 - Certain types of cancer are also associated with defects in DNA repair pathways.

DNA Recombination Mechanisms

Recombination between homologous DNA molecules involves the activity of numerous enzymes that cleave, unwind, stimulate single-strand invasions of double helices, repair, and join strands of DNA. We discussed the main features of recombination between homologous chromosomes in Chapter 7, but we did not consider the molecular details of the process. Because many of the gene products involved in the repair of damaged DNA also are required for recombination between homologous chromosomes, or crossing over, we will

now examine some of the molecular aspects of this important process. Moreover, recombination usually, perhaps always, involves some DNA repair synthesis. Thus, much of the information discussed in the preceding sections is relevant to the process of recombination.

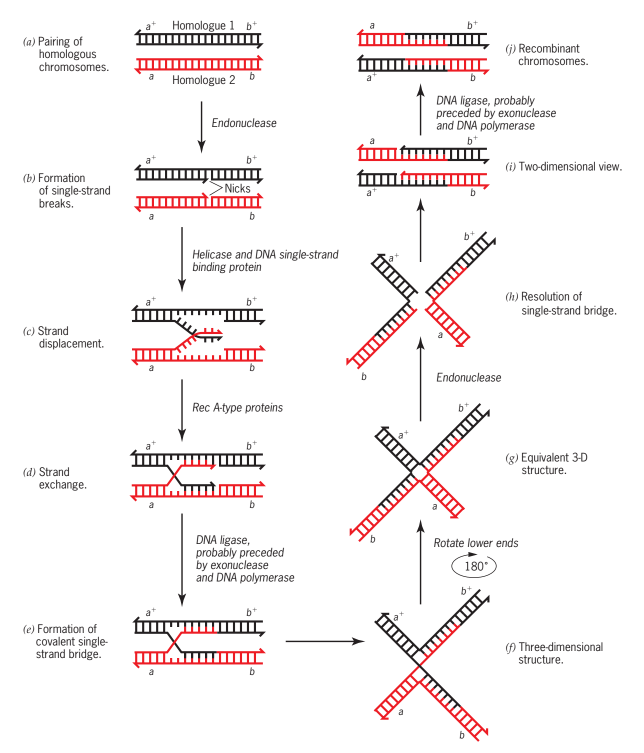
RECOMBINATION: CLEAVAGE AND REJOINING OF DNA MOLECULES

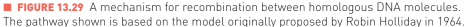
In Chapter 7, we discussed the experiment of Creighton and McClintock showing that crossing over occurs by breakage of parental chromosomes and rejoining of the parts in new combinations. Evidence demonstrating that recombination occurs by breakage and rejoining has also been obtained by autoradiography and other techniques. Indeed, the main features of the process of recombination are now well established, even though specific details remain to be elucidated.

Much of what we know about the molecular details of crossing over is based on the study of *recombination-deficient mutants* of *E. coli* and *S. cerevisiae*. Biochemical studies of these mutants have shown that they are deficient in various enzymes and other proteins required for recombination. Together, the results of genetic and biochemical studies have provided a fairly complete picture of recombination at the molecular level.

Many of the popular models of crossing over were derived from a model proposed by Robin Holliday in 1964. Holliday's model was one of the first that explained most of the genetic data available at the time by a mechanism involving the breakage, reunion, and repair of DNA molecules. An updated version of the Holliday model is shown in **Figure 13.29**. This mechanism, like many others that have been invoked, begins when an endonuclease cleaves single strands of each of the two parental DNA molecules (breakage). Segments of the single strands on one side of each cut are then displaced from their complementary strands with the aid of DNA helicases and single-strand binding proteins. The helicases unwind the two strands of DNA in the region adjacent to single-strand incisions. In *E. coli*, the *RecBCD complex* contains both an endonuclease activity that makes single-strand breaks in DNA and a DNA helicase activity that unwinds the complementary strands of DNA in the region adjacent to each nick.

The displaced single strands then exchange pairing partners, base-pairing with the intact complementary strands of the homologous chromosomes. This process is stimulated by proteins like the *E. coli* RecA protein. RecA-type proteins have been characterized in many species, both prokaryotic and eukaryotic. RecA protein and its homologues stimulate *single-strand assimilation*, a process by which a single strand of DNA displaces its homologue in a DNA double helix. RecA-type proteins promote reciprocal exchanges of DNA single strands between two DNA double helices in two steps. In the first step, a single strand of one double helix is assimilated by a





second, homologous double helix, displacing the identical or homologous strand and base-pairing with the complementary strand. In the second step, the displaced single strand is similarly assimilated by the first double helix. The RecA protein mediates these exchanges by binding to the unpaired strand of DNA, aiding in the search for a homologous DNA sequence, and, once a homologous double helix is found, promoting the replacement of one strand with the unpaired strand. If complementary sequences already exist as single strands, the presence of RecA protein increases the rate of renaturation by over 50-fold.

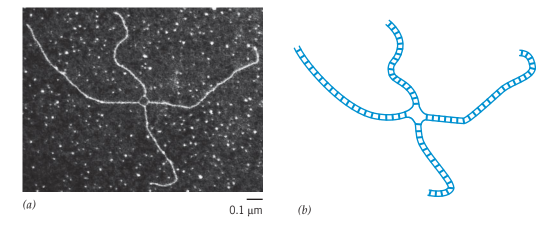


FIGURE 13.30 Electron micrograph (a) and diagram (b) of a *chi form*. Two DNA molecules have been caught in the process of genetic recombination using the electron microscope. This electron micrograph provides direct physical evidence for the existence of the Holliday recombination intermediate. Note how this molecule corresponds exactly to the theoretical structure that arises in *panel (g)* of the prototype Holliday recombination model, shown in Figure 13.29. *Micrograph Courtesy of H. Potter, University of South Florida and D. Dressler, Harvard University.*

The cleaved strands are then covalently joined in new combinations (reunion) by DNA ligase. If the original breaks in the two strands do not occur at exactly the same site in the two homologues, some tailoring will be required before DNA ligase can catalyze the reunion step. This tailoring involves the excision of nucleotides by an exonuclease and repair synthesis by a DNA polymerase. The sequence of events described so far will produce X-shaped recombination intermediates called *chi forms*, which have been observed by electron microscopy in several species (**Figure 13.30**). The chi forms are resolved by enzyme-catalyzed breakage and rejoining of the complementary DNA strands to produce two recombinant DNA molecules. In *E. coli*, chi structures can be resolved by the product of either the *recG* gene or the *ruvC* gene (repair of *UV*-induced damage). Each gene encodes an endonuclease that catalyzes the cleavage of single strands at chi junctions (see Figure 13.29).

A substantial body of evidence indicates that homologous recombination occurs by more than one mechanism-probably by several different mechanisms. In S. cerevisiae, the ends of DNA molecules produced by double-strand breaks are highly recombinogenic. This fact and other evidence suggest that recombination in yeast often involves a double-strand break in one of the parental double helices. Thus, in 1983, Jack Szostak, Franklin Stahl, and colleagues proposed a double-strand break model of crossing over. According to their model, recombination involves a double-strand break in one of the parental double helices, not just single-strand breaks as in the Holliday model. The initial breaks are then enlarged to gaps in both strands. The two single-stranded termini produced at the double-stranded gap of the broken double helix invade the intact double helix and displace segments of the homologous strand in this region. The gaps are then filled in by repair synthesis. This process yields two homologous chromosomes joined by two single-strand bridges. The bridges are resolved by endonucleolytic cleavage, just as in the Holliday model. Both the double-strand-break model and the Holliday model nicely explain the production of chromosomes that are recombinant for genetic markers flanking the region in which the crossover occurs.

GENE CONVERSION: DNA REPAIR SYNTHESIS ASSOCIATED WITH RECOMBINATION

Up to this point, we have discussed only recombination events that can be explained by breakage of homologous chromatids and the reciprocal exchange of parts. However,

analysis of tetrads of meiotic products of certain fungi reveals that genetic exchange is not always reciprocal. For example, if crosses are performed between two closely linked mutations in the mold *Neurospora*, and asci containing wild-type recombinants are analyzed, these asci frequently do not contain the reciprocal, double-mutant recombinant.

Consider a cross involving two closely linked mutations, m_1 and m_2 . In a cross of $m_1 m_2^+$ with $m_1^+ m_2$, asci of the following type are observed:

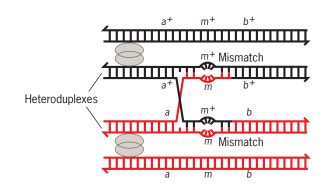
Spore pair 1: $m_1^+ m_2$ Spore pair 2: $m_1^+ m_2^+$ Spore pair 3: $m_1^- m_2^+$ Spore pair 4: $m_1^- m_2^+$

Wild-type $m_1^+ m_2^+$ spores are present, but the $m_1 m_2$ double-mutant spores are not present in the ascus. Reciprocal recombination would produce an $m_1 m_2$ chromosome whenever an $m_1^+ m_2^+$ chromosome was produced. In this ascus, the $m_2^+:m_2$ ratio is 3:1 rather than 2:2 as expected. One of the m_2 alleles appears to have been "converted" to the m_2^+ allelic form. Thus, this type of nonreciprocal recombination is called **gene conversion**. We might assume that gene conversion results from mutation, except that it occurs at a higher frequency than the corresponding mutation events, always produces the allele present on the homologous chromosome, not a new allele, and is correlated about 50 percent of the time with reciprocal recombination of flanking markers. The last observation strongly suggests that gene conversion is now believed to result from DNA repair synthesis associated with the breakage, excision, and reunion events of crossing over.

With closely linked markers, gene conversion occurs more frequently than reciprocal recombination. In one study of the *bis1* gene of yeast, 980 of 1081 asci containing *bis*⁺ recombinants exhibited gene conversion, whereas only 101 showed classical reciprocal recombination.

The most striking feature of gene conversion is that the input 1:1 allele ratio is not maintained. This can be explained easily if short segments of parental DNA are degraded and then resynthesized with template strands provided by DNA carrying the other allele. Given the mechanisms of excision repair discussed earlier in this chapter, the Holliday model of crossing over explains gene conversion for genetic markers located in the immediate vicinity of the crossover. In Figure 13.29*d*–*i*, there is a segment of DNA between the a^+ and b^+ loci where complementary strands of DNA from the two homologous chromosomes are base-paired. If a third pair of alleles located within this segment were segregating in the cross, mismatches in the two double helices would be present. DNA molecules containing such mismatches, or different alleles in the two complementary strands of a double helix, are called **heteroduplexes.** Such heteroduplex molecules occur as intermediates in the process of recombination.

If Figure 13.29*e* were modified to include a third pair of alleles, and the other two chromatids were added, the tetrad would have the following composition:



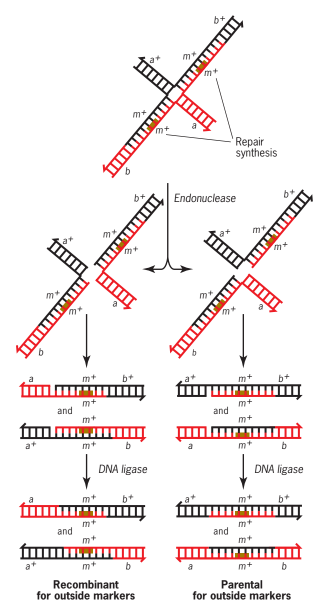
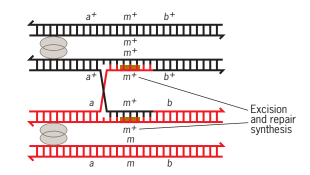


FIGURE 13.31 Formation of either the recombinant (bottom left) or parental (bottom right) combinations of flanking markers in association with gene conversion. The recombination intermediate at the top is equivalent to that illustrated in Figure 13.29g, but shows the mismatch-repaired chromatids of the tetrad diagrammed in the text. This tetrad produces an ascus showing $3 m^+$ to 1 m gene conversion. Cleavage of the single-strand bridge in the vertical plane (left) produces the recombinant ($a^+ b$ and $a b^+$) arrangement of flanking markers, whereas cleavage in the horizontal plane yields the parental ($a^+ b^+$ and a b) arrangement of the flanking markers.

If the mismatches are resolved by nucleotide excision repair (see Figure 13.27), in which the *m* strands are excised and resynthesized with the complementary m^+ strands as templates, the following tetrad will result:



As a result of semiconservative DNA replication during the subsequent mitotic division, this tetrad will yield an ascus containing six m^+ ascospores and two *m* ascospores, the 3:1 gene conversion ratio.

Suppose that only one of the two mismatches in the tetrad just described is repaired prior to the mitotic division. In this case, the semiconservative replication of the remaining heteroduplex will yield one m^+ homoduplex and one *m* homoduplex, and the resulting ascus will contain a $5m^+$:3m ratio of ascospores. Such 5:3 gene conversion ratios do occur. They result from postmeiotic (mitotic) segregation of unrepaired heteroduplexes.

Gene conversion is associated with the reciprocal recombination of flanking markers approximately 50 percent of the time. This correlation is nicely explained by the Holliday model of recombination presented in Figure 13.29. If the two recombinant chromatids of the tetrad just diagrammed are drawn in a form equivalent to that shown in Figure 13.29g,

the association of gene conversion with reciprocal recombination of flanking markers can easily be explained (**■** Figure 13.31). The single-strand bridge connecting the two chromatids must be resolved by endonucleolytic cleavage to complete the recombination process. This cleavage may occur either horizontally or vertically on the chi form drawn in Figure 13.31. Vertical cleavage will yield an ascus showing both gene conversion and reciprocal recombination of flanking markers. Horizontal cleavage will yield an ascus showing gene conversion and the parental combination of flanking markers. Thus, if cleavage occurs in the vertical plane half of the time and in the horizontal plane half of the time, gene conversion will be associated with reciprocal recombination of flanking markers about 50 percent of the time, as observed.

KEY POINTS

- Crossing over involves the breakage of homologous DNA molecules and the rejoining of parts in new combinations.
- When genetic markers are closely linked, nonreciprocal recombination, or gene conversion, often occurs, yielding 3:1 ratios of the segregation alleles.
- Gene conversion results from DNA repair synthesis that occurs during the recombination process.

Basic Exercises

Illustrate Basic Genetic Analysis

- **1.** Consider the role of mutation in evolution. Could species evolve in the absence of mutation?
- **Answer:** No. Mutation is the essential first step in the evolutionary process; it is the ultimate source of all new genetic variation. Recombination mechanisms produce new combinations of this genetic variation, and natural (or artificial) selection preserves the combinations that produce organisms that are the best adapted to the environments in which they live. Without mutation, evolution could not occur.
- 2. Consider a short segment of a wild-type gene with the following nucleotide-pair sequence:

5'-ATG TCC GCA TGG GGA -3' 3'-TAC AGG CGT ACC CCT -5'

Transcription of this gene segment yields the following mRNA nucleotide sequence:

5'-AUG UCC GCA UGG GGA-3'

and translation of this mRNA produces the amino acid sequence:

methionine-serine-alanine-tryptophan-glycine

If a single nucleotide-pair substitution occurs in this gene, changing the G:C at position 7 to A:T, what effect will this mutation have on the polypeptide produced by this gene?

Answer: The mRNA produced by the gene segment with the mutation will now be:

5'-AUG UCC ACA UGG GGA-3'

and will encode the amino acid sequence:

methionine-serine-threonine-tryptophan-glycine

Note that the third amino acid of the mutant polypeptide is threonine instead of alanine as in the wild-type polypeptide. Thus, this base-pair substitution, like most base-pair substitutions, results in a single amino acid substitution in the polypeptide encoded by the gene.

3. If a single nucleotide-pair substitution occurs in the gene segment shown in Exercise 2, changing the G:C at position 12 to A:T, what effect will this mutation have on the polypeptide produced by this gene?

Testing Your Knowledge

Integrate Different Concepts and Techniques

1. Charles Yanofsky isolated a large number of auxotrophic mutants of *E. coli* that could grow only on medium containing the amino acid tryptophan. How could such mutants be identi-

Answer: The resulting mRNA sequence will be: 5'-AUG UCC GCA UGA GGA-3'

termination codon

with the fourth codon changed from UGG, a tryptophan codon, to UGA, one of the three chain-termination codons. As a result, the mutant polypeptide will be prematurely terminated at this position, yielding a truncated protein.

4. If a single A:T base pair is inserted between nucleotide pairs 6 and 7 in the gene segment shown in Exercise 2, what effect will this change have on the polypeptide specified by this gene?

Answer: The nucleotide sequence of the mRNA specified by the mutant gene segment will be:

5'-AUG UCC AGC AUG GGG A-J'

and the polypeptide produced from the altered mRNA will be:

methionine-serine-serine-methionine-glycline

altered amino acid sequence

The base-pair insertion will alter the reading frame of the mRNA (trinucleotides read as codons) distal to the site of the mutation. As a result, all of the amino acids specified by codons downstream from the site of the insertion will be changed, producing an abnormal (usually nonfunctional) protein. In many cases, an insertion will shift a termination codon into the proper reading frame for translation, causing a truncated polypeptide to be produced.

5. If the two DNA molecules shown in the following diagram, where the arrowhead indicates the 3' end of each strand, undergo crossing over by breakage and reunion, will both of the recombinants shown be produced with equal frequency?



Answer: No. During recombination, only DNA strands with the same polarity can be joined. The second recombinant will not be produced.

fied? If a specific tryptophan auxotroph resulted from a nitrous acid-induced mutation, could it be induced to revert back to prototrophy by treatment with 5-bromouracil (5–BU)?

- Answer: The culture of mutagenized bacteria must be grown in medium containing tryptophan so that the desired mutants can survive and reproduce. The bacteria should then be diluted, plated on agar medium containing tryptophan, and incubated until visible colonies are produced. The colonies are next transferred to plates lacking tryptophan by the replica-plating technique developed by the Lederbergs (see Figure 13.15). The desired tryptophan auxotrophs will grow on the plates containing tryptophan, but not on the replica plates lacking tryptophan. Because nitrous acid and 5–BU produce transition mutations in both directions, A:T \leftrightarrow G:C, any mutation induced with nitrous acid should be induced to backmutate with 5–BU.
- 2. Assume that you recently discovered a new species of bacteria and named it *Escherichia mutaphilium*. During the last year, you have been studying the *mutA* gene and its polypeptide product, the enzyme trinucleotide mutagenase, in this bacterium. *E. mutaphilium* has been shown to use the established, nearly universal genetic code and to behave like *Escherichia coli* in all other respects relevant to molecular genetics.

The sixth amino acid from the amino terminus of the wild-type trinucleotide mutagenase is histidine, and the wild-type *mutA* gene has the triplet nucleotide-pair sequence

3'-GTA-5' 5'-CAT-3'

at the position corresponding to the sixth amino acid of the gene product. Seven independently isolated mutants with single nucleotide-pair substitutions within this triplet have also been characterized. Furthermore, the mutant trinucleotide mutagenases have all been purified and sequenced. All seven are different: they contain, respectively, glutamine, tyrosine, asparagine, aspartic acid, arginine, proline, and leucine as the sixth amino acid from the amino terminus.

Mutants *mutA1*, *mutA2*, and *mutA3* will not recombine with each other, but each will recombine with each of the other four mutants (*mutA4*, *mutA5*, *mutA6*, and *mutA7*) to yield true wild-type recombinants. Similarly, mutants A4, A5, and A6 will not recombine with each other but will each yield true wild-type recombinants in crosses with each of the other four mutants. Finally, crosses between *mutA1* and *mutA7* yield about twice as many true wild-type recombinants as do crosses between *mutA6* and *mutA7*.

Mutants A1 and A6 are induced to back-mutate to wildtype by treatment with 5-bromouracil (5-BU), whereas mutants A2, A3, A4, A5, and A7 are not induced to backmutate by treatment with 5–BU. Mutants A2 and A4 grow slowly on minimal medium, whereas mutants A3 and A5 carry null mutations (producing completely inactive gene products) and are incapable of growth on minimal medium. This difference has been used to select for mutation events from genotypes mutA3 and mutA5 to genotypes mutA2and mutA4. Mutants A3 and A5 can be induced to mutate to A2 and A4, respectively, by treatment with 5-bromouracil or hydroxylamine. However, mutant A3 cannot be induced to mutate to A4, nor A5 to A2, by treatment with either mutagen.

Use the information given above and the nature of the genetic code (Table 12.1) to deduce which mutant allele specifies the mutant polypeptide with each of the seven different amino acid substitutions at position 6 of trinucleotide mutagenase, and describe the rationale behind each of your deductions.

Answer: The following deductions can be made from the information given.

- (a) The wild-type His codon must be CAU based on the nucleotide-pair sequence of the gene.
- (b) The codons for the seven amino acids found at position 6 in the mutant polypeptides must be connected to CAU by a single-base change because the mutants were all derived from wild-type by a single nucleotide-pair substitution. Thus, the degeneracy of the genetic code is not a factor in deducing specific codon assignments.
- (c) Because of the nature of the genetic code—specifically the degeneracy at the third (3') position in each codon there are three possible amino acid substitutions due to single-base substitutions (caused by single base-pair substitutions in DNA) at each of the first two positions (the 5' base and the middle base), but only one possible amino acid change due to a single-base change at position 3 (the 3' base in the codon). For ease of discussion, the three nucleotide-pair positions in the triplet under consideration will be referred to as position 1 (corresponding to the 5' base in the codon), position 2 (the middle nucleotide pair), and position 3 (corresponding to the 3' base in the mRNA codon).
- (d) Since A1, A2, and A3 do not recombine with each other, they must all result from base-pair substitutions at the same position in the triplet, at either position 1 or position 2. The same is true for A4, A5, and A6. Since A7 recombines with each of the other six mutant alleles, it must result from the single base-pair substitution at position 3 that leads to an amino acid change.
- (e) The only amino acid with codons connected to the His codon CAU by single-base changes at position 3 is Gln (codons CAA and CAG). Thus, the *mutA*7 polypeptide must have glutamine as the sixth amino acid.
- (f) Since mutA7 (the third position substitution) yields about twice as many wild-type recombinants in crosses with mutA1 as in crosses with mutA6, the A1 substitution must be at position 1 and the mutA6 substitution must be at position 2. Combined with (d) above, this places the A2 and A3 substitutions at position 1 and the A4 and A5 substitutions at position 2.
- (g) Since mutA1 and mutA6 are induced to revert to wildtype by 5-BU, they must be connected to the triplet of nucleotide pairs encoding His by transition mutations that is,

$$(mutA1) \stackrel{\text{ATA}}{\text{TAT}} \stackrel{\text{5-BU}}{\longleftrightarrow} \stackrel{\text{GTA}}{\underset{\text{CAT}}{\text{CAT}}} \stackrel{\text{5-BU}}{\longleftarrow} \stackrel{\text{GCA}}{\underset{\text{CGT}}{\text{CGT}}} (mutA6)$$

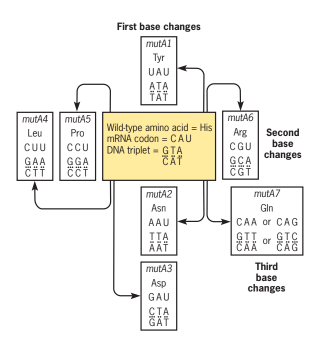
(h) Since *mutA3* and *mutA5* are induced to mutate to *mutA2* and *mutA4*, respectively, by hydroxylamine, *A3* must be connected to *A2*, and *A5* to *A4*, specifically by G:C \rightarrow A:T transitions—that is,

$$(mutA3) \stackrel{\text{CTA}}{\text{GAT}} \xrightarrow{\text{HA}} \stackrel{\text{TTA}}{\longrightarrow} \stackrel{\text{TTA}}{\underset{\text{AAT}}{\text{TTA}}} (mutA2)$$

and

$$(mutA5) \xrightarrow{\text{GGA}}_{\text{CCT}} \xrightarrow{\text{HA}} \xrightarrow{\text{GAA}}_{\text{CTT}} (mutA4)$$

Collectively, these deductions establish that the following relationships between the amino acids, codons, and nucleotide-pair triplets are present at the position of interest in the trinucleotide mutagenase polypeptides, mRNAs, and genes in the seven different mutants:



Questions and Problems

Enhance Understanding and Develop Analytical Skills

- 13.1 Identify the following point mutations represented in DNA and in RNA as (1) transitions, (2) transversions, or (3) reading frameshifts. (a) A to G; (b) C to T; (c) C to G; (d) T to A; (e) UAU ACC UAU to UAU AAC CUA; (f) UUG CUA AUA to UUG CUG AUA.
- **13.2 GO** Of all possible missense mutations that can occur in a segment of DNA encoding the amino acid tryptophan, what is the ratio of transversions to transitions if all single base-pair substitutions occur at the same frequency?
- **13.3** Both lethal and visible mutations are expected to occur in fruit flies that are subjected to irradiation. Outline a method for detecting (a) X-linked lethals and (b) X-linked visible mutations in irradiated *Drosophila*.
- **13.4** How can mutations in bacteria causing resistance to a particular drug be detected? How can it be determined whether a particular drug causes mutations or merely identifies mutations already present in the organisms under investigation?
- **13.5** Published spontaneous mutation rates for humans are generally higher than those for bacteria. Does this indicate that individual genes of humans mutate more frequently than those of bacteria? Explain.
- **13.6** A precancerous condition (intestinal polyposis) in a particular human family group is determined by a single dominant gene. Among the descendants of one woman who died with cancer of the colon, 10 people

have died with the same type of cancer and 6 now have intestinal polyposis. All other branches of the large kindred have been carefully examined, and no cases have been found. Suggest an explanation for the origin of the defective gene.

- **13.7** Juvenile muscular dystrophy in humans depends on an X-linked recessive gene. In an intensive study, 33 cases were found in a population of some 800,000 people. The investigators were confident that they had found all cases that were well enough advanced to be detected at the time the study was made. The symptoms of the disease were expressed only in males. Most of those with the disease died at an early age, and none lived beyond 21 years of age. Usually, only one case was detected in a family, but sometimes two or three cases occurred in the same family. Suggest an explanation for the sporadic occurrence of the disease and the tendency for the gene to persist in the population.
- **13.8** Products resulting from somatic mutations, such as the navel orange and the Delicious apple, have become widespread in citrus groves and apple orchards. However, traits resulting from somatic mutations are seldom maintained in animals. Why?
- **13.9** If a single short-legged sheep should occur in a flock, suggest experiments to determine whether the short legs are the result of a mutation or an environmental effect. If due to a mutation, how can one determine whether the mutation is dominant or recessive?

- **13.10** How might enzymes such as DNA polymerase be involved in the mode of action of both mutator and antimutator genes (mutant genes that increase and decrease, respectively, mutation rates)?
- **13.11** How could spontaneous mutation rates be optimized by natural selection?
- **13.12** A mutator gene Dt in maize increases the rate at which the gene for colorless aleurone (*a*) mutates to the dominant allele (*A*), which yields colored aleurone. When reciprocal crosses were made (i.e., seed parent dt/dt, $a/a \times Dt/Dt$, a/a and seed parent Dt/Dt, $a/a \times dt/dt$, a/a), the cross with Dt/Dt seed parents produced three times as many dots per kernel as the reciprocal cross. Explain these results.
- 13.13 A single mutation blocks the conversion of phenylalanine to tyrosine. (a) Is the mutant gene expected to be pleiotropic? (b) Explain.
- **13.14** How can normal hemoglobin (hemoglobin A) and hemoglobin S be distinguished?
- **13.15** If CTT is a DNA triplet (transcribed strand of DNA) specifying glutamic acid, what DNA and mRNA base triplet alterations could account for valine and lysine in position 6 of the β -globin chain?
- **13.16** The bacteriophage T4 genome contains about 50 percent A:T base pairs and 50 percent G:C base pairs. The base analog 2-aminopurine induces A:T \rightarrow G:C and G:C \rightarrow A:T base-pair substitutions by undergoing tautomeric shifts. Hydroxylamine is a mutagenic chemical that reacts specifically with cytosine and induces only G:C \rightarrow A:T substitutions. If a large number of independent mutations were produced in bacteriophage T4 by treatment with 2-aminopurine, what percentage of these mutations should you expect to be induced to mutate back to the wild-type genotype by treatment with hydroxylamine?
- 13.17 Assuming that the β -globin chain and the α -globin chain shared a common ancestor, what mechanisms might explain the differences that now exist in these two chains? What changes in DNA and mRNA codons would account for the differences that have resulted in dissimilar amino acids at corresponding positions?
- **13.18** In a given strain of bacteria, all of the cells are usually killed when a specific concentration of streptomycin is present in the medium. Mutations that confer resistance to streptomycin occur. The streptomycin-resistant mutants are of two types: some can live with or without streptomycin; others cannot survive unless this drug is present in the medium. Given a streptomycin-sensitive strain of this species, outline an experimental procedure by which streptomycin-resistant strains of the two types could be established.
- 13.19 One stock of fruit flies was treated with 1000 roentgens (r) of X rays. The X-ray treatment increased the mutation rate of a particular gene by 2 percent. What percentage increases in the mutation rate of this gene would be

expected if this stock of flies was treated with X-ray doses of 1500 r, 2000 r, and 3000 r?

- **13.20** Why does the frequency of chromosome breaks induced by X rays vary with the total dosage and not with the rate at which it is delivered?
- **13.21** A reactor overheats and produces radioactive tritium (H³), radioactive iodine (I¹³¹), and radioactive xenon (Xn¹³³). Why should we be more concerned about radioactive iodine than the other two radioactive isotopes?
- 13.22 One person was in an accident and received 50 roent-gens (r) of X rays at one time. Another person received 5 r in each of 20 treatments. Assuming no intensity effect, what proportionate number of mutations would be expected in each person?
- 13.23 A cross was performed in *Neurospora crassa* between a strain of mating type A and genotype $x^+ m^+ z$ and a strain of mating type a and genotype $x m z^+$. Genes x, m, and z are closely linked and present in the order x-m-z on the chromosome. An ascus produced from this cross contained two copies ("identical twins") of each of the four products of meiosis. If the genotypes of the four products of meiosis showed that gene conversion had occurred at the m locus and that reciprocal recombination had occurred at the xand z loci, what might the genotypes of the four products look like? In the parentheses that follow, write the genotypes of the four haploid products of meiosis in an ascus showing gene conversion at the m locus and reciprocal recombination of the flanking markers (at the x and z loci).

Ascus Spore Pairs

	1–2			3-4			5-6		7–8
()	()	()	()

- **13.24** How does nitrous acid induce mutations? What specific end results might be expected in DNA and mRNA from the treatment of viruses with nitrous acid?
- **13.25** Are mutational changes induced by nitrous acid more likely to be transitions or transversions?
- **13.26** You are screening three new pesticides for potential mutagenicity using the Ames test. Two *his*⁻ strains resulting from either a frameshift or a transition mutation were used and produced the following results (number of revertant colonies):

Strain 1	Transition Mutant Control (no chemical)	Transition Mutant + Chemical	Transition Mutant + Chemical + Rat Liver Enzymes
Pesticide #1	21	180	19
Pesticide #2	18	19	17
Pesticide #3	25	265	270

Strain 2	Frameshift Mutant Control (no chemical)	Frameshift Mutant + Chemical	Frameshift Mutant + Chemical + Rat Liver Enzymes
Pesticide #1	5	4	5
Pesticide #2	7	5	93
Pesticide #3	6	9	7

What type of mutations, if any, do the three pesticides induce?

- **13.27** How does the action and mutagenic effect of 5-bromouracil differ from that of nitrous acid?
- **13.28** Sydney Brenner and A. O. W. Stretton found that nonsense mutations did not terminate polypeptide synthesis in the *rII* gene of the bacteriophage T4 when these mutations were located within a DNA sequence interval in which a single-nucleotide insertion had been made on one end and a single nucleotide deletion had been made on the other. How can this finding be explained?
- 13.29 Seymour Benzer and Ernst Freese compared spontaneous and 5-bromouracil-induced mutants in the *rII* gene of the bacteriophage T4; the mutagen increased the mutation rate $(rII^+ \rightarrow rII)$ several hundred times above the spontaneous mutation rate. Almost all (98 percent) of the 5-bromouracil-induced mutants could be induced to revert to wild-type $(rII \rightarrow rII^+)$ by 5-bromouracil treatment, but only 14 percent of the spontaneous mutants could be induced to revert to wild-type by this treatment. Discuss the reason for this result.
- **13.30** How do acridine-induced changes in DNA result in inactive proteins?

Use the known codon-amino acid assignments given in Chapter 12 to work the following problems.

13.31 Mutations in the genes encoding the α and β subunits of hemoglobin lead to blood diseases such as thalassemias and sickle-cell anemia. You have found a family in China in which some members suffer from a new genetic form of anemia. The DNA sequences at the 5' end of the non-template strand of the normal and mutant DNA encoding the α subunit of hemoglobin are as follows:

Normal 5'-ACGTTATGCCGTACTGCCAGCTAACT-GCTAAAGAACAATTA......-3'

Mutant 5'-ACGTTATGCCCGTACTGCCAGCTAACT-GCTAAAGAACAATTA......-3'

- (a) What type of mutation is present in the mutant hemoglobin gene?
- (b) What are the codons in the translated portion of the mRNA transcribed from the normal and mutant genes?
- (c) What are the amino acid sequences of the normal and mutant polypeptides?

- 13.32 Go Bacteriophage MS2 carries its genetic information in RNA. Its chromosome is analogous to a polygenic molecule of mRNA in organisms that store their genetic information in DNA. The MS2 minichromosome encodes four polypeptides (i.e., it has four genes). One of these four genes encodes the MS2 coat protein, a polypeptide 129 amino acids long. The entire nucleotide sequence in the RNA of MS2 is known. Codon 112 of the coat protein gene is CUA, which specifies the amino acid leucine. If you were to treat a replicating population of bacteriophage MS2 with the mutagen 5-bromouracil, what amino acid substitutions would you expect to be induced at position 112 of the MS2 coat protein (i.e., Leu \rightarrow other amino acid)? (*Note*: Bacteriophage MS2 RNA replicates using a complementary strand of RNA and base-pairing like DNA.)
- **13.33** Would the different amino acid substitutions induced by 5-bromouracil at position 112 of the coat polypeptide that you indicated in Problem 13.32 be expected to occur with equal frequency? If so, why? If not, why not? Which one(s), if any, would occur more frequently?
- **13.34** Would such mutations occur if a nonreplicating suspension of MS2 phage was treated with 5-bromouracil?
- 13.35 Recall that nitrous acid deaminates adenine, cytosine, and guanine (adenine \rightarrow hypoxanthine, which base-pairs with cytosine; cytosine \rightarrow uracil, which base-pairs with adenine; and guanine \rightarrow xanthine, which base-pairs with cytosine). Would you expect nitrous acid to induce any mutations that result in the substitution of another amino acid for a glycine residue in a wild-type polypeptide (i. e., glycine \rightarrow another amino acid) if the mutagenesis were carried out on a suspension of mature (nonreplicating) T4 bacteriophage? (*Note:* After the mutagenic treatment of the phage suspension, the nitrous acid is removed. The treated phage are then allowed to infect *E. coli* cells to express any induced mutations.) If so, by what mechanism? If not, why not?
- **13.36** Keeping in mind the known nature of the genetic code, the information given about phage MS2 in Problem 13.32, and the information you have learned about nitrous acid in Problem 13.35, would you expect nitrous acid to induce any mutations that would result in amino acid substitutions of the type glycine \rightarrow another amino acid if the mutagenesis were carried out on a suspension of mature (nonreplicating) MS2 bacteriophage? If so, by what mechanism? If not, why not?
- **13.37** Would you expect nitrous acid to induce a higher frequency of Tyr \rightarrow Ser or Tyr \rightarrow Cys substitutions? Why?
- 13.38 Which of the following amino acid substitutions should you expect to be induced by 5-bromouracil with the highest frequency? (a) Met → Leu; (b) Met → Thr; (c) Lys → Thr; (d) Lys → Gln; (e) Pro → Arg; or (f) Pro → Gln? Why?
- 13.39 The wild-type sequence of part of a protein is

NH₂-Trp-Trp-Trp-Met-Arg-Glu-Trp-Thr-Met

Each mutant in the following table differs from wild-type by a single point mutation. Using this information, determine the mRNA sequence coding for the wild-type polypeptide. If there is more than one possible nucleotide, list all possibilities.

Mutant	Amino Acid Sequence of Polypeptide
1	Trp-Trp-Trp Met
2	Trp-Trp-Trp-Met-Arg-Asp-Trp-Thr-Met
3	Trp-Trp-Trp-Met-Arg-Lys-Trp-Thr-Met
4	Trp-Trp-Trp-Met-Arg-Glu-Trp-Met-Met

13.40 Acridine dyes such as proflavin are known to induce primarily single base-pair additions and deletions. Suppose that the wild-type nucleotide sequence in the mRNA produced from a gene is

5'-AUGCCCUUUGGGAAAGGGUUUCCCUAA-3'

Also, assume that a mutation is induced within this gene by proflavin and, subsequently, a revertant of this mutation is similarly induced with proflavin and shown to result from a second-site suppressor mutation within the same gene. If the amino acid sequence of the polypeptide encoded by this gene in the revertant (double mutant) strain is

NH₂-Met-Pro-Phe-Gly-Glu-Arg-Phe-Pro-COOH

what would be the most likely nucleotide sequence in the mRNA of this gene in the revertant (double mutant)?

13.41 Eight independently isolated mutants of *E. coli*, all of which are unable to grow in the absence of histidine (his⁻), were examined in all possible cis and trans heterozygotes (partial diploids). All of the *cis* heterozygotes were able to grow in the absence of histidine. The trans heterozygotes yielded two different responses: some of them grew in the absence of histidine; others did not. The experimental results, using "+" to indicate growth and "0" to indicate no growth, are given in the accompanying table. How many genes are defined by these eight mutations? Which mutant strains carry mutations in the same gene(s)?

Growth of Trans Heterozygotes (without Histidine)

Mutant	1	2	3	4	5	6	7	8
8	0	0	0	0	0	0	+	0
7	+	+	+	+	+	+	0	
6	0	0	0	0	0	0		
5	0	0	0	0	0			
4	0	0	0	0				
3	0	0	0					
2	0	0						
1	0							

13.42 Assume that the mutants described in Problem 13.41 vielded the following results. How many genes would they have defined? Which mutations would have been in the same gene(s)?

Mutant	1	2	3	4	5	6	7	8
8	+	+	+	+	+	+	0	0
7	+	+	+	+	+	+	0	
6	+	+	+	+	0	0		
5	+	+	+	+	0			
4	+	+	0	0				
3	+	+	0					
2	0	0						
1	0							

Growth of Trans Heterozygotes (without Histidine)

- 13.43 In Drosophila, white, white cherry, and vermilion are all X-linked mutations affecting eye color. All three mutations are recessive to their wild-type allele(s) for red eyes. A white-eyed female crossed with a vermilioneved male produces white-eved male offspring and redeved (wild-type) female offspring. A white-eved female crossed with a white cherry-eyed male produces whiteeyed sons and light cherry-eyed daughters. Do these results indicate whether or not any of the three mutations affecting eye color are located in the same gene? If so, which mutations?
- 13.44 The loz (lethal on Z) mutants of bacteriophage X are conditional lethal mutants that can grow on E. coli strain Y but cannot grow on E. coli strain Z. The results shown in the following table were obtained when seven loz mutants were analyzed for complementation by infecting E. coli strain Z with each possible pair of mutants. A "+" indicates that progeny phage were produced in the infected cells, and a "0" indicates that no progeny phage were produced. All possible cis tests were also done, and all cis heterozygotes produced wild-type yields of progeny phage.

Mutant	1	2	3	4	5	6	7
7	+	+	0	+	0	0	0
6	+	+	+	+	+	0	
5	+	+	0	+	0		
4	0	0	+	0			
3	+	+	0				
2	0	0					
1	0						

Propose three plausible explanations for the apparently anomalous complementation behavior of *loz* mutant number 7. (b) What simple genetic experiments can be used to distinguish between the three possible explanations? (c) Explain why specific outcomes of the proposed experiments will distinguish between the three possible explanations.

Genomics on the Web at http://www.ncbi.nlm.nih.gov

Sickle-cell disease is caused by a single base-pair substitution in the human β -globin gene. This mutation changes the sixth amino acid in the mature polypeptide from glutamic acid to valine (see Figure 1.9). This single amino acid change, in turn, causes all the symptoms of this painful and eventually fatal disease.

- What other mutations in the human β-globin gene have changed the glutamic acid at position 6 to some other amino acid? What are these hemoglobin variants called? Are there β-globin variants with an amino acid substitution at position 6 and another amino acid substitution elsewhere in the polypeptide?
- 2. Proline is present at position 5 in normal human β-globin. What amino acid substitutions have occurred at this position in mutant β-globins? How about the glutamic acid present at position 7? Are there mutations that change this amino acid to something else?

- **3.** Mutations have been documented at a large number of the 146 base-pair triplets (specifying mRNA codons) in the human β-globin gene. How many of these triplets have mutated to produce an amino acid substitution in the polypeptide?
- **4.** What genes are located next to the β-globin gene on human chromosome 11? What are the functions of the delta-, gamma A-, gamma G-, and epsilon-globin genes? Is there any significance to their arrangement on the chromosome?

Hint: At the NCBI web site, search all databases with the query "beta-globin variants." Start with the results in OMIM (Online Medical Inheritance in Man), click HBB (online symbol for the human β -globin gene), on the left bar, click HbVar for a list of all the human β -globin variants characterized to date, then return to the HBB page and click "Gene Map" for a list of the genes next to *HBB*.



The Techniques of Molecular Genetics

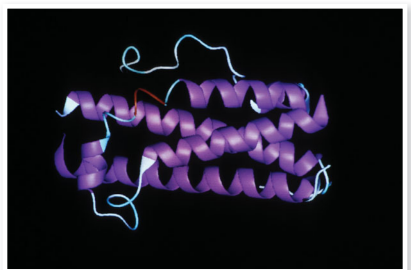
CHAPTER OUTLINE

- Basic Techniques Used to Identify, Amplify, and Clone Genes
- Construction and Screening of DNA Libraries
- The Molecular Analysis of DNA, RNA, and Protein
- The Molecular Analysis of Genes and Chromosomes

Treatment of Pituitary Dwarfism with Human Growth Hormone

Kathy was a typical child in most respects—happy, playful, a bit mischievous, and intelligent. Indeed, the only thing unusual about Kathy was her small stature. She was born with pituitary dwarfism, which results from a deficiency of human growth hormone (hGH). Kathy seemed destined to remain abnormally small throughout her life. Then, at age 10, Kathy began receiving treatments of hGH synthesized in bacteria. She grew five inches during her first year of treatments. By continuing to receive hGH during maturation, Kathy reached the short end of the normal height distribution for adults. Without these treatments, she would have remained abnormally small in stature.

The hGH that allowed Kathy to grow to near-normal size was one of the first products of genetic engineering, the use of designed or modified genes to synthesize desired products. hGH was initially produced in *E. coli* cells harboring a modified gene composed of the coding sequence for hGH fused to synthetic bacterial regulatory elements. This chimeric gene was constructed *in vitro* and introduced into *E. coli* by transformation. In 1985, hGH produced in *E. coli* became the second pharmaceutical



Computer-generated model of the structure of human growth hormone.

product of genetic engineering to be approved for use in humans by the U.S. Food and Drug Administration. Human insulin, which was the first such product, was approved in 1982. How do scientists construct a gene that will produce hGH or human insulin in *E. coli*? They accomplish this feat by combining the coding sequence of the human growth hormone or human insulin gene with regulatory sequences that will ensure its expression in *E. coli* cells. Once they have pieced together the gene in the test tube, they must introduce it into living bacteria so that it can be expressed. In the past, the synthesis of human proteins in bacteria seemed like science fiction. Today, human proteins are routinely produced in bacteria or eukaryotic cells growing in culture. In this chapter, we focus on the powerful tools of molecular genetics that allow researchers to construct genes from components derived from different species and to express these novel genes in both bacteria and eukaryotic cells.

Much of what we know about the structure of genes has been obtained by molecular studies of genes and chromosomes made possible by the development of **recombinant DNA technologies.** Recombinant DNA approaches begin with the **cloning** of specific genes. The cloning of a gene involves its isolation, its insertion into a small self-replicating genetic element such as a plasmid or viral chromosome, and its amplification during the replication of the plasmid or viral chromosome in an appropriate host cell (usually an *E. coli* cell). The small self-replicating genetic elements used to clone genes are called **cloning vectors.** Gene cloning—the isolation and amplification of a given gene—should not be confused with the cloning of organisms—the production of an organism, such as the lamb named Dolly, from a single cell obtained from an adult organism.

The isolation and cloning of a specific gene is a complex process. However, after a gene has been cloned, it can be subjected to a whole array of manipulations that allow investigations of gene structure–function relationships. Usually, a cloned gene is sequenced; that is, the nucleotide-pair sequence of the gene is determined. If the function of the gene is unknown, its nucleotide sequence can be compared with thousands of gene sequences stored in three large computer gene banks—one in Germany, a second in Japan, and a third in the United States (see Focus on GenBank in Chapter 15). Sometimes the function of a gene can be deduced based on its similarity to other genes whose functions are known. Given the nucleotide sequence of a gene and knowledge of the genetic code, the amino acid sequence of the polypeptide encoded by the gene can be predicted. The predicted amino acid sequence of the polypeptide can then be searched for amino acid sequences that may provide clues about its function. Nucleic acid and protein sequence databases have become important resources for research in molecular genetics, and they will become increasingly important in both basic biological research and the diverse applications of this research (Chapters 15 and 16).

Basic Techniques Used to Identify, Amplify, and Clone Genes

The haploid genome of a mammal contains about 3×10^9 nucleotide pairs. If the combined exons of the average gene are 3000 nucleotide pairs long (many are larger), the coding region of the gene will represent one of a million such sequences in the genome.

Recombinant DNA, gene cloning, and DNA amplification techniques allow scientists to isolate and characterize essentially any DNA sequence from any organism.

Although most of the DNA in mammalian genomes does not consist of genes, still, isolating any one gene is like searching for the proverbial needle in a haystack. Most techniques used in the analysis of genes and other DNA sequences require that the sequence be available in significant quantities in pure or essentially pure form. How can one identify the segment of a DNA molecule that carries a single gene and isolate enough of this sequence in pure form to permit molecular analyses of its structure and function?

The development of recombinant DNA and gene-cloning technologies has provided molecular geneticists with methods by which genes or other segments of large chromosomes can be isolated, replicated, and studied by nucleic acid sequencing techniques, electron microscopy, and other analytical techniques. Indeed, genes or other DNA sequences can be amplified by two distinct approaches—one with amplification of the sequence occurring *in vivo* and the other *in vitro*. The second approach can only be used when short nucleotide sequences on either side of the DNA sequence of interest are known.

Solve It!

How Many *Not*l Restriction Fragments in Chimpanzee DNA?

The genome of the chimpanzee (Pan troglodytes) is about the same size as the human (Homo sapiens) genome, but the diploid chromosome number in chimpanzees is 48, rather than 46 as in humans. Sex determination in chimpanzees occurs by the XX-XY mechanism just as in humans. All chimps contain 23 pairs of autosomes; in addition, females contain 2 X chromosomes and males an X chromosome and a Y chromosome. The haploid nuclear genome of the chimpanzee contains 2,928,563,828 nucleotide pairs. The mitochondrial genome of the chimpanzee consists of a circular molecule of DNA 16,600 nucleotide pairs long. If you assume that G, C, A, and T are present in equal amounts and are distributed randomly throughout both the nuclear and mitochondrial genomes of the chimp, how many restriction fragments would be produced by cleaving total DNA from a male chimpanzee with Notl, a restriction endonuclease that cleaves a specific eight nucleotide-pair sequence?

To see the solution to this problem, visit the Student Companion site. In the first approach, a minichromosome carrying the gene of interest is produced in the test tube and is then introduced into an appropriate host cell. This gene-cloning procedure involves two essential steps: (1) the incorporation of the gene of interest into a small self-replicating chromosome (*in vitro*) and (2) the amplification of the recombinant minichromosome by its replication in an appropriate host cell (*in vivo*). Step 1 involves the joining of two or more different DNA molecules *in vitro* to produce **recombinant DNA molecules**, for example, a human gene inserted into an *E. coli* plasmid or other self-replicating minichromosome. Step 2 is really the gene-cloning event in which the recombinant DNA molecule is replicated or "cloned" to produce many identical copies for subsequent biochemical analysis. In step 2, the recombinant minichromosome is introduced into *E. coli* cells where it replicates to produce many copies of the recombinant DNA molecule. Although the entire procedure is often referred to as the recombinant DNA or gene-cloning technique, these terms actually refer to two separate steps in the process.

In the second approach, short DNA strands that are complementary to DNA sequences on either side of the gene or DNA sequence of interest are synthesized and used to initiate its amplification *in vitro* by a special (heat-stable) DNA polymerase. This procedure—called the polymerase chain reaction (PCR)—is an extremely powerful gene-amplification tool. The amplified products can then be analyzed and sequenced, and, if desired, they can be inserted into cloning vectors and replicated *in vivo* for additional studies. Amplification of a DNA sequence by PCR frequently eliminates the need to clone the sequence by replication *in vivo*. Thus, procedures involving the amplification of DNA sequences by PCR have commonly replaced earlier *in vivo* amplification protocols. However, PCR can only be used when nucleotide sequences flanking the gene or DNA sequence of interest are known.

THE DISCOVERY OF RESTRICTION ENDONUCLEASES

The ability to clone and sequence essentially any gene or other DNA sequence of interest from any species depends on a special class of enzymes called restriction endonucleases (from the Greek term éndon meaning "within"; endonucleases make internal cuts in DNA molecules). Many endonucleases make random cuts in DNA, but the restriction endonucleases are site-specific, and Type II restriction enzymes cleave DNA molecules only at specific nucleotide sequences called restriction sites. Type II restriction enzymes cleave DNA at these sites regardless of the source of the DNA. Different restriction endonucleases are produced by different microorganisms and recognize different nucleotide sequences in DNA (Table 14.1). The restriction endonucleases are named by using the first letter of the genus and the first two letters of the species that produces the enzyme. If an enzyme is produced only by a specific strain, a letter designating the strain is appended to the name. The first restriction enzyme identified from a bacterial strain is designated I, the second II, and so on. Thus, restriction endonuclease EcoRI is produced by Escherichia coli strain RY13. Hundreds of restriction enzymes have been characterized and purified; thus, restriction endonucleases that cleave DNA molecules at many different DNA sequences are available. For an extensive list of restriction enzymes, see http://en.wikipedia.org/ wiki/List_of_restriction_enzyme_cutting_sites:_A#Whole_list_navigation.

Restriction endonucleases were discovered in 1970 by Hamilton Smith and Daniel Nathans (see A Milestone in Genetics: Restriction Endonucleases on the Student Companion site). They shared the 1986 Nobel Prize in Physiology or Medicine with Werner Arber, who carried out pioneering research that led to the discovery of restriction enzymes. The biological function of restriction endonucleases is to protect the genetic material of bacteria from "invasion" by foreign DNAs, such as DNA molecules from another species or viral DNAs. As a result, restriction endonucleases are sometimes referred to as the immune systems of prokaryotes.

All cleavage sites in the DNA of an organism must be protected from cleavage by the organism's own restriction endonucleases; otherwise the organism would commit suicide by degrading its own DNA. In many cases, this protection of endogenous cleavage sites is accomplished by **methylation** of one or more nucleotides in each

TABLE 14.1

Recognition Sequences and Cleavage Sites of Representative Restriction Endonucleases							
Enzyme	Source	Recognition Sequence ^a and Cleavage Sites ^b	Type of Ends Produced				
<i>Eco</i> RI	Escherichia coli strain RY13	5'-GAA TTC -3' 3'-CTT AAG-5' digest 3'-CTT.	+ 5'-AATTC-3' 5' Overhang: AA-5' G-5' 5' Overhang:				
Hincll	Haemophilus influenzae strain R_{c}	5'-GTPy PuAC-3' → 5'-GTP 3'-CAPu PyTG-5' → 3'-CAP	² y-3' + 5'-PuAC-3' Blunt ² u-5' + 3'-PyTG-5'				
HindIII	Haemophilus influenzae strain R_{d}	5'-AAG CTT-3' 3'-TTC GAA-5' → 5'-A 3'-TTC	CGA-5' + ^{5'-AGCTT-3'} 5' Overhang:				
Hpall	Haemophilus parainfluenzae	5'-CC GG-3' 3'-GG CC-5' → 5'-C 3'-GG CC-5'					
Alul	Arthrobacter luteus	5'-AG [↓] CT-3' → 5'-AG- 3'-TC [↓] GA-5' → 3'-TC-	3' + 5'-CT-3' Blunt 5' 3'-GA-5' Blunt				
Pstl	Providencia stuartii	5'-CTG CÅG-3' 3'-GAC GTC-5' $\longrightarrow 5'$ -CTG 3'-G	GCA-3' + G-3' 3' Overhang: 3'-ACGTC-5' 3' Overhang:				
Clal	Caryophanon latum	5'-ATC GAT-3' → 5'-AT 3'-TAG CTA-5' → 3'-TAG	C-5' ⁺ 5'-CGAT-3' 5' Overhang: C-5' TA-5' 5' Overhang:				
Sacl	Streptomyces achromogenes	5'-GAG CTC-3' 5'-GAG 3'-CTC GAG-5' 3'-C	GCT-3' + C-3' 3' Overhang 3'-TCGAG-5' 3' Overhang				
Notl	Nocardia otitidis	5'-GCGG CCGC-3' 3'-CGCC GGCG-5'	CCGG-5' + ^{5'-GGCCGC-3'} 5' Overhang:				

Recognition Sequences and Cleavage Sites of Representative Restriction Endonucleases

^a The axis of dyad symmetry in each palindromic recognition sequence is indicated by the red dot; the DNA sequences are the same reading in opposite directions from this point and switching the top and bottom strands to correct for their opposite polarity. Pu indicates that either purine (adenine or guanine) may be present at this position; Py indicates that either pyrimidine (thymine or cytosine) may be present. ^b The position of each bond cleaved is indicated by an arrow. Note that with some restriction endonucleases the cuts are staggered (at different positions

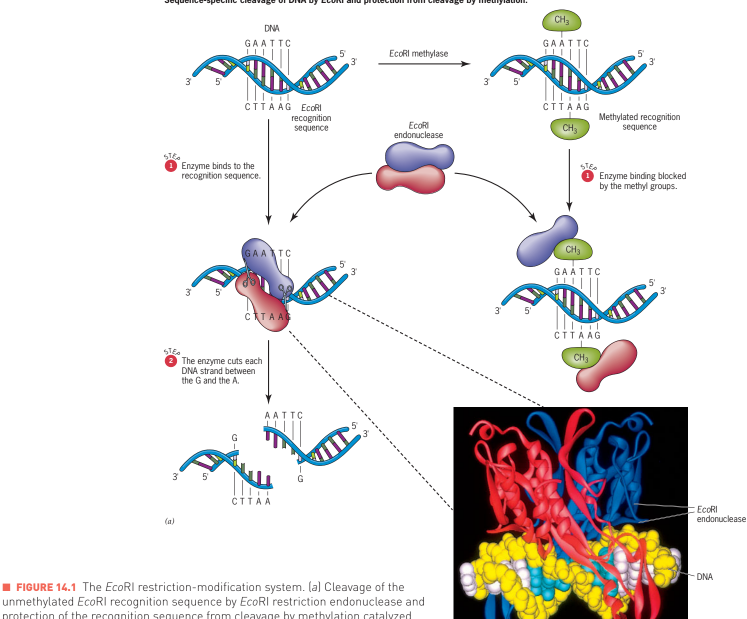
in the two complementary strands).

nucleotide sequence that is recognized by the organism's own restriction endonuclease (**Figure 14.1**). Methylation occurs rapidly after replication, catalyzed by site-specific methylases produced by the organism. Each restriction endonuclease will cleave a foreign DNA molecule into a fixed number of fragments, the number depending on the number of restriction sites in the particular DNA molecule. Test your understanding of restriction endonucleases by working through Solve It: How Many *Not*I Restriction Fragments in Chimpanzee DNA?

An interesting feature of restriction endonucleases is that they commonly recognize DNA sequences that are **palindromes**—that is, nucleotide-pair sequences that read the same forward or backward from a central axis of symmetry, as in the nonsense phrase

AND MADAM DNA

In addition, a useful feature of many restriction nucleases is that they make staggered cuts; that is, they cleave the two strands of a double helix at different points (Figure 14.1). (Other restriction endonucleases cut both strands at the same place and produce bluntended fragments.) Because of the palindromic nature of the restriction sites, the staggered



Sequence-specific cleavage of DNA by EcoRI and protection from cleavage by methylation.

unmethylated *Eco*RI recognition sequence by *Eco*RI restriction endonuclease and protection of the recognition sequence from cleavage by methylation catalyzed by the *Eco*RI methylase. (*b*) Diagram of the structure of the *Eco*RI-DNA complex based on X-ray diffraction data. The two subunits of the *Eco*RI endonuclease are shown in red and blue.

(b) Structure of an EcoRI-DNA complex based on X-ray diffraction data.

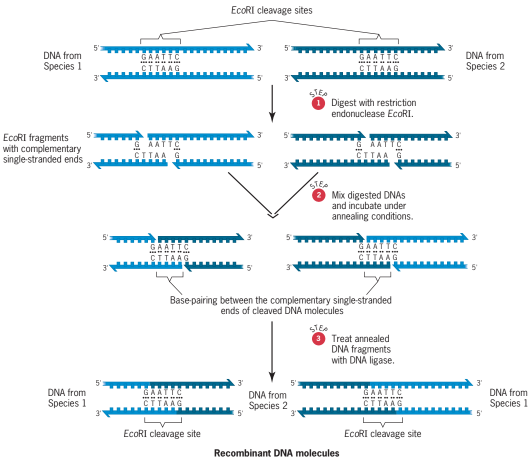
cuts produce segments of DNA with complementary single-stranded ends. For example, cleaving a DNA molecule of the following type:



with the restriction endonuclease EcoRI will yield



Because all the resulting DNA fragments will have complementary singlestranded termini, they will hydrogen bond with each other and can be rejoined under the appropriate renaturation conditions by using the enzyme **DNA ligase** to re-form the missing phosphodiester linkages in each strand (see Chapter 10). Thus, DNA



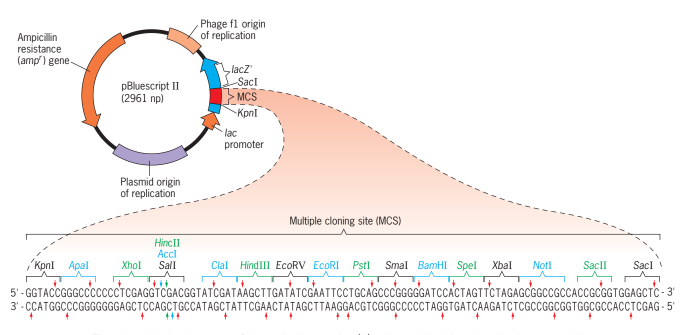
■ FIGURE 14.2 The construction of recombinant DNA molecules *in vitro*. DNA molecules isolated from two different species are cleaved with a restriction enzyme, mixed under annealing conditions, and are covalently joined by treatment with DNA ligase. The DNA molecules can be obtained from any species—animal, plant, or microbe. The digestion of DNA with the restriction enzyme *Eco*RI produces the same complementary single-stranded 5'-AATT-3' ends regardless of the source of the DNA.

molecules can be cut into pieces, called **restriction fragments**, and the pieces can be joined together again with DNA ligase, almost at will.

THE PRODUCTION OF RECOMBINANT DNA MOLECULES IN VITRO

A restriction endonuclease catalyzes the cleavage of a specific sequence of nucleotide pairs regardless of the source of the DNA. It will cleave phage DNA, *E. coli* DNA, corn DNA, human DNA, or any other DNA, as long as the DNA contains the nucleotide sequence that it recognizes. Thus, restriction endonuclease *Eco*RI will produce fragments with the same complementary single-stranded ends, 5'-AATT-3', regardless of the source of DNA, and two *Eco*RI fragments can be covalently fused regardless of their origin; that is, an *Eco*RI fragment from human DNA can be joined to an *Eco*RI fragment from *E. coli* DNA just as easily as two *Eco*RI fragments from *E. coli* DNA or two *Eco*RI fragments from human DNA can be joined. A DNA molecule of the type shown in **Figure 14.2**, containing DNA fragments from two different sources, is referred to as a recombinant DNA molecule. The ability of geneticists to construct such recombinant DNA molecules at will is the basis of the recombinant DNA technology that has revolutionized molecular biology in the last three decades.

The first recombinant DNA molecules were produced in Paul Berg's laboratory at Stanford University in 1972. Berg's research team constructed recombinant DNA molecules that contained phage lambda genes inserted into the small circular DNA molecule of simian virus 40 (SV40). In 1980, Berg was a co-recipient of the Nobel Prize in Chemistry as a result of this accomplishment. Shortly thereafter, Stanley Cohen and colleagues, also at Stanford, inserted an *Eco*RI restriction fragment from one DNA molecule into the cleaved, unique *Eco*RI restriction site of a self-replicating plasmid. When this recombinant plasmid was introduced into *E. coli* cells by transformation, it exhibited autonomous replication, just like the original plasmid.



■ FIGURE 14.3 The plasmid cloning vector Bluescript II contains (1) a plasmid origin of replication controlling doublestranded DNA synthesis, (2) a phage f1 origin of replication controlling single-stranded DNA synthesis, (3) an ampicillinresistance gene (*amp'*) that serves as a dominant selectable marker, (4) the promoter for the *lac* genes and the promoterproximal segment (*Z'*) of the *lacZ* gene, and (5) a polylinker or multiple cloning site (MCS) containing a cluster of unique restriction enzyme cleavage sites (18 are shown). The MCS is located within the *lacZ'* gene segment; therefore, when foreign DNA is inserted into the MCS, it disrupts LacZ' function. The designators and brackets showing the locations of recognition sequences for the restriction enzymes are above the MCS DNA sequence. The cleavage sites are marked with red arrows except for *Accl* and *HinclI*, where they are marked with blue and green arrows, respectively.

AMPLIFICATION OF RECOMBINANT DNA MOLECULES IN CLONING VECTORS

The various applications of recombinant DNA techniques require not only the construction of recombinant DNA molecules, as shown in Figure 14.2, but also the **amplification** of these recombinant molecules; that is, the production of many copies or **clones** of these molecules. This is accomplished by making sure that one of the parental DNAs incorporated into the recombinant DNA molecule is capable of self-replication. In practice, the gene or DNA sequence of interest is inserted into a specially chosen cloning vector. Most of the commonly used cloning vectors have been derived from plasmids or bacteriophage chromosomes (Chapter 8).

A cloning vector has three essential components: (1) an origin of replication, (2) a **dominant selectable marker gene**, usually a gene that confers drug resistance to the host cell, and (3) at least one *unique restriction endonuclease cleavage site*—a cleavage site that is present only once in a region of the vector that does not disrupt either the origin of replication or the selectable marker gene (**Trigure 14.3**). Modern cloning vectors contain a cluster of unique restriction sites called a **polylinker** or a **multiple cloning site** (Figure 14.3).

Many cloning vectors are modified versions of plasmids, the extrachromosomal, double-stranded circular molecules of DNA present in bacteria (Chapter 8). Plasmids range from about 1 kb (1 kilobase = 1000 base pairs) to over 200 kb in size, and many replicate autonomously. Many plasmids also carry antibiotic-resistance genes, which are ideal selectable markers.

A limiting factor in using plasmid vectors is that they will only accept relatively small foreign DNA inserts—maximum sizes of 10–15 kb. Thus, scientists searched for vectors that could replicate even when very large inserts were present. Some of these vectors are listed in Table 14.2, along with the maximum sizes of inserts that they would accept. Phage lambda vectors were widely used for several years; then more sophisticated vectors were constructed by combining components from viruses and plasmids. Phagemids combine components of *phage* such as M13 with parts of plasmids. Cosmids

TABLE 14.2							
Selected Cloning Vectors and Maximum Insert Sizes							
Vector	Maximum Insert Size						
Plasmids Phagemids Phage lambda Cosmids Bacterial artificial chromosomes (BACs) Phage P1 artificial chromosomes (PACs) Yeast artificial chromosomes (YACs)	15 kb 15 kb 23 kb 44 kb 300 kb 300 kb 600 kb						

contain the cohesive ends (*cos* sites) of lambda (see Figure 10.8) in plasmids. Yeast artificial chromosomes (YACs) are linear minichromosomes containing just the essential parts of yeast chromosomes—the origin of replication, centromere, and telomeres—along with a selectable marker and a multiple cloning site. Bacterial artificial chromosomes (BACs) and P1 artificial chromosomes (PACs) combine multiple cloning sites and selectable marker genes with the essential components of *bac*terial fertility (F) factors and phage *P*1 chromosomes, respectively. YACs, BACs, and PACs accept much larger foreign DNA inserts than plasmids and phage lambda cloning vectors (Table 14.2).

Bluescript (Figure 14.3) is a phagemid vector with a multiple cloning site (MCS) that contains many unique restriction enzyme cleavage sites, two distinct origins of replication, and a good selectable marker—a gene that makes the host bacterium resistant to ampicillin. The MCS is located within the 5' portion of the coding region of the *lacZ* gene, which encodes β -galactosidase, the enzyme that catalyzes the first step in the catabolism of lactose (Chapter 18). When foreign DNA is inserted into one of the restriction sites in the MCS, it disrupts the function of the plasmid-encoded *lacZ* product. This inactivation of the amino-terminal segment of β -galactosidase provides a good visual test for determining whether or not the Bluescript plasmid in a cell contains a foreign DNA insert.

The basis for this visual test is as follows. The presence of β -galactosidase in cells can be monitored based on its ability to cleave the substrate 5-bromo-4-chloro-3-

indolyl- β -D-galactoside (usually called X-gal) to galactose and 5-bromo-4chloroindigo. X-gal is colorless; 5-bromo-4-chloroindigo is blue. Thus, cells containing active β -galactosidase produce blue colonies on agar medium containing X-gal, whereas cells lacking β -galactosidase activity produce white colonies on X-gal plates (**\blacksquare Figure 14.4**).

The molecular basis of the β -galactosidase activity that provides the color indicator test for Bluescript vectors is somewhat more complex. The *lacZ* gene of *E. coli* is over 3 kb long, and placing the entire gene in the plasmid would make the vector larger than desired. The Bluescript vector contains only a small part of the *lacZ* gene. This *lacZ'* gene segment encodes only the amino-terminal portion of β -galactosidase. However, the presence of a functional copy of the *lacZ'* gene segment can be detected because of a unique type of complementation. When a functional copy of the *lacZ'* gene segment in a cell that contains a particular *lacZ* mutant allele on the chromosome or on an F' plasmid, the two defective *lacZ* sequences yield polypeptides that together have β -galactosidase activity. The mutant allele, designated *lacZ* $\Delta M15$, synthesizes a Lac protein that lacks amino acids 11 through 14 from the amino terminus. The absence of these amino acids prevents the mutant polypeptides from interacting to produce the active tetrameric form of the enzyme.

The presence of the amino-terminal fragment (the first 147 amino acids) of the lacZ polypeptide encoded by the *lacZ'* gene fragment on Bluescript plasmids facilitates tetramer formation by the $\Delta M15$ deletion polypeptides. This yields active β -galactosidase, which permits the X-gal color test to be utilized without placing the entire *lacZ* gene in the pBluescript vector.



FIGURE 14.4 Photograph illustrating the use of X-gal to identify *E. coli* colonies containing (blue) or lacking (white) β -galactosidase activity. In this case, the cells in the white colonies harbor Bluescript plasmids with foreign DNA fragments inserted into the multiple cloning site, and the cells in the blue colonies contain Bluescript plasmids with no insert.

CLONING LARGE GENES AND SEGMENTS OF GENOMES IN BACs, PACs, AND YACs

Some eukaryotic genes are very large. For example, the gene for human dystrophin (a protein that links filaments to membranes in muscle cells) is over 2000 kb in length. Research on large genes and chromosomes is much easier using vectors that accept large foreign DNA inserts, namely, BACs, PACs, and YACs (see Table 14.2). These vectors accept inserts of size 300 to 600 kb. BACs and PACs are less complex and easier to construct and work with than YACs. In addition, BACs and PACs replicate in E. coli like plasmid vectors. Thus, BAC and PAC vectors have largely replaced YAC vectors in the studies of large genes and genomes such as those of mammals and flowering plants.

PAC vectors have been constructed that permit negative selection against vectors lacking foreign DNA inserts. These PAC vectors contain the sacB gene of Bacillus subtilis. This gene encodes the enzyme levan sucrase, which catalyzes the transfer of fructose groups to various carbohydrates. The presence of this enzyme is lethal to E. coli cells when grown in medium containing 5 percent sucrose. The inactivation of the sacB gene by the insertion of foreign DNA in a BamHI restriction site in the gene can be used to select vectors containing inserts. Cells containing vectors with inserts can grow on medium containing 5 percent sucrose; cells with vectors lacking inserts cannot grow on this medium. Cells containing vectors lacking inserts lyse during the first hour of growth in the presence of 5 percent sucrose. As a result, all surviving cells contain vectors with inserts located within the *sacB* gene—inserts that eliminate levan sucrase activity.

PAC and BAC vectors have been modified to produce shuttle vectors that can replicate both in E. coli and in mammalian cells. The structure of one of these vectors is shown in Figure 14.5. This shuttle vector, pJCPAC-Mam1, contains the *sacB* gene, which allows for positive selection of cells carrying vectors with inserts, plus the origin of replication (oriP) and the gene encoding nuclear antigen 1 of the Epstein-Barr virus, which facilitate replication of the vector in mammalian cells. In addition, the pur^r

(puromycin-resistance) gene has been added so that mammalian cells carrying the vector can be selected on medium containing the antibiotic

plasmid replication regulatory puromycin. Similar BAC shuttle vectors have also been constructed.

AMPLIFICATION OF DNA SEQUENCES BY THE POLYMERASE CHAIN REACTION (PCR)

Today, we have complete or nearly complete nucleotide sequences of many genomes, including the human genome. The availability of these sequences in GenBank and other databases allows researchers to isolate genes or other DNA sequences of interest without using cloning vectors or host cells. The amplification of the DNA sequence is performed entirely in vitro, and the sequence can be amplified a millionfold or more in just a few hours. All that is required to use this procedure is knowledge of short nucleotide sequences flanking the sequence of interest. This in vitro amplification of genes and other DNA sequences is accomplished by the polymerase chain reaction (usually referred to as **PCR**). PCR involves using synthetic oligonucleotides complementary to known sequences flanking the sequence of interest to prime enzymatic amplification of the intervening segment of DNA in the test tube. The PCR procedure for amplifying DNA sequences was developed by Kary Mullis, who received the 1993 Nobel Prize in Chemistry for this work.

The PCR procedure involves three steps, each repeated many times (**Figure 14.6**). In step 1, the genomic DNA containing the sequence to be amplified is denatured by heating to 92-95°C for about 15 seconds. In step 2, the denatured DNA is annealed to an excess of the synthetic oligonucleotide primers by incubating

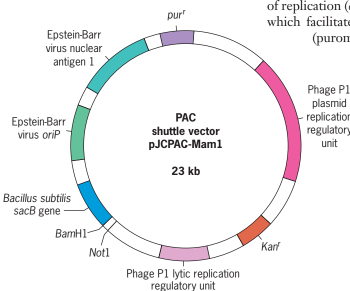


FIGURE 14.5 Structure of the PAC mammalian shuttle vector pJCPAC-Mam1. The vector can replicate in either E. coli or mammalian cells. It can replicate in *E. coli* at low copy number under the control of the bacteriophage P1 plasmid replication unit or be amplified by inducing the phage P1 lytic replication unit (under the control of the lac inducible promoter; see Chapter 18). It can replicate in mammalian cells by using the origin of replication (oriP) and nuclear antigen 1 of the Epstein-Barr virus. Genes kan^r and *pur^r* provide dominant selectable markers for use in *E. coli* and mammalian cells, respectively. The sacB gene (derived from Bacillus subtilis) is used for negative selection against vectors lacking DNA inserts (see text for details). BamHI and NotI are cleavage sites for these two restriction endonucleases.

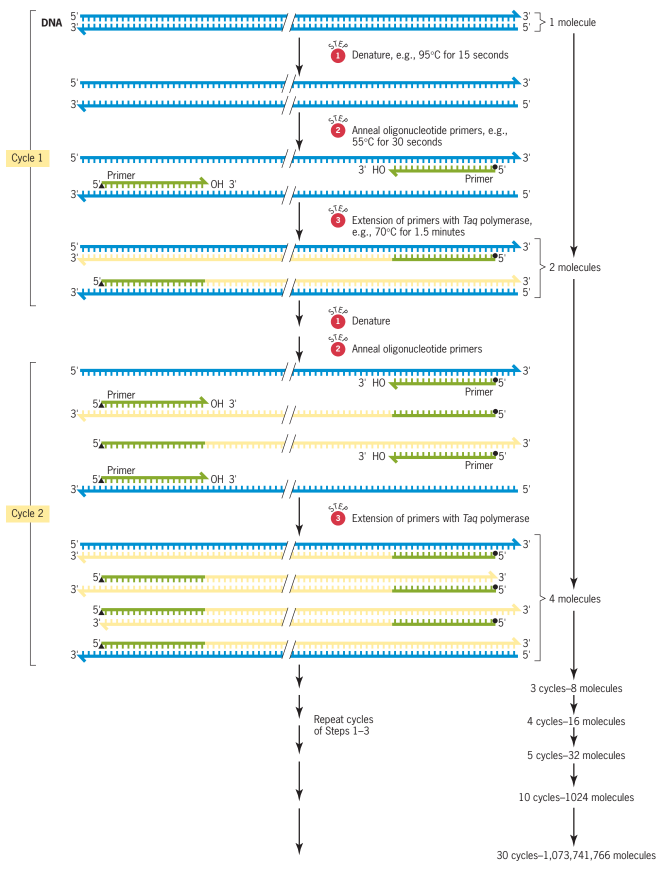


FIGURE 14.6 The use of PCR to amplify DNA molecules *in vitro*. Each cycle of amplification involves three steps: (1) denaturation of the genomic DNA being analyzed, (2) annealing of the denatured DNA to chemically synthesized oligonucleotide primers with sequences complementary to sites on opposite sides of the DNA region of interest, and (3) enzymatic replication of the region of interest by *Taq* polymerase.

them together at 50–60°C for 30 seconds. The ideal annealing temperature depends on the base composition of the primer. In step 3, DNA polymerase is used to replicate the DNA segment between the sites complementary to the oligonucleotide primers. The primer provides the free 3'-OH required for covalent extension, and the denatured genomic DNA provides the required template function (Chapter 10). Polymerization is usually carried out at 70–72°C for 1.5 minutes. The products of the first cycle of replication are then denatured, annealed to oligonucleotide primers, and replicated again with DNA polymerase. The procedure is repeated many times until the desired level of amplication is achieved. Note that *amplification occurs geometrically*. One DNA double helix will yield 2 double helices after one cycle of replication, 4 after two cycles, 8 after three cycles, 16 after four cycles, 1024 after ten cycles, and so on. After 30 cycles of amplification, more than a billion copies of the DNA sequence will have been produced.

Initially, PCR was performed with DNA polymerase I of *E. coli* as the replicase. Because this enzyme is heat-inactivated during the denaturation step, fresh enzyme had to be added at step 3 of each cycle. A major improvement in PCR amplification of DNA came with the discovery of a heat-stable DNA polymerase in the thermophilic bacterium, *Thermus aquaticus*. This polymerase, called *Taq* polymerase (*<u>T</u>. <u>aquaticus</u> polymerase), remains active during the heat denaturation step. As a result, polymerase does not have to be added after each cycle of denaturation. Instead, excess <i>Taq* polymerase and oligonucleotide primers can be added at the start of the PCR process, and amplification cycles can be carried out by sequential alterations in temperature. PCR machines or thermal cyclers change the temperature automatically and hold large numbers of samples, making PCR amplification of specific DNA sequences a relatively simple task.

One disadvantage of PCR is that errors are introduced into the amplified DNA copies at low but significant frequencies. Unlike most DNA polymerases, *Taq* polymerase does not contain a built-in $3' \rightarrow 5'$ proofreading activity, and, consequently, it produces a higher than normal frequency of replication errors. If an incorrect nucleotide is incorporated during an early PCR cycle, it will be amplified just like any other nucleotide in the DNA sequence. When high fidelity is required, PCR is performed using heat-stable polymerases—such as *Pfu* (from *Pyrococcus furiosus*) or *Thi* (from *Thermococcus litoralis*)—that possess $3' \rightarrow 5'$ proofreading activity. A second disadvantage of *Taq* polymerase is that it amplifies long tracts of DNA—greater than a few thousand nucleotide pairs—inefficiently. If long segments of DNA need to be amplified, the more processive *Tfl* polymerase from *Thermus flavus* is used in place of *Taq* polymerase. *Tfl* polymerase that 35 kb cannot be efficiently amplified by PCR.

PCR technologies provide shortcuts for many applications that require large amounts of a specific DNA sequence. These procedures permit scientists to obtain definitive structural data on genes and DNA sequences when very small amounts of DNA are available. One important application occurs in the diagnosis of inherited human diseases, especially in cases of prenatal diagnosis, where limited amounts of fetal DNA are available. A second major application occurs in forensic cases involving the identification of individuals by using DNA isolated from very small tissue samples. Few criteria can provide more definitive evidence of identity than DNA sequences. By using PCR amplification, DNA sequences can be obtained from minute amounts of DNA isolated from a few drops of blood, semen, or even individual human hairs. Thus, *PCR DNA profiling (fingerprinting)* experiments play important roles in legal cases involving uncertain identity. Some of the applications of PCR are discussed in Chapter 16.

KEY POINTS

 The discovery of restriction endonucleases—enzymes that recognize and cleave DNA in a sequence-specific manner—allowed scientists to produce recombinant DNA molecules in vitro.

- DNA sequences can be inserted into small, self-replicating DNA molecules called cloning vectors and amplified by replication in vivo after being introduced into living cells by transformation.
- The polymerase chain reaction (PCR) can be used to amplify specific DNA sequences in vitro.

Construction and Screening of DNA Libraries

The first step in cloning a gene from an organism usually involves the construction of a **genomic DNA library**—a set of DNA clones collectively containing the entire genome. Sometimes, individual chromo-

somes of an organism are isolated by a procedure that sorts chromosomes based on size and DNA content. The DNAs from the isolated chromosomes are then used to construct chromosome-specific DNA libraries. The availability of chromosomespecific DNA libraries facilitates the search for a gene that is known to reside on a particular chromosome, especially for organisms like humans with large genomes. After their construction, libraries are amplified by replication and used to identify individual genes or DNA sequences of interest to the researcher.

An alternative approach to gene cloning restricts the search for a gene to DNA sequences that are transcribed into mRNA copies. The RNA retroviruses (Chapter 17) encode an enzyme called reverse transcriptase, which catalyzes the synthesis of DNA molecules complementary to single-stranded RNA templates. These DNA molecules are called **complementary DNAs (cDNAs)**. They can be converted to

double-stranded cDNA molecules with DNA polymerases (Chapter 10), and the double-stranded cDNAs can be cloned in plasmid vectors. By starting with mRNA, geneticists are able to construct **cDNA** libraries that contain only the coding regions of the expressed genes of an organism.

CONSTRUCTION OF GENOMIC LIBRARIES

Genomic DNA libraries are usually prepared by isolating total DNA from an organism, digesting the DNA with a restriction endonuclease, and inserting the restriction fragments into an appropriate cloning vector. If the restriction enzyme that is used makes staggered cuts in DNA, producing complementary singlestranded ends, the restriction fragments can be ligated directly into vector DNA molecules cut with the same enzyme (**Figure 14.7**). When this procedure is used, the foreign DNA inserts can be excised from the vector DNA by cleavage with the restriction endonuclease used to prepare the genomic DNA fragments for cloning.

Once the genomic DNA fragments are ligated into vector DNA, the recombinant DNA molecules must be introduced into host cells for amplification by replication *in vivo*. This step usually involves transforming antibiotic-sensitive recipient cells under conditions where a single recombinant DNA molecule is introduced per cell (for most cells) (Chapter 8). When *E. coli* is used, the bacteria must first be made permeable to DNA by treatment with chemicals or a short pulse of electricity. Transformed cells are then selected by growing the cells under conditions where the selectable marker gene of the vector is essential for growth.

A good genomic DNA library contains essentially all of the DNA sequences in the genome of interest. For large genomes, complete libraries contain hundreds of thousands of different recombinant clones.

DNA libraries can be constructed and screened for genes and other sequences of interest.

Isolate *E.coli* plasmid Bluescript II DNA and mouse genomic DNA.

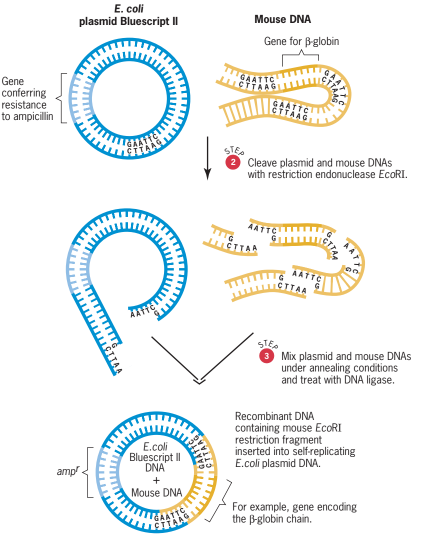
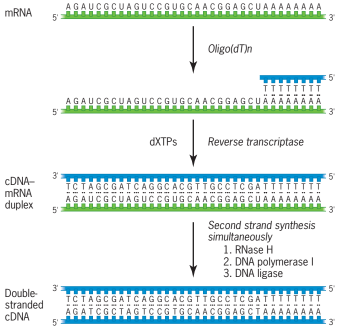
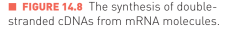


FIGURE 14.7 Procedure used to clone DNA restriction fragments with complementary single-stranded ends.





CONSTRUCTION OF cDNA LIBRARIES

Most of the DNA sequences present in the large genomes of higher animals and plants do not encode proteins. Thus, expressed DNA sequences can be identified more easily by working with complementary DNA (cDNA) libraries. Because most mRNA molecules contain 3' poly(A) tails, poly(T) oligomers can be used to prime the synthesis of complementary DNA strands by reverse transcriptase (**Figure 14.8**). Then, the RNA–DNA duplexes are converted to doublestranded DNA molecules by the combined activities of ribonuclease H, DNA polymerase I, and DNA ligase. Ribonuclease H degrades the RNA template strand, and short RNA fragments produced during degradation serve as primers for DNA synthesis. DNA polymerase I catalyzes the synthesis of the second DNA strand and replaces RNA primers with DNA strands, and DNA ligase seals the remaining single-strand breaks in the double-stranded DNA molecules. These double-stranded cDNAs can be inserted into plasmid or phage λ cloning vectors by adding complementary single-stranded tails to the cDNAs and vectors.

SCREENING DNA LIBRARIES FOR GENES OF INTEREST

The genomes of higher plants and animals are very large. For example, the human genome contains 3×10^9 nucleotide pairs. Thus, searching genomic DNA or cDNA libraries of multicellular eukaryotes for a specific gene or other DNA sequence of interest requires the identification of a single DNA sequence in a library that contains a million or more different sequences. The most powerful screening procedure is genetic selection: searching for a DNA sequence in the library that can restore the wild-type phenotype to a mutant organism. When genetic selection cannot be employed, more laborious molecular screens must be carried out. Molecular screens usually involve the use of DNA or RNA sequences as hybridization probes or the use of antibodies to identify gene products encoded by cDNA clones.

Genetic Selection

The simplest procedure for identifying a clone of interest is **genetic selection**. For example, the *Salmonella typhimurium* gene that confers resistance to penicillin can be easily cloned. A genomic library is constructed from the DNA of a *pen^r* strain of *S. typhimurium*. Penicillin-sensitive *E. coli* cells are transformed with the recombinant DNA clones in the library and are plated on medium containing penicillin. Only the transformed cells harboring the *pen^r* gene will be able to grow in the presence of penicillin.

When mutations are available in the gene of interest, genetic selection can be based on the ability of the wild-type allele of a gene to restore the normal phenotype to a mutant organism. Although this type of selection is called **complementation screening**, it really depends on the dominance of wild-type alleles over mutant alleles that encode inactive products. For example, the genes of *S. cerevisiae* that encode histidinebiosynthetic enzymes were cloned by transforming *E. coli* histidine auxotrophs with yeast cDNA clones and selecting transformed cells that could grow on histidine-free medium. Indeed, many plant and animal genes have been identified based on their ability to complement mutations in *E. coli* or yeast.

Complementation screening has limitations. Eukaryotic genes contain introns, which must be spliced out of gene transcripts prior to their translation. Because *E. coli* cells do not possess the machinery required to excise introns from eukaryotic genes, complementation screening of eukaryotic clones in *E. coli* is restricted to cDNAs, from which the intron sequences have already been excised. In addition, the complementation screening procedure depends on the correct transcription of the cloned gene in the new host. Eukaryotes have signals that regulate gene expression that are different from those in prokaryotes; therefore, the complementation approach is more

likely to work with prokaryotic genes in prokaryotic organisms, and eukaryotic genes in eukaryotic organisms. For this reason, researchers often use *S. cerevisiae* to screen eukaryotic DNA libraries by the complementation procedure.

Molecular Hybridization

The first eukaryotic DNA sequences to be cloned were genes that are highly expressed in specialized cells. These genes included the mammalian α - and β -globin genes and the chicken ovalbumin gene. Red blood cells are highly specialized for the synthesis and storage of hemoglobin. Over 90 percent of the protein molecules synthesized in red blood cells during their period of maximal biosynthetic activity are globin chains. Similarly, ovalbumin is a major product of chicken oviduct cells. As a result, RNA transcripts of the globin and ovalbumin genes can be easily isolated from reticulocytes and oviduct cells, respectively. These RNA transcripts can be employed to synthesize radioactive cDNAs, which, in turn, can be used to screen genomic DNA libraries by *in situ* colony or plaque hybridization (**I Figure 14.9**). Colony hybridization is used with libraries constructed in plasmid and cosmid vectors; plaque hybridization is used with

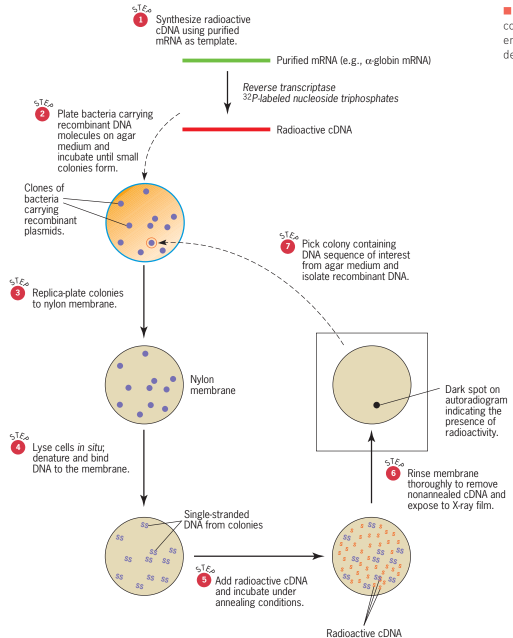


FIGURE 14.9 Screening DNA libraries by colony hybridization. A radioactive cDNA is employed as a hybridization probe. See text for details.

Solve It!

How Can You Clone a Specific Notl Restriction Fragment from the Orangutan Genome?

You are studying what appears to be an inherited disorder in the Sumatran orangutan (*Pongo abelii*), and you want to clone a 95-kb *Not*l restriction fragment from the orangutan that cross-hybridizes with a specific human gene. You have pBluescript II and pJCPAC-Mam1 DNAs available to use as cloning vectors. Which vector would you use to clone the *Not*I fragment of interest, and how would you proceed to construct and identify the clone of interest?

To see the solution to this problem, visit the Student Companion site. libraries in phage lambda vectors. We will focus on *in situ* colony hybridization here, but the two procedures are virtually identical.

The colony hybridization screening procedure involves transfer of the colonies formed by transformed cells onto nylon membranes, hybridization with a radioactively labeled DNA or RNA probe, and autoradiography (Figure 14.9). The labeled DNA or RNA is employed as a probe for hybridization (see Appendix C: In Situ Hybridization) to denatured DNA from colonies grown on the nylon membranes. The DNA from the lysed cells is bound to the membranes before hybridization so that it won't come off during subsequent steps in the procedure. After time is allowed for hybridization between complementary strands of DNA, the membranes are washed with buffered salt solutions to remove nonhybridized cDNA and are then exposed to X-ray film to detect the presence of radioactivity on the membrane. Only colonies that contain DNA sequences complementary to the radioactive cDNA will yield radioactive spots on the autoradiographs (Figure 14.9). The locations of the radioactive spots are used to identify colonies that contain the desired sequence on the original replicated plates. These colonies are used to purify DNA clones harboring the gene or DNA sequence of interest. Test your comprehension of the methods used to prepare and screen genomic libraries by working Solve It: How Can You Clone a Specific NotI Restriction Fragment from the Orangutan Genome?

KEY POINTS

- DNA libraries can be constructed that contain complete sets of genomic DNA sequences or DNA copies (cDNAs) of mRNAs in an organism.
- Specific genes or other DNA sequences can be isolated from DNA libraries by genetic complementation or by hybridization to labeled nucleic acid probes containing sequences of known function.

The Molecular Analysis of DNA, RNA, and Protein

DNA, RNA, or protein molecules can be separated by gel electrophoresis, transferred to membranes, and analyzed by various procedures.

The development of recombinant DNA techniques has spawned many new approaches to the analysis of genes and gene products. Questions that were totally unapproachable just 25 years ago can now be investigated with relative ease. Geneticists can isolate and

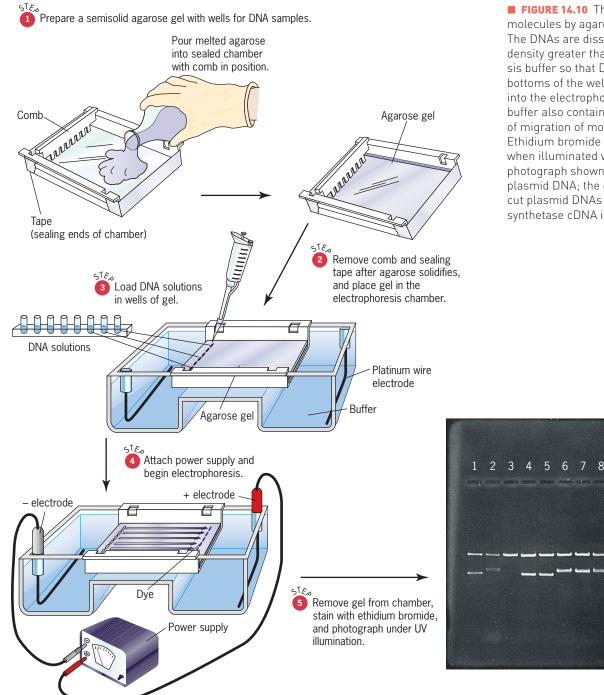
characterize essentially any gene from any organism; however, the isolation of genes from large eukaryotic genomes is sometimes a long and laborious process (Chapter 16). Once a gene has been cloned, its expression can be investigated in even the most complex organisms such as humans.

Is a particular gene expressed in the kidney, the liver, bone cells, hair follicles, erythrocytes, or lymphocytes? Is this gene expressed throughout the development of the organism or only during certain stages of development? Is a mutant allele of this gene similarly expressed, spatially and temporally, during development? Or does the mutant allele have an altered pattern of expression? If the latter, is this altered pattern of expression responsible for an inherited syndrome or disease? These questions and many others can now be routinely investigated using well-established methodologies.

A comprehensive discussion of the techniques used to investigate gene structure and function is far beyond the scope of this text. However, let's consider some of the most important methods used to investigate the structure of genes (DNA), their transcripts (RNA), and their final products (usually proteins).

ANALYSIS OF DNAs BY SOUTHERN BLOT HYBRIDIZATIONS

Gel electrophoresis is a powerful tool for the separation of macromolecules with different sizes and charges. DNA molecules have an essentially constant charge per unit mass; thus, they separate in agarose and acrylamide gels almost entirely on the basis of size or conformation. Agarose or acrylamide gels act as molecular sieves, retarding the passage of large molecules more than small molecules. Agarose gels are better sieves for large molecules (larger than a few hundred nucleotides); acrylamide gels are better for separating small DNA molecules. **■** Figure 14.10 illustrates the separation of DNA restriction fragments by agarose gel electrophoresis. The procedures used



■ FIGURE 14.10 The separation of DNA molecules by agarose gel electrophoresis. The DNAs are dissolved in loading buffer with density greater than that of the electrophoresis buffer so that DNA samples settle to the bottoms of the wells, rather than diffusing into the electrophoresis buffer. The loading buffer also contains a dye to monitor the rate of migration of molecules through the gel. Ethidium bromide binds to DNA and fluoresces when illuminated with ultraviolet light. In the photograph shown, lane 3 contained *Eco*RI-cut plasmid DNA; the other lanes contained *Eco*RIcut plasmid DNAs carrying maize glutamine synthetase cDNA inserts.

> - 3.2 kb - 1.7 kb `1.5 kb

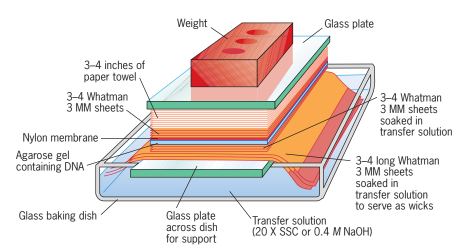


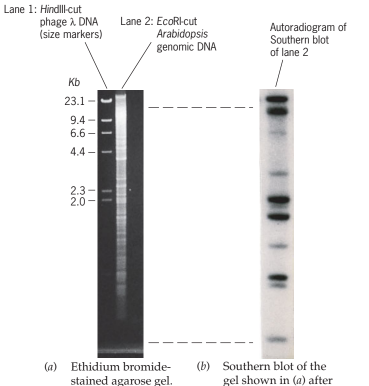
FIGURE 14.11 Procedure used to transfer DNAs separated by gel electrophoresis to nylon membranes. The transfer solution carries the DNA from the gel to the membrane as the dry paper towels on top draw the salt solution from the reservoir through the gel to the towels. The DNA binds to the membrane on contact. The membrane with the DNA bound to it is dried and baked under vacuum to affix the DNA firmly prior to hybridization. SSC is a solution containing sodium chloride and sodium citrate.

to separate RNA and protein molecules are largely the same in principle but involve slightly different techniques because of the unique properties of each class of macromolecule (see the section Variation in Protein Structure in Chapter 24).

In 1975, E. M. Southern published an important procedure that allowed investigators to identify the locations of genes and other DNA sequences on restriction fragments separated by gel electrophoresis. The essential feature of this technique is the transfer of the DNA molecules that have been separated by gel electrophoresis onto nitrocellulose or nylon membranes (**Figure 14.11**). Such transfers of DNA to membranes are called Southern blots after the scientist who developed the technique. The DNA is denatured either prior to or during transfer by placing the gel in an alkaline solution. After transfer, the DNA is immobilized on the membrane by drying or UV irradiation. A radioactive DNA probe containing the sequence of interest is then hybridized (see Appendix C: In Situ Hybridization) with the immobilized DNA on the membrane. The

probe will hybridize only with DNA molecules that contain a nucleotide sequence complementary to the sequence of the probe. Nonhybridized probe is then washed off the membrane, and the washed membrane is exposed to X-ray film to detect the presence of the radioactivity. After the film is developed, the dark bands show the positions of DNA sequences that have hybridized with the probe (**■** Figure 14.12).

The ability to transfer DNA molecules that have been separated by gel electrophoresis to nylon membranes for hybridization studies and other types of analyses has proven to be extremely useful (see Focus on Detection of a Mutant Gene Causing Cystic Fibrosis).



hybridization to a labeled

β-tubulin cDNA.

ANALYSIS OF RNAs BY NORTHERN BLOT HYBRIDIZATIONS

If DNA molecules can be transferred from agarose gels to nylon membranes for hybridization studies, we might expect that RNA molecules separated by agarose gel electrophoresis could be similarly transferred and analyzed. Indeed, such RNA transfers are used routinely in genetics laboratories. RNA blots are called **northern blots** in recognition of the fact that the procedure is analogous to the Southern blotting technique, but with RNA molecules being separated and transferred to a membrane. As we will discuss in the next section, this terminology has been extended to the transfer of proteins from gels to membranes, a procedure called western blotting.

FIGURE 14.12 Identification of genomic restriction fragments harboring specific DNA sequences by the Southern blot hybridization procedure. (a) Photograph of an ethidium bromide-stained agarose gel containing phage λ DNA digested with *Hin*dIII (left lane), and *Arabidopsis thaliana* DNA digested with *Eco*RI (right lane). The λ DNA digest provides size markers. The *A. thaliana* DNA digest was transferred to a nylon membrane by the Southern procedure (Figure 14.11) and hybridized to a radioactive DNA fragment of a cloned β -tubulin gene. The resulting Southern blot is shown in (*b*); nine different *Eco*RI fragments hybridized with the β -tubulin probe.

FOCUS ON

bases in the coding

amino acids in

DETECTION OF A MUTANT GENE CAUSING CYSTIC FIBROSIS

ystic fibrosis is characterized by the accumulation of mucus in the lungs, pancreas, and liver, and the subsequent malfunction of these organs. It is the most common inherited disease in humans of northern European descent. In Chapter 16, we discuss cystic fibrosis and the identification and characterization of the gene that causes it. Here, we will focus on the use of PCR to amplify the *CF* alleles in genomic DNA from members of families afflicted with this disease and the detection of the most common mutant allele by Southern blot hybridization to labeled oligonucleotide probes.

Approximately 70 percent of the cases of CF result from a specific mutant allele of the *CF* gene. This mutant allele, *CF* Δ *F508*, contains a three-base deletion that eliminates a phenylalanine residue at position 508 in the polypeptide product. Because the nucleotide sequence of the *CF* gene is known and since the *CF* Δ *F508* allele differs from the wild-type allele by three base pairs, it was possible to design oligonucleotide probes that hybridize specifically with the wild-type *CF* allele or the Δ *F508* allele under the appropriate conditions.

The wild-type CF gene and gene product have the following nucleotide and amino acid sequences in the region altered by the $\Delta F508$ mutation:

strand: 5'-AAA GAA AAT ATC ATC TTT GGT GTT-3'

product: NH2-Lys Glu Asn Ile Ile Phe Gly Val-COOH

deleted in $\Delta F508$

whereas the Δ *F508* allele and product have these sequences:

bases in the coding deletion strand: 5'-AAA GAA AAT ATC AT. . . T GGT GTT-3'

amino acids in product: NH_2 -Lys Glu Asn Ile Ile — Gly Val-COOH

Phe absent

Based on these nucleotide sequences, Lap-Chee Tsui and colleagues synthesized oligonucleotides spanning this region of the mutant and wild-type alleles of the *CF* gene and tested their specificity. They demonstrated that at 37°C under a standard set of conditions, one oligonucleotide probe (oligo-N: 3'-CTTTTATAG-TAGAAACCAC-5') hybridized only with the wild-type allele, whereas another (oligo- Δ F: 3'-TTCTTTATAGTA... ACCACAA-5') hybridized only with the Δ F508 allele. Their results showed that the oligo- Δ F probe could be used to detect the Δ F508 allele in either the homozygous or heterozygous state. When Tsui and coworkers used these allele-specific oligonucleotide probes to analyze CF patients and their parents for the presence of the Δ F508 mutation, they found that many of the patients were homozygous for this mutation, whereas most of their parents were heterozygous, as is expected. Some of their results are shown in **Figure 1**.

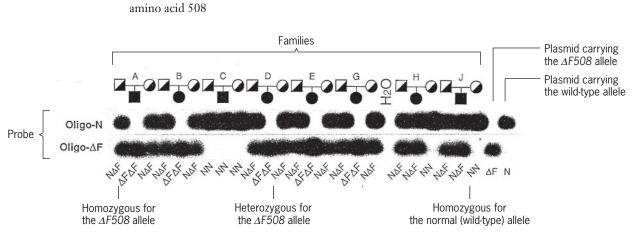


FIGURE 1 Detection of *CF* wild-type and $\Delta F508$ alleles by hybridization of labeled allele-specific oligonucleotide probes to genomic DNAs transferred to nylon membranes by the Southern blotting procedure (Figure 14.11). PCR was used to amplify the *CF* loci in genomic DNAs isolated from individual family members. The PCR products were separated by gel electrophoresis, transferred to membranes, denatured, and hybridized to the radioactive oligonucleotide probes (described above). Duplicate Southern blots were prepared; one blot was hybridized to the probe specific for the wild-type *CF* allele (top lane), and the other was hybridized to the probe specific for the $\Delta F508$ allele (bottom lane). The family pedigrees shown at the top represent offspring with *CF* and their heterozygous parents. Note that the $\Delta F508$ allele is present in families A, B, D, E, and G. Family C carries a different *CF* allele, and families H and J have one parent with the $\Delta F508$ allele and the other parent with a different *CF* allele. The lane labeled H₂O is a control containing only water. In the pedigrees at the top, filled symbols represent individuals who carry two mutant *CF* alleles, and half-filled symbols represent individuals who carry mutant and wild-type *CF* alleles.

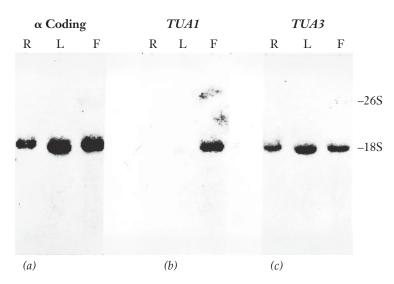


FIGURE 14.13 Typical northern blot hybridization data. Total RNAs were isolated from roots (R), leaves (L), and flowers (F) of *A. thaliana* plants, separated by agarose gel electrophoresis, and then transferred to nylon membranes. The autoradiogram shown in (*a*) is of a blot that was hybridized to a radioactive probe containing an α -tubulin coding sequence. This probe hybridizes to the transcripts of all six α -tubulin genes in *A. thaliana*. The autoradiograms shown in (*b*) and (*c*) are of RNA blots that were hybridized to DNA probes specific for the α 1- and α 3-tubulin genes (*TUA1* and *TUA3*, respectively). The results show that the α 3-tubulin transcript is present in all organs analyzed, whereas the α 1-tubulin transcript is present only in flowers. The 18S and 26S ribosomal RNAs provide size markers. Their positions were determined from a photograph of the ethidium-bromide stained gel prior to transfer of the RNAs to the nylon membrane.

The northern blot procedure is essentially identical to that used for Southern blot transfers (Figure 14.11). However, RNA molecules are very sensitive to degradation by RNases. Thus, care must be taken to prevent contamination of materials with these extremely stable enzymes. Furthermore, most RNA molecules contain considerable secondary structure and must therefore be kept denatured during electrophoresis in order to separate them on the basis of size. Denaturation is accomplished by adding formaldehyde or some other chemical denaturant to the buffer used for electrophoresis. After transfer to an appropriate membrane, the RNA blot is hybridized to either RNA or DNA probes just as with a Southern blot.

Northern blot hybridizations (**■** Figure 14.13) are extremely helpful in studies of gene expression. They can be used to determine when and where a particular gene is expressed. However, we must remember that northern blot hybridizations only measure the accumulation of RNA transcripts. They provide no information about why the observed accumulation has occurred. Changes in transcript levels may be due to changes in the rate of transcription or to changes in the rate of transcript degradation. More sophisticated procedures must be used to distinguish between these possibilities.

ANALYSIS OF RNAs BY REVERSE TRANSCRIPTASE-PCR (RT-PCR)

The enzyme reverse transcriptase catalyzes the synthesis of DNA strands that are complementary to RNA templates. It can be used *in vitro* to synthesize DNAs that are complementary to RNA template strands. The resulting DNA strands can

then be converted to double-stranded DNA by several different procedures (for example, see Figure 14.8), including the use of a second primer and the heat-stable *Taq* DNA polymerase. The resulting DNA molecules can then be amplified by standard PCR [see the section Amplification of DNA Sequences by the Polymerase Chain Reaction (PCR) earlier in this chapter].

The first strand of DNA, often called a **cDNA** because it is complementary to the mRNA under study, can be synthesized by using an oligo(dT) primer that will anneal to the 3'-poly(A) tails of all mRNAs, or by using gene-specific primers (sequences complementary to the RNA molecule of interest). Gene-specific oligonucleotide primers are usually chosen to anneal to sequences in the 3'-noncoding regions of the mRNAs. **Figure 14.14** illustrates how such primers can be used in RT-PCR to amplify a specific gene transcript. The products of these amplifications are analyzed by gel electrophoresis. Wherever a product appears in the gel, the investigator knows that the sample from which it was generated contained the mRNA under study. This procedure is therefore a quick and easy way of ascertaining whether or not a particular gene is being transcribed.

Many modifications of the RT-PCR procedure have been developed, with a major emphasis on making it more quantitative. For example, known amounts of the RNA under study can be analyzed to determine the relationship between RNA input and DNA output. By knowing this relationship, an investigator can use the quantity of DNA generated by an experimental sample to extrapolate back to the amount of RNA that was initially present in that sample.

ANALYSIS OF PROTEINS BY WESTERN BLOT TECHNIQUES

Polyacrylamide gel electrophoresis is an important tool for the separation and characterization of proteins. Because many functional proteins are composed of two or

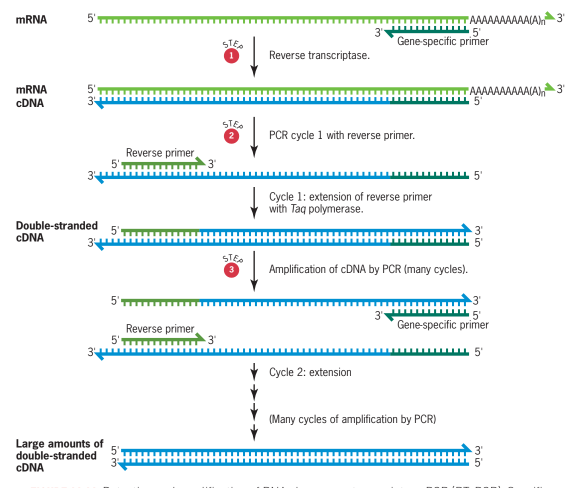


FIGURE 14.14 Detection and amplification of RNAs by reverse transcriptase PCR (RT-PCR). Specific gene transcripts are amplified by first using reverse transcriptase to synthesize a single-stranded DNA that is complementary to the mRNA of interest. The synthesis is initiated with a gene-specific oligonucleotide primer (a primer that will only anneal to the mRNA of interest). The complementary DNA strand is then synthesized by using a reverse primer and *Taq* polymerase. Large quantities of double-stranded cDNA are subsequently synthesized by standard PCR reactions in the presence of both the gene-specific and reverse PCR primers.

more subunits, individual polypeptides are separated by electrophoresis in the presence of the detergent sodium dodecyl sulfate (SDS), which denatures the proteins. After electrophoresis, the proteins are detected by staining with Coomassie blue or silver stain. However, the separated polypeptides also can be transferred from the gel to a nitrocellulose membrane, and individual proteins can be detected with antibodies. This transfer of proteins from acrylamide gels to nitrocellulose membranes, called **western blotting**, is performed by using an electric current to move the proteins from the gel to the surface of the membrane.

After transfer, a specific protein of interest is identified by placing the membrane with the immobilized proteins in a solution containing an antibody to the protein. Nonbound antibodies are then washed off the membrane, and the presence of the initial (primary) antibody is detected by placing the membrane in a solution containing a secondary antibody. This secondary antibody reacts with immunoglobulins (the group of proteins comprising all antibodies) in general (Chapter 20). The secondary antibody is conjugated to either a radioactive isotope (permitting autoradiography) or an enzyme that produces a visible product when the proper substrate is added.

KEY POINTS • DNA restriction fragments and other small DNA molecules can be separated by agarose or acrylamide gel electrophoresis and transferred to nylon membranes to produce DNA gel blots called Southern blots.

- The DNAs on Southern blots can be hybridized to labeled DNA probes to detect sequences of interest by autoradiography.
- When RNA molecules are separated by gel electrophoresis and transferred to membranes for analysis, the resulting RNA gel blots are called northern blots.
- RNA molecules can be detected and analyzed by reverse transcriptase-PCR (RT-PCR).
- When proteins are transferred from gels to membranes and detected with antibodies, the products are called western blots.

The Molecular Analysis of Genes and Chromosomes

The sites at which restriction enzymes cleave DNA molecules can be used to construct physical maps of the molecules; however, nucleotide sequences provide the ultimate physical maps of DNA molecules.

Recombinant DNA techniques allow geneticists to determine the structure of genes, chromosomes, and entire genomes. Indeed, molecular geneticists have constructed detailed genetic and physical maps of the genomes of many organisms (Chapter 15).

The ultimate physical map of a genetic element is its nucleotide sequence, and the complete nucleotide sequences of the genomes of thousands of viruses, bacteria, mitochondria, chloroplasts, and numerous eukaryotic organisms have already been determined. In

October 2004, the International Human Genome Sequencing Consortium published a "nearly complete" sequence of the human genome. That sequence contained only 341 gaps and covered 99 percent of the gene-rich chromatin in the human genome (Chapter 15). In the following sections, we discuss the construction of restriction enzyme cleavage site maps of genes and chromosomes and the determination of DNA sequences.

PHYSICAL MAPS OF DNA MOLECULES BASED **ON RESTRICTION ENZYME CLEAVAGE SITES**

Most restriction endonucleases cleave DNA molecules in a site-specific manner (see Table 14.1). As a result, they can be used to generate **physical maps** of chromosomes that are of great value in assisting researchers in isolating DNA fragments carrying genes or other DNA sequences of interest. The sizes of the restriction fragments can be determined by polyacrylamide or agarose gel electrophoresis (see Figure 14.10). Because of the nucleotide subunit structure of DNA, with one phosphate group per nucleotide, DNA has an essentially constant charge per unit of mass. Thus, the rates of migration of DNA fragments during electrophoresis provide accurate estimates of their lengths, with the rate of migration inversely proportional to length.

The procedure that is used to map the restriction enzyme cleavage sites is illustrated in Figure 14.15. The sizes of DNA restriction fragments are estimated by using a set of DNA markers of known size. In Figure 14.15, a set of DNA molecules that differ in length by 1000 nucleotide pairs are used as size markers. Consider a DNA molecule approximately 6000 nucleotide pairs (6 kb) in length. When the 6-kb DNA molecule is cut with EcoRI, two fragments of sizes 4000 and 2000 nucleotide pairs are produced. The possible positions of the single *Eco*RI cleavage site in the molecule are shown in Figure 14.15b. When the same DNA molecule is cleaved with HindIII, two fragments of sizes 5000 and 1000 nucleotide pairs result.

The possible locations of the single *Hin*dIII cleavage site are shown in Figure 14.15*c*. Note that at this stage of the analysis no deductions can be made about the relative positions of the *Eco*RI and *Hin*dIII cleavage sites. The *Hin*dIII cleavage site may be located in either of the two *Eco*RI restriction fragments. The molecule is then

■ FIGURE 14.15 Procedure used to map restriction enzyme cleavage sites in DNA molecules. (*a*-*d*) Structures of the DNA molecule or of restriction fragments of the molecule either (*a*) uncut or cut with (*b*) *Eco*RI, (*c*) *Hin*dIII, or (*d*) *Eco*RI and *Hin*dIII. (*e*) The separation of these DNA molecules and fragments by agarose gel electrophoresis. The left lane on the gel contains a set of molecular size markers, a set of DNA molecules of size 1000 nucleotide pairs and multiples thereof.

simultaneously digested with both *Eco*RI and *Hin*dIII, and three fragments of sizes 3000, 2000, and 1000 nucleotide pairs are produced. This result establishes the positions of the two cleavage sites relative to one another on the molecule. Since the 2000-nucleotide-pair *Eco*RI restriction fragment is still present (not cut by *Hin*dIII), the *Hin*dIII cleavage site must be at the opposite end of the molecule from the *Eco*RI cleavage site (Figure 14.15*d*). By extending this type of analysis to include the use of several different restriction enzymes, more extensive maps of restriction sites can be constructed. When large numbers of restriction enzymes are employed, detailed maps of entire chromosomes can be constructed. An important aspect of these **restriction maps** is that, unlike genetic maps (Chapter 7), they reflect true physical distances along the DNA molecule.

By combining computer-assisted restriction mapping with other molecular techniques, it is possible to construct physical maps of entire genomes. The first multicellular eukaryote for which this was accomplished is *Caenorhabditis elegans*, a worm that is important for studies on the genetic control of development (Chapter 20). Moreover, the physical map of the *C. elegans* genome has been correlated with its genetic map. Thus, when an interesting new mutation is identified in *C. elegans*, its position on the genetic map often can be used to obtain clones of the wild-type gene from a large international *C. elegans* clone bank.

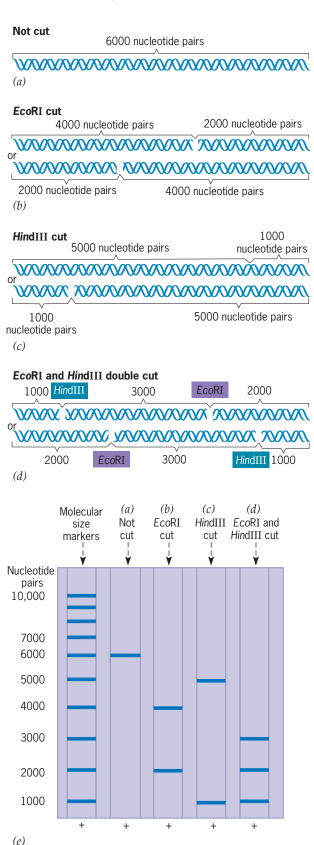
NUCLEOTIDE SEQUENCES OF GENES AND CHROMOSOMES

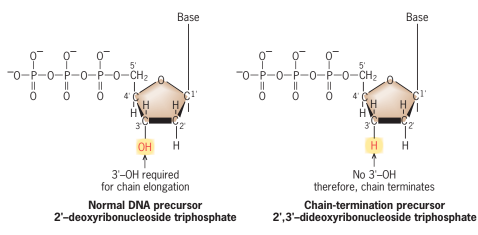
The ultimate physical map of a specific gene or chromosome is its nucleotide-pair sequence, complete with a chart of all nucleotide-pair changes that alter the function of that gene or chromosome. Prior to 1975, the thought of trying to sequence entire chromosomes was barely conceivable—at best, it was a laborious task requiring years of work. By late 1976, however, the entire 5386-nucleotide-long chromosome of phage ϕ X174 had been sequenced. Today, sequencing is a routine laboratory procedure. The complete nucleotide sequences of the genomes of 2442 viruses, 1372 bacteria, 93 archaea, and 40 eukaryotes are known, and genome sequencing projects are in progress for another 3670 bacteria, 84 archaea, and 612 eukaryotes. In addition, the sequence of 99 percent of the euchromatin in the human genome is known (Chapter 15).

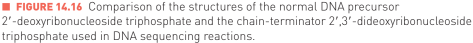
Our initial ability to sequence essentially any DNA molecule was the result of four major developments. The most important breakthrough was the discovery of restriction enzymes and their use in preparing homogeneous samples of specific segments of chromosomes. Another major advance was the improvement of gel electrophoresis procedures to the point

where DNA chains that differ in length by a single nucleotide could be resolved. Gene-cloning techniques to facilitate the preparation of large quantities of a particular DNA molecule were also important. Finally, researchers invented efficient procedures by which the nucleotide sequences of DNA molecules can be determined.

DNA sequencing protocols depend on the generation of a population of DNA fragments that all have one end in common (all end at exactly the same nucleotide)







and terminate at all possible positions (every consecutive nucleotide) at the other end. The common end is the 5'-terminus of the sequencing primer. The 3'-terminus of the primer contains a free —OH, which is the site of chain extension by DNA polymerase. Chain extension produces fragments with variable 3' ends—with ends at every possible nucleotide position along the DNA strand. These fragments are then separated on the basis of chain length by polyacrylamide gel electrophoresis.

Today, all DNA sequencing is performed using automated DNA sequencing machines. Initially, sequencing machines utilized a DNA sequencing protocol published in 1977 by Frederick Sanger and colleagues. Sanger shared the 1980 Nobel Prize in Chemistry for this work; he also received the 1958 Nobel Prize in Chemistry for determining the amino acid sequence of insulin. Today, new and faster DNA sequencing methods are replacing the Sanger procedure.

The Sanger procedure uses *in vitro* DNA synthesis in the presence of specific chain-terminators to generate populations of DNA fragments that end at As, Gs, Cs, and Ts, respectively. 2',3'-Dideoxyribonucleoside triphosphates (ddXTPs) (**Figure 14.16**) are the chain-terminators most frequently used in the Sanger sequencing protocol. Recall that DNA polymerases have an absolute requirement for a free 3'—OH on the DNA primer strand (Chapter 10). If a 2',3'-dideoxynucleotide is added to the end of a chain, it will block subsequent extension of that chain since the 2',3'-dideoxynucleotides have no 3'-OH. By using (1) 2',3'-dideoxythymidine triphosphate (ddTTP), (2) 2',3'-dideoxycytidine triphosphate (ddCTP), (3) 2',3' -dideoxyadenosine triphosphate (ddATP), and (4) 2',3'-dideoxyguanosine triphosphate (ddGTP), each labeled with a dye that fluoresces a different color, as chainterminators in a DNA synthesis reaction, a population of nascent fragments will be generated that includes chains with 3' termini at every possible position. Moreover, all chains that terminate with ddG will fluoresce one color; those that terminate with ddA will fluoresce a second color; chains that terminate with ddC will fluoresce a third color; and those that terminate with ddT will fluoresce a fourth color (**Figure 14.17**).

In the reaction tube, the ratio of dXTP:ddXTP (where X can be any one of the four bases) is kept at approximately 100:1, so that the probability of termination at a given X in the nascent chain is about 1/100. This yields a population of fragments terminating at all potential (X) termination sites within a distance of a few hundred nucleotides from the original primer terminus.

After the DNA chains generated in the reaction are released from the template strands by denaturation, they are separated by polyacrylamide capillary gel electrophoresis; their positions in the gel are detected with a scanning laser and a fluorescence detector, and recorded on a computer. The computer prints out the sequence of fluorescence peaks recorded as each nascent chain moves past the laser beam. The shortest

Set up a DNA polymerization reaction containing the following: 1 **DNA** polymerase 3' - AGTGTCAAGA - 5' Template strand dGTP, dATP, dTTP, and dCTP Primer strand 5' MOH 3' All four 2',3'-dideoxyribonucleoside triphosphate chain terminators, each labeled with a different fluorescent dye: ddGTP, ddATP, ddCTP, and ddTTP. STE 2 Incubate reaction mixture. Synthesized chains terminating with: ddG ddC ddT ddA 3' - AGTGTCAAGA - 5' 3' - AGTGTCAAGA - 5' 3' - AGTGTCAAGA - 5' 3' - AGTGTCAAGA 5 ww TCdd ww Tdd MW TCACAG^{dd} ww TCAdd [≁] TCACAGT^{dd} MW TCACAdd MW TCACdd Products **ww TCACAGTT^{dd} MW TCACAGTTCdd www_TCACAGTTCTdd** STED Denature products and separate by polyacrylamide capillary gel electrophoresis. STES Record sequence with a scanning laser, a fluorescence Load detector, and a computer A - 50 $_{\rm T}$ c ^A Primer TCACAGT 3 M T C A C A G 5' T C A C A 5' TCAC 5' 5' • T C A 5 • T C 5' 3' luorescence GACACI Laser detector \square Computer Capillary tube С Read DNA sequence from computer printout. 20 10 30 40 50 5' 3'

FIGURE 14.17 Sequencing DNA by the 2',3'-dideoxynucleoside triphosphate chain-termination procedure. *In vitro* DNA synthesis is performed in the presence of the four 2',3'-dideoxy chain-terminators: ddGTP, ddATP, ddCTP, and ddTTP, each labeled with a different fluorescent dye. The reaction mixture contains all the components required for DNA synthesis (see text for details). The dideoxy terminator at the 3' end of each chain is determined by the fluorescence of the attached dye. In the example shown, ddG fluoresces dark blue (appears black), ddC fluoresces light blue, ddA fluoresces green, and ddT fluoresces red. Because the shortest chain migrates the greatest distance, the nucleotide sequence of the longest chain (shown reading 5' \rightarrow 3' at the top of the computer printout) is obtained by reading the sequence starting with the first chain to pass the laser beam and continuing with each chain one nucleotide longer through to the longest chain.

PROBLEM-SOLVING SKILLS

Determining the Nucleotide Sequences of Genetic Elements

THE PROBLEM

Ten micrograms of a decanucleotide-pair *Hpa*l restriction fragment were isolated from the double-stranded DNA chromosome in the chloroplast of *Arabidopsis thaliana*. Octanucleotide poly (A) tails were then added to the 3' ends of both strands using the enzyme terminal transferase and dATP as shown in the following sequence:

> 5'-X X X X X X X X X X X -3' 3'-X'X'X'X'X'X'X'X'X'X'-5'

terminal transferase dATP

5'-X X X X X X X X X X A A A A A A A A-3' 3'-A A A A A A A A X'X'X'X'X'X'X'X'X'-5'

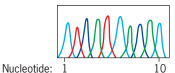
where X and X' can be any of the four standard nucleotides, but X' is always complementary to X.

The two complementary strands were then separated, and each strand was sequenced by the 2',3'-dideoxyribonucleoside triphosphate chain-termination method. Reaction 1 contained strand 1, primer, DNA polymerase, and all the other components required for DNA synthesis *in vitro*, plus the four standard dideoxynucleoside triphosphate chain-terminators—ddTTP, ddCTP, ddATP, and ddGTP each labeled with a dye that fluoresces at a different wavelength. The structure of the template-primer used in reaction 1 is as follows:

Strand 1: 3'-AAAAAAAAX'X'X'X'X'X'X'X'X'X'X'X'X'X'5' 5'-TTTTTTT-OH

Sequencing reaction 2 contained the same components as reaction 1 with the exception of the template-primer complex. Reaction 2 contained complementary strand 2; thus, the template-primer complex used in reaction 2 had the following structure:

After incubating the two reactions to allow time for DNA synthesis, the DNAs in each reaction were denatured, and the reaction products were separated by capillary gel electrophoresis using an automated DNA sequencing machine. The dyes used to label the chain-terminators fluoresce at different wavelengths, which are recorded by a photocell as the products of the reactions are separated in the capillary tube (see Figure 14.17). In the standard sequencing reactions, the chains terminating with ddG fluoresce dark blue, those terminating with ddC fluoresce light blue, those terminating with ddA fluoresce green, and those terminating with ddT fluoresce red. The computer printout for sequencing reaction 1 is as follows.



Draw the expected computer printout for sequencing reaction 2 (complementary strand 2 as template) in the following box. (Use the format shown above.)



FACTS AND CONCEPTS

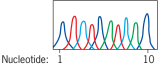
1. All DNA polymerases have an absolute requirement for a free 3'-hydroxyl on the end of the primer strand that will be extended by DNA polymerization reactions.

10

- **2.** All DNA synthesis occurs 5' to 3'; that is, all synthesis occurs by the addition of nucleotides to the 3' end of the primer strand.
- **3.** The addition of a 2', 3'-dideoxyribonucleoside monophosphate to the 3' end of a primer strand will block its extension.
- **4.** Polyacrylamide gel electrophoresis separates DNA strands on the basis of size and conformation.
- **5.** DNA chains have a constant charge per unit mass; that is, they have one negative charge per nucleotide.
- **6.** Because of their constant charge per unit mass, polynucleotide chains can be separated based on their size (length in nucleotides or nucleotide pairs).
- **7.** Linear DNA molecules that differ in length by one nucleotide can be separated by polyacrylamide gel electrophoresis for chains up to a few hundred nucleotides long.
- **8.** The shortest chains will migrate the largest distance during gel electrophoresis.
- **9.** Polyacrylamide gel electrophoresis performed in thin capillary tubes yields excellent separation of DNA chains differing in length by one nucleotide.
- **10.** The two strands of a double helix have opposite chemical polarity; if one strand has 5' to 3' polarity, the complementary strand has 3' to 5' polarity.

ANALYSIS AND SOLUTION

Because all DNA synthesis occurs by the addition of nucleotides to the 3'-OH terminus of the primer strand, all synthesis occurs in the $5' \rightarrow 3'$ direction. Therefore, the sequence of the nascent DNA chain synthesized with strand 1 as template is read 5' to 3' from the left to the right on the computer printout. The shortest nascent DNA fragment fluoresced light blue, indicating that it terminated with ddC, which means there was a G at this position in the template strand. Reading the ladder of bands from the left (shortest chain) to the right (longest chain) reveals that the sequence of the nascent strand is 5'-CTGATCAGAC-3'. Therefore, the sequence of the complementary template strand (strand 1) is 5'-GTCTGATCAG-3'. Now, if strand 2 is used as the template strand in the sequencing reaction, the nascent strand will have the sequence of strand 1, so the sequence of nucleotides (indicated by the fluorescent peaks) will be as shown in the following. The sequence of the nascent strand will be 5'-GTCTGATCAG-3', reading the peaks from the left (shortest chain) to the right (longest chain), and the sequence of the complementary template strand will be 5'-CTGATCAGAC-3'.



For further discussion visit the Student Companion site.



chain moves through the gel first, and each chain thereafter is one nucleotide longer than the preceding one. The dideoxynucleotide at the end of each chain will determine the color of fluorescence. Thus, the sequence of the longest newly synthesized DNA chain can be determined by simply reading the sequence of fluorescence peaks from the shortest chain to the longest chain (Figure 14.17). See Problem-Solving Skills: Determining the Nucleotide Sequences of Genetic Elements to test your understanding of automated DNA sequencing machines that use the Sanger procedure.

New approaches to DNA sequencing are now replacing the Sanger chainterminator method, and new—so-called second generation—DNA sequencing machines can sequence up to 25 billion nucleotide pairs per day. Many of the new sequencing procedures utilize sequencing-by-synthesis protocols in which the primer strands of immobilized primer-template complexes are extended by DNA polymerase by adding deoxyribonucleoside triphosphates one at a time and recording the sequence of nucleotide additions based on light signals recorded by a CCD (charge-coupled device) sensor. One such procedure is called *pyrosequencing* because it relies on the detection of the pyrophosphate released when a nucleotide is added to the end of a primer strand.

Another procedure utilizes a laser beam to record the addition of fluorescently labeled nucleotides during the extension of primer strands bound to tiny beads in a water–oil mixture. This procedure is called 454 sequencing. Yet another procedure, called Illumina sequencing (formerly Solexa sequencing), uses reversible terminators to detect single nucleotides as they are added to growing DNA strands. In the sequencing machines that use this procedure, large numbers of reactions occur simultaneously; thus, it is often called massively parallel sequencing. All of these systems are extremely fast, and new sequencing strategies are currently being developed. Although we are not there yet, the goal of sequencing an entire human genome for \$1000 has gone from being science fiction to a reasonable possibility.

- Detailed physical maps of DNA molecules can be prepared by identifying the sites that are cleaved by various restriction endonucleases.
- The nucleotide sequences of DNA molecules provide the ultimate physical maps of genes and chromosomes.

KEY POINTS

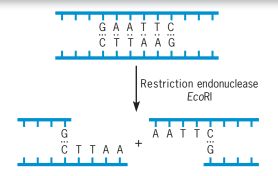
Basic Exercises

Illustrate Basic Genetic Analysis

- 1. What is a recombinant DNA molecule?
- **Answer:** A recombinant DNA molecule is constructed *in vitro* from portions of two different DNA molecules, often DNA molecules from two different species.

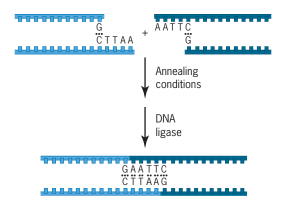


- 2. What are restriction endonucleases?
- **Answer:** Restriction endonucleases are enzymes that cleave DNA molecules in a sequence-specific manner such that all of the fragments produced have the same nucleotide sequences at their ends. Many restriction enzymes make staggered cuts in palindromic DNA sequences, yielding fragments with complementary single-stranded termini, as shown here.



- **3.** How are restriction endonucleases used to construct recombinant DNA molecules *in vitro*?
- **Answer:** If DNA molecules from two different sources (perhaps different species) are both digested with a restriction endonuclease that recognizes a palindromic DNA sequence and makes staggered cuts in the two strands, the resulting fragments will have complementary single-stranded ends.

If these DNA fragments are mixed, the complementary ends will pair, and the addition of DNA ligase will produce recombinant DNA molecules, as shown here.



- **4.** Why is the polymerase chain reaction (PCR) such a powerful tool for use in analyses of DNA?
- Answer: Because PCR amplifies DNA sequences geometrically large quantities of specific sequences can be obtained

Testing Your Knowledge

Integrate Different Concepts and Techniques

- The human genome (haploid) contains about 3 × 10° nucleotide pairs of DNA. If you digest a preparation of human DNA with *Not*I, a restriction endonuclease that recognizes and cleaves the octameric sequence 5'-GCGGCCGC-3', how many different restriction fragments would you expect to produce? Assume that the four bases (G, C, A, and T) are equally prevalent and randomly distributed in the human genome.
- Answer: Assuming that the four bases are present in equal amounts and are randomly distributed, the chance of a specific nucleotide occurring at a given site is 1/4. The chance of a specific dinucleotide sequence (e.g., AG) occurring is $1/4 \times 1/4 = (1/4)^2$ and the probability of a specific octanucleotide sequence is $(1/4)^8$ or 1/65,536. Therefore, NotI will cleave such DNA molecules an average of once in every 65,536 nucleotide pairs. If a linear DNA molecule is cleaved at *n* sites, n + 1 fragments will result. A genome of 3×10^9 nucleotide pairs should contain about 45,776 (3 \times 10%/65,536) NotI cleavage sites. If the entire human genome consisted of a single molecule of DNA, NotI would cleave it into 45,776 + 1 fragments. Given that these cleavage sites are distributed on 24 different chromosomes, complete digestion of the human genome with NotI should yield about 45,776 + 24 restriction fragments.
- 2. The maize gene *gln2*, which encodes the chloroplastic form of the enzyme glutamine synthetase, contains a single cleavage site for *Hin*dIII, but no cleavage site for *Eco*RI. You are given an *E. coli* plasmid cloning vector that

starting with just one or a few molecules. If one begins with a single molecule of DNA, 10 cycles of replication will yield 1024 DNA double helices, and 20 cycles will yield 1,048,576.

- 5. How are 2',3'-dideoxyribonucleoside triphosphates used in DNA sequencing protocols?
- **Answer:** The 2',3'-dideoxyribonucleoside triphosphates function as specific terminators of DNA synthesis. When a 2',3'-dideoxyribonucleoside monophosphate is added to the end of a nascent DNA chain, that chain can no longer be extended by DNA polymerase because of the absence of the 3'—OH required for chain extension. By using the appropriate ratios of 2'-deoxyribonucleoside triphosphates to 2',3'-dideoxyribonucleoside triphosphates in DNA synthesis reactions *in vitro*, DNA chains are produced that terminate at all possible nucleotide positions. Separation of these nascent DNA chains by gel electrophoresis and detection of their positions in the gel with fluorescent dyes are then used to determine their nucleotide sequences (see Figure 14.17).

contains a unique *Hin*dIII cleavage site within the gene *amp*^r, which confers resistance to the antibiotic ampicillin on the host cell, and a unique *Eco*RI cleavage site within a second gene *tet*^r, which makes the host cell resistant to the antibiotic tetracycline. You are also given an *E. coli* strain that is sensitive to both ampicillin and tetracycline (*amp*^s *tet*^s). How would you go about constructing a maize genomic DNA library that includes clones carrying a complete *gln2* gene?

Answer: Maize genomic DNA should be purified and digested with *Eco*RI. Vector DNA should be similarly purified and digested with *Eco*RI. The maize *Eco*RI restriction fragments and the *Eco*RI-cut plasmid DNA molecules will now have complementary single-stranded ends (5'-AATT-3'). The maize restriction fragments should next be mixed with the *Eco*RI-cut plasmid molecules and covalently inserted into the linearized vector molecules in an ATP-dependent reaction catalyzed by DNA ligase. The ligation reaction will produce circular recombinant plasmids, some of which will contain maize *Eco*RI fragment inserts. Insertion of maize DNA fragments into the *Eco*RI site of the plasmid disrupts the *tet*^r gene so that the resulting recombinant plasmids will no longer confer tetracycline resistance to host cells.

amp^s tet^s E. coli cells should then be transformed with the recombinant plasmid DNAs, and the cells should be plated on medium containing ampicillin to select for transformed cells harboring plasmids. The majority of the cells will not be transformed and, thus, will not

grow in the presence of ampicillin. The cells that grow on ampicillin-containing medium should be retained for analysis. This collection of cells harboring different *Eco*RI fragments of the maize genome represents a clone library that should contain clones with an intact *gln2* gene since this gene contains no *Eco*RI cleavage site. Note that the *Hin*dIII site of the vector could be used to construct a similar maize genomic *Hin*dIII fragment library, but such a library would not contain intact *gln2* genes because of the *Hin*dIII cleavage site in *gln2*.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **14.1** (a) In what ways is the introduction of recombinant DNA molecules into host cells similar to mutation? (b) In what ways is it different?
- 14.2 **(constraint)** Listed in this question are four different single strands of DNA. Which of these, in their double-stranded form, would you expect to be cleaved by a restriction endonuclease?
 - (a) ACTCCAGAATTCACTCCG(b) GCCTCATTCGAAGCCTGA(c) CTCGCCAATTGACTCGTC(d) ACTCCACTCCCGACTCCA
- **14.3** If the sequence of base pairs along a DNA molecule occurs strictly at random, what is the expected frequency of a specific restriction enzyme recognition sequence of length (a) four and (b) six base pairs?
- **14.4** In what ways do restriction endonucleases differ from other endonucleases?
- **14.5** Of what value are recombinant DNA and gene-cloning technologies to geneticists?
- **14.6** What determines the sites at which DNA molecules will be cleaved by a restriction endonuclease?
- 14.7 Restriction endonucleases are invaluable tools for biologists. However, genes encoding restriction enzymes obviously did not evolve to provide tools for scientists. Of what possible value are restriction endonucleases to the microorganisms that produce them?
- **14.8** Why is the DNA of a microorganism not degraded by a restriction endonuclease that it produces, even though its DNA contains recognition sequences normally cleaved by the endonuclease?
- **14.9** One of the procedures for cloning foreign DNA segments takes advantage of restriction endonucleases like *Hin*dIII (see Table 14.1) that produce complementary single-stranded ends. These enzymes produce identical complementary ends on cleaved foreign DNAs and on the vector DNAs into which the foreign DNAs are inserted. Assume that you have inserted your favorite gene into the *Hin*dIII site in the polycloning region of the Bluescript cloning vector with DNA ligase, have amplified the plasmid containing your gene in *E. coli*, and have isolated a large

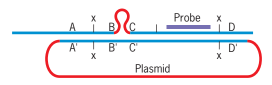
quantity of gene/Bluescript DNA. How could you excise your favorite gene from the Bluescript vector?

14.10 You are working as part of a research team studying the structure and function of a particular gene. Your job is to clone the gene. A restriction map is available for the region of the chromosome in which the gene is located; the map is as follows:

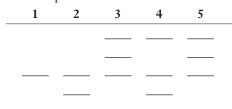


Your first task is to prepare a genomic DNA library that contains clones carrying the entire gene. Describe how you would prepare such a library in the plasmid vector Bluescript (see Figure 14.3), indicating which restriction enzymes, media, and host cells you would use.

- 14.11 Compare the nucleotide-pair sequences of genomic DNA clones and cDNA clones of specific genes of higher plants and animals. What is the most frequent difference that you would observe?
- 14.12 Most of the genes of plants and animals that were cloned soon after the development of recombinant DNA technologies were genes encoding products that are synthesized in large quantities in specialized cells. For example, about 90 percent of the protein synthesized in mature red blood cells of mammals consists of α - and β -globin chains, and the globin genes were among the first mammalian genes cloned. Why were genes of this type so prevalent among the first eukaryotic genes that were cloned?
- **14.13** Genomic clones of the chloroplastic glutamine synthetase gene (*gln2*) of maize are cleaved into two fragments by digestion with restriction endonuclease *Hind*III, whereas full-length maize *gln2* cDNA clones are not cut by *Hind*III. Explain these results.
- 14.14 💿 In the following illustration, the upper line shows a gene composed of segments A–D. The lower circle shows a mutant version of this gene, consisting of two fused pieces (A'-B', C'-D'), carried on a plasmid. You attempt a directed mutagenesis of a diploid cell by transforming cells with the cloned mutant gene. The following diagram shows the desired pairing of the plasmid and chromosome just prior to recombination.

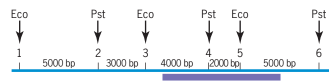


You prepare DNA from the cells, digest it with an enzyme that cuts at x, and hybridize the cleaved DNA with the probe shown above. The following diagram shows a Southern blot of possible results.



(a) Which lane shows fragments produced from DNA in the cell before transformation? (b) Which lane shows fragments produced from DNA in the cell in which the anticipated targeted mutagenesis occurred? (c) Which of these blot patterns might be expected if two crossovers occurred, one between A and B, and the other between C and D?

- 14.15 (a) What experimental procedure is carried out in Southern, northern, and western blot analyses? (b) What is the major difference between Southern, northern, and western blot analyses?
- 14.16 What major advantage does the polymerase chain reaction (PCR) have over other methods for analyzing nucleic acid structure and function?
- 14.17 The cloning vectors in use today contain an origin of replication, a selectable marker gene (usually an antibioticresistance gene), and one additional component. What is this component, and what is its function?
- 14.18 The drawing in this problem shows a restriction map of a segment of a DNA molecule. Eco refers to locations where the restriction endonuclease *Eco*RI cuts the DNA, and Pst refers to locations where the restriction enzyme *Pst*I cuts the DNA. Potential restriction sites are numbered 1–6. Distances between restriction sites are shown on the bottom scale in base pairs (bp). The thick line represents the part of the molecule that has homology with a probe.



- (a) Assume that individual 1 has restriction sites 1 through 6. If DNA is digested with *Pst*I, what are the expected sizes of the DNA fragments that will hybridize with the probe?
- (b) Assume that individual 2 has a mutation that eliminates site 4. If DNA is digested with *PstI*, what are the expected sizes of the DNA fragments that will hybridize with the probe?
- (c) Assume that individual 3 has a mutation that eliminates site 5. If the DNA is digested with *Pst*I, what are the

expected sizes of the DNA fragments that will hybridize with the probe?

- (d) If the DNA of individual 1 is digested with both *PstI* and *Eco*RI, what are the expected sizes of the DNA fragments that will hybridize with the probe?
- (e) If the DNA of individual 3 is digested with both *Pst*I and *Eco*RI, what are the expected sizes of the DNA fragments that will hybridize with the probe?
- 14.19 The cystic fibrosis (*CF*) gene (location: chromosome 7, region q31) has been cloned and sequenced, and studies of CF patients have shown that about 70 percent of them are homozygous for a mutant *CF* allele that has a specific three-nucleotide-pair deletion (equivalent to one codon). This deletion results in the loss of a phenylalanine residue at position 508 in the predicted CF gene product. Assume that you are a genetic counselor responsible for advising families with CF in their pedigrees regarding the risk of CF among their offspring. How might you screen putative CF patients and their parents and relatives for the presence of the *CF*\Delta*F*508 mutant gene? What would the detection of this mutant gene in a family allow you to say about the chances that CF will occur again in the family?
- 14.20 Cereal grains are major food sources for humans and other animals in many regions of the world. However, most cereal grains contain inadequate supplies of certain of the amino acids that are essential for monogastric animals such as humans. For example, corn contains insufficient amounts of lysine, tryptophan, and threonine. Thus, a major goal of plant geneticists is to produce corn varieties with increased kernel lysine content. As a prerequisite to the engineering of high-lysine corn, molecular biologists need more basic information about the regulation of the biosynthesis and the activity of the enzymes involved in the synthesis of lysine. The first step in the anabolic pathway unique to the biosynthesis of lysine is catalyzed by the enzyme dihydrodipicolinate synthase. Assume that you have recently been hired by a major U.S. plant research institute and that you have been asked to isolate a clone of the nucleic acid sequence encoding dihydrodipicolinate synthase in maize. Briefly describe four different approaches you might take in attempting to isolate such a clone and include at least one genetic approach.
- **14.21** You have just isolated a mutant of the bacterium *Shigella dysenteriae* that is resistant to the antibiotic kanamycin, and you want to characterize the gene responsible for this resistance. Design a protocol using genetic selection to identify the gene of interest.
- **14.22** You have isolated a cDNA clone encoding a protein of interest in a higher eukaryote. This cDNA clone is *not* cleaved by restriction endonuclease *Eco*RI. When this cDNA is used as a radioactive probe for blot hybridization analysis of *Eco*RI-digested genomic DNA, three radioactive bands are seen on the resulting Southern blot. Does this result indicate that the genome of the eukaryote in question contains three copies of the gene encoding the protein of interest?

14.23 A linear DNA molecule is subjected to single and double digestions with restriction endonucleases, and the following results are obtained:

Enzymes	Fragment Sizes (in kb)	
EcoRI	2.9, 4.5, 7.4, 8.0	
HindIII	3.9, 6.0, 12.9	
EcoRI and HindIII	1.0, 2.0, 2.9, 3.5, 6.0, 7.4	

Draw the restriction map defined by these data.

14.24 A circular DNA molecule is subjected to single and double digestions with restriction enzymes, and the products are separated by gel electrophoresis. The results are as follows (fragment sizes are in kb):

EcoRI	EcoRI and HindIII	HindIII	<i>Bam</i> HI		<i>Hin</i> dIII and <i>Bam</i> HI
8	5	12	6	6	6
4	4		6	4	5
	3			2	1

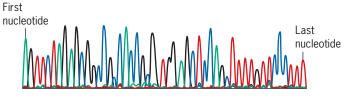
Draw the restriction map of this DNA molecule.

14.25 You are studying a circular plasmid DNA molecule of size 10.5 kilobase pairs (kb). When you digest this plasmid with restriction endonucleases *Bam*HI, *Eco*RI, and *Hin*dIII, singly and in all possible combinations, you obtain linear restriction fragments of the following sizes:

Enzymes	Fragment Sizes (in kb)
BamHI	7.3, 3.2
EcoRI	10.5
HindIII	5.1, 3.4, 2.0
BamHI + EcoRI	6.7, 3.2, 0.6
BamHI + HindIII	4.6, 2.7, 2.0, 0.7, 0.5
EcoRI + HindIII	4.0, 3.4, 2.0, 1.1
BamHI + EcoRI + HindIII	4.0, 2.7, 2.0, 0.7, 0.6, 0.5

Draw a restriction map for the plasmid that fits your data.

14.26 The automated DNA sequencing machines utilize fluorescent dyes to detect the nascent DNA chains synthesized in the presence of the four dideoxy (ddX) chain-terminators, each labeled with a different fluorescent dye. The dyes fluoresce at different wavelengths, which are recorded by a photocell as the products of the reactions are separated based on length by capillary gel electrophoresis (see Figure 14.17). In the standard sequencing reaction, the chains terminating with ddG fluoresce dark blue (peaks appear black in computer printout), those terminating with ddC fluoresce light blue, those terminating with ddA fluoresce green, and those terminating with ddT fluoresce red. The computer printout for the sequence of a short segment of DNA is as follows.



What is the nucleotide sequence of the nascent strand of DNA?

What is the nucleotide sequence of the DNA template strand?

14.27 Ten micrograms of a decanucleotide-pair *Hpa*I restriction fragment were isolated from the double-stranded DNA chromosome of a small virus. Octanucleotide poly(A) tails were then added to the 3' ends of both strands using terminal transferase and dATP; that is,

5'-X X X X X X X X X X X X-3' 3'-X'X'X'X'X'X'X'X'X'-5'

 \downarrow terminal transferase, dATP

5-X X X X X X X X X X X X AAAAAAAA-3' 3'-AAAAAAAX'X'X'X'X'X'X'X'X'X'-5'

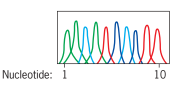
where X and X' can be any of the four standard nucleotides, but X' is always complementary to X.

The two complementary strands ("Watson" strand and "Crick" strand) were then separated and sequenced by the 2',3'-dideoxyribonucleoside triphosphate chaintermination method. The reactions were primed using a synthetic poly(T) octamer; that is,

Watson strand

Crick strand

Two DNA sequencing reactions were carried out. Reaction 1 contained the Watson strand template/primer shown above; reaction 2 contained the Crick strand template/primer. Both sequencing reactions contained DNA polymerase and all other substrates and components required for DNA synthesis in vitro plus the standard four 2',3'-dideoxyribonucleoside triphosphate chainterminators-ddGTP, ddCTP, ddATP, and ddTTPeach labeled with a different fluorescent dye. The dyes fluoresce at different wavelengths, which are recorded by a photocell as the products of the reactions are separated by capillary gel electrophoresis (see Figure 14.17). In the standard sequencing reaction, the chains terminating with ddG fluoresce dark blue (peaks appear black in the computer printouts), those terminating with ddC fluoresce light blue, those terminating with ddA fluoresce green, and those terminating with ddT fluoresce red. The computer printout for sequencing reaction 1, which contained the Watson strand as template, is as follows.



Draw the predicted computer printout for reaction 2, which contained the Crick strand as template, in the following box. Remember that all DNA synthesis occurs in the 5' \rightarrow 3' direction and that the sequence of the nascent strand reads 5' to 3' from left to right in the printout.



Genomics on the Web at http://www.ncbi.nlm.nih.gov

In this chapter, we have discussed a DNA test for one of the most prevalent mutant alleles that causes cystic fibrosis, and in Chapter 16 (Figure 16.2), we will examine a DNA test for mutant genes that result in Huntington's disease.

- 1. Are DNA tests available for mutant genes that cause other inherited human diseases? If so, what are some of the diseases for which DNA tests are currently available?
- 2. What are some of the molecular techniques used in these DNA tests? gel electrophoresis? PCR? Southern blots?
- **3.** How reliable are these tests? Can they be performed on fetal cells obtained by amniocentesis? on single cells obtained from eight-cell pre-embryos?

Hint: At the NCBI web site, go to Human Genome Resources and then to OMIM (Online Mendelian Inheritance in Man), and search for "DNA tests for mutant alleles." Also visit http://www.genetests.org for information on 607 laboratories providing tests for 1549 different human genetic diseases.

Genomics



CHAPTER OUTLINE

- Genomics: An Overview
- Correlated Genetic, Cytological, and Physical Maps of Chromosomes
- Map Position-Based Cloning of Genes
- The Human Genome Project
- RNA and Protein Assays of Genome Function
- Comparative Genomics

first recognized their uniqueness after studying a skull and other bones found by miners in Germany's Neander Valley (Neander Tal in German). The Neanderthals coexisted with our ancestors in Europe from 45,000 to 30,000 years ago, and perhaps in the Middle East as early as 80,000 years ago, according to the archaeological records from caves in the area. Indeed, the Neanderthals and early humans both lived in caves, had similar tools, and used spears to hunt deer and gazelles. So, the big question has always been: did our ancestors and Neanderthals mate and exchange genes? Paleoanthropologists were never able to agree on the answer to that question—until very recently.

In May 2010, that question was answered in the affirmative when an international research team led by Svante Pääbo published about two-thirds of the sequence of the Neanderthal genome. How can scientists sequence DNA from an extinct species? One possibility is to find specimens frozen and highly preserved in ice such as the woolly mammoths discovered in Siberia. The other possibility—used to sequence the Neanderthal genome—is to use DNA extracted from bones. Bones contain DNA that remains intact long after an animal dies, and the researchers were able to sequence fragments of DNA extracted from the bones of three female Neanderthals who lived in the Vindija cave in Croatia about 40,000 years ago. By repeatedly sequencing short fragments of DNA and carefully removing contaminating microbial DNA sequences, Pääbo and his colleagues were able to splice together almost two-thirds of the Neanderthal genome.

After assembling over 60 percent of the Neanderthal genome, the research team compared the sequence with the sequences of five living humans—from China, France, Papua New Guinea, South Africa, and West Africa, respectively. What they found was somewhat of a surprise, because an earlier comparison of the mitochondrial DNA sequences of the two species showed no traces of human DNA sequences in the Neanderthal mitochondrial DNA, and vice versa. What they discovered was that Europeans and Asians, but not

The Neanderthal Genome: What It Reveals about Our Ancestors

The Neanderthals (*Homo neanderthalensis*) are believed to be our closest evolutionary relatives. They lived in Europe and Asia from about 130,000 years ago until perhaps 28,000 years ago, when they became extinct. They are called Neanderthals because scientists



Photo of a man walking his dog through the cutout silhouette of a Neanderthal man in a monument at Mettmann, Germany, where the first Neanderthal fossils were discovered.

Africans, have inherited from 1 to 4 percent of their genes from Neanderthals. These results indicate that Neanderthals and humans interbred some 80,000 years ago after humans left Africa but before they spread throughout Europe and Asia. The genomes of humans who remained in Africa until after this period of interbreeding, therefore, do not contain Neanderthal sequences.

Now that the question about mating between Neanderthals and humans has been answered, can the sequence of the Neanderthal genome provide answers to other questions about human evolution? What genes make humans "human"? What caused the Neanderthals to go extinct and humans to become the dominant species on our planet? What human genes have evolved since the split between humans and Neanderthals? The Neanderthal genome research team has identified a few genes that may be important in distinguishing the two species. They include genes involved in cognitive and skeletal development; however, more research is needed to evaluate their significance.

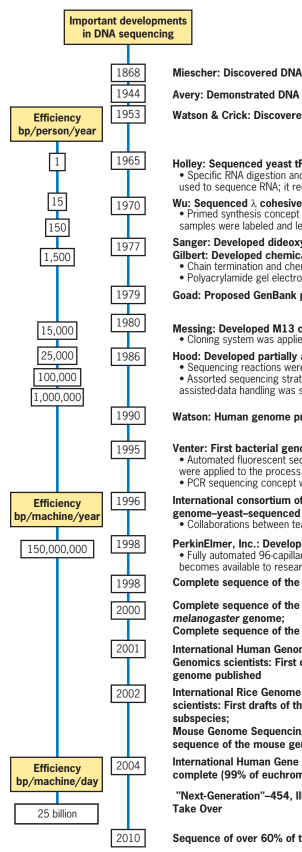
Is this the end of the story? No! Scientists are now trying to piece together the genome of the Denisovans, cousins of the Neanderthals, who lived in Asia from approximately 400,000 to 50,000 years ago. Paleoanthropologists believe that the Denisovans interbred with the ancestors of the current inhabitants of New Guinea. Now, scientists have extracted Denisovan DNA from a finger bone and a tooth discovered in a Siberian cave. Their results to date indicate that 4.8 percent of the DNA in people from New Guinea is derived from the Denisovan genome. Where do we go from here? Will "Lucy's" genome be sequenced next?

Gregor Mendel studied the effects of seven genes on traits in peas, but he studied no more than three genes in any one cross. Today's geneticists can study the expression of all the genes—the entire genome—of an organism in a single experiment. As of February 2011, the complete nucleotide sequences of the genomes of 2585 viruses, defective viruses, and viroids, 735 plasmids, 2362 mitochondria, 201 chloroplasts, 94 archaea, 1318 true bacteria, and 41 eukaryotes had been determined. In addition, the genomes of another 370 eukaryotes have been sequenced, and the sequences are currently being assembled into complete genome sequences. Sequencing projects for another 630 eukaryotic genomes are underway, and the complete genome sequences of several humans are now available. Finally, as discussed in Chapter 9, the goal of the 1000 Genomes Project is to sequence at least 2500 genomes from people representing ancestral groups from around the world. Indeed, some scientists are predicting that it may be possible to sequence an entire human genome for as little as \$1000 in the near future.

The list of eukaryotes whose genomes have been sequenced includes important model organisms in genetics: baker's yeast *Saccharomyces cerevisae*, the fruit fly *Drosophila melanogaster*; and the plant *Arabidopsis thaliana*. It also includes the protozoan *Plasmodium falciparum*, which causes the most dangerous form of malaria, and the mosquito *Anopheles gambiae*, which is the host organism most responsible for the spread of this disease. The silkworm (*Bombyx mori*), an economically important insect, is on the list, and so are several vertebrates: the mouse (*Mus musculus*), the Norwegian rat (*Rattus norvegicus*), the Red Jungle Fowl—an ancestor of domestic chickens (*Gallus gallus*), the puffer fish (*Fugu rubripes*), our closest living relative the chimpanzee (*Pan troglodytes*), and our own species (*Homo sapiens*).

One of the original goals of the Human Genome Project was to determine the complete nucleotide sequence of the human genome by the year 2005. As it turned out, two first drafts of the sequence—one by a public consortium and the other by a private company—were published in February 2001. Indeed, a nearly complete sequence of the human genome comprising 99 percent of the euchromatic DNA was released in October 2004, a full year ahead of the original goal. The sequence of the genome of our closest living relative, the chimpanzee (*Pan troglodytes*), was completed in 2006, and about two-thirds of the sequence of the genome of our close evolutionary relative, the extinct Neanderthal (*Homo neanderthalensis*), was published in 2010.

The rapid improvements in DNA sequencing technology that occurred during the last two decades have allowed researchers to collect large amounts of sequence data. The new second-generation sequencing machines now make it possible to sequence entire human genomes in one day (see Chapter 14). However, sequencing was not always so easy. It took Robert Holley, 1968 Noble Prize recipient, several years to determine the 77-nucleotide sequence of the alanine tRNA from yeast (see Figure 12.12). A few of the major advances in sequencing technology, as well as some of the landmarks in the study of genomes, are highlighted in **E Figure 15.1**.



Miescher: Discovered DNA

Avery: Demonstrated DNA as "genetic material"

Watson & Crick: Discovered double-helix structure of DNA

Holley: Sequenced yeast tRNA^{Ala}

 Specific RNA digestion and chromatography methods were used to sequence RNA; it required large quantities of sample.

Wu: Sequenced λ cohesive end DNA

• Primed synthesis concept and 2-D electrophoresis were used; samples were labeled and less material was required.

Sanger: Developed dideoxy termination sequencing procedure;

- Gilbert: Developed chemical degradation sequencing protocol
- Chain termination and chemical degradation concepts were developed.
- Polyacrylamide gel electrophoresis was used to separate DNA tracts.

Goad: Proposed GenBank prototype

Messing: Developed M13 cloning vectors Cloning system was applied.

Hood: Developed partially automated sequencing system

- Sequencing reactions were optimized.
- Assorted sequencing strategies were applied and computer assisted-data handling was started.

Watson: Human genome project initiated

Venter: First bacterial genomes sequenced

 Automated fluorescent sequencing instruments and robotic operations were applied to the process.

PCR sequencing concept was introduced.

International consortium of scientists: First eukaryotic

Collaborations between teams of scientists.

PerkinElmer, Inc.: Developed 96-capillary sequencer

• Fully automated 96-capillary electrophoresis sequencing system becomes available to research laboratories.

Complete sequence of the Caenorhabditis elegans genome

Complete sequence of the euchromatic portion of the Drosophila

Complete sequence of the Arabidopsis thaliana genome

International Human Genome Sequencing Consortium and Celera Genomics scientists: First drafts of the sequence of the human

International Rice Genome Sequencing Project and Syngenta scientists: First drafts of the genomic sequences of two rice

Mouse Genome Sequencing Consortium: First draft of the sequence of the mouse genome

International Human Gene Sequencing Consortium: Nearly complete (99% of euchromatin) sequence of the human genome

"Next-Generation"-454, Illumina, SOLiD-Sequencing Machines

Sequence of over 60% of the Neanderthal Genome Is Published

FIGURE 15.1 Advances in DNA sequencing efficiency, some of the technological developments that enhanced the productivity of sequencers, and some landmarks in DNA sequencing. Initially, all the steps in DNA sequencing were performed manually, making it a very labor-intensive process. Today, fully automated sequencing machines have greatly increased efficiency.

In the present era, vast amounts of sequence data accumulate daily. Most of these data are the results of research projects funded by government agencies—the National Institutes of Health (NIH), the National Science Foundation (NSF), and the Department of Energy (DOE) in the United States—and comparable agencies in other countries. Thus, these data are public information and are available to anyone who wants to use them. Making the sequences public has been accomplished by establishing sequence databases that are available free on the web at http://www.ncbi.nlm. nih.gov/entrez/query.fcgi (see Focus on GenBank).

Of course, just making the databases available is not enough. We must be able to extract information from them—that is, to "mine" the databases—and then analyze the extracted information efficiently and accurately. This process requires computer software that can search the vast DNA sequences in genomes of interest. The need for such software has spawned a new scientific discipline called bioinformatics. Mathematicians, computer scientists, and molecular biologists who work in this discipline develop computer-search algorithms that can extract information from DNA and protein sequence data.

The availability of entire genome sequences has opened the door to bioinformatic analyses and to functional studies of the genes contained in these sequences. Microarrays—including the so-called gene chips—allow scientists to investigate the expression of all the genes in an organism simultaneously (see the section RNA and Protein Assays of Genome Function in this chapter). Other procedures use known nucleotide sequences to dissect metabolic pathways by "knocking out" or turning off the expression of genes (see the section Reverse Genetics in Chapter 16).

In this chapter, we will discuss some of the tools and techniques that are used to study the structure and function of genomes, we will examine the spectacular progress of the Human Genome Project, and we will see how comparisons of genomes can contribute to our understanding of evolution. In the following chapter, we will examine other technical advances—DNA profiling, human gene therapy, and the production of transgenic microorganisms, plants, and animals. We will also see how geneticists have identified the defective genes that are responsible for two tragic human conditions, Huntington's disease and cystic fibrosis. The procedures used to identify these genes have become methodological paradigms for the identification of many other disease-related genes in humans.

FOCUS ON

GENBANK

n 1979 Walter Goad, a physicist working at the Los Alamos National Laboratory (LANL) in New Mexico, came up with the idea for a database that would contain all available DNA sequences. From 1982 until 1992, Goad and his colleagues incorporated sequences into the database—now named GenBank and maintained it at LANL. Today, this database is maintained by the National Center for Biotechnology Information (NCBI), which is part of the National Library of Medicine (NLM) at the National Institutes of Health (NIH) in Bethesda, Maryland. The content of the database has grown enormously since Goad and his colleagues created it. At the end of 1982, GenBank contained 680,338 nucleotide pairs of sequenced DNA, but by January 2011, it contained over 117 billion nucleotide pairs (**■ Figure 1**).

Databases comparable to GenBank also were established in Europe and Japan. The European Molecular Biology Laboratory (EMBL) Data Library was set up in Germany in 1980, and the DNA DataBank of Japan (DDBJ) was established in 1984. GenBank, EMBL, and DDBJ subsequently joined forces and formed the International Nucleotide Sequence Database Collaboration, which allows researchers to search all three databases simultaneously.

The development of search and retrieval programs that screen databases for sequences similar to input sequences has provided scientists with an important research tool. In particular, NCBI's Entrez retrieval system has proven invaluable. This system is available free at http://www.ncbi.nlm.nih.gov/entrez. The amount of information available at the Entrez web site has increased every year. It encompasses not only DNA and protein sequence databases, but also a huge bibliographic database called PubMed that covers most of the journals in medicine and biology. Today, you can search all these databases simultaneously by using NCBI's global cross-database search engine, and the search page will give you the number of items found (that is, the "hits") in each database. For example, a "Search across databases" using the query "HBB" (abbreviation for the human beta-globin gene)



will yield 948 hits in PubMed Central (free, full-text journal articles), 57 Books, 1725 Nucleotide hits (sequences in GenBank), 726 SNP (single-nucleotide polymorphisms) hits, and so on.

A discussion of all the databases that can be searched with Entrez is far beyond the scope of this textbook. You are encouraged to visit the site and explore its databases. They include the PubMed and DNA databases mentioned above, and databases of protein sequences, three-dimensional macromolecular structures, cancer chromosomes and genes, expressed sequences, single-nucleotide polymorphisms, whole-genome sequences, and many more.

Let's perform one Entrez search to illustrate how it works. Assume that you have just determined the nucleotide sequence of a segment of DNA from an organism of interest, and you want to know if that DNA has already been sequenced or if it is similar to sequences in any of the current databases. One of the quickest ways to obtain this information is to perform a BLAST (*Basic Local Alignment Search Tool*) search with your sequence as the input, or *query*, sequence. Let's start at the NCBI home page: http:// www.ncbi.nlm.nih.gov. First, you will need to sign in or register if this is your initial use of the software at the site. Then, select "BLAST" from the Popular Resources list, and select "nucleotide blast." Next, paste the following sequence into the query box.

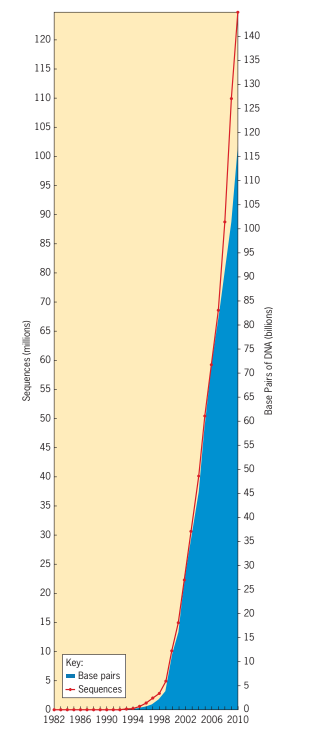
5'-ATGAGAGAAATTCTTCATATTCAAGGAGGTCAGTGCGGAAACCAGATCGG AGCTAAGTTCTGGGAAGTTATTTGCGGCGAGCACGGTATTGATCAAACCG-3'.

Before you click the "BLAST!" button, give your job a title (e.g., your name) and choose "Nucleotide collection (nr/nt)" as your "Database." Now, click the "BLAST!" button. Your results should appear in about 10 seconds. They should include a list of "Sequences producing significant alignments" and the alignment of each sequence with your query sequence.

The first six sequences are all independently obtained sequences of the same gene, the β 9 tubulin gene of *Arabidopsis thaliana*; the rest are independent sequences of closely related genes in the same and related species. Note that the query sequence is a perfect match with the first six sequences and differs from sequences 9 through 12 (the *A. thaliana* β 8 tubulin gene) at 12 nucleotide positions. These two sequences are members of a gene family that encodes a set of very closely related proteins with the same or very similar functions.

Suppose you want to know more about the sequences identified in your search. Let's select sequence number 6 with accession number M84706; click on the accession number. That will take you to the sequence submitted to GenBank along with information about the sequence and the original publication (Snustad et al., 1992). To obtain a copy of that publication, just click on the PUBMED article number (1498609). The Abstract of that paper will appear first. If you then click on "Free Full Text," you will be able to download a copy of the entire paper.

This brief exploration of the Entrez web site illustrates the power and convenience of the software and the databases now available. Without these tools, geneticists would be hard-pressed to make much sense out of the vast number of DNA sequences currently available.



■ FIGURE 1 Growth of GenBank from its origin in 1982 to 2011. The left and right ordinates show the size of the collection in number of DNA sequences (red) and number of nucleotide pairs (blue), respectively. The number of different sequences has grown from 606 at the end of 1982 to 122.9 million at the beginning of 2011.

Genomics: An Overview

Genomics is the subdiscipline of genetics that focuses on the structure and function of entire genomes. Geneticists have used the term *genome* for over seven decades to refer to one complete copy of the genetic information or one complete set of chromosomes (monoploid or haploid) of an organism. In contrast, **genomics** is a relatively new term. The word *genomics* appears to have been coined by Thomas Roderick in 1986 to refer

to the genetics subdiscipline of mapping, sequencing, and analyzing the functions of entire genomes, and to serve as the name of a new journal—*Genomics*—dedicated to the communication of new information in this subdiscipline.

As more detailed maps and sequences of genomes became available, the genomics subdiscipline was divided into *structural genomics*—the study of the genome structure, *functional genomics*—the study of the genome function; and **comparative genomics**—the study of genome evolution. Functional genomics includes analyses of the **transcriptome**, the complete set of RNAs transcribed from a genome, and the **proteome**, the complete set of proteins encoded by a genome. Indeed, functional genomics has spawned an entirely new discipline, **proteomics**, which has as its goal the determination of the structures and functions of all the proteins in an organism.

Whereas structural genomics is quite advanced with the complete nucleotide sequences available for many organisms, functional genomics is presently in an explosive growth phase. New array hybridization and gene-chip technologies allow researchers to monitor the expression of entire genomes—all the genes in an organism—at various stages of growth and development or in response to environmental changes. These powerful new tools promise to provide a wealth of information about genes and how they interact with each other and with the environment.

KEY POINT

 Genomics is the subdiscipline of genetics devoted to the mapping, sequencing, and functional and comparative analyses of genomes.

Correlated Genetic, Cytological, and Physical Maps of Chromosomes

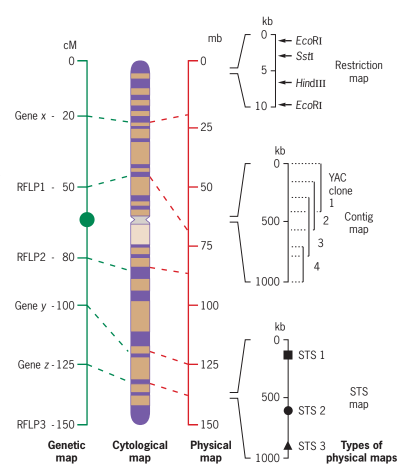
The chromosomal locations of genes and other molecular markers can be mapped based on recombination frequencies, positions relative to cytological features, or physical distances. The ability of scientists to identify and isolate genes based on information about their location in the genome was one of the first major contributions of genomics research. In principle, this approach, called *positional cloning*, can be used to identify and clone any gene with a known phenotypic effect in any species. Positional cloning has been used extensively in many species, including humans. Indeed, in Chapter 16, we will consider the use of posi-

tional cloning to identify the human genes responsible for Huntington's disease and cystic fibrosis.

Because the utility of positional cloning depends on the availability of detailed maps of the regions of the chromosomes where the genes of interest reside, major efforts have focused on developing detailed maps of the human genome and the genomes of important model organisms such as *D. melanogaster*; *C. elegans*, and *A. thaliana*. The goal of this research is to construct correlated genetic and physical maps with markers distributed at relatively short intervals throughout the genome. In the case of the human and *Drosophila* genomes, the genetic and physical maps can also be correlated with cytological maps (banding patterns) of the chromosomes (**Figure 15.2**). We will discuss the construction of these maps in the following sections of this chapter.

Recall that genetic maps (Figure 15.2, left) are constructed from recombination frequencies, with 1 centiMorgan (cM) equal to the distance that yields an average frequency of recombination of 1 percent (Chapter 7). Genetic maps with markers spaced at short intervals—high-density genetic maps-are often constructed by using molecular markers such as restriction fragments of different lengths (restriction fragment-length polymorphisms, or RFLPs). Cytological maps (Figure 15.2, center) are based on the banding patterns of chromosomes observed with the microscope after treatment with various stains (Chapter 6). Physical maps (Figure 15.2, right), such as the restriction maps (Figure 15.2, top right) discussed in Chapter 14, are based on the molecular distancesbase pairs (bp), kilobases (kb, 1000 bp), and megabases (mb, 1 million bp)-separating sites on the giant DNA molecules present in chromosomes. Physical maps often contain the locations of overlapping genomic clones or contigs (Figure 15.2, right center) and unique nucleotide sequences called sequence-tagged sites, or STSs (Figure 15.2, bottom right).

Physical maps of a chromosome can be correlated with the genetic and cytological maps in several ways. Genes that have been cloned can be positioned on the cytological map by *in situ* hybridization (see Appendix C). Correlations between the genetic and physical maps can be established by locating clones of genetically mapped genes or RFLPs on the physical map. Markers that are mapped both genetically and physically are called *anchor markers*; they anchor the physical map to the genetic map and vice versa. Physical maps of chromosomes can also be correlated with genetic and cytological maps by using (1) PCR (see Figure 14.6) to amplify short—usually 200 to 500 bp—unique genomic DNA sequences, (2) Southern blots to relate these sequences to overlapping clones on physical maps, and (3) *in situ* hybridization to determine their chro-



■ FIGURE 15.2 Correlation of the genetic, cytological, and physical maps of a chromosome. Genetic map distances are based on crossing over frequencies and are measured in percentage recombination, or centiMorgans (cM), whereas physical distances are measured in kilobase pairs (kb) or megabase pairs (mb). Restriction maps, contig maps, and STS (sequence-tagged site) maps are described in the text.

mosomal locations (cytological map positions). These short, unique anchor sequences are called *sequence-tagged sites* (*STSs*). Another approach uses short cDNA sequences (DNA copies of mRNAs), or *expressed-sequence tags* (*ESTs*), as hybridization probes to anchor physical maps to RFLP maps (genetic maps) and cytological maps.

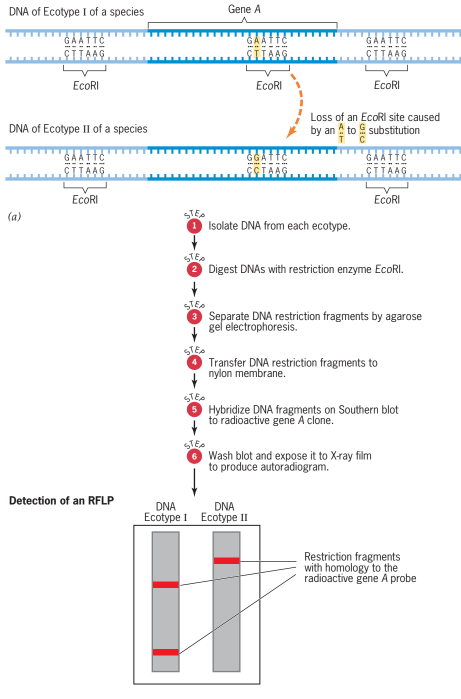
Physical distances do not correlate directly with genetic map distances because recombination frequencies are not always proportional to molecular distances. However, the two are often reasonably well correlated in euchromatic regions of chromosomes. In humans, 1 cM is equivalent, on average, to about 1 mb of DNA.

RESTRICTION FRAGMENT-LENGTH POLYMORPHISM (RFLP) AND SHORT TANDEM REPEAT (STR) MAPS

When mutations change the nucleotide sequences in restriction enzyme cleavage sites, the enzymes no longer recognize them (**Figure 15.3***a*). Other mutations may create new restriction sites. These mutations result in variations in the lengths of the DNA fragments produced by digestion with various restriction enzymes (**Figure 15.3***b*). Such **restriction fragment-length polymorphisms**, or **RFLPs**, have proven invaluable in constructing detailed genetic maps for use in positional cloning. The RFLPs are mapped just like other genetic markers; they segregate in crosses like codominant alleles.

The DNAs of different geographical isolates, different ecotypes (strains adapted to different environmental conditions), and different inbred lines of a species contain **FIGURE 15.3** The mutational origin (a) and detection (b) of RFLPs in different ecotypes of a species. In the example shown, an $A:T \rightarrow G:C$ base-pair substitution results in the loss of the central *Eco*RI recognition sequence present in gene *A* of the DNA of ecotype I. This mutation might have occurred in an ecotype II ancestor during the early stages of its divergence from ecotype I.

Mutational origin of an RFLP



(b)

many RFLPs that can be used to construct detailed genetic maps. Indeed, the DNAs of different individuals—even relatives—often exhibit RFLPs. Some RFLPs can be visualized directly when the fragments in DNA digests are separated by agarose gel electrophoresis, stained with ethidium bromide, and viewed under ultraviolet light. Other RFLPs can be detected only by using specific cDNA or genomic clones as radioactive hybridization probes on genomic Southern blots (Figure 15.3*b*). The RFLPs themselves are the phenotypes used to classify the progeny of crosses as parental or recombinant. RFLPs segregate as codominant markers in crosses, with the restriction fragments from both of the homologous chromosomes visible in gels or detected on autoradiograms of Southern blots produced from the gels.

RFLP markers have proven especially valuable in mapping the chromosomes of humans, where researchers must rely on the segregation of spontaneously occurring mutant alleles in families to estimate map distances. Pedigree-based mapping of this type is done by comparing the probabilities that the genetic markers segregating in the pedigree are unlinked or linked by various map distances. In 1992, geneticists used this procedure to construct an early map of about 2000 RFLPs on the 24 human chromosomes. **■** Figure 15.4 shows the correlation between an RFLP map and the cytological map of human chromosome 1.

In humans, the most useful RFLPs involve short sequences that are present as tandem repeats. The number of copies of each sequence present at a given site on a chromosome is highly variable. These sites, called variable number tandem repeats (VNTRs, also called minisatellites) and short tandem repeats (STRs, also called microsatellites) are therefore highly polymorphic. VNTRs and STRs vary in length not because of differences in the positions of restriction enzyme cleavage sites, but because of differences in the number of copies of the repeated sequence between the restriction sites. The use of VNTRs and STRs in humans is discussed in more detail in Chapter 16 (see the section DNA Profiling).

STRs have proven extremely valuable in constructing high-density maps of eukaryotic chromosomes. STRs are polymorphic tandem repeats of sequences only two to five nucleotide pairs long. STRs composed of polymorphic tandem repeats of the dinucleotide sequence AC/TG (AC in one strand; TG in the complementary strand) provide especially useful markers in humans. In 1996, a group of French and Canadian researchers published a comprehensive map of 5264 AC/TG STRs in the human genome. These STRs defined 2335 sites with an average distance of 1.6 cM or about 1.6 mb between adjacent markers.

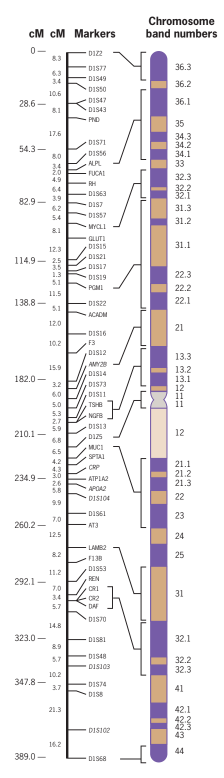
By 1997, a large international consortium had used RFLPs to map over 16,000 human genes (ESTs and cloned genes) and had integrated their map with the physical map of the human genome. In this collaborative study, over 20,000 STSs were mapped to 16,354 distinct loci. These genetic maps, composed primarily of RFLP markers, VNTRs, and STRs, have made it possible to identify and characterize mutant genes that are responsible for many human diseases (Chapter 16).

CYTOGENETIC MAPS

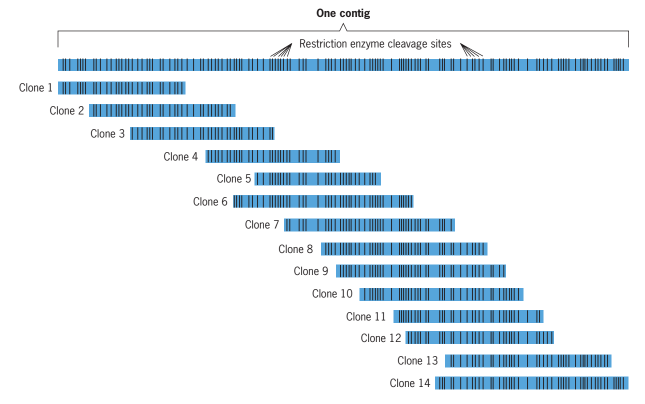
In some species, genes and clones can be positioned on the cytological maps of the chromosomes by *in situ* hybridization (Appendix C). For example, in *Drosophila*, the banding patterns of the giant polytene chromosomes in the salivary glands provide high-resolution maps of the chromosomes (Chapter 6). Thus, a clone of unknown genetic content can be positioned on the cytological map with considerable precision. In mammals, including humans, fluorescent *in situ* hybridization (FISH; see Appendix C: *In Situ* Hybridization) can be used to position clones on chromosomes stained by any of several chromosome-banding protocols. Figure 1 in Appendix C illustrates how FISH can be used to determine the chromosomal location of specific DNA sequences on human chromosomes. If RFLPs can be identified that overlap with these sequences, they can be used as STS sites that anchor genetic maps of chromosomes to cytological maps by Southern blot hybridization experiments (Chapter 14), they can also be used to tie the physical maps to both the genetic and cytological maps of the chromosomes.

PHYSICAL MAPS AND CLONE BANKS

The RFLP mapping procedure has been used to construct detailed genetic maps of chromosomes, which, in turn, have made positional cloning feasible. These genetic maps have been supplemented with physical maps of chromosomes. By isolating and preparing restriction maps of large numbers of genomic clones, overlapping clones can be identified and used to construct physical maps of chromosomes and even entire genomes. In principle, this procedure is simple (**■** Figure 15.5). However, in practice,



■ FIGURE 15.4 Correlation of the RFLP map (left) and the cytological map (right) of human chromosome 1. Molecular markers and a few genes are shown in the center. Distances are in centiMorgans (cM), with the uppermost marker set at position 0 on the left and distances between adjacent markers shown in the second column from the left. The brackets on the left of the cytological map show the chromosomal locations of the indicated genes and molecular markers.



■ FIGURE 15.5 A contig map produced from overlapping genomic clones. Large—200 to 500 kb—genomic clones, such as those present in PAC and BAC vectors (Chapter 14), are used to construct contig maps. Restriction maps of individual clones are prepared and searched by computer for overlaps. Overlapping clones are then organized into contig maps like the one shown here. When the physical map of a genome is complete, each chromosome will be represented by a single contig map.

it is a formidable task, especially for large genomes. The restriction maps of large genomic clones in PAC and BAC vectors (Chapter 14) are analyzed by computer and organized in overlapping sets of clones called **contigs**. As more data are added, adjacent contigs are joined; when the physical map of a genome is complete, each chromosome will correspond to a single contig map.

The construction of physical maps of entire genomes requires that vast amounts of data be searched for overlaps. Nevertheless, detailed physical maps are available for several genomes, including the human, *C. elegans, D. melanogaster*, and *A. thaliana* genomes. These physical maps have been used to prepare clone banks that contain catalogued clones collectively spanning entire chromosomes. Thus, if a researcher needs a clone of a particular gene or segment of a chromosome, that clone may already have been catalogued in the clone bank, or clone library, and be available on request. Obviously, the availability of such clone banks and the correlated physical maps of entire genomes are dramatically accelerating genetic research. Indeed, searching for a specific gene with and without the aid of a physical map is like searching for a book in a huge library with and without a computer catalog giving the locations of the books in the library.

KEY POINTS

- **Solution** Genetic maps of chromosomes are based on recombination frequencies between markers.
 - Cytogenetic maps are based on the location of markers within, or near, cytological features of chromosomes observed by microscopy.
 - Physical maps of chromosomes are based on distances in base pairs, kilobase pairs, or megabase pairs separating markers.
 - High-density maps that integrate the genetic, cytological, and physical maps of chromosomes have been constructed for many chromosomes, including all of the human chromosomes.

Map Position-Based Cloning of Genes

The first eukaryotic genes to be cloned were genes that are expressed at very high levels in specialized tissues or cells. For example, about 90 percent of the protein synthesized in mammalian reticulocytes is hemoglobin. Thus, α - and β -globin mRNAs could Detailed genetic, cytogenetic, and physical maps of chromosomes allow scientists to isolate genes by chromosome walks and chromosome jumps.

be easily isolated from reticulocytes and used to prepare radioactive cDNA probes for genomic library screens. However, most genes are not expressed at such high levels in specialized cells. Thus, how are genes that are expressed at moderate or low levels cloned? One important approach has been to map the gene precisely and to search for a clone of the gene by using procedures that depend on its location in the genome. This approach, called **positional cloning**, can be used to identify any gene, given an adequate map of the region of the chromosome in which it is located.

The steps in positional cloning are illustrated in **Figure 15.6**. The gene is first mapped to a specific region of a given chromosome by genetic crosses or, in the case of humans, by pedigree analysis, which usually requires large families. The gene is next localized on the physical map of this region of the chromosome. Candidate genes in the segment of the chromosome identified by physical mapping are then isolated from mutant and wild-type individuals and sequenced to identify mutations that would result in a loss of gene function. As we will discuss in Chapter 16, the human genes responsible for inherited disorders such as Huntington's disease

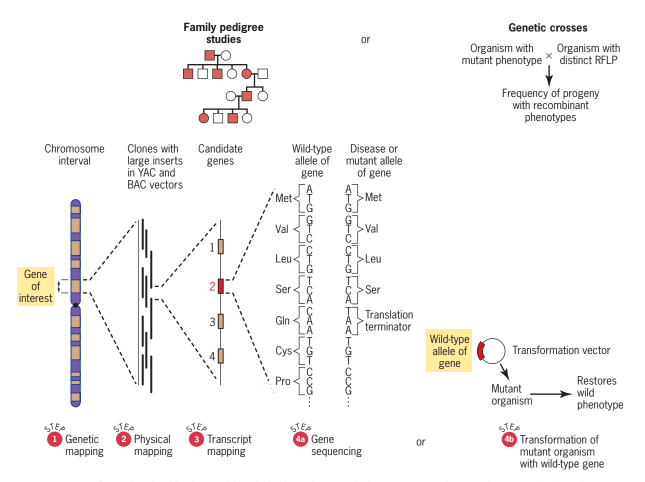
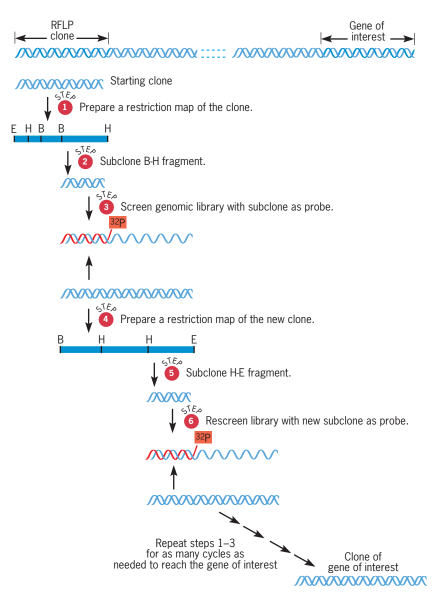


FIGURE 15.6 Steps involved in the positional cloning of genes. In humans, genetic mapping must be done by pedigree analysis, and candidate genes must be screened by sequencing wild-type and mutant alleles (step 4a). In other species, the gene of interest is mapped by appropriate genetic crosses, and the candidate genes are screened by transforming the wild-type alleles into mutant organisms and determining whether or not they restore the wild-type phenotype (step 4b).



■ FIGURE 15.7 Positional cloning of a gene by chromosome walking. A chromosome walk starts with the identification of a molecular marker (such as the RFLP shown at top) close to the gene of interest and proceeds by repeating steps 1 through 3 as many times as is required to reach the gene of interest (bottom).

and cystic fibrosis have been identified by using the positional cloning approach. In species where transformation is possible, copies of the wild-type alleles of candidate genes are introduced into mutant organisms to determine whether the wild-type genes will restore the wild phenotype. Restoration of the wild phenotype to a mutant organism provides strong evidence that the introduced wild-type gene is the gene of interest.

CHROMOSOME WALKS AND JUMPS

Positional cloning is accomplished by mapping the gene of interest, identifying an RFLP, VNTR, STR, or other molecular marker near the gene, and then "walking" or "jumping" along the chromosome until the gene is reached.

Chromosome walks are initiated by the selection of a molecular marker (RFLP or known gene clone) close to the gene of interest and the use of this clone as a hybridization probe to screen a genomic library for overlapping sequences. Restriction maps are constructed for the overlapping clones identified in the library screen, and the restriction fragment farthest from the original probe is used to screen a second genomic library constructed by using a different restriction enzyme or to rescreen a library prepared from a partial digest of genomic DNA. Repeating this procedure several times and isolating a series of overlapping genomic clones allow a researcher to walk along the chromosome to the gene of interest (**Figure 15.7**). Without information about the orientation of the starting clone on the linkage map, the initial walk will have to proceed in both directions until another RFLP is identified and it is determined whether the new RFLP is closer to or farther away from the gene of interest than is the starting RFLP.

Verification that a clone of the gene of interest has been isolated is accomplished in various ways. In experimental organisms such as *Drosophila* and *Arabidopsis*, verification is achieved by introducing the wild-type allele of the gene into a mutant organism and showing that it restores the wild-type phenotype. In humans, verification usually involves determining the nucleotide

sequences of the wild-type gene and several mutant alleles and showing that the coding sequences of the mutant genes are defective and unable to produce functional gene products.

Chromosome walking is very difficult in species with large genomes (the walk is usually too far) and an abundance of dispersed repetitive DNA (each repeated sequence is a potential roadblock). Chromosome walking is easier in organisms such as *A. thaliana* and *C. elegans*, which have small genomes and little repetitive DNA.

When the distance from the closest molecular marker to the gene of interest is large, a technique called **chromosome jumping** can be used to speed up an otherwise long walk. Each jump can cover a distance of 100 kb or more. Like a walk, a jump is initiated by using a molecular probe such as an RFLP, VNTR, or STR as a starting point. However, with chromosome jumps, large DNA fragments are prepared by partial digestion of genomic DNA with a restriction endonuclease. The large genomic fragments are then circularized with DNA ligase. A second restriction endonuclease is used to excise the junction fragment from the circular molecule. This junction fragment will contain both ends of the long fragment; it can be identified by hybridizing

the DNA fragments on Southern blots to the initial molecular probe. A restriction map of the junction fragment is prepared, and a restriction fragment that corresponds to the distal end of the long genomic fragment is cloned and used to initiate a chromosome walk or a second chromosome jump. Chromosome jumping has proven especially useful in work with large genomes such as the human genome. Chromosome jumps played a key role in identifying the human cystic fibrosis gene (Chapter 16).

- Detailed genetic, cytogenetic, and physical maps of chromosomes permit researchers to isolate genes based on their location in the genome.
- If a molecular marker such as a restriction fragment-length polymorphism (RFLP), variable number tandem repeat (VNTR), or short tandem repeat (STR) maps close to a gene, the gene can usually be isolated by chromosome walks or chromosome jumps.

KEY POINTS

The Human Genome Project

As the recombinant DNA, gene cloning, and DNA sequencing technologies improved in the 1970s and early 1980s, scientists began discussing the possibility of sequencing all 3×10^{9} nucleotide pairs in the human genome. These discussions led to the launching of the **Human Genome Project** in 1990. The initial

Detailed genetic, cytogenetic, and physical maps are available for all 24 human chromosomes, and complete, or nearly complete, nucleotide sequences are available for the genomes of many species, including *Homo sapiens*.

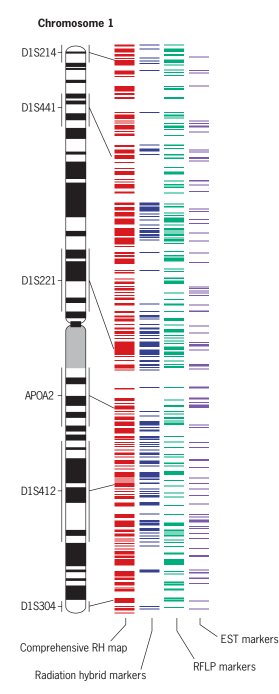
goals of the Human Genome Project were (1) to map all of the human genes, (2) to construct a detailed physical map of the entire human genome, and (3) to determine the nucleotide sequences of all 24 human chromosomes by the year 2005. Scientists soon realized that this huge undertaking should be a worldwide effort. Therefore, an international Human Genome Organization (HUGO) was established to coordinate the efforts of human geneticists around the world.

James Watson, who, with Francis Crick, discovered the double-helix structure of DNA, was the first director of this ambitious project, which was expected to take nearly two decades to complete and to cost in excess of \$3 billion. In 1993, Francis Collins, who, with Lap-Chee Tsui, led the research teams that identified the cystic fibrosis gene, replaced Watson as director of the Human Genome Project. In addition to work on the human genome, the Human Genome Project has served as an umbrella for similar mapping and sequencing projects on the genomes of several other organisms, including the bacterium *E. coli*, the yeast *S. cerevisiae*, the fruit fly *D. melanogaster*; the plant *A. thaliana*, and the worm *C. elegans*.

MAPPING THE HUMAN GENOME

Rapid progress was made in mapping the human genome from the launching of the Human Genome Project. Complete physical maps of chromosomes Y and 21 and detailed RFLP maps of the X chromosome and all 22 autosomes were published in 1992. By 1995, the genetic map contained markers separated by, on average, 200 kb. A detailed STR map of the human genome was published in 1996, and a comprehensive map of 16,354 distinct loci was released in 1997. All of these maps have proven invaluable to researchers cloning genes based on their locations in the genome.

Unfortunately, the resolution of genetic mapping in humans is quite low—in the range of 1–10 mb. The resolution of fluorescent *in situ* hybridization (FISH) is also approximately 1 mb. Higher resolution mapping (down to 50 kb) can be achieved by radiation hybrid mapping, a modification of the somatic-cell hybridization mapping procedure. Standard somatic-cell hybridization involves the fusion of human cells and rodent cells growing in culture and the correlation of human gene products with human chromosomes retained in the hybrid cells.



■ FIGURE 15.8 A high-resolution map of human chromosome 1. The cytogenetic map of chromosome 1 is shown on the left, along with the locations of six anchor markers. To the right of the cytogenetic map are four genetic maps that show the locations of the comprehensive radiation hybrid markers (red lines), the high-confidence radiation hybrid markers (blue lines), the RFLP markers (green lines), and the ESTs (purple lines).

Radiation hybrid mapping is performed by fragmenting the chromosomes of the human cells with heavy irradiation prior to cell fusion. The irradiated human cells are then fused with Chinese hamster (or other rodent) cells growing in culture, usually in the presence of a chemical such as polyethylene glycol to increase the efficiency of cell fusion. The human–Chinese hamster somatic-cell hybrids are then identified by growth in an appropriate selection medium.

Many of the human chromosome fragments become integrated into the Chinese hamster chromosomes during this process and are transmitted to progeny cells just like the normal genes in the Chinese hamster chromosomes. The polymerase chain reaction (PCR; see Chapter 14) is then used to screen a large panel of the selected hybrid cells for the presence of human genetic markers. Chromosome maps are constructed based on the assumption that the probability of an X ray-induced break between two markers is directly proportional to the distances separating them in chromosomal DNA.

Several groups used the radiation hybrid mapping procedure to construct highdensity maps of the human genome. In 1997, Elizabeth Stewart and coworkers published a map of 10,478 STSs based on radiation hybrid mapping data; their map of human chromosome 1 is shown in **Figure 15.8**.

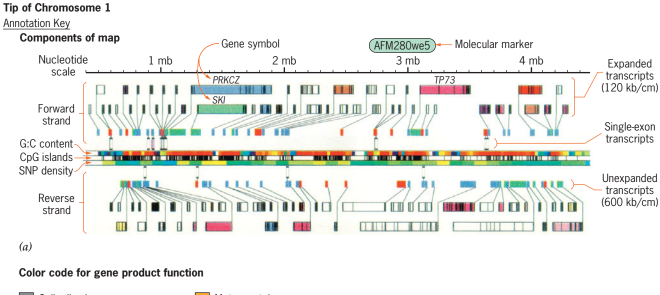
SEQUENCING THE HUMAN GENOME

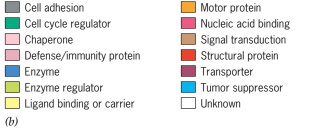
Whereas the gene-mapping work advanced quickly, progress toward sequencing the human genome initially lagged behind schedule. However, that all changed rapidly beginning in 1998. During May of 1998, J. Craig Venter announced that he had formed a private company, Celera Genomics, with the goal of sequencing the human genome in just three years. (For details, see A Milestone in Genetics: Two Drafts of the Sequence of the Human Genome on the Student Companion site.) Shortly thereafter, the leaders of the public Human Genome Project's sequencing laboratories announced that they had revised their schedule and planned to complete the sequence of the human genome by 2003—two years earlier than originally proposed. From that point in time, everything accelerated.

The complete sequence of the first human chromosome—small chromosome 22—was published in December 1999. The complete sequence of human chromosome 21 followed in May of 2000. Then, with the intervention of the White House, Venter, of Celera Genomics, and Francis Collins, director of the public Human Genome Project, agreed to publish first drafts of the sequence of the human genome at the same time. The Celera and public sequences were both published in February 2001. **Figure 15.9** shows an annotated, sequence-based map of a 4-mb segment at the tip of the short arm of human chromosome 1. This map illustrates the positions and orientations of known and predicted genes in one small portion of the human genome. For similar maps of the entire human genome, see the February 15, 2001, issue of *Nature* and the February 16, 2001, issue of *Science*.

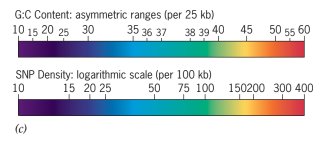
The amount of information in these first drafts of the human genome was overwhelming, including the sequence of over 2650 megabase pairs of DNA (over 2,650,000,000 bp). The human genome is more than 25 times the size of the previously sequenced *Drosophila* and *Arabidopsis* genomes, and more than eight times the sum of all the genomes sequenced before it.

The sequence of the human genome provided one surprise: there appeared to be only about 25,000 to 30,000 genes rather than the estimated 50,000 to 120,000 genes suggested by earlier studies. The distribution of functions for the 26,383 genes predicted by the Celera sequence is shown in **Figure 15.10**. About 60 percent of the predicted proteins have similarities with proteins of other species whose genomes have been sequenced (**Figure 15.11**). Over 40 percent of the predicted human proteins share similarities with *Drosophila* and *C. elegans* proteins. Indeed, only 94 of 1278 protein families predicted by the sequence of the human genome are specific to vertebrates. The rest have evolved from domains of proteins in distant ancestors, including prokaryotes and unicellular eukaryotes.





Color code for G:C content and single-nucleotide polymorphism (SNP) density



■ FIGURE 15.9 Annotated, sequence-based map of a 4-mb segment of DNA at the tip of human chromosome 1, assembled by researchers at Celera Genomics. (a) The top line gives distances in mb. The next three panels show predicted transcripts from one strand of DNA (the "forward strand"), whereas the bottom three panels show transcripts specified by the other strand of DNA (the "reverse strand"). The middle three panels give the G:C content, the positions of CpG islands, which occur upstream of genes, and the density of single-nucleotide polymorphisms (SNPs), respectively. (b) The color code for gene-product functions, and (c) the color codes for G:C content and SNP density.

On average, there is one gene per 145 kb in the human genome, although there is some clustering of highly expressed genes in euchromatic regions of specific chromosomes. The average human gene is about 27,000 bp in length and contains 9 exons. Exons make up only 1.1 percent of the genome, whereas introns account for 24 percent, with 75 percent of the genome being intergenic DNA. Of the intergenic DNA, at least 44 percent is derived from transposable genetic elements (see Chapter 17 for details).

The two first drafts of the sequence of the human genome were incomplete, containing over 100,000 gaps. Therefore, the International Human Genome Sequencing Consortium continued to work on filling in these gaps and completing the sequence. By October 2004, they had reduced the number of gaps to 341 and had completed the sequence of 99 percent of the euchromatic DNA in the human genome. Surprisingly, the estimated number of genes in the genome had decreased again—to just 22,287

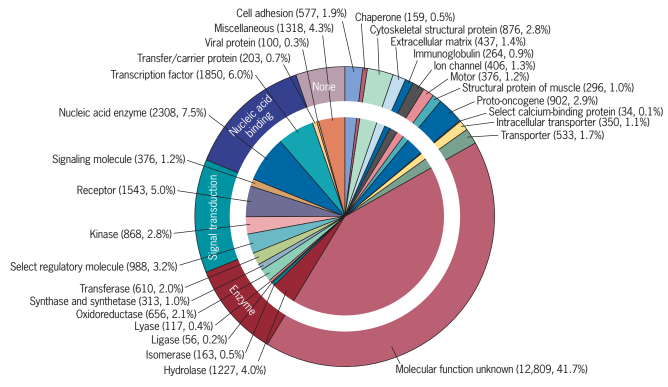
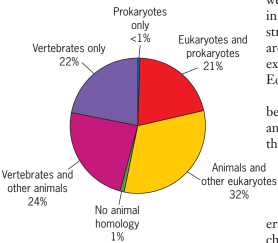


FIGURE 15.10 Functional classification of the 26,383 genes predicted by Celera Genomics' first draft of the sequence of the human genome. Each sector gives the number and percentage of gene products in each functional class in parentheses. Note that some classes overlap: a proto-oncogene, for example, may encode a signaling molecule.



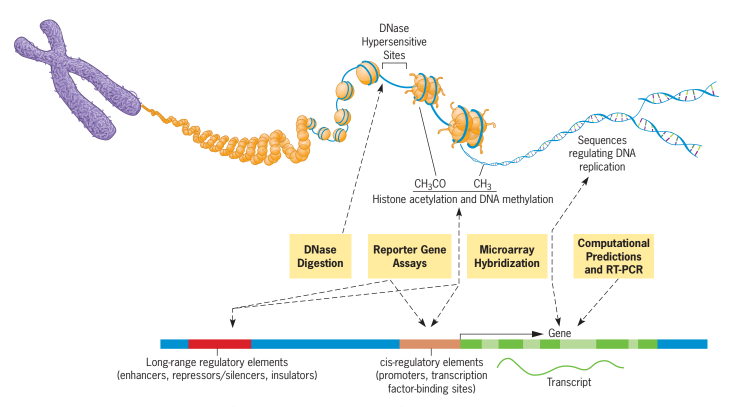
■ FIGURE 15.11 Pie chart showing homology of predicted human proteins to proteins of other species where homologues were detected by computer searches of public databases.

protein-coding genes—in the more complete sequence. There are, of course, other genes that specify RNA products—rRNAs, tRNAs, snRNAs, and miRNAs—that will increase the total number of genes significantly.

With the development of new sequencing technology that enables large genomes like the human genome to be sequenced quickly and at much lower costs, it has become feasible to sequence individual human genomes (Chapter 14). James D. Watson and J. Craig Venter were the first two individuals to have their genomes sequenced. Then, Jeffrey M. Kidd and coworkers mapped and sequenced structural variation in eight human genomes of diverse origin. The individuals selected for study were of African, Asian, and European ancestry. The researchers focused on changes in the genomes in the range of 1 kb to 1 mb and documented a large amount of structural diversity—especially deletions, inversions, and insertions. Today, scientists are busy sequencing 2,500 human genomes from around the world to determine the extent of sequence variability in human genomes of diverse origin (see On the Cutting Edge: The 1000 Genomes Project in Chapter 9).

The wealth of information provided by the sequences of human genomes is just beginning to be exploited. Given that only about 1.1 percent of the genome encodes amino acid sequences in polypeptides, the big question is what are the functions of the rest of the components of the human genome. Francis Collins and other leaders of the Sequencing Consortium are focusing on this question. They have organized a new consortium, ENCODE (*ENCyclopedia Of DNA Elements*), whose goal is to identify all of the nongenic functional elements in the human genome. These elements will include regulatory sequences such as promoters, enhancers, silencers, sites of methylation and acetylation, and other factors involved in the control of chromatin structure and gene expression (**■ Figure 15.12**).

The identification and functional characterization of the nongenic elements in the human genome has already become a fascinating story. The functions of the vast noncoding "dark matter" in the genome is proving to be a complex and intriguing subject. Large segments of noncoding DNA are conserved between mice and humans,



■ FIGURE 15.12 The goal of the ENCODE (*ENC*yclopedia *Of D*NA *E*lements) Project Consortium is to identify the nongenic functional elements in the human genome. The elements will include regulatory sequences such as promoters, enhancers, silencers, repressor-binding sites, transcription factor-binding sites, and sites of chemical modifications such as acetylation and methylation. They will also include sequences that alter chromatin structure by interacting with DNA-binding proteins and the histones that package DNA into nucleosomes. Some of these elements will alter chromatin structure producing DNase hypersensitive sites (characteristic of chromatin that is transcriptionally active—see Chapter 19). Tools to be used in these studies will include reporter gene assays and microarray hybridizations (discussed in subsequent sections of this chapter) and reverse-transcript PCR (RT-PCR)—polymerase chain reactions using RNAs as templates to identify transcribed regions of the genome.

which suggests they must contain important functional elements. When researchers look for sequences that increase the risks of human diseases, about 40 percent are located in intergenic regions. Moreover, recent studies indicate that about 80 percent of the genome is actually transcribed into RNAs of unknown function. Some of this noncoding RNA consists of important small regulatory miRNAs (see Chapter 19). Another component includes "large intervening noncoding RNAs" with regulatory and unknown functions. This "dark matter" in the genome also includes chemically modified sequences that control the epigenetic expression of genes from one generation to the next by modifying chromatin structure (see Chapter 19). Clearly, we have a lot to learn about the noncoding components of the human genome.

Another international consortium—the *Hu*man *P*roteome *O*rganization (HUPO)—has been formed with the goal of determining the structures and functions of all the proteins encoded by the human genome. Despite the wealth of data provided by the sequence of the human genome, the functional dissection of the genome is just beginning. We still have a long way to go before we will really understand the structure and function of the 3×10^{9} nucleotide pairs in the human genome.

The availability of the sequence of the human genome raises a whole new set of questions about the proper use of this new knowledge. Many of these questions focus on an individual's right to privacy. For example, if a mutation that causes a lateonset disorder such as Huntington's disease (Chapter 16) is discovered in a family, who should have access to this information? If such information were available to the public, widespread discrimination might occur. Employers might not hire members of the family, and medical schools might not admit talented young scholars to their M.D. programs. Will insurance companies provide health and life insurance to someone who carries a mutant gene that increases the risk of cancer or leads to a lateonset disorder like Huntington's disease? If they do, will the insurance be affordable, or will it be priced beyond the reach of all but the wealthy? Given the recent increase in the amount of genetic information available, it seems clear that laws protecting the privacy of this information will be needed in the future. Indeed, in the United States, the Genetic Information Nondiscrimination Act (GINA) was passed in 2008, and it should protect individuals from discrimination by employers and insurance companies based on information obtained by genetic studies and DNA tests.

THE HUMAN HAPMAP PROJECT

Human genomes contain a large amount of genetic variation. In Chapter 6 we discussed gross changes—deletions, duplications, inversions, and translocations—in the structures of genomes. In the preceding section, we discussed changes of intermediate size—deletions, insertions, and inversions in the range of 1 kb to 1 mb—in eight human genomes. Small changes—insertions or deletions of one or a few nucleotide pairs—are even more frequent. The most common changes in human genomes are single nucleotide-pair substitutions, for example, A:T to G:C or G:C to A:T substitutions (Chapter 13). Basepair substitutions of this type have produced a large number of *single-nucleotide polymorphisms* (SNPs, pronounced "snips") in human genomes. Most of these SNPs are not located in the coding regions of genes and do not result in mutant phenotypes. When the nucleotide sequences of the same chromosomes of two individuals are compared, one SNP is present, on average, in every 1200 nucleotide pairs.

SNPs can be detected in human genomes by the microarray hybridization or "gene-chip" technology described in a later section of this chapter (see Microarrays and Gene Chips). In brief, hybridization probes can be synthesized that can detect single-nucleotide differences in DNA molecules. If a DNA molecule matches a probe exactly, it will bind to that probe; if it does not match exactly, it will not bind. Thus, if a segment of DNA from one individual has an A:T base pair at a specific position, and the corresponding segment of DNA from another individual has a G:C base pair at this position, it is possible to distinguish these two individuals genetically by hybridizing their DNA to probes that will bind to one or the other of the two DNA segments. These and thousands of other diagnostic probes can be arrayed systematically on a silicon wafer (see Figure 15.16) to screen for single-nucleotide differences in genomic DNA collected from a sample of individuals. Usually the DNA from each individual is amplified by PCR using primers that flank genomic regions of interest, and the amplified DNA is labeled in some way before hybridizing it with the diagnostic array of probes. In a study conducted at Perlegen Sciences, Inc., researchers used this microarray technology to determine the genotypes of 71 people at more than 1.5 million sites in the human genome—an amazing accomplishment! By studying these polymorphisms in different subpopulations, it may be possible to trace important genetic events in the evolutionary history of our species and to predict a person's susceptibility to diseases like cancer and heart disease.

Individual SNPs may be present in one human population and absent from another. When present, they may vary in frequency from one population to another. Most SNPs present in human populations were produced by a single mutation in one individual that subsequently spread through the population. Each SNP is associated with other SNPs that were present on the ancestral chromosome at the time that the mutation generating the SNP occurred. SNPs that are closely linked tend to be passed on to progeny as a unit because there is little chance for crossovers to shuffle them into new combinations. The SNPs on a chromosome or a segment of a chromosome that tend to be inherited together define a genetic unit called a haplotype (**Figure 15.13**). Of course, mutation will modify haplotypes, and crossing over will generate new haplotypes during the course of evolution.

Because of their frequency and distribution throughout the human genome, SNPs have proven to be valuable genetic markers. The study of haplotypes defined by SNPs is providing important information about the relationships among different

■ FIGURE 15.13 Haplotypes are sets of linked SNPs and other genetic markers that tend to

be inherited as a unit.

DNA from:	SNP1	SNP2	SNP3	SNP4
Individual 1	TACA <mark>G</mark> GATC ·	· · · TTCG <mark>G</mark> GCC/	A AATC <mark>C</mark> ATGC	···· AAGC <mark>T</mark> CGGA ····
Individual 2	TACA <mark>A</mark> GATC ·	TTCG <mark>A</mark> GCCA	a aatc <mark>c</mark> atgc	AAGC <mark>T</mark> CGGA
Individual 3	taca <mark>g</mark> gatc ∙	ttcg <mark>a</mark> gcc <i>i</i>	a aatc <mark>t</mark> atge	AAGC <mark>T</mark> CGGA
Individual 4	TACA <mark>G</mark> GATC ·	TTCG <mark>A</mark> GCC#	a aatc <mark>c</mark> atgc	AAGC <mark>C</mark> CGGA
		 *	-,: `` ' ' ':	i
		Haplotype 1 G	ĠĊŤ	
		Haplotype 2 A	АСТ	
		Haplotype 3 G	A <mark>T</mark> T	
		Haplotype 4 G	A C C	

ethnic groups and about human evolution (see Chapter 24). The study of SNPs and haplotypes is also helping researchers identify genes that are involved in susceptibility to diseases such as breast cancer, glaucoma, amyotrophic lateral sclerosis (ALS, also known as Lou Gehrig's disease), and rheumatoid arthritis. The strategy in these studies has been to determine the SNP genotypes of large samples of people and then search for associations between the SNPs (or the haplotypes defined by linked SNPs) and particular diseases. Once an association has been found, the SNP or haplotype can be used to help predict the risk that an individual will develop the disease, and in favorable cases, it may help to identify the actual disease-causing gene.

Because of the value of SNP haplotypes in studying ancestry and evolution in human populations and in finding disease associations, researchers from around the world have initiated the International HapMap Project. The goal of this collaborative enterprise is to identify and map SNPs using DNA samples from many different human populations. The data collected by the Project are being made available as a resource for all genomic researchers.

- Researchers collaborating on the Human Genome Project have constructed detailed maps of all 24 human chromosomes.
- Other participants of the Human Genome Project have determined the complete, or nearly complete, nucleotide sequences of the genomes of several important model organisms.
- A nearly complete sequence of the euchromatic DNA in the human genome was released in October 2004.
- Scientists throughout the world have initiated the International Human HapMap Project with the goal of characterizing the similarities and differences in human genomes worldwide.

RNA and Protein Assays of Genome Function

Knowing the complete sequence of the human genome will help identify genes responsible for human diseases and should lead to successful gene therapies for some of these diseases. However, it will not tell us what these genes do or how they control biological processes. Indeed, by itself, the nucleotide sequence of a gene, a chromosome, or an entire genome is uninformative. Only when supplemented with information about their The availability of the nucleotide sequences of entire genomes has led to the development of microarray, gene-chip, and reporter gene technologies that permit researchers to study the expression of all the genes of an organism simultaneously.

functions do sequences become truly meaningful. Thus, information about the functions of nucleotide sequences must still be obtained by traditional genetic studies and by molecular analyses. If geneticists want to understand the genetic control of the growth and development of a mature human from a single fertilized egg (Chapter 20), they will need to know much more than the sequence of the human genome. But the availability of KEY POINTS

the ultimate map of the human genome, its nucleotide sequence, will certainly accelerate progress toward understanding the programs of gene expression that control morphogenesis. Indeed, the development of new technologies such as "gene-chip" hybridizations is designed to take advantage of the availability of the sequences of complete genomes (see the section Microarrays and Gene Chips).

EXPRESSED SEQUENCES

In large eukaryotic genomes, only a small proportion of the DNA encodes proteins. In the yeast *S. cerevisiae*, almost 70 percent of the genome encodes proteins, and there is one gene for every 2 kb of sequence. In humans, only about 1 percent of the genome encodes amino acid sequences, and there is one gene for every 130 kb of sequence. Thus, in order to focus on the protein-coding content of genomes, many scientists have analyzed cDNA clones (DNAs complementary to RNA molecules; see Chapter 14) or ESTs rather than genomic clones. By 1996, the public databases contained more than 600,000 cDNA sequences, about 450,000 of which were human cDNAs. The number of human cDNA sequences nearly doubled to about 800,000 by late 1997. However, many of these cDNA sequences are derived from the same gene transcripts. Multiple cDNAs can be obtained from different segments of a single gene transcript or from alternative splicing of a gene transcript. For example, the human gene that encodes serum albumin is represented by more than 1300 EST sequences in public databases.

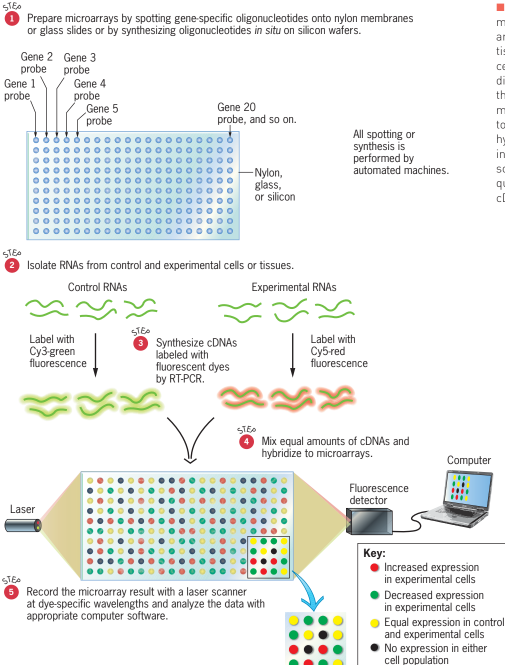
The transcripts of different genes can usually be recognized by distinct 3' untranslated regions—different nucleotide sequences in the region between the 3' translation-termination codon and the 3' terminus of the transcript. When sequence comparisons were made between the regions of cDNAs corresponding to the 3' untranslated regions of the transcripts, the cDNAs were grouped into 49,625 clusters (with 97 percent sequence identity within clusters). Prior to the publication of the two drafts of the human genome, the number of clusters was considered a good estimate of the number of distinct genes. Of the sequence clusters, 4563 corresponded to known human genes. One problem with estimating gene numbers from EST sequences is that the ESTs may be derived from nonoverlapping regions of a gene transcript. In any case, it now seems clear that human gene number estimates based on EST databases gave gene numbers that were too high.

MICROARRAYS AND GENE CHIPS

Given the sequence of an entire genome, geneticists can immediately begin to study the expression of every gene in the organism. Oligonucleotide hybridization probes can be synthesized that are complementary to segments of the transcripts of every open reading frame (ORF), or PCR can be used to make millions of copies of each gene in a genome. Thus, scientists can monitor changes in total genome expression over time, throughout development, or in response to changes in the environment. Such knowledge should prove invaluable in understanding human diseases such as cancer and, perhaps, even the aging process.

New technologies now allow scientists to produce **microarrays** that contain thousands of hybridization probes on a single membrane or other solid support. Oligonucleotides can be synthesized that are complementary to the RNA transcripts of every gene in an organism's genome, and they can be attached to solid supports such as nylon membranes, glass slides, or silicon surfaces for use as hybridization probes. Or the oligonucleotide chains can be synthesized in microarrays on silicon surfaces or on arrays of microbeads. In the case of **gene chips**, thousands of probes are synthesized on silicon wafers 1–2 square centimeters in size. Thus, a single gene chip can be used to study the expression of thousands of genes.

RNAs to be analyzed are isolated from the cells or tissues of interest—for example, normal cells and cancer cells—and used to synthesize fluorescent dye-labeled cDNAs by RT-PCR (see Chapter 14). These labeled cDNAs are then hybridized to the probes on microarrays to compare the levels of expression of genes of interest or of all the



■ FIGURE 15.14 Preparation and use of microarrays to study gene expression. RNAs are isolated from control and experimental tissues, for example, normal cells and cancer cells, and used to prepare cDNAs labeled with different fluorescent dyes. Equal amounts of the cDNA samples are mixed and hybridized to microarrays containing probes complementary to the cDNAs of the genes of interest. After hybridization, the microarrays are analyzed using sophisticated laser scanners and computer software that remove background noise and quantify the signals from the two fluorescent cDNA populations.

genes in the genome (**■** Figure 15.14). After hybridization is complete, the arrays are washed and then scanned with lasers and fluorescence detectors with micrometer resolution, and the results are analyzed and recorded using computer software designed to remove background noise and amplify positive signals (**■** Figure 15.15). The gene chip shown in **■** Figure 15.16 contains a microarray of over 10,000 oligonucleotide probes on a single silicon wafer.

The genome sequencing projects and the microarray hybridization technologies have spawned the new subdiscipline of functional genomics, which focuses on the expression of entire genomes. However, some geneticists have argued that this has been the goal of the science of genetics since its inception. As knowledge in the field has advanced, geneticists have been able to study the expression of more and more genes. Now, for the first time, they can study the expression of all the genes of

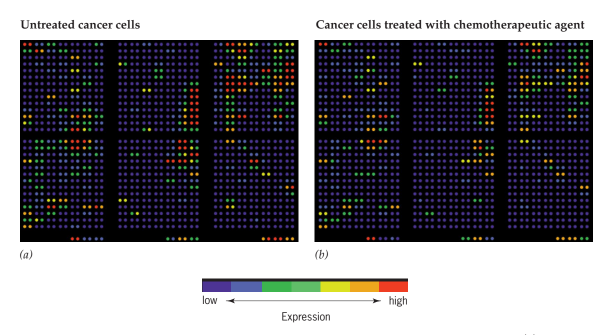


FIGURE 15.15 Microarray hybridization data comparing the levels of expression of 588 genes in (*a*) untreated human cancer cells and (*b*) human cancer cells treated with a chemotherapeutic agent. The photographs were produced using a scanner to measure the intensities of the hybridization signals on the microarrays and converting them to visual images with the appropriate computer software. Changes in levels of gene expression induced by the chemotherapeutic agent can be detected by comparing the two arrays.

an organism simultaneously. Probe microarrays are available that allow biologists to analyze the expression of the nearly 6000 genes of budding yeast. In addition, DNA chips that permit scientists to study the expression of the approximately 17,000 genes of *Drosophila melanogaster*; the roughly 26,000 genes of *Arabidopsis thaliana*, and the approximately 20,500 human genes are now available to the research community. The

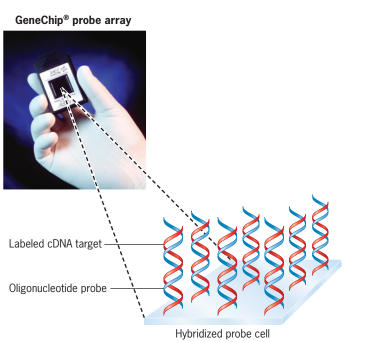


Image of a hybridized microarray

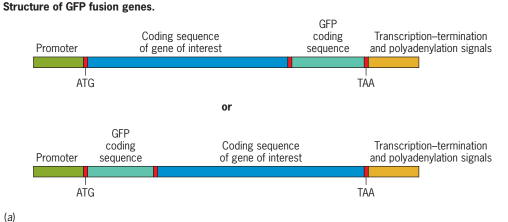
FIGURE 15.16 Photograph of a gene chip (top left) and a photograph of a hybridized microarray (top right). Gene chips and other types of microarrays allow researchers to analyze the expression of all the genes of an organism simultaneously. The gene chips contain thousands of oligonucleotide hybridization probes that allow scientists to detect the transcripts of thousands of genes in one experiment.

ability to analyze the expression of entire genomes has resulted in a vast amount of new information in biology and will eventually lead to an understanding of the normal process of human development and the causes of at least some human diseases.

THE GREEN FLUORESCENT PROTEIN AS A REPORTER OF PROTEIN SYNTHESIS

Array hybridizations and gene chips can be used to determine whether genes are transcribed, but they provide no information about the translation of the gene transcripts. Thus, biologists often use antibodies to detect the protein products of genes of interest. Western blots are used to detect proteins separated by electrophoresis (Chapter 14), and antibodies coupled to fluorescent compounds are used to detect the location of proteins *in vivo*. However, both of these approaches provide only a single time-point assay of a protein in a cell, tissue, or organism.

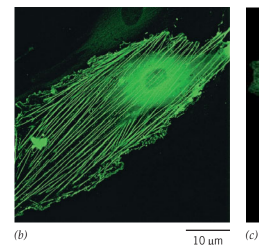
The discovery of a naturally occurring fluorescent protein, the **green fluorescent protein (GFP)** of the jellyfish *Aequorea victoria*, has provided a powerful tool that can be used to study gene expression at the protein level. GFP is now being used to monitor the synthesis and localization of specific proteins in a wide variety of living cells. These studies entail constructing fusion genes that contain the nucleotide sequence encoding GFP, coupled in frame to the nucleotide sequence encoding the protein of interest; introducing the chimeric gene into cells by transformation; and studying the fluorescence of the fusion protein in transgenic cells exposed to blue or UV light (**Transformation** Figure 15.17). Because GFP is a small protein, it can often be coupled to proteins without interfering with their activity or interaction with other cellular components.



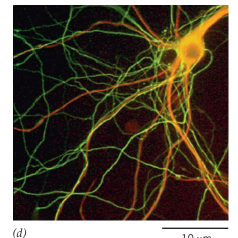
GFP-tagged tubulin

■ FIGURE 15.17 Use of the green fluorescent protein (GFP) of the jellyfish to study protein localization in living cells. (a) Structure of GFP fusion genes. The GFP coding sequence may be placed at either end of the gene of interest or at internal positions. (b-d) Immunofluorescence localization of GFP-tagged proteins: (b) smooth-muscle actin in a fibroblast cell; (c) the microtubule structural protein tubulin in Chinese hamster ovary cells; and (d) double labeling of two microtubule-binding proteins. MAP2 labeled with blue light-emitting GFP and tau labeled with green light-emitting GFP, in a rat neuron. With the light filters used for microscopy, MAP2 and tau appear red and green, respectively.

GFP-tagged actin



Supplied art is low(72)DPI. GFP-tagged MAP2 (red) plus GFP-tagged tau (green)



το μπ

10 µm

As the name implies, GFP fluoresces bright green when exposed to blue or ultraviolet light. The chromophore of GFP is produced by the posttranslational cyclization and oxidation of an encoded serine/tyrosine/glycine tripeptide. This chromophore is largely protected from ion and solvent effects by encasement in a barrel-like fold of the mature protein. Unlike other bioluminescent proteins, GFP does not require the addition of substrates, cofactors, or any other substances to fluoresce—only exposure to blue or UV light. Thus, GFP can be used to study gene expression in living cells and to study protein localization and movement in cells over time. By mutagenizing the GFP gene, molecular biologists have produced variant forms of GFP that emit blue or yellow light, variants that fluoresce up to 35 times more intensely than the wild-type GFP, and variants whose fluorescence depends on the pH of the microenvironment. These GFP variants can be used to study the synthesis and intracellular localization of two or more proteins simultaneously (Figure 15.17*d*).

Some geneticists are using GFP fusions to study changes in the expression of all the genes encoding proteins that are involved in a particular metabolic pathway in response to treatment of cells or tissues with a specific drug or potential therapeutic agent. They construct an entire set of chimeric genes containing the GFP coding region fused in frame to the coding regions of other genes, introduce them into host cells, and monitor their expression by quantifying the fluorescence of fusion proteins separated by electrophoresis or other techniques. Technologies are being developed that will allow scientists to observe changes in the levels of large arrays of GFP fusion proteins by capillary electrophoresis (electrophoresis performed in small capillary tubes), monitored by sensitive microphotodetectors and sophisticated computer software. Indeed, in the not-too-distant future, functional genomics may involve the use of DNA chips to detect gene transcripts and "protein chips" to detect the polypeptides encoded by these transcripts.

KEY POINTS

- Once the complete nucleotide sequence of a genome has been determined, scientists can study the temporal and spatial patterns of expression of all the genes of the organism.
- Microarrays of gene-specific hybridization probes on gene chips allow researchers to study the transcription of thousands of genes simultaneously.
- Chimeric genes that contain the coding region of the green fluorescent protein of the jellyfish fused with the coding regions of the genes of experimental organisms can be used to study the localization of proteins in living cells.

Comparative Genomics

Comparisons of the nucleotide sequences of the genomes of organisms have resulted in a better understanding of taxonomic relationships and of the changes responsible for the evolution of species from common ancestors. Now that we know the complete nucleotide sequences of the genomes of more than 2500 viruses, over 1400 archaea and eubacteria, and 41 eukaryotes (plus another 370 currently being assembled and 630 being sequenced)—over 122 billion nucleotide pairs of DNA in total—how can we use this information? By itself, the sequence of a DNA molecule is totally uninformative. In genetics, we use mutational dissection to determine the functions of various units of DNA. But how can we extract information from the vast amounts

of sequence information in the databanks? In this section, we will briefly discuss a few of the tools that are used to "mine" information from DNA sequences—those currently available and the vast number of new sequences that are accumulating daily.

We have been promised that the sequence of the human genome will lead to new approaches in the practice of medicine and that treatments will be tailored to the genotype of each individual. For example, physicians probably monitor women with a mutation in the *BRCA1* (*BR*east *CA*ncer 1) gene for breast cancer more vigilantly than other women (see Chapter 21). However, for the most part, medical treatment based on an individual's genotype, other than for classical hereditary disorders, is

still largely futuristic. So, what were the immediate contributions of the genomics era? The answer to this question seems clear—an enhanced knowledge of the evolutionary relationships between species and other taxonomic groups. The evidence of interbreeding between early humans and Neanderthals discussed at the beginning of this chapter is one example. By comparing the nucleotide sequences of the genomes of organisms—a subdiscipline called **comparative genomics**—scientists have documented many of the changes responsible for the divergence of species from common ancestors. Comparative genomics is a powerful new tool for studies of evolution. Evolutionary "trees"—phylogenies that show the relationships between species or other taxonomic groups—can be constructed from DNA sequences (see the section Molecular Phylogenies in Chapter 24). We will briefly examine some of the changes that have occurred in the genomes of the cereal grasses and selected mammals in the last two sections of this chapter. Chapter 24 presents a more detailed discussion of evolutionary processes.

BIOINFORMATICS

Knowledge of the nucleotide sequences of entire genomes has provided a wealth of information and a new challenge. How can we extract information from these sequences? This challenge has spawned a new scientific discipline called **bioinformatics (bio**logy + **informatics)**. You know that biology is the study of life. Informatics is the science of gathering, manipulating, storing, retrieving, and classifying recorded information. Bioinformatics involves doing all these things with biological information—most notably DNA and protein sequences. A comprehensive discussion of bioinformatics is beyond the scope of this textbook. Nevertheless, let's briefly examine some of the tools used to study the nucleotide sequences of DNAs and the amino acid sequences of proteins.

Suppose that you have just sequenced a DNA restriction fragment isolated from your favorite organism. How would you begin to analyze the function of this DNA molecule? First, you might ask whether anyone had previously sequenced this DNA or similar DNA molecules. To answer this question, you need a computer program that can search large DNA databases for similar sequences. Software programs designed to search sequence databases were first developed in the 1980s, and today there are programs designed to do almost anything that you can imagine. Some of the more popular programs were developed by the Genetics Computer Group (GCG) at the University of Wisconsin. We will use a couple of their programs to illustrate how nucleotide sequences can be studied.

Earlier in this chapter (see Focus on GenBank), we discussed the use of the Entrez web site (http://www.ncbi.nlm.nih.gov/entrez) to search for DNA sequences similar to a query sequence by using BLAST software. The other program that is used for rapidly searching huge databases is called FASTA. In the search that we performed in the Focus on GenBank exercise, we discovered that our sequence encoded part of the β 9-tubulin of *Arabidopsis* and that it was closely related to the β 8-tubulin gene of this plant. Indeed, the BLAST alignment tool showed us exactly how similar the query sequence was to each of the two genes. See Problem-Solving Skills: Using Bioinformatics to Investigate DNA Sequences to gain more experience using the tools discussed here.

Now, let's assume that our new sequence is not represented in GenBank, and let's ask how we might begin to analyze it. The most elementary step in trying to identify genes within nucleotide sequences is to look for open reading frames (ORFs)—sequences that can be translated into amino acid sequences without encountering any in-frame translation-termination codons. "Map" is a GCG program that can be used to translate double-stranded DNA in all six reading frames (three in each strand of DNA). Let's use the Map software to look for ORFs in a short segment of DNA from the green alga *Chlamydomonas reinhardti* (**■** Figure 15.18). The standard three-letter amino acid abbreviations are quite cumbersome when dealing with large protein databases; therefore, the single-letter code (see Figure 12.1) is used in bioinformatic analyses.

When the short segment of *Chlamydomonas* DNA is translated in all three reading frames, only reading frame 5 lacks termination codons (Figure 15.18). Thus, any ORF

PROBLEM-SOLVING SKILLS



Using Bioinformatics to Investigate DNA Sequences

THE PROBLEM

You have decided to follow the lead of Craig Venter and James Watson and have your genome sequenced. The first 100 nucleotides had the sequence acatttgctt ctgacacaac tgtgttcact agcaacctca aacagacacc atggtgcatc tgactcctga ggagaagtct gccgttactg ccctgtgggg. What is the function of this DNA? On what chromosome is it located? Is the sequence unique, or are there similar sequences present elsewhere in your genome? Is this sequence present in the genomes of other species?

FACTS AND CONCEPTS

- 1. The entire human genome—excluding some regions of highly repetitive DNA in heterochromatin—has been sequenced, and the sequences have been deposited in GenBank.
- 2. The sequences of the genomes of several other mammals including our closest living relative—the chimpanzee—are also available in GenBank.
- **3.** The NCBI web site (http://www.ncbi.nlm.nih.gov) contains bioinformatic tools that can be used to search GenBank for specific DNA sequences and/or for the proteins encoded by these sequences.
- **4.** The BLAST (*B*asic *L*ocal Alignment Search *T*ool) software allows you to search through specific genome sequences or all of the sequences in GenBank for similar sequences.

5. The NCBI web site can also be searched for publications that report the results of studies on specific DNA sequences and their products.

ANALYSIS AND SOLUTION

A BLAST search of the "Human genomic + transcript" sequences in the GenBank nucleotide database informs us that the 100-nucleotide sequence is part of the human β -globin gene (*HBB*) on chromosome 11. The 100-nucleotide sequence is identical to the sequence of one strand of the human *HBB* gene. The sequence is also very similar (93 percent identical) to the sequence of the human δ -globin gene located adjacent to the β -globin gene. A BLAST search of all NCBI Genomes (Chromosomes) shows that the sequence differs at only 1 nucleotide from the homologous sequence on chromosome 11 of the chimpanzee (*Pan troglodytes*) and at only 7 nucleotides from the homologous sequence on chromosome 14 of the rhesus monkey (*Macaca mulatta*). Clearly, the sequences of the β -globin genes are highly conserved in all primates. Indeed, a more detailed analysis would show that the β -globin genes of all vertebrates are highly conserved.

For further discussion visit the Student Companion site.

spanning this segment of DNA would have to be in reading frame 5. Of course, if we were searching for genes, we would usually be looking for ORFs that are much longer than the short DNA sequence in Figure 15.18. The software developed to search nucleotide sequences for genes can also screen for promoters, ribosome-binding sites, and other conserved sequences.

DNA sequence



Translation products

Reading Frame 1:	G V Y * I G * L F I A W C C R * M R D G
2:	ACIKLGNSSLRGVVDR * GMA
3:	R V L N W V T L H C V V L * I D E G W Q
	+++++++
4:	L R T N F Q T V R * Q T T N Y I S S P H
5:	A H I L N P L E E N R P T T S L H P I A
6:	P T Y * I P Y S K M A H H O L Y I L S P

FIGURE 15.18 Illustration of the use of the Wisconsin GCG "Map" program to identify ORFs by translating a 60-base-pair segment of DNA from *C. reinhardtii* in all six reading frames. Note the presence of translation–termination codons in all reading frames except number 5. Thus, it is the only reading frame that could be part of a larger ORF. Recall that translation is always $5' \rightarrow 3'$; thus, the amino-termini are on the left for translation products 1–3 and on the right for products 4–6. The translation–termination sites are designated by asterisks. They correspond to the termination triplets shown underlined in red, green, or blue, depending on the reading frame.

The presence of introns in eukaryotic genes makes gene searches more difficult than in prokaryotes. Eukaryotic gene searches involve scanning for intron splice sites in addition to ORFs and other regulatory sequences. Today, the gene-search programs are tailored to individual species by factoring in features of their genomes—for instance, base composition, codon usage, and preferences for certain sequences in regulatory elements. The only way to be absolutely certain that a predicted gene is real and that its introns have been identified correctly is to isolate and sequence a full-length cDNA clone and compare its sequence with the sequence of a genomic clone. There are currently dozens of gene-prediction programs; they go by names such as GRAIL (Gene Recognition and Analysis Internet Link), GeneMark (one of the first used to search for genes in *E. coli*), GeneScan, and GeneFinder.

One common feature of eukaryotic genomes is the presence of gene families sets of genes that encode very similar proteins (often called isoforms). These proteins usually have redundant or overlapping functions. In order to compare all of the genes in a gene family, bioinformaticians have developed programs that will align multiple nucleotide or amino acid sequences, allowing a direct visual comparison of all members of a gene or protein family. Multiple alignments are especially useful in identifying conserved DNA sequences that are important regulatory elements, such as protein-binding sites. They are also useful in identifying important, and thus conserved, functional domains within proteins. **■** Figure 15.19 shows the alignment of the amino-terminal regions of the eight β -tubulins of *Arabidopsis thaliana*. Note that within the 60 or 61 amino acids shown there are five regions of four or more amino acids that are conserved in all eight proteins.

Genes with very similar nucleotide sequences, such as the nine genes encoding the eight β -tubulins shown in Figure 15.19, often—but not always—owe their similarity to having evolved from a common ancestral gene. Such genes are said to be **homologous**; note that "similar" and "homologous" are not synonyms. The nine β -tubulin genes of *Arabidopsis* are homologous; that is, they are **homologues**. Indeed,

Tubulin	Amin acid		Amino acid
β6:	1	MREILHIQGGQCGNQIGsKFWEVVCdEHGIDpTGRYvG nsDLQLERVNVYYNEASCGRyV	60
β8:	1	eq:meilhiggggggggggggggggggggggggggggggggggg	60
β2 and β3:	1	MREILHIQGGQCGNQIGAKFWEVVCAEHGIDpTGRYtGD SDLQLERiNVYYNEASCGRFV	60
β7:	1	MREILHIQGGQCGNQIGSKFWEVVnlEHGIDqTGRYvGD SeLQLERvNVYYNEASCGRYV	60
β5:	1	MREILHIQGGQCGNQIGSKFWEVICDEHGIDsTGRYsGDtADLQLERINVYYNEASGGRYV	61
β1:	1	MREILHvQGGQCGNQIGSKFWEVICDEHGvDpTGRYnGDSADLQLERINVYYNEASGGRYV	61
β4:	1	MREILHIQGGQCGNQIGAKFWEVICDEHGIDbTGQYvGDS pLQLERIdVYFNEASGGKYV	60
β9:	1	MREILHIQGGQCGNQIGAKFWEVICgEHGIDqTGQscGD tDLQLERINVYFNEASGGKYV	60
0			
Conserve	a seqs	. 1 2 3 4 5	

FIGURE 15.19 Illustration of a multi-sequence protein alignment using the single-letter amino acid code (see Figure 12.1). The alignment—generated with the GCG "PileUp" program—compares the amino-terminal regions of the eight β -tubulins of *Arabidopsis*, which are encoded by nine genes. Genes *TUB2* (*TU*bulin Beta number 2) and *TUB3* encode identical products (the β 2- and β 3-tubulins): these two genes differ only at positions corresponding to the degenerate third bases of codons. The most similar sequences are grouped together. The single-letter codes for amino acids are in uppercase when adjacent sequences are identical and are in lowercase when adjacent sequences are different.

two of the genes were produced by a recent (on the evolutionary time scale) geneduplication event; they differ at only 30 base pairs (corresponding to degenerate bases in codons) and encode the same polypeptide. These genes are also called **paralogous genes** or **paralogues**—homologous genes within a species. Homologous genes present in different species are called **orthologous genes** or **orthologues**. The tubulin genes of *Arabidopsis* and *Chlamydomonas* are orthologues.

PROKARYOTIC GENOMES

TABLE 15.1

Haemophilus influenzae was the first cellular organism to have its entire genome sequenced; the sequence was published in 1995. By February 2011, the complete sequences of the genomes of 1412 archaea and bacteria were available in the public databases, and sequencing projects were underway for another 3695 species. The sequenced genomes range in size from 490,885 bp for *Nanoarchaeum equitans*—an obligate symbiont; to 580,076 bp for *Mycoplasma genitalium*—thought to have the smallest genome of any non-symbiotic bacterium; to 4,403,837 bp for *Mycobacterium tuberculosis*—the cause of more human deaths than any other infectious bacterium; to 4,639,675 bp for *Bradyrbizobium japonicum*—a soil bacterium capable of colonizing plant root nodules. The size and predicted gene content of a few prokaryotic genomes are shown in **Table 15.1**; for a complete list, see http://www.ncbi.nlm.nih.gov/genomes/lproks.cgi.

One of the striking features of bacterial genomes is their variability in size within a species. Studies on *E. coli, Prochlorococcus marinus*, and *Streptococcus coelicolor* have documented variations in genome size of up to a million nucleotide pairs between different strains of the same species.

Of the bacterial genomes sequenced to date, the sequence of the *E. coli* strain K12 genome created the most excitement among biologists. *E. coli* is the most studied and best understood cellular organism on our planet. Geneticists, biochemists, and molecular biologists have utilized *E. coli* as the preferred model organism for decades. Most of what is known about bacterial genetics was learned from research on *E. coli*. Thus, the 1997 publication of the complete sequence of the *E. coli* genome with its 4467 putative protein-coding genes was an important milestone in genetics. Known and putative genes specifying proteins and stable RNAs make up 87.8 percent and 0.8 percent of the genome, respectively, and noncoding repetitive elements account for 0.7 percent of the genome. Thus, 10.7 percent of the genome must involve regulatory sequences and sequences with unknown functions.

Species	Genome Size in Nucleotide Pairs	Predicted Numbe of Genes
Archaea		
Nanoarchaeum equitans	490,885	582
Sulfolobus solfataricus	2,992,245	3,033
Eubacteria		
Bradyrhizobium japonicum	9,105,828	8,373
<i>Escherichia coli</i> , strain K12 MG1655	4,639,675	4,467
<i>Escherichia coli</i> , strain 0157 EDL933	5,528,970	5,463
<i>Legionella pneumophila</i> , strain Paris	3,503,610	3,136
Mycobacterium tuberculosis, strain CDC	4,403,837	4,293
Mycobacterium genitalium	580,076	525
Yersinia pestis, strain KIM	4,600,755	4,240

The genomes of *M. tuberculosis* (tuberculosis), *Legionella pneumophila* (Legionnaire's disease), *Yersinia pestis* (bubonic plague), and other infectious bacteria are also of great interest because of the pathogenicity of these organisms and the hope that a complete understanding of their metabolism will suggest ways to prevent these often fatal diseases.

The genome of *M. genitalium* is of special interest because it may approximate the "minimal gene set" for a cellular organism—the smallest set of genes that will allow a cell to reproduce. The genome of *M. genitalium* contains only 525 predicted genes, and engineered mutations have shown that at least 100 of these genes are not essential to survival. By comparing the 525 genes of *M. genitalium* with those of other bacteria, and using information about the functions of these genes in other bacteria, researchers have estimated that the minimal number of genes required for the reproduction of a cellular organism is somewhere between 265 and 350.

A LIVING BACTERIUM WITH A CHEMICALLY SYNTHESIZED GENOME

After sequencing the small genome of *Mycoplasma genitalium* with its 525 predicted genes, J. Craig Venter and colleagues became interested in the "minimal gene set"— the minimum number of genes that would support life—of a single-celled organism. To prepare for testing the hypothesis that the "minimal gene set" consisted of about 300 genes, researchers at the J. Craig Venter Institute in Maryland decided to construct a totally synthetic bacterial genome. Because of the slow growth rate and parasitic lifestyle of *M. genitalium*, they decided to synthesize the genome of its faster growing relative *M. mycoides*.

The starting points for their work were the published nucleotide sequences of the genomes of two strains of *M. mycoides*. They began by synthesizing oligonucleotides, which they spliced together into 1080-bp cassettes with 80-bp overlaps. Venter and associates verified the accuracy of their synthetic processes by sequencing all of the cassettes. The cassettes were designed with *Not*I restriction sites at their termini. The key strategy in assembling the cassettes into a complete genome was to transform yeast cells and select for the products of homologous recombination *in vivo*. They first assembled 1078 1080-bp cassettes, which were spliced together to produce 109 10,080-bp assemblies, which in turn were joined together to produce eleven 100,000-bp mega-assemblies. The 100-kb genome segments were subsequently joined to produce the complete 1,077,947-bp genome. The synthesis and assembly process is illustrated in **Figure 15.20**.

The research team inserted four marker DNAs to use in distinguishing their synthetic genome from wild-type *M. mycoides* genomes and deleted one 4-kb unessential region. The marker DNAs included the *E. coli lacZ* gene, which permitted the identification of cells carrying the synthetic genome as blue colonies on X-gal plates (see Figure 14.4), and a tetracycline-resistance gene, which made it possible to select cells carrying the synthetic genome after transplantation into tetracycline-sensitive cells.

They used PCR carried out with primer pairs that spanned each of the eleven 100-kb genome segments to screen for the complete synthetic genome, comparing the results with those obtained using the genome of a wild-type strain. Once they had the complete genome assembled, they had to determine whether it was functional. They did this by transplanting the synthetic genome to cells of a closely related species *M. capricolum*. The restriction system of recipient cells had previously been inactivated by an insertion mutation so that the foreign DNA would not be degraded (see Figure 14.1). After transplantation of the intact synthetic genome to *M. capricolum* cells, the transiently "binucleate cells" were plated on X-gal medium containing tetracycline. The tetracyline selected for cells carrying the synthetic *M. mycoides* genome, and X-gal allowed the desired bacteria—blue colonies—to be distinguished from the remaining recipient cells—white colonies.

Now that Venter and coworkers have developed the technology required to synthesize complete bacterial genomes and transfer them from yeast cells to bacterial

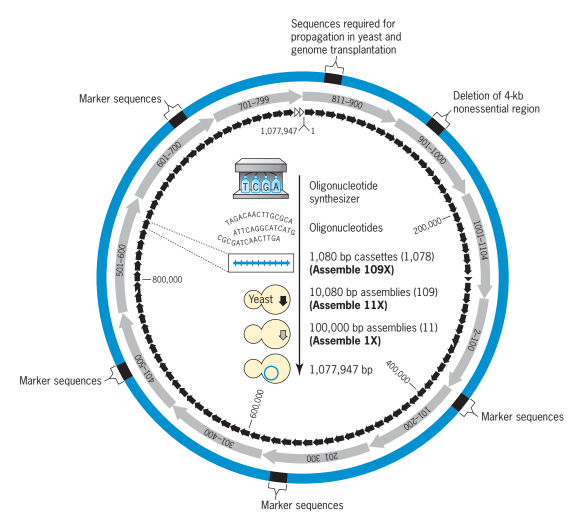


FIGURE 15.20 Strategy used to create a completely synthetic bacterial genome. The construction of the synthetic bacterial genome started with the synthesis of oligonucleotide sequences corresponding to established sequences in the genome of wild-type strains of *M. mycoides*. These sequences were then spliced together in the series of assemblies shown to produce a complete 1,077,947 bp *M. mycoides* genome. The genome was shown to be functional by transplanting it into cells of a closely related species *M. capricolum*.

cells, they can pursue the question of the "minimal gene set" by deleting genes and testing for viability. They can also attempt to produce bacteria with synthetic genomes that synthesize valuable products or that degrade environmental pollutants, and so on. Although the procedure is still very costly, technical improvements should lead to more efficient and less expensive synthetic genomes in the future.

THE GENOMES OF CHLOROPLASTS AND MITOCHONDRIA

Eukaryotic cells contain membrane-bounded compartments or organelles that play important roles in energy metabolism. Mitochondria convert organic molecules into energy by aerobic, or oxidative, metabolism, and chloroplasts use energy from sunlight to synthesize organic material from water and carbon dioxide—a process called photosynthesis. Both of these organelles almost certainly developed from prokaryotic cells that established symbiotic—mutually beneficial—relationships with host cells. These prokaryotes brought their genomes with them, along with their ability to carry out aerobic metabolism and photosynthesis. As a result, mitochondria and chloroplasts contain their own genomes. In both cases, however, these organelles utilize some imported proteins encoded by nuclear genes to supplement gene products specified by organellar genes. Today, eukaryotic cells have become highly dependent on these former prokaryotic invaders. Plants could not perform photosynthesis without chloroplasts, and neither plants nor animals could carry out aerobic respiration without mitochondria.

Mitochondrial Genomes

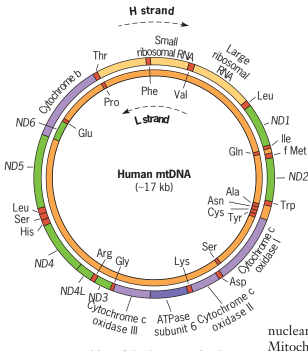
Mitochondrial genetic systems consist of DNA and the molecular machinery needed to replicate and express the genes contained in this DNA. This machinery includes the macromolecules needed for transcription and translation. Mitochondria even possess their own ribosomes. Many of these macromolecules are encoded by mitochondrial genes, but some are encoded by nuclear genes and therefore are imported from the cytosol.

Mitochondrial DNA, or **mtDNA** as it is usually abbreviated, was discovered in the 1960s, initially through electron micrographs that revealed DNA-like fibers within the mitochondria. Later, these fibers were extracted and characterized by physical and chemical procedures. The advent of recombinant DNA techniques made it possible to analyze mtDNA in great detail. In fact, the complete nucleotide sequences of mtDNA molecules from many different species have now been determined. Representative mtDNAs are shown in **Table 15.2**. For a complete list of the mitochondrial genomes sequenced to date, see http://www.ncbi.nlm.nih.gov/genomes/genlist.cgi?taxid= 2759&type=4&name=Eukaryota Organelles. Mitochondrial DNA molecules vary enormously in size, from about 6 kb in the malaria-causing parasite *Plasmodium* to 2500 kb in certain flowering plants. Each mitochondrion appears to contain several copies of the DNA, and because each cell usually has many mitochondria, the number of mtDNA molecules per cell can be very large. A vertebrate oocyte, for example, may contain as many as 10⁸ copies of the mtDNA. Somatic cells, however, have fewer copies, perhaps less than 1000.

Most mtDNA molecules are circular, but in some species, such as the alga *Chlamydomonas reinhardtii* and the ciliate *Paramecium aurelia*, they are linear. The circular mtDNA molecules, which have been studied the most thoroughly, appear to be organized in many different ways. In the vertebrates 37 distinct genes are packed into a 16- to 17-kb circle, leaving little or no space between genes. In some of the flowering plants, an unknown number of genes are dispersed over a very large circular DNA molecule hundreds or thousands of kilobases in size.

TABLE 15.2

Size and Gene Content of Selected Mitochondrial and Chloroplast Genomes						
Species	Common Name	Genome Size in Nucleotide Pairs	Predicted Number of Genes			
Mitochondrial Genomes						
Arabidopsis thaliana	mouse ear cress	366,924	57			
Caenorhabditis elegans	roundworm	13,794	12			
Drosophila melanogaster	fruit fly	19,517	37			
Homo sapiens	human	16,571	37			
<i>Oryza sativa</i> Indica	rice	491,515	96			
Saccharomyces cerevisae	baker's yeast	85,779	43			
Zea mays subsp. mays	corn	569,630	218			
Chloroplast Genomes						
Arabidopsis thaliana	mouse ear cress	154,478	129			
Chlamydomonas reinhardtii	green alga	203,828	109			
Marchantia polymorpha	liverwort	121,024	134			
<i>Oryza sativa</i> Japonica	rice	134,525	159			
Zea mays subsp. mays	corn	140,384	158			
Data are from the NCBI web site (http://www.ncbi.nim.nih.gov/Genomes).						



■ FIGURE 15.21 Map of the human mitochondrial genome. *ND1-6* are genes encoding subunits of the enzyme NADH reductase; the tRNA genes in the mtDNA are indicated by abbreviations for the amino acids. Arrows show the direction of transcription. Genes on the inner circle are transcribed from the L (light) strand of the DNA, whereas genes on the outer circle are transcribed from the H (heavy) strand of the DNA.

Animal mtDNA is small and compact. In humans, for example, the mtDNA is 16,571 base pairs long and contains 37 genes (**Figure 15.21**), including two that encode ribosomal RNAs, 22 that encode transfer RNAs, and 13 that encode polypeptides involved in oxidative phosphorylation, the process that mitochondria use to recruit energy. In mice, cattle, and frogs, the mtDNA is similar to that of human beings-an indication of a basic conservation of structure within the vertebrate subphylum. Invertebrate mtDNAs are about the same size as vertebrate mtDNAs, but their genetic organization is somewhat different. In fungi, the mtDNA is considerably larger than it is in animals. Yeast, for example, possesses circular mtDNA molecules 78 kb long. Plant mtDNA is much larger than the mtDNA of other organisms (Table 15.2). It is also more variable in structure. One of the first plant mtDNAs to be sequenced was from the liverwort, Marchantia polymorpha. The mtDNA from this primitive, nonvascular plant is a 186-kb circular molecule with 94 substantial open reading frames (ORFs). In vascular plants, the mtDNA is larger than it is in *Marchantia*; for example, it is a 570-kb circular molecule in maize. Higher plant mtDNA molecules contain many noncoding sequences, including some that are duplicated.

Most—perhaps all—mitochondrial gene products function solely within the mitochondrion. However, they do not function alone. Many nuclear gene products are imported to augment or facilitate their function. Mitochondrial ribosomes, for example, are constructed with ribosomal RNA transcribed from mitochondrial genes and with ribosomal proteins encoded by nuclear genes. The ribosomal proteins are synthesized in the cytosol and imported into the mitochondria for assembly into ribosomes.

Many of the polypeptides needed for aerobic metabolism are also synthesized in the cytosol. These include subunits of several proteins involved in oxidative phosphorylation—for example, the ATPase that is responsible for binding the energy of aerobic metabolism into ATP. However, because some of the subunits of this protein are synthesized in the mitochondria, the complete protein is actually a mixture of nuclear and mitochondrial gene products. This dual composition suggests that the nuclear and mitochondrial genetic systems are coordinated in some way so that equivalent amounts of their products are made. To expand your comprehension of the structures of mitochondrial genomes, answer the questions posed in Solve It: What Do We Know about the Mitochondrial Genome of the Extinct Woolly Mammoth?

Chloroplast Genomes

Chloroplasts are specialized forms of a general class of plant organelles called **plastids**. Botanists distinguish among several kinds of plastids, including chromoplasts (plastids containing pigments), amyloplasts (plastids containing starch), and elaioplasts (plastids containing oil or lipid). All three types seem to develop from small membranebounded organelles called proplastids, and, within a particular plant species, all seem to contain the same DNA. This DNA is generally referred to as **chloroplast DNA**, abbreviated simply as **cpDNA**.

In higher plants, cpDNAs typically range from 120 to 160 kb in size, and in algae, from 85 to 292 kb (Table 15.2). In a few species of green algae in the genus *Acetabularia*, the cpDNA is much larger, about 2000 kb. The plant cpDNAs that have been sequenced are circular molecules.

The number of cpDNA molecules in a cell depends on two factors: the number of chloroplasts and the number of cpDNA molecules within each chloroplast. For example, in the unicellular alga *Chlamydomonas reinhardtii*, there is only one chloroplast per cell, and it contains about 100 copies of the cpDNA. In *Euglena gracilis*, another unicellular organism, there are about 15 chloroplasts per cell, and each contains about 40 copies of the cpDNA.

All cpDNA molecules carry basically the same set of genes, but in different species these genes are arranged in different ways. The basic gene set includes genes for ribosomal RNAs, transfer RNAs, some ribosomal proteins, various polypeptide

■ FIGURE 15.22 Genetic organization of the chloroplast genome in the liverwort *Marchantia polymorpha*. Symbols: *rpo*, RNA polymerase; *rps*, ribosomal proteins of small subunit; *rpl* and *secX*, ribosomal proteins of large subunit; *4.55*, *55*, *16S*, *23S*, rRNAs of the indicated size; *rbs*, ribulose bisphosphate carboxylase; *psa*, photosystem I; *psb*, photosystem II; *pet*, cytochrome b/f complex; *atp*, ATP synthesis; *infA*, initiation factor A; *frx*, iron-sulfur proteins; *ndh*, putative NADH reductase; *mpb*, chloroplast permease; tRNA genes are indicated by abbreviations for the amino acids.

components of the photosystems involved in capturing solar energy, the catalytically active subunit of the enzyme ribulose 1,5-bisphosphate carboxylase, and four subunits of a chloroplast-specific RNA polymerase. More than 200 cpDNA molecules have been sequenced in their entirety. Table 15.2 lists a few examples; the complete list is available at http://www.ncbi.nlm.nih.gov/genomes/GenomesGroup. cgi?taxid=2759&opt=plastid.

Two of the first cpDNAs sequenced were from the liverwort, *Marchantia polymorpha* (**■** Figure 15.22), and from the tobacco plant, *Nicotiana tabacum*. The tobacco cpDNA is larger (155,844 bp) and contains about 150 genes. Most cpDNAs have a pair of large inverted repeats that contain the genes for the ribosomal RNAs.

As mentioned earlier, the development of functional chloroplasts depends on the expression of both nuclear and chloroplast genes. The nuclear genes are transcribed in the nucleus and translated in the cytosol. The products of nuclear genes that function in the chloroplast must be imported from the cytosol. Once imported, these proteins must act in concert with cpDNA-encoded proteins. Functional chloroplasts thus depend on the coordinated activities of both nuclear and chloroplast gene products.

EUKARYOTIC GENOMES

Baker's yeast, *Saccharomyces cerevisiae*, was the first eukaryotic organism to have its entire genome sequenced. The complete 12,068-kb sequence of the *S. cerevisiae* genome was assembled in 1996 through an international collaboration of about 600 scientists working in Europe, North America, and Japan. The yeast genome contains 5885 potential protein-coding genes, about 140 genes specifying ribosomal RNAs, 40 genes for small nuclear RNA molecules, and over 200 tRNA genes. Researchers have systematically generated deletions of essentially all (5916, or 96.5 percent) of the

Solve It!

What Do We Know about the Mitochondrial Genome of the Extinct Woolly Mammoth?

The woolly mammoth, Mammuthus primigenius, disappeared from most of its range about 10,000 years ago, with a small population surviving on the Wrangel Island in the Arctic Ocean until about 4700 years ago. Given that the species has been extinct for almost 5000 years, how can scientists have sequenced major portions of its nuclear genome and its entire mitochondrial genome? How large is the woolly mammoth's mitochondrial genome (mtDNA)? How many proteincoding genes does it contain? How many noncoding RNA molecules does it specify? Is the sequence of the woolly mammoth's mtDNA more similar to that of human mtDNA or to that of elephant mtDNA? (1) If you compare the mtDNAs of (1) the woolly mammoth and the elephant and (2) the Neanderthals and humans, which sequences are the more closely related?

To see the solution to this problem, visit the Student Companion site.

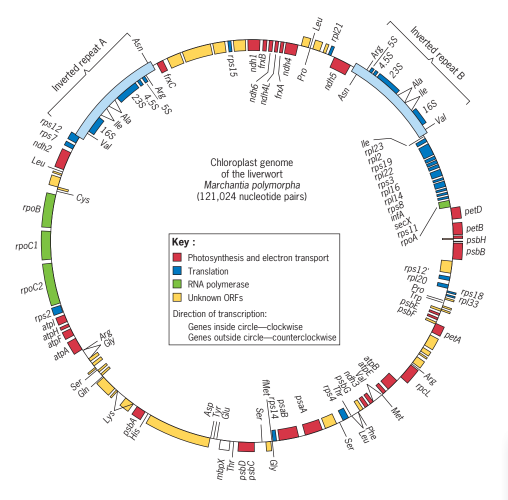


TABLE 15.3

Size and Predicted Gene Content of Selected Eukaryotic Genomes

Size and Frederica Serie Solite	int of Detected Edital Jotte Dellon			
Species	Common Name	Genome Size in Nucleotide Pairs	Predicted Number of Genes*	Gene Density (bp/gene) ⁺⁺
Protist				
Plasmodium falciparum	malaria protozoan	22,820,308	5,361	4,300
Fungus				
Saccharomyces cerevisae	baker's yeast	12,057,909	6,268	1,900
Nematode				
Caenorhabditis elegans	roundworm	100,291,841	20,516	4,900
Insect	f	121 000 000	10 700	0 500
Drosophila melanogaster Plant	fruit fly	131,000,899	13,792	9,500
Arabidopsis thaliana	mouse ear cress	119,186,496	28,152	4,200
Vertebrates				
Danio rerio	zebra fish	1,571,018,465	23,524	66,800
Homo sapiens	human	2,851,330,913	22,287	127,900
Mus musculus	mouse	2,932,368,526	25,396	115,500
Pan troglodytes	chimpanzee	2,928,563,828	21,098	139,000

Data are from the NCBI web site (http://www.ncbi.nim.nih.gov/Genomes), the Ensembl web site (http://www.ensembl.org), or the CBS Genome Atlas Database (http://www.cbs.dtu.dk/services/GenomeAtlas).

*Gene numbers are Ensembl predictions (minus pseudogenes when data are available).

⁺⁺Values are rounded to the nearest 100 bp.

predicted 6268 genes in the yeast genome. Of the genes tested, 1105 (18.7 percent) were found to be essential for growth in rich glucose medium—that is, deletions in these genes were lethal. Some deletions did not cause lethality because the yeast genome contains many duplicated genes. Both copies of these genes must be deleted to have a lethal effect. Many other yeast genes can be deleted without killing the organism. However, knockouts of these genes are often associated with changes in morphology or with impaired growth.

The sequences of the genomes of other eukaryotic model systems soon followed. The sequence of 99 percent of the genome of the worm *Caenorhabditis elegans* was published in 1998, and nearly complete sequences of the genomes of the fruit fly *Drosophila melanogaster* and the model plant *Arabidopsis thaliana* followed in 2000. The release of two first drafts of the sequence of the human genome in 2001 probably received more coverage by the international news media than any other event in the history of biology. As mentioned earlier, a nearly complete sequence of the human genome was released in 2004.

What have we learned from all these sequences? In contrast to the genomes of archaea and eubacteria, gene density varies widely among different eukaryotic species, ranging from one gene per 1900 bp in baker's yeast to one gene per 127,900 bp (145,000 bp if the unsequenced heterochromatin is included) in humans. Genome size and gene content are shown for the genomes of selected eukaryotes in **Table 15.3**; for a complete list of sequenced genomes and ongoing sequencing projects, see http://www.ncbi.nlm.nih.gov/genomes/leuks.cgi. The genomes of the single-celled eukaryotes are like yeast, with one gene for every 1000 to 2000 bp. Gene density decreases to one gene per 4000 to 5000 bp for *Arabidopsis* and *C. elegans*, to one gene per 9500 bp in *D. melanogaster*; and is the lowest in mammals at one gene for every 115,000 to 129,000 bp. The observed decrease in gene density with increased developmental complexity raises questions about the functions of the noncoding DNA. As mentioned earlier (see Figure 15.12), an international *ENC*yclopedia *Of DNA Elements* (ENCODE) Project Consortium is investigating the functions of noncoding DNAs. Test your ability to

use some of the tools on the web to analyze eukaryotic DNA sequences by working through Solve It: What Can You Learn about DNA Sequences Using Bioinformatics?

One reason for the lower gene density in the larger eukaryotic genomes is that these genomes contain considerable amounts of repetitive DNA (see Chapter 9). Baker's yeast contains very little repetitive DNA, although about 30 percent of its genes are duplicated. By contrast, the genomes of multicellular eukaryotes contain lots of repetitive DNA, and the amount of this material is, in most cases, directly related to genome size. For example, only about 10 percent of the small genome of *C. elegans* consists of moderately repetitive DNA, whereas about 45 percent of the large genomes of mammals is composed of moderately repeated DNA sequences. Most of these moderately repetitive sequences are derived from transposable genetic elements (see Chapter 17).

Highly repetitive DNA is also more abundant in the larger genomes; however, there is no direct correlation between the amount of highly repetitive DNA and genome size. Indeed, closely related species sometimes differ significantly in the amount of highly repetitive DNA. For example, 18 percent of the genome of *D. melanogaster* consists of highly repetitive DNA, whereas 45 percent of the DNA in *D. virilis* is highly repetitive. Much of the highly repetitive DNA in most species, including humans, is present in the regions of chromosomes that flank the centromeres (centromeric heterochromatin) and in the telomeres. This DNA is difficult to sequence. In fact, most of the unsequenced DNA in the human genome—472 million base pairs—consists of highly repetitive sequences, and 24 of the gaps in the human genome sequence correspond to blocks of centromeric heterochromatin in the 24 chromosomes.

Introns are a significant component of eukaryotic DNA, and they are more prevalent and longer in the larger eukaryotic genomes. Intergenic regions are also longer in the larger eukaryotic genomes. In contrast, the number of distinct protein domains functional regions of proteins—encoded by genes does not seem to vary much with genome size. The predicted numbers of protein domains encoded by the *A. thaliana*, *D. melanogaster*, and human genomes are 1012, 1035, and 1262, respectively. However, humans and other vertebrates make greater use of alternate pathways of transcript splicing (see Chapter 19) to shuffle these domains into more combinations, increasing protein diversity.

GENOME EVOLUTION IN THE CEREAL GRASSES

The cereal crops provide much of the food for humans and their domestic livestock, which provide additional human foodstuff. Therefore, enhanced cereal productivity is an important component of the effort to feed the constantly expanding human population on our planet. Increased knowledge of the genomes of these agronomic species is key to achieving this objective. Recent comparative analyses of the genomes of several cereal species indicate that much of the information obtained by mapping and sequencing the smallest of these genomes—the 400-mb genome of rice—will be directly applicable to other cereal grasses because of the conservation of genome structure in these species.

When Graham Moore and colleagues compared the high-density genetic maps of the chromosomes of several cereal grasses, they discovered that despite large differences in genome size and chromosome number, the linkage relationships of blocks of unique DNA sequences and known genes were remarkably conserved. In contrast, the quantities and locations of repetitive DNA sequences were highly variable.

The striking conservation of genome structure in the cereal grasses is illustrated most clearly by drawing the rice genome as a circular array and aligning the conserved blocks of genes in the other species with the rice genome (**Figure 15.23**). This circular display of the cereal genomes does not imply any circularity of ancestral chromosomes; it simply permits maximal alignment of homologous blocks of genes. The alignment also emphasizes the presence of duplicate copies of each block of genes in the maize genome, indicating that maize has evolved from a tetraploid ancestor. Interestingly, one set of genes is present largely in the small chromosomes of maize, and the second set is present primarily in the large chromosomes.

Solve It!

What Can You Learn about DNA Sequences Using Bioinformatics?

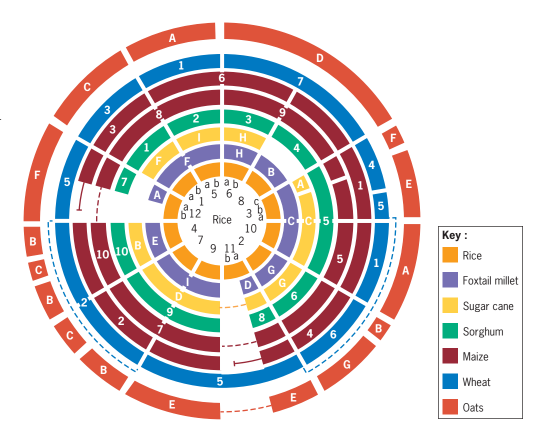
Translate the following two DNA sequences in all six reading frames using the software available at http://www.expasy.org/tools/ dna.html.

Sequence 1	: atggtgctgt	ctcctgccga
caagaccaac	gtcaaggccg	cctggggtaa
ggtcggcgcg	cacgctggcg	agtatggtgc
ggaggccctg	gagaggatgt	tcctgtcctt
ccccaccacc		

Sequence 2: aatatgctta ccaagctgt gattccaaat attacgtaaa tacacttgca aaggaggatg tttttagtag caatttgtac tgatggtatg gggccaagag atatatctta gagggagggc

Which sequence is likely to be part of the coding sequence of a gene? Which sequence is clearly not part of a coding sequence? Why? Next perform a BLAST search at PubMed's Entrez web site (http:// www.ncbi.nlm.nih.gov/entrez) using the potential coding sequence as your query. Is the sequence present in GenBank? Is it a coding sequence? in what organism? in what gene?

To see the solution to this problem, visit the Student Companion site. ■ FIGURE 15.23 Simplified comparative map of the genomes of seven cereal grasses. Chromosomes and segments of chromosomes (denoted by capital letters) of the various cereal grasses are aligned with the chromosomes of rice, the grass species with the smallest genome (center). The maize genome has two similar copies of each block of genes and thus occupies two rings of the circle. The outer dashed lines connect adjacent segments of chromosomes. Similar segments of chromosomes in the oats genome are not connected by dashed lines for the sake of simplicity.



The conserved structures of the cereal grass genomes should assist plant breeders in their attempts to produce varieties with increased yield, pest resistance, drought tolerance, and other desired traits. Because of the conserved genome structure, information obtained from sequencing the relatively small rice genome should be more easily applied to breeding and genetic engineering projects on the other cereal crop species.

GENOME EVOLUTION IN MAMMALS

Mammalian genomes exhibit conservation of chromosome structure similar to that observed in the cereal grasses. Although genetic mapping is being done in over 200 mammalian species, high-density maps are currently available for only human, mouse, dog, rat, and a few agriculturally important farm animals such as swine and cattle. These detailed chromosome maps can be used to demonstrate the conserved linkage relationships of genes in species where such maps are available. For other species, a procedure called **chromosome painting** has been used for comparative genome analyses. Chromosome painting is a variation of fluorescent *in situ* hybridization (FISH; see Appendix C: *In Situ* Hybridization) in which chromosomes are "painted" different colors by using DNA hybridization probes labeled with fluorescent dyes that emit light of different wavelengths.

In comparative genomic studies, DNA sequences from one species are used to paint the chromosomes of a related species. Such cross-species chromosome painting experiments are called Zoo-FISH experiments. They are usually done under conditions of reduced hybridization stringency, which allows the detection of cross-hybridization between the partially complementary strands of homologous genes.

Some of the most interesting chromosome painting studies used the sequences in a chromosome-specific genomic library (Chapter 14) from one species to paint the chromosomes of related species. Indeed, different fluorescent dyes can be used to label sequences from two or more different chromosomes of one species, and these fluorescence-tagged sequences can be used to "paint" the chromosomes of related species. Chromosome-specific libraries are available for all 24 human chromosomes, and sequences from these

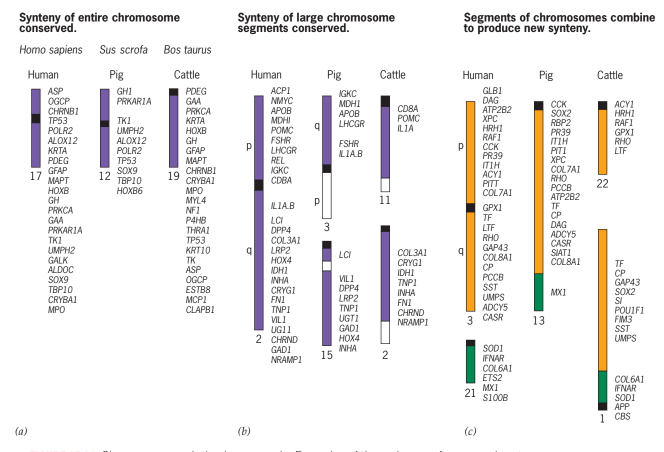


FIGURE 15.24 Chromosome evolution in mammals. Examples of three classes of conserved synteny (conserved blocks of linked genes) are illustrated. (a) Human chromosome 17 is an example of the conservation of entire chromosomes. Note that even with conserved synteny inversions occur, changing the order of some genes. (b) Human chromosome 2 provides an example of the conservation of large chromosomal segments.
 (c) Human chromosomes 3 and 21 illustrate the formation of new synteny by chromosome fusion. The chromosomal locations of a few genes are shown to the right of the chromosomes. Fewer genes have been mapped in pigs than in cattle or humans. Species-specific chromosome designators are given below the chromosomes.

libraries have been used to "paint" the chromosomes of several related species, including most of the primates and a few more distantly related mammals.

Upon reviewing comparative linkage and chromosome painting data, Bhanu Chowdhary and colleagues concluded that the evolution of mammalian chromosomes involved three classes of conserved synteny. (Synteny is the presence of genes on the same chromosome.) The three classes are (1) conservation of entire chromosomes, (2) conservation of large segments of chromosomes, and (3) the joining of segments of different chromosomes to produce new synteny. Each type of chromosome evolution is illustrated in **Figure 15.24**.

The synteny of genes on human chromosome 17 is conserved on chromosome 12 of the pig and chromosome 19 of cattle (Figure 15.24*a*). Similar patterns of conservation of human chromosomes 13 and 20 have been observed in these species.

Human chromosome 2 provides an example of the second class of conserved synteny, the conservation of large segments of chromosomes (Figure 15.24*b*). Major portions of the long and short arms of human chromosome 2 are conserved on two chromosomes of the pig and cattle (also in the horse and cat—not shown). Human chromosomes 4, 5, 6, 9, and 11 also have large chromosome segments conserved in the other mammals studied.

The third pattern of conserved synteny—the joining of segments of chromosomes to produce new synteny—has occurred with human chromosomes 3 and 21 (Figure 15.24c). Chromosome 13 of the pig appears to contain the genes of both human chromosomes arranged as if they had simply fused together. In cattle, one block of genes present on human chromosome 3 is present on a separate chromosome (22), whereas most of the rest of the genes from human chromosome 3 are on a chromosome (1) with the genes from human chromosome 21. Other examples of this pattern of conserved synteny involve human chromosomes 12 and 22, 14 and 15, and 16 and 19.

Despite the extensive conservation of blocks of genes illustrated by the examples discussed above, more detailed comparisons of chromosome structure in mammals show that numerous chromosome rearrangements have occurred during the evolution of even closely related species. A comparison of human chromosomes with the chromosomes of the white-cheeked gibbon, *Hylobates concolor*; revealed that at least 21 translocations have occurred during the evolution of these two species from their common ancestor. Inversions and other intrachromosomal rearrangements are especially prevalent in the genomes of closely related species.

KEY POINTS • Comparative genomics—comparing the nucleotide sequences of genomes—has provided new information about the relationships between various taxonomic groups.

- Bioinformatics is the science of storing, comparing, and extracting information from biological systems, especially DNA and protein sequences.
- The nucleotide sequences of prokaryotic genomes have provided evidence of the diversity and plasticity of species and of different strains of the same species.
- Mitochondrial genomes are usually circular and range in size from 6 kb to 2500 kb, whereas chloroplast genomes also are usually circular and are typically 120 to 292 kb in size with more than 100 genes.
- As eukaryotic organisms have increased in complexity, the proportion of their genomes that encodes proteins has decreased.
- Comparative genomics has revealed a remarkable conservation of synteny in related eukaryotic species, such as mammals and the cereal grasses.

Basic Exercises

Illustrate Basic Genetic Analysis

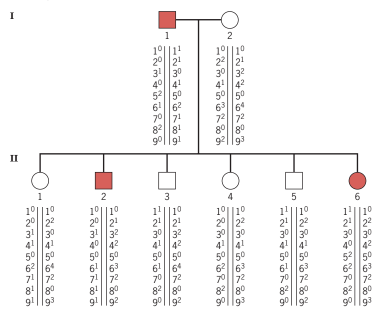
- 1. What is a genetic map?
- **Answer:** A genetic map shows the positions of genes and other markers such as restriction fragment-length polymorphisms (RFLPs) on a chromosome based on recombination frequencies.
- 2. What is a cytological map?
- **Answer:** A cytological map shows the positions of genes and other genetic markers relative to the banding patterns of chromosomes.
- **3.** What is a physical map of a DNA molecule or chromosome?
- **Answer:** A physical map of a DNA molecule or chromosome gives the positions of genes or other markers based on the actual distances in base pairs (bp), kilobase pairs (kb), or megabase pairs (mb) separating them. Restriction maps, contig maps, and sequence-tagged site (STS) maps are examples of physical maps.
- 4. How can genetic maps, cytological maps, and physical maps of chromosomes be correlated?

- **Answer:** If a gene is cloned and positioned on all three maps, it provides an anchor marker that can be used to relate the genetic, cytological, and physical maps to each other. All three types of maps are colinear arrays showing the locations of nucleotide sequences on the chromosome. They differ in the units that are used to assign the positions of markers along the linear arrays.
- 5. How can the map position of a gene on a chromosome be used to identify and clone the gene?
- **Answer:** Once a gene has been positioned on the genetic, cytological, or physical map of a chromosome, molecular markers such as RFLPs close to the gene can be used to initiate chromosome walks and jumps starting at the linked marker and progressing along the chromosome to the position of the gene of interest. The identity of the gene must be established by transforming a mutant organism with a wild-type copy of the gene and showing that it restores the wild-type phenotype or, in humans, by comparing the nucleotide sequences of the gene in a number of affected and unaffected individuals (see Figure 15.6).

Testing Your Knowledge

Integrate Different Concepts and Techniques

1. Best disease is a form of blindness in humans that develops gradually in adults. It is caused by an autosomal dominant mutation on chromosome 11. Nine RFLPs, designated 1 through 9, map on chromosome 11 in numerical order. The polymorphisms at each site are designated by superscripts 0 through N, where N + 1 is the number of polymorphisms present at a site in the family represented by the accompanying pedigree. DNA was obtained from each member of the family, digested with the appropriate restriction enzyme, subjected to gel electrophoresis, transferred to a nylon membrane by Southern blotting, denatured, and hybridized to radioactive probes that detect all the RFLPs. After hybridization, the membranes were exposed to X-ray film, and the autoradiograms were used to determine which RFLP(s) was present in each member of the family. The results are shown in the pedigree. Circles represent females; squares represent males; red symbols indicate individuals with Best disease.



Which RFLP site is closest to the mutation that causes Best disease? Which allele of this RFLP is present on the chromosome that carries the Best disease mutation?

Questions and Problems

Enhance Understanding and Develop Analytical Skills

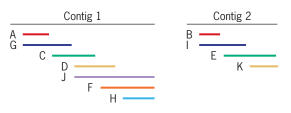
- **15.1** Distinguish between a genetic map, a cytogenetic map, and a physical map. How can each of these types of maps be used to identify a gene by positional cloning?
- **15.2** What is the difference between chromosome walking and chromosome jumping? Why must chromosome jumps,

- **Answer:** RFLP site 4 is closest to the Best disease mutation, which is present on the copy of chromosome 11 carrying the 4^o allele of the polymorphism. Of the polymorphisms on chromosome 11, only the 4^o allele is present in all three family members with Best disease and absent from all five members with normal vision.
- 2. Eleven genomic clones, each containing DNA from chromosome 4 of *Drosophila melanogaster*; were tested for crosshybridization in all pairwise combinations. The clones are designated A through K, and the results of the hybridizations are shown in the accompanying table. A plus sign indicates that hybridization occurred; a minus sign indicates that no hybridization was observed.

	<u>A</u>	<u>B</u>	<u>C</u>	<u>D</u>	<u>E</u>	<u>F</u>	<u>G</u>	<u>H</u>	<u> </u>	J	<u>K</u>
K:	-	-	-	-	+	-	-	-	-	-	+
J:	-	-	+	+	-	+	-	+	-	+	
I:	-	+	-	-	+	-	-	-	+		
H:	-	-	-	-	-	+	-	+			
G:	+	-	+	-	-	-	+				
F:	-	-	-	+	-	+					
E:	-	-	-	-	+						
D:	-	-	+	+							
C:	-	-	+								
B:	-	+									
A:	+										

Based on the hybridization results shown in the table, how many contigs do these clones define? Draw the contig map(s) defined by these data.

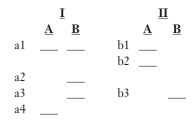
Answer: Maps of the two contigs defined by the 11 mutations are as follows:



rather than just chromosome walks, sometimes be used to identify a gene of interest?

15.3 What is a contig? an RFLP? a VNTR? an STS? an EST? How is each of these used in the construction of chromosome maps?

15.4 The following is a Southern blot of *Eco*RI-digested DNA of rye plants from two different inbred lines, A and B. Developed autoradiogram I shows the bands resulting from probing the blot with ³²P-labeled cDNA1. Autoradiogram II shows the same Southern blot after it was stripped of probe and reprobed with ³²P-labeled cDNA2.



(a) Which bands would you expect to see in the autoradiogram of a similarly probed Southern blot prepared using *Eco*RI-digested DNA from F_1 hybrid plants produced by crossing the two inbred lines? (b) What can you conclude about the gene(s) represented by band a1 on blot I in the two inbreds? (c) The F_1 plants were crossed to plants possessing only bands a1, a4, and b3. DNA was isolated from several individual progeny and digested with *Eco*RI. The resulting DNA fragments were separated by gel electrophoresis, transferred to a nylon membrane, and hybridized with radioactive cDNA1 and cDNA2 probes. The following table summarizes the bands present in autoradiograms obtained using DNA from individual progeny.

	Bands Present								
Plant No.	a1	a2	a3	A4	b1	b2	b3		
1	+	+	+	+			+		
2	+	+	+	+			+		
3	+	+	+	+			+		
4	+	+	+	+			+		
5	+	+	+	+	+	+	+		
6	+			+	+	+	+		
7	+			+	+	+	+		
8	+			+	+	+	+		
9	+			+	+	+	+		
10	+			+			+		

Interpret these data. Do the data provide evidence for RFLPs? at how many loci? Are any of the RFLPs linked? If so, what are the linkage distances defined by the data?

15.5 As part of the Human Genome Mapping Project, you are trying to clone a gene involved in colon cancer. Your first step is to localize the gene using RFLP markers. In the following table, RFLP loci are defined by STS (sequence-tagged site) number (for example, STS1), and the gene for colon cancer is designated *C*.

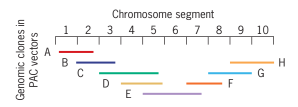
Loci	% Recombination	Loci	% Recombination
<i>C</i> , STS1	50	STS1, STS5	10
<i>C</i> , STS2	15	STS2, STS3	30
<i>C</i> , STS3	15	STS2, STS4	14
<i>C</i> , STS4	1	STS2, STS5	50
<i>C</i> , STS5	40	STS3, STS4	16
STS1, STS2	50	STS3, STS5	25
STS1, STS3	35	STS4, STS5	41
STS1, STS4	50		

(a) Given the percentage recombination between different RFLP loci and the gene for colon cancer shown in the table, draw a genetic map showing the order and genetic distances between adjacent RFLP markers and the gene for colon cancer. (b) Given that the human genome contains approximately $3.3 \times 10^{\circ}$ base pairs of DNA and that the human genetic map contains approximately 3300 centiMorgans, approximately how many base pairs of DNA are located along the stretch of chromosome defined by this RFLP map? (*Hint:* First figure how many base pairs of DNA are present per cM in the human genome.) (c) How many base pairs of DNA are present in the region between the colon cancer gene and the nearest STS?

- **15.6** What are STRs? Why are they sometimes called micro-satellites?
- **15.7** You have cloned a previously unknown human gene. What procedure will allow you to position this gene on the cytological map of the human genome without performing any pedigree analyses? Describe how you would carry out this procedure.
- **15.8** You have identified a previously unknown human EST. What must be done before this new EST can be called an STS?
- **15.9** VNTRs and STRs are specific classes of polymorphisms. What is the difference between a VNTR and a STR?
- **15.10** An RFLP and a mutant allele that causes albinism in humans cannot be shown to be separated by recombination based on pedigree analysis or by radiation hybrid mapping. Do these observations mean that the RFLP occurs within or overlaps the gene harboring the mutation that causes albinism? If so, why? If not, why not?
- **15.11** Why is the resolution of radiation hybrid mapping of human genes higher than the resolution of standard somatic-cell hybrid mapping?
- **15.12** An RFLP and a mutation that causes deafness in humans both map to the same location on the same chromosome. How can you determine whether or not the RFLP overlaps with the gene containing the deafness mutation?
- **15.13** What were the goals of the Human Genome Project? What impact has achieving these goals had on the practice

of medicine to date? What are some of the predicted future impacts? What are some of the possible misuses of human genome data?

- **15.14** To be useful as a genetic marker for positional cloning of a mutant allele that causes an inherited abnormality in humans, an RFLP must be present on the same homologue as the mutation. Why?
- **15.15** Which type of molecular marker, RFLP or EST, is most likely to mark a disease-causing mutant gene in humans? Why?
- **15.16** Bacteriophage Φ X174 contains 11 genes in a genome of 5386 bp; *E. coli* has a predicted 4467 genes in a genome of about 4.639 kb; *S. cerevisiae* has about 6000 genes in a genome of size 12.1 mb; *C. elegans* has about 22,000 genes present in a genome of about 100 mb; and *H. sapiens* has an estimated 20,500 genes in its 3000-mb genome. Which genome has the highest gene density? the lowest gene density? Does there appear to be any correlation between gene density and developmental complexity? If so, describe the correlation.
- **15.17** A contig map of one segment of chromosome 3 of *Arabidopsis* is shown as follows.



(a) If an EST hybridizes with genomic clones C, D, and E, but not with the other clones, in which segment of chromosome 3 is the EST located? (b) If a clone of gene ARA hybridizes only with genomic clones C and D, in which chromosome segment is the gene located? (c) If a restriction fragment hybridizes with only one of the genomic clones shown above, in which chromosome segment(s) could the fragment be located?

15.18 💿 Eight human–Chinese hamster radiation hybrids were tested for the presence of six human ESTs designated A through F. The results are shown in the following table, where a plus indicates that a marker was present and a minus indicates that it was absent.

		1	2	Rac 3	diatio 4	n hyt 5	orid 6	7	8
	А	_	+	_	_	+	+	_	+
	В	+	-	+	-	-	+	-	+
Marker	С	-	+	+	+	-	-	-	+
Ma	D	+	-	+	+	-	-	+	-
	Е	+	-	+	+	-	-	+	-
	F	-	+	-	+	+	+	+	-

Based on these data, do any of the ESTs appear to be closely linked? Which ones? What would be needed for you to be more certain of your answer?

- **15.19** What is the major advantage of gene chips as a microarray hybridization tool?
- **15.20** What major advantage does the green fluorescent protein of the jellyfish have over other methods for studying protein synthesis and localization?
- **15.21** You are given chromosome-specific cDNA libraries for all 24 human chromosomes. How might these libraries be used to study chromosome evolution in primates?
- **15.22** Of the cereal grass species, only maize contains two copies of each block of linked genes. What does this duplication of sets of maize genes indicate about the origin of this agronomically important species?
- **15.23** Five human genomic DNA clones present in PAC vectors were tested by hybridization for the presence of six sequence-tagged sites designated STS1 through STS6. The results are given in the following table; a plus indicates the presence of the STS, and a minus indicates the absence of the STS.

			STS					
		1	2	3	4	5	6	
PAC clone	А	+	-	+	+	-	-	
	В	+	-	-	-	+	-	
	С	-	-	+	+	-	+	
	D	-	+	-	-	+	-	
	Е	-	-	+	-	-	+	

(a) What is the order of the STS sites on the chromosome?

- (b) Draw the contig map defined by these data.
- 15.24 At the beginning of this chapter, we discussed the sequence of two-thirds of the nuclear genome of Homo neanderthalensis. The complete sequences of six mitochondrial genomes of H. neanderthalensis have been available for some time; the first H. neanderthalensis mtDNA sequence was published in 2008. How similar are the sequences of the mtDNAs of H. neanderthalensis and H. sapiens? Are the genomes similar in size? Is the amount of diversity observed in the mtDNAs of Neanderthals and humans the same? If not, what might this tell us about the sizes of Neanderthal and human populations? How many genes are present in the H. neanderthalensis mitochondrial genome? How many of these genes encode proteins? How many specify structural RNA molecules? Are there any pseudogenes in H. neanderthalensis mtDNA? All of these questions can be answered by visiting the http://www.ncbi.nlm.nih.gov web site.
- **15.25** Assume that you have just sequenced a small fragment of DNA that you had cloned. The nucleotide sequence of this segment of DNA is as follows.

aagtagtcgaaaccgaattccgtagaaacaactcgcacgctccggttcgtgttgcaacaaaataggcattcccatcgcggcagttagaatcaccgagtgcccagagtcacgttcgtaagcaggcgcagtttacaggcagcagaaaaatcgattgaacagaaatggctggcggtaaagcaggccaggattcgggcaaggccaaggcgaaggcggtatcgcgttccgcgcgcgcggg In an attempt to learn something about the identity or possible function of this DNA sequence, you decide to perform a BLAST (nucleotide blast) search on PubMed's Entrez web site (http://www.ncbi.nlm.nih.gov/entrez). Paste or type this sequence into the query sequence box. Run the search and examine the sequences most closely related to your query sequence. Are they coding sequences? What proteins do they encode? The first sequence listed is NM 079795.2. Go to the Entrez "Search across databases" tool (click the box at the bottom of the Entrez home page), type or paste NM_079795.2 in as the query, and click on Go. You will get results ("hits") in five databases: Nucleotide, Gene, HomoloGene, Probe, and UniGene. Examine the information in all five databases and see what you can learn about the function of your DNA or the protein that it encodes. What have you learned about your DNA sequence from these searches? Repeat the BLAST search with only half of your sequence as the query sequence. Do you still identify the same sequences in the databases? If you use one-fourth of your sequence as a query, do you still retrieve the same sequences? What is the shortest DNA sequence that you can use as a query and still identify the same sequences in the databanks?

15.26 PubMed's Entrez web site (http://www.ncbi.nlm.nih.gov/ entrez) can also be used to search for protein sequences. Instead of performing a BLAST search with a nucleic acid query, one performs a protein blast with a polypeptide (amino acid sequence) query. Assume that you have the following partial sequence of a polypeptide:

GYDVEKNNSRIKLGLKSLVSKGILVQTKGT-GASGSFKLNKKAASGEAKPQAKKAGAAKA

Go to the Entrez web site and click BLAST on the left. Then click on protein blast and enter your query sequence in the box at the top. Then click BLAST. Your

results should be available in 10 to 15 seconds. The first two sequences listed should be identical to your query sequence; the third sequence should differ from your query by a single amino acid. What is the identity of your query sequence?

15.27 The sequence of a gene in *Drosophila melanogaster* that encodes a histone H2A polypeptide is:

aagtagtcgaaaccgaattccgtagaaacaactcgcacgctccggtttcgtgttgcaacaaaataggcattcccatcgcggcagttagaatcaccgagtgcccagagtcacgttcgtaagcaggcgcagtttacaggcagcagaaaaatcgattgaacagaaatggctggcggtaaagcaggcaaggattcgggcaaggccaaggcgaaggcggtatcgcgttccgcgcgcgcgggtcttcagttccccgtgggtcgcatccatcgtcatctcaagagccgcactacgtcacatggacgcgtcggagccactgcagccgtgtactccgct-catcgaaggacttgaaagtgaaacgtatcactcctcgccacttacagctcgccattcgcggagacgaggagctggacagcctgatcaaggcaaccatcgctggtggcggtgtcattccgcacatacacaagtcgctgatcggcaaaaaggaggaaacggtgcaggatccgcagcggaagggcaacgtcattetgtcgcaggcctactaagccagtcggcaatcggacgccttcgaaacatgcaacactaatgtttaattcagatttcagcagagacaagctaaaacaccgacgagttgtaatcatttctgtgcgccagcatatatttcttatatacaacg-gttgagtatgaaacaaaaacaaaacttaatttagagcaaagtaattacacgaataaatttaataaaaaaaactataataaaaacgcc.

Let's use the translation software available on the Internet at http://www.expasy.org/tools/dna.html to translate this gene in all six possible reading frames and see which reading frame specifies histone H2A. Just type or paste the DNA sequence in the "ExPASy Translate" tool box, and click TRANSLATE SEQUENCE. The results will show the products of translation in all six reading frames with **Met**'s and **Stop**'s boldfaced to highlight potential open reading frames. Which reading frame specifies histone H2A?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

The chimpanzee, *Pan troglodytes*, is our closest living relative. Humans and chimps evolved from a common ancestor that lived approximately 6 million years ago.

- 1. How similar are the chimpanzee and human genomes?
- 2. If you compare some important proteins—for example, the α and β -globins—of humans and chimps, how similar are their amino acid sequences?
- 3. If you compare the nucleotide sequences of the genes encoding the α and β -globins, how similar are they?
- **4.** Are the amino acid sequences of the proteins or the nucleotide sequences of the genes more similar? Why might this be expected?
- **5.** Given the striking similarities between the human and chimpanzee genomes, what kinds of differences do you think are

likely to explain the behavioral differences between humans and chimps?

Hint: At the NCBI web site, go to Genome Biology, then under Genome Resources, Genome Projects Database, click on Mammals and go to *Pan troglodytes*. To compare sequences, go back to Genome Biology and click on HomoloGene. Search using *HBB* (the gene symbol for β -hemoglobin), and click on 1. HomoloGene:68066. Scroll down to Show Pairwise Alignments and do a BLAST comparison of the *Pan* and *Homo* β -globins. To compare the nucleotide sequences of the genes, return to the Entrez home page and perform the search in the Nucleotide database and carry out a similar BLAST search using the nucleotide sequence.

Applications of Molecular Genetics



CHAPTER OUTLINE

- Use of Recombinant DNA Technology to Identify Human Genes and Diagnose Human Diseases
 - Human Gene Therapy
 - DNA Profiling
 - Production of Eukaryotic Proteins in Bacteria
 - Transgenic Animals and Plants
 - Reverse Genetics: Dissecting Biological Processes by Inhibiting Gene Expression

Leber's congenital amaurosis is caused by autosomal recessive mutations in any of at least 12 different genes. Type II, the most severe form of the disease, is caused by mutations in a gene called *RPE64*, which is expressed in retinal pigment epithelial (RPE) cells that provide the pigment rhodopsin to the photoreceptors in the back of the eye. In the absence of the *RPE64* gene product, the photoreceptors degenerate leading to blindness.

This type of blindness is not restricted to humans; it also occurs in other mammals, especially dogs. Indeed, mutations in the canine version of *RPE64* are common in the Briard breed of dogs and cause a very similar type of blindness. In 2001, scientists at the University of Pennsylvania demonstrated that they could restore some vision in blind dogs by injecting copies of functional *RPE64* genes into retinal cells. This work set the stage for similar gene therapy trials in humans with the inherited disorder.

The first such gene therapy trials in humans were done at the Children's Hospital of Philadelphia and in the United Kingdom in 2008. The goals of these studies were to test the safety of the gene therapy procedure being used. In all cases, one eye was treated and the other eye was untreated. The initial results showed that four of the six young adults who were treated with good *RPE64* genes exhibited improved vision in their treated eye. However, based on their results with dogs, the researchers predicted that performing the gene therapy on children would yield a larger response because they have more intact retinal cells than adults.² Nine more patients were treated, including four children ages 8 to 11, and the results

²Kaiser, J., October 24, 2009. Gene Therapy Helps Blind Children See. http:// news.sciencemag.org/sciencenow/2009/10/24-01.html.

Gene Therapy Improves Sight in Child with Congenital Blindness

The first unusual thing that Nancy and Ethan Haas noticed about their son Corey was that he rarely made eye contact with them as an infant. Then, as a toddler, he had a tendency to bump into things; however, his most unusual trait was his attraction to bright lights. According to his dad, Corey "was constantly staring at lights." He started wearing glasses when he was 10 months old. When Corey was six years old, his doctors discovered that he had a rare inherited disorder called Leber's congenital amaurosis type II. Corey's doctors told his parents that he would probably be completely blind by age 40, so he started learning Braille in preparation for blindness.¹



Corey Haas playing a video game prior to the gene therapy that improved his vision. Corey has a rare inherited disorder called Leber's congenital amaurosis type II, which limits his vision and would have led to blindness without gene therapy.

¹Waters, R., October 24, 2009. Gene Therapy Gives Sight to Blind Children with Rare Disorder. www.bloomberg.com/apps/news?pid=newsarchive&sid=arwB5NT9QXeg.

were impressive. The children showed improved ability to maneuver through an obstacle course and increased sensitivity to light.³

One of the treated children was Corey Haas. Corey told reporters at a press conference in October 2008 that he can now recognize faces, read large-print books, ride his bike around his neighborhood,

³The Children's Hospital of Philadelphia Press Release. December 15, 2010. One Shot of Gene Therapy and Children with Congenital Blindness Can Now See. http://multivu.prnewswire.com/mnr/chop/40752. and even play baseball.⁴ For more details on Corey's gene therapy and its impact on his life, watch the video entitled "New Hope for Gene Therapy: A Young Boy's Fight against Blindness" at www.youtube.com/ watch?v=FyR99anGBqE. Hopefully, Corey's improved vision will continue, and he will not face blindness at age 40. And, hopefully, there will be more success stories for gene therapy in the future.

⁴Kaiser, J., October 24, 2009.

Just as geneticists now know the complete pathway of morphogenesis of bacteriophage T4 (Chapter 13), in the future they will know the complete pathway of morphogenesis of a yeast cell, a fruit fly, an *Arabidopsis* plant, or, indeed, even a human being. Moreover, at some point, biologists will understand the molecular basis of learning and memory and will know what molecular events underlie the aging process. Most important, they will understand the complex mechanisms that regulate cell division in humans and should be able to use this knowledge to prevent or cure at least some types of human cancer and life-threatening viral infections.

Use of Recombinant DNA Technology to Identify Human Genes and Diagnose Human Diseases

The mutant genes that cause Huntington's disease and cystic fibrosis were identified by positional cloning. These genes and other mutant genes that cause human diseases can be detected using DNA probes. Recombinant DNA techniques have revolutionized the search for defective genes that cause human diseases. Indeed, numerous major "disease genes" have already been identified by positional cloning (Chapter 15). In addition, the mutations responsible for the diseases have been determined by comparing the nucleotide sequences of wild-type and mutant alleles of the genes. The coding sequences of the wild-type alleles have been translated by computer to predict the amino acid sequences of the gene products. Oligopeptides have been synthesized based on the predicted amino acid sequences and

used to produce antibodies, which, in turn, have been used to localize the gene products and to investigate their functions *in vivo*. The results of these studies will allow future treatment of some of these diseases by gene therapy.

HUNTINGTON'S DISEASE

Huntington's disease (HD) is genetic disorder caused by an autosomal dominant mutation, which occurs in about one of every 10,000 individuals of European descent. Individuals with HD undergo progressive degeneration of the central nervous system, usually beginning at age 30 to 50 years and terminating in death 10 to 15 years later. To date, HD is untreatable. However, identification of the gene and the mutational defect responsible for HD has kindled hope for an effective treatment in the future. Because of the late age of onset of the disease, most HD patients already have children before the disease symptoms appear. Since the disorder is caused by a dominant mutation, each child of a heterozygous HD patient has a 50 percent chance of being afflicted with the disease. These children observe the degeneration and death of their HD parent, knowing that they have a 50:50 chance of suffering the same fate.

The gene responsible for HD (*HTT*, for *huntingtin*) was one of the first human genes shown to be tightly linked to a restriction fragment-length polymorphism (RFLP). In 1983, James Gusella, Nancy Wexler, and coworkers demonstrated that the *HTT* gene cosegregated with an RFLP that mapped near the end of the short arm of

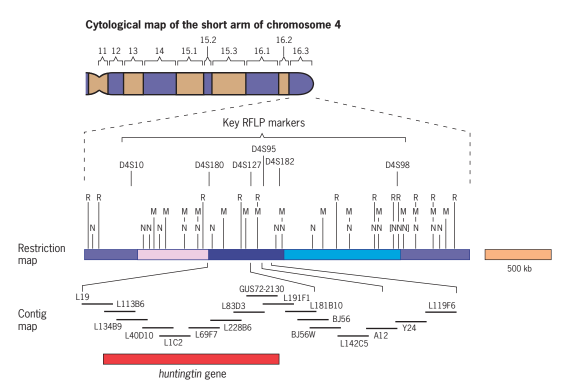


FIGURE 16.1 Identification of the gene responsible for Huntington's disease by positional cloning. The cytological map of the short arm of chromosome 4 is shown at the top. The RFLP markers, restriction map, and contig map used to locate the *huntingtin* gene are shown below the cytological map. M, N, and R represent *Mlul*, *Notl*, and *Nrul* restriction sites, respectively.

chromosome 4. They based their findings mainly on data from studies of two large families, one in Venezuela and one in the United States. Subsequent research showed that the linkage was about 96 percent complete; 4 percent of the offspring of *HTT* heterozygotes were recombinant for the RFLP and the mutant *HTT* allele. Given this early localization of the *HTT* gene to a relatively short segment of chromosome 4, some geneticists predicted that the *HTT* gene would soon be cloned and characterized. However, the task was more difficult than anticipated and took a full 10 years to accomplish.

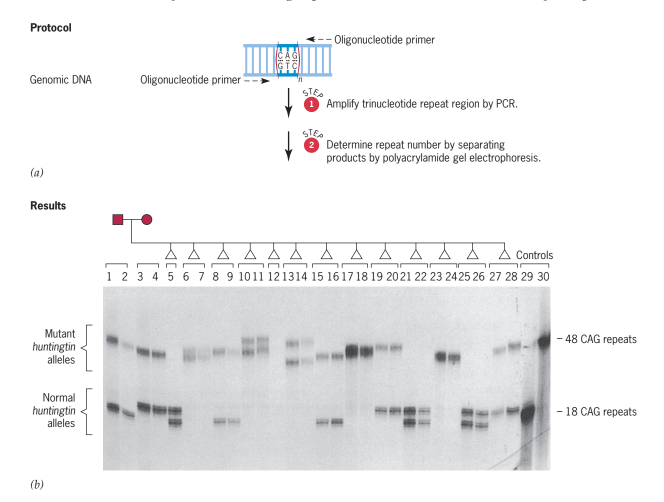
By using positional cloning procedures, Gusella, Wexler, and coworkers identified a gene, first called *IT15* (for Interesting Transcript number 15) and subsequently named *huntingtin*, that spans about 210 kb near the end of the short arm of chromosome 4 (**Figure 16.1**). This gene contains a trinucleotide repeat, $(CAG)_n$, which is present in 11 to 34 copies on each chromosome 4 of healthy individuals. In individuals with HD, the chromosome carrying the *HTT* mutation contains 42 to 100 or more copies of the CAG repeat in this gene. Moreover, the age of onset of HD is negatively correlated with the number of copies of the trinucleotide repeat. Rare juvenile onset of the disease occurs in children with an unusually high repeat copy number. The trinucleotide repeat regions of *HTT* genes are unstable, with repeat numbers often expanding and sometimes contracting between generations. Gusella, Wexler, and collaborators detected expanded CAG repeat regions in chromosomes from 72 different families with HD, leaving little doubt that they had identified the correct gene.

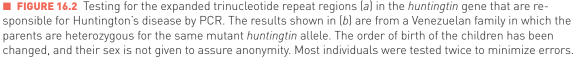
The *buntingtin* gene is expressed in many different cell types, producing a large 10- to 11-kb mRNA. The coding region of the *buntingtin* mRNA predicts a protein 3144 amino acids in length. Unfortunately, the predicted amino acid sequence of the huntingtin protein has provided little information about its function. It exhibits no sequence homology with other proteins. In cells, huntingtin protein is found associated with microtubules and vesicles, suggesting that it might be involved in transport or cytoskeletal attachments of some type. The dominance of the *HTT* mutation indicates that the mutant protein causes the disease.

The expanded CAG repeat region in the mutant *huntingtin* gene encodes an abnormally long polyglutamine region near the amino terminus of the protein. The elongated polyglutamine region fosters protein–protein interactions that lead to the accumulation of aggregates of the huntingtin protein in brain cells. These protein aggregates are thought to cause the clinical symptoms of HD, and current approaches to treatment involve attempts to disrupt or eliminate these protein aggregates.

HD was the fourth human disease to be associated with an unstable trinucleotide repeat. In 1991, fragile X syndrome—the second most common form of mental retardation in humans—was the first human disorder to be associated with an expanded trinucleotide repeat. We discuss fragile X syndrome and the expanded trinucleotide repeat responsible for that disorder in Focus on Fragile X Syndrome and Expanded Trinucleotide Repeats. Shortly thereafter, myotonic dystrophy and spinobulbar muscular atrophy (both diseases associated with loss of muscle control) were shown to result from expanded trinucleotide repeats. Today, over 40 different human disorders—many associated with neurodegenerative abnormalities—have been shown to result from expanded trinucleotide repeats. They include several types of spinocerebellar ataxia, dentatorubro-pallidoluysian atrophy (Haw River syndrome), Friedreich ataxia, and fragile X syndrome. The high frequency of human disorders caused by the expansion of trinucleotide repeats indicates that this may be a common mutational event in our species.

Although the identification of the genetic defect, the expanded trinucleotide repeat in the *huntingtin* gene, has not led to a treatment of the disorder, it has provided a simple and accurate DNA test for the huntingtin mutation (**■** Figure 16.2). Once the nucleotide sequences of the *huntingtin* gene on either side of the trinucleotide repeat region were





FOCUS ON

FRAGILE X SYNDROME AND EXPANDED TRINUCLEOTIDE REPEATS

ragile X syndrome is the second (after Down syndrome; see Chapter 6) most common form of inherited mental retardation in humans. Individuals with fragile X syndrome show significant mental impairment; they may also exhibit facial and behavioral abnormalities. The fragile X syndrome occurs in about 1 in 4000 males and in about 1 in 7000 females. Pedigree studies indicate that the fragile X syndrome is caused by a dominant, X-linked mutation that is incompletely penetrant. About 20 percent of hemizygous males and about 30 percent of heterozygous females do not show symptoms. Fragile X syndrome was the first human disorder to be associated with an unstable trinucleotide repeat (see the section Huntington's Disease at the beginning of this chapter).

Early studies demonstrated that the fragile X syndrome is associated with a cytological anomaly detectable in cells cultured in the absence of thymidine and folic acid. This anomaly—a constriction near the tip of the long arm of the X chromosome—gives the impression that the tip is ready to detach from the rest of the chromosome (**Figure 1***a*), hence the name fragile X chromosome. Molecular analysis subsequently showed that this chromosome contains an unstable trinucleotide repeat, $(CGG)_n$, at the fragile site. This repeat is located in the 5'-untranslated region of a gene designated as *FMR-1*, for fragile X mental retardation gene 1 (**Figure 1***b*). The protein product of this gene, denoted FMRP, accumulates in the dendrites of neurons, which are long extensions of the neuronal cell body that make connections with other cells.

FMRP is an RNA-binding protein. It is found in complexes with mRNAs and other components of the translation apparatus, and it may play a role in transporting mRNA molecules or in regulating their translation. Transcription of the *FMR-1* gene is turned off in fragile X patients, and the absence of FMRP protein seems to be the cause of the observed mental deficiencies.

How is the loss of FMRP expression related to the unstable trinucleotide repeat in the 5'-region of the *FMR-1* gene? Normal—that is, expressed—*FMR-1* genes contain 6 to 59 copies of this

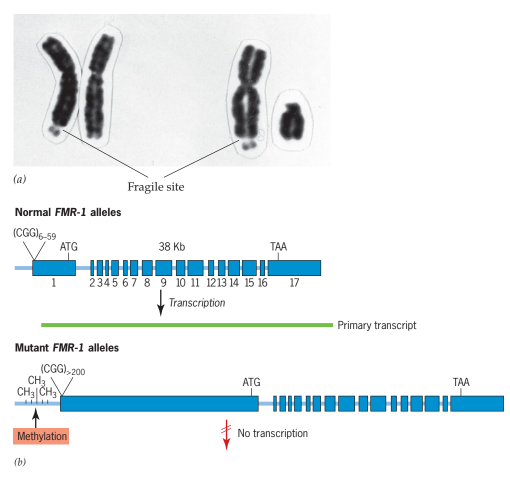


FIGURE 1 (a) The fragile X and a normal X chromosome from a female (left), and the fragile X and a normal Y chromosome from a male (right). (b) The location and number of CGG trinucleotide repeats in normal (top) and mutant alleles (bottom) of the *FMR-1* gene. The promoters of the mutant alleles are heavily methylated, which blocks transcription.

FOCUS ON (continued)

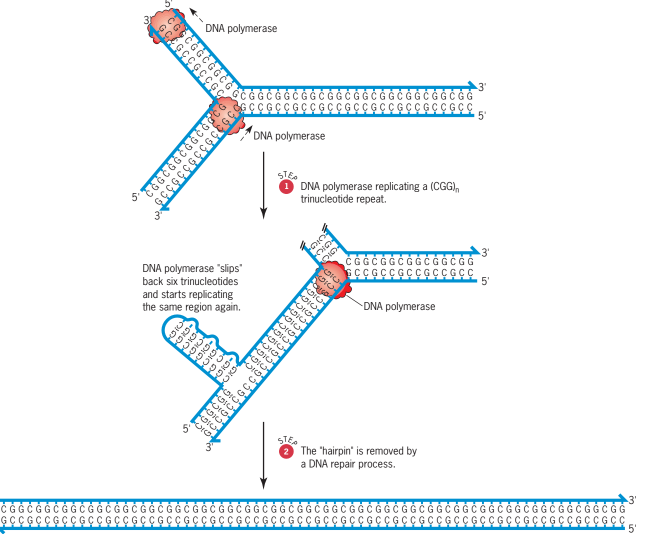


repeat. By contrast, abnormal—that is, unexpressed—*FMR-1* genes, which are found in people who have the fragile X syndrome, contain from 200 to 1500 copies. Somehow an increase in the number of trinucleotide repeats interferes with the expression of the *FMR-1* gene. One hypothesis is that the increased number of repeats leads to chemical modification of the DNA in the promoter of the *FMR-1* gene. This promoter is highly methylated in individuals who have the fragile X syndrome. Various studies have shown that hypermethylation of DNA, especially in and around promoters, silences gene expression (see Chapter 19).

What causes the trinucleotide repeats in chromosomes to increase their copy number from 6-59 to 200-1500? One hypothesis is that during DNA replication, DNA polymerase may "slip," or

"stutter," when it passes through a region containing lots of short, tandem repeats (**Figure 2**). After repair systems clean up the resulting hairpin structures, the repeat region may be significantly expanded. If so, this would explain why the repeated regions tend to be unstable from generation to generation.

Within a year of the discovery of the unstable trinucleotide repeat in the *FMR-1* gene, another neurodegenerative disorder, spinobulbar muscular atrophy (also known as Kennedy's disease), was linked to an unstable trinucleotide repeat, this time (CAG)_n. Other neurodegerative disorders have since been shown to result from expanded trinucleotide repeats. The best known of these is Huntington's disease. Mutations involving unstable trinucleotide repeats therefore seem to be a significant type of genetic defect in our species.



Expanded trinucleotide repeat

FIGURE 2 A possible mechanism for the expansion of trinucleotide repeats. During the replication of the tandem repeat, DNA polymerase falls off the template strand, slips backwards, and then reinitiates synthesis in a previously replicated region. The hairpin formed as a result of the slippage is recognized as a defect by a DNA repair enzyme, which initiates the repair process. A DNA polymerase involved in the repair pathway catalyzes the synthesis of a strand complementary to the unfolded hairpin, producing an expanded trinucleotide repeat region.

PROBLEM-SOLVING SKILLS



Testing for Mutant Alleles That Cause Fragile X Mental Retardation

THE PROBLEM

The second most common inherited type of mental retardation in humans is caused by expanded CGG trinucleotide repeats in the *FMR-1* (for fragile X mental retardation gene 1) gene. See Focus on Fragile X Syndrome and Expanded Trinucleotide Repeats for details. Design a DNA test for the presence of *FMR-1* mutant alleles. How will the results of the test tell you whether an individual is homozygous or heterozygous for the mutant allele when present?

FACTS AND CONCEPTS

- **1.** Normal individuals usually have 6 to 59 copies of the CGG trinucleotide present in the region between the promoter and the translation start site of the *FMR-1* gene.
- **2.** Individuals with fragile X syndrome usually have more than 200 copies of this trinucleotide.
- **3.** The entire euchromatic portion of the human genome has been sequenced. Thus, the sequence of the *FMR-1* gene and the genomic sequences flanking it are known.
- **4.** PCR can be used to amplify the region of the *FMR-1* gene that contains the CGG trinucleotide repeats.
- **5.** Polyacrylamide gel electrophoresis can be used to determine the sizes of small DNA molecules.

ANALYSIS AND SOLUTION

- 1. Synthesize forward and reverse oligonucleotide PCR primers (see Figure 14.6) that are complementary to sequences flanking the trinucleotide repeat region of the *FMR-1* gene.
- Use these primers to amplify the trinucleotide repeat region in genomic DNA samples from the individuals to be tested. Genomic DNAs from individuals with a known number of CGG trinucleotide repeats—both normal and expanded—should be included as controls.
- **3.** Use polyacrylamide gel electrophoresis to determine the sizes of the amplified DNAs (see Figure 14.10). The controls will serve as size markers in this analysis.
- 4. DNA samples from individuals who are heterozygous for normal and expanded *FMR-1* alleles will yield two amplified DNA fragments—a smaller fragment containing 6 to 59 copies of the repeat and a larger fragment containing more than 200 copies of the repeat. DNA samples from individuals who are homozygous for an *FMR-1* allele will yield one amplified DNA fragment—small if two normal alleles are present, larger if two mutant alleles are present.

For further discussion visit the Student Companion site.

known, oligonucleotide primers could be synthesized and used to amplify the region by PCR, and the number of CAG repeats could be determined by polyacrylamide gel electrophoresis. Thus, individuals at risk of carrying the mutant *huntingtin* gene can easily be tested for its presence. Because the PCR procedure requires little DNA, the test for HD also can be performed prenatally on fetal cells obtained by amniocentesis or chorionic biopsy (see Focus on Amniocentesis and Chorionic Biopsy in Chapter 6). Test your understanding of the DNA test for mutant *HTT* alleles by devising a similar DNA test for the mutant alleles containing expanded CGG trinucleotide repeats that cause the second most common form of mental retardation in humans (see Problem-Solving Skills: Testing for Mutant Alleles that Cause Fragile X Mental Retardation).

Given the availability of the DNA test for the *huntingtin* mutation, individuals who are at risk of transmitting the defective gene to their children can determine whether they carry it before starting a family. Each person with a heterozygous parent has a 50 percent chance of not carrying the defective gene. If the test is negative, she or he can begin a family with no concern about transmitting the mutation. If the test is positive, the fetus can be tested prenatally, or the couple can consider *in vitro* fertilization and DNA tests on eight-cell pre-embryos prior to implantation (see On the Cutting Edge: Screening Eight-Cell Pre-Embryos for Tay-Sachs Mutations in Chapter 13). If the tests are negative for the *HTT* mutation, the embryo can be implanted in the mother's uterus with the knowledge that it carries two normal copies of the *huntingtin* gene. If used conscientiously, the DNA test for the *HTT* mutation should diminish human suffering from this dreaded disease.

CYSTIC FIBROSIS

Cystic fibrosis (CF) is one of the most common inherited diseases in humans, affecting 1 in 2000 newborns of northern European heritage. CF is inherited as an autosomal recessive mutation, and the frequency of heterozygotes is estimated to be about 1 in

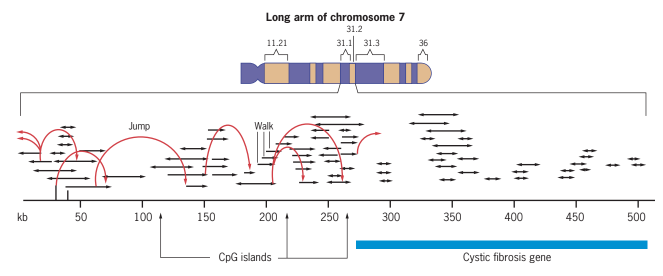


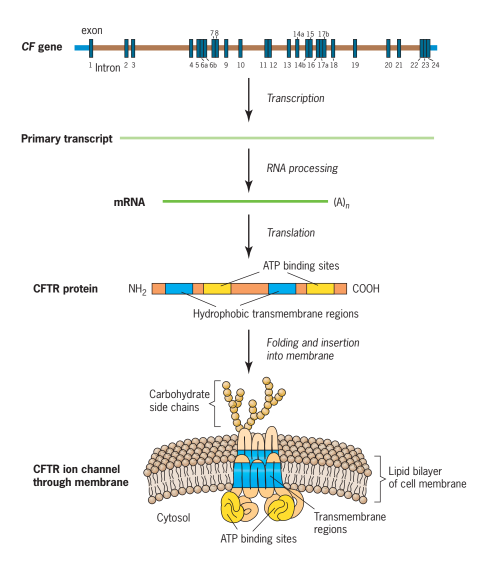
FIGURE 16.3 The sequence of chromosome walks and jumps used to locate and characterize the cystic fibrosis gene. The positions of CpG islands used as landmarks in locating the 5' end of the gene are also shown.

25 in Caucasian populations. In the United States alone, over 30,000 people suffer from this devastating disease. One easily diagnosed symptom of CF is excessively salty sweat, a largely benign effect of the mutant gene. Other symptoms are anything but benign. The lungs, pancreas, and liver become clogged with mucus, which results in chronic infections and the eventual malfunction of these vital organs. In addition, mucus often builds up in the digestive tract, causing individuals to be malnourished no matter how much they eat. Lung infections are recurrent, and patients often die from pneumonia or other infections of the respiratory system. In 1940, the average life expectancy for a newborn with CF was less than two years. With improved methods of treatment, life expectancy has gradually increased. Today, the life expectancy for someone with CF is about 32 years, but the quality of life is poor.

Identification of the *CF* gene is one of the major successes of positional cloning (Chapter 15). Biochemical analyses of cells from CF patients had failed to identify any specific metabolic defect or mutant gene product. Then, in 1989, Francis Collins and Lap-Chee Tsui and their coworkers identified the *CF* gene and characterized some of the mutations that cause this tragic disease. The cloning and sequencing of the *CF* gene quickly led to the identification of its product, which in turn has suggested approaches to clinical treatment of the disease and hope for successful gene therapy in the future.

The *CF* gene was first mapped to the long arm of chromosome 7 by its cosegregation with RFLPs. Further RFLP mapping localized the gene to a 500-kb region of chromosome 7. The two RFLP markers closest to the *CF* gene were then used to initiate chromosome walks (see Figure 15.7) and jumps and to begin construction of a detailed physical map of the region (**■** Figure 16.3). Three kinds of information were used to narrow the search for the *CF* gene.

- 1. Human genes are often preceded by clusters of cytosines and guanines called CpG islands (Chapter 19). Three such clusters are present just upstream from the *CF* gene (Figure 16.3).
- **2.** Important coding sequences usually are conserved in related species. When exon sequences from the *CF* gene were used to probe Southern blots containing restriction fragments from human, mouse, hamster, and bovine genomic DNAs (often called *zoo blots*), the exons were found to be highly conserved.
- **3.** As previously mentioned, CF is known to be associated with abnormal mucus in the lungs, pancreas, and sweat glands. A cDNA library was prepared from mRNA isolated from sweat gland cells growing in culture and screened by colony hybridization using exon probes from the *CF* gene (candidate *CF* gene at the time).



Use of the sweat gland cDNA library proved to be critical in identifying the *CF* gene because northern blot experiments subsequently showed that this gene is expressed only in epithelial cells of the lungs, pancreas, salivary glands, sweat glands, intestine, and reproductive tract. Thus, cDNA clones of the *CF* gene would not have been identified using cDNA libraries prepared from other tissues and organs. The northern blot results also showed that the putative *CF* gene is expressed in the appropriate tissues.

Identification of a candidate gene as a disease gene hinges on comparisons of normal and mutant alleles from several different families. CF is unusual in that 70 percent of the mutant alleles contain the same three-base deletion, $\Delta F 508$, which eliminates the phenylalanine residue at position 508 in the CF gene product. Unlike the *buntingtin* gene, the nucleotide sequence of the CF gene proved very informative. The gene is huge, spanning 250 kb and containing 24 exons (**Figure 16.4**). The CF mRNA is about 6.5 kb in length and encodes a protein of 1480 amino acids. A computer search of the protein data banks quickly showed that the CF gene product is similar to several ion channel proteins, which form pores between cells through which ions pass. The CF gene product, called the cystic fibrosis transmembrane conductance regulator; or CFTR protein, forms ion channels (Figure 16.4) through the membranes of cells that line the respiratory tract, pancreas, sweat glands, intestine, and other organs and regulates the flow of salts and water in and out of these cells. Because the mutant CFTR protein does not function properly in CF patients, salt accumulates in epithelial cells and mucus builds up on the surfaces of these cells.

■ FIGURE 16.4 The structures of the *CF* gene and its product, the CFTR protein. The CFTR protein forms ion channels through the membranes of epithelial cells of the lungs, intestine, pancreas, sweat glands, and some other organs.

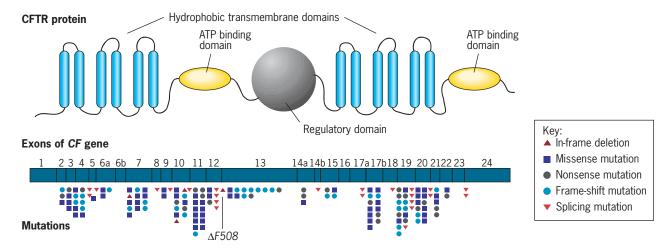


FIGURE 16.5 Mutations in the *CF* gene that cause cystic fibrosis. The distribution and classification of the mutations that cause cystic fibrosis are shown below the exons of the *CF* gene. A schematic diagram of the CFTR protein is shown above the exon map to illustrate the domains of the protein that are altered by the mutations. About 70 percent of all cases of *CF* result from mutation $\Delta F508$, which deletes the phenylalanine present at position 508 of the normal CFTR protein.

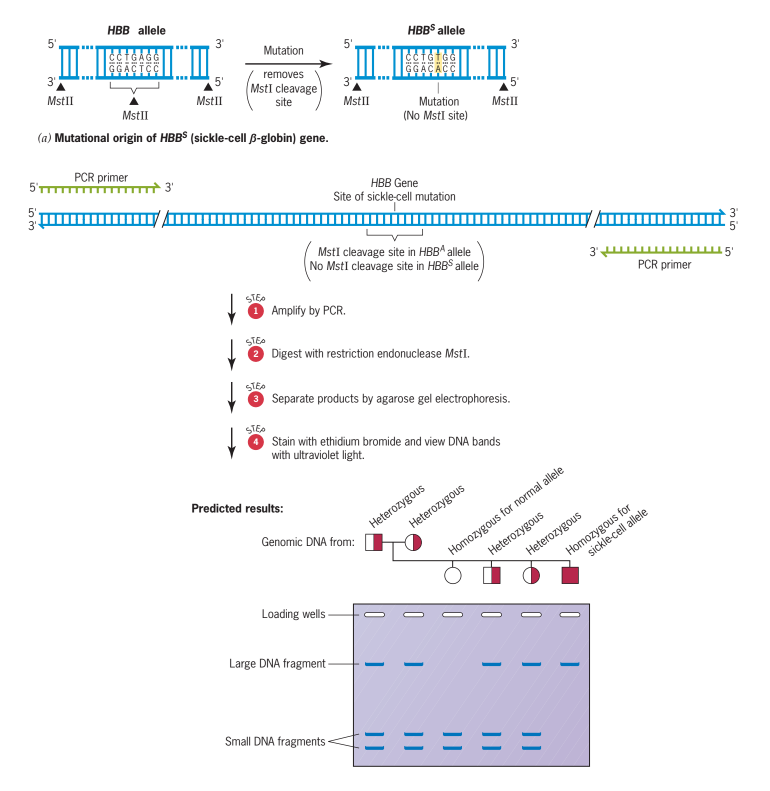
The presence of mucus on the lining of the respiratory tract leads to chronic, progressive infections by *Pseudomonas aeruginosa, Staphylococcus aureus*, and related bacteria. These infections, in turn, frequently result in respiratory failure and death. However, the mutations in the *CF* gene are pleiotropic; they cause a number of distinct phenotypic effects. Malfunctions of the pancreas, liver, bones, and intestinal tract are common in individuals with CF. Although CFTR forms chloride channels (Figure 16.4), it also regulates the activity of several other transport systems such as potassium and sodium channels. Some work suggests that CFTR may play a role in regulating lipid metabolism and transport. CFTR interacts with a number of other proteins and undergoes phosphorylation/dephosphorylation by kinases and phosphatases. Thus, CFTR should be considered multifunctional. Indeed, some of the symptoms of CF may result from the loss of CFTR functions other than the chloride channels.

Although 70 percent of the cases of CF are due to the $\Delta F508$ trinucleotide deletion, over 900 different *CF* mutations have been identified (representative mutations are shown in **Figure 16.5**). About 20 of these mutations are quite common; others are rare, and many have been identified in only one individual. Several of these mutations can be detected by DNA screens such as the test for the $\Delta F508$ deletion illustrated in the Focus on Detection of a Mutant Gene Causing Cystic Fibrosis in Chapter 14. These tests can be performed on fetal cells obtained by amniocentesis or chorionic biopsy. They have also been done successfully on eight-cell pre-implantation embryos produced by *in vitro* fertilization. The diversity of the mutations that cause CF (see Figure 16.5) makes it very difficult to devise DNA tests for all of the mutant *CF* alleles.

MOLECULAR DIAGNOSIS OF HUMAN DISEASES

Once the gene responsible for a human disease has been cloned and sequenced and the mutations that cause the disorder are known, molecular tests for the mutant alleles usually can be designed. These tests can be performed on small amounts of DNA by using PCR to amplify the DNA segment of interest (see Figure 14.6). Thus, they can be performed prenatally on fetal cells obtained by amniocentesis or chorionic biopsy, or even on a single cell from a pre-embryo produced by *in vitro* fertilization.

Some molecular diagnoses involve simply testing for the presence or absence of a specific restriction enzyme cleavage site in DNA. For example, the mutation that causes sickle-cell anemia (Chapter 13) removes a cleavage site for the restriction enzyme MstII (**Theorem 16.6**). The HBB^{s} (sickle-cell) allele can be distinguished from



(b) Distinguishing the HBB and HBB^S alleles by simple molecular techniques.

FIGURE 16.6 (a) The mutation that produces the sickle-cell β -globin (*HBB*^s) allele from the normal β -globin (*HBB*^A) allele removes an *Mst*l cleavage site from the gene. That change can be used to distinguish the two alleles by simple molecular techniques. (b) Detection of the sickle-cell β -globin mutation in the *HBB*^s allele by amplification of fragments of the *HBB* gene from genomic DNA and cleavage with restriction enzyme *Mst*II.

the normal β -globin allele (*HBB^A*) by synthesizing PCR primers that are complementary to DNA sequences flanking the sickle-cell mutation in the *HBB^S* gene and using them to amplify this segment from genomic DNA. The amplified DNA can then be treated with *Mst*II and the products of the reaction separated by agarose gel electrophoresis and stained with ethidium bromide. If the amplified DNA is cleaved by *Mst*II to produce two fragments, it contains the normal *HBB^A* allele; if it isn't cleaved, it contains the mutant *HBB^S* allele. If the genomic DNA was isolated from an individual who is heterozygous for these *HBB* alleles, half will be cleaved and half will remain intact (Figure 16.6). Thus, the presence of the *HBB^S* allele can be diagnosed by a simple molecular test.

For inherited disorders such as Huntington's disease and fragile X syndrome, which result from expanded trinucleotide repeat regions in genes, PCR and Southern blots can be used to detect the mutant alleles. The DNA test for the *huntingtin* gene is illustrated in Figure 16.2. Other types of mutations can be detected by using allele-specific oligonucleotides to probe genomic Southern blots. This procedure is illustrated for the $\Delta F508$ mutation in the CF gene—the most frequent cause of cystic fibrosis—in Focus on Detection of a Mutant Gene Causing Cystic Fibrosis in Chapter 14. Indeed, once the mutations responsible for a disease have been characterized, the development of DNA tests to detect the most common ones is usually routine. The availability of diagnostic tests for mutations that cause human diseases has contributed greatly to the field of genetic counseling, providing invaluable information to families in which the genetic defects occur.

- **KEY POINTS** *The mutant genes responsible for Huntington's disease and cystic fibrosis were identified by positional cloning.*
 - The nucleotide sequences of the huntingtin and CF genes were used to predict the amino acid sequences of their polypeptide products and to obtain information about the functions of the gene products.
 - The characterization of the huntingtin and CF genes has led to the development of DNA tests that detect some of the mutations that cause Huntington's disease and cystic fibrosis.
 - Mutant genes that are responsible for inherited human disorders can often be diagnosed by DNA tests.
 - The results of DNA tests for mutant genes that cause inherited diseases allow genetic counselors to inform families of the risks of having affected children.

Human Gene Therapy

Gene therapy—introducing functional copies of a gene into an individual with two defective copies of the gene—is a potential tool for treating inherited human diseases. Of the over 6000 inherited human diseases catalogued to date, only a few are currently treatable. For many of these diseases, the missing or defective gene product cannot be supplied exogenously, as insulin is supplied to diabetics. Most enzymes are unstable and cannot be delivered in functional form to their sites of action in the body, at least not in a form that provides for long-term activity. Cell membranes are impermeable to large macromolecules such

as proteins; thus, enzymes must be synthesized in the cells where they are needed. Therefore, treatment of inherited diseases is largely restricted to those cases where the missing metabolite is a small molecule that can be distributed to the appropriate tissues of the body through the circulatory system, or the symptoms can be controlled by modifying the individual's diet. For many other inherited diseases, **gene therapy** offers the most promising approach to successful treatment. Gene therapy involves adding a normal (wild-type) copy of a gene to the genome of an individual carrying defective copies of the gene. A gene that has been introduced into a cell or organism is called a **transgene** (for *trans*ferred *gene*) to distinguish it from endogenous genes, and the organism carrying the introduced gene is said to be **transgenic**. If gene therapy is successful, the transgene will synthesize the missing gene product and restore the normal phenotype.

Before considering specific examples, we need to discuss two types of gene therapy: **somatic-cell** or **nonheritable gene therapy**, and **germ-line** or **heritable gene therapy**. In higher animals such as humans, the reproductive or germ-line cells are produced by a cell lineage separate from all somatic-cell lineages. Thus, somatic-cell gene therapy will treat the disease symptoms of the individual but will not cure the disease. That is, the defective gene(s) will still be present in the germ-line cells of the patient after somatic-cell gene therapy and may be transmitted to his or her children. All of the gene-therapy treatments of human diseases that we will discuss here are somaticcell gene therapies. Germ-line gene therapy is being performed on mice and other animals, but not on humans.

The distinction between somatic-cell and germ-line gene therapy is important when we discuss humans. The frequently expressed concerns about humankind's "tinkering with nature" or "playing God" apply to germ-line gene transfers, not to somatic-cell gene therapy. Major moral and ethical considerations are involved in any decision to perform germ-line modifications of human genes. In contrast, somatic-cell gene therapy is no different from enzyme (gene-product) therapy or cell, tissue, and organ transplants. In transplants, entire organs, with all the foreign genes present in the genome of every cell in the organ, are implanted in patients. In current somatic-cell gene therapies, some of the patient's own cells are removed, repaired, and reimplanted in the patient. Thus, somatic-cell gene therapy is less complex and less life-threatening for an individual than an organ transplant.

To perform somatic-cell gene therapy, wild-type genes must be introduced into and expressed in cells homozygous or hemizygous for a mutant allele of the gene. In principle, the wild-type gene could be delivered to the mutant cells by any of several different procedures. Most commonly, viruses are used to carry the wild-type gene into cells. In the case of retroviral vectors, the wild-type transgene is integrated along with the retroviral DNA—into the DNA of the host cell. Thus, when retroviral vectors are used, the transgene is transmitted to all progeny cells in the affected cell lineage.

With other viral vectors, such as those derived from adenoviruses, the transgenes are present only transiently in host cells because the genomes of these viruses replicate autonomously and persist only until the immune system eliminates the viruses along with the infected cells. The advantage of these vectors over retroviral vectors is that no potentially harmful mutations are induced during the integration step (Chapter 13). However, they have two major disadvantages: (1) transgene expression is transient, lasting only as long as the viral infection persists, and (2) most humans exhibit strong immune responses to these viruses, presumably because of prior exposure to the same or closely related viruses. For example, in early attempts to treat cystic fibrosis by somatic-cell gene therapy, an adenoviral vector carrying the CF gene was inhaled by patients, with the hope that lung cells would become infected and synthesize enough of the CF gene product to alleviate some of the symptoms of the disease. Unfortunately, these treatments proved ineffective, at least in part because of rapid immune responses to these viruses in the individuals receiving the treatments.

With diseases such as cystic fibrosis, where effective gene therapy will require long-term transgene expression, the standard adenovirus vectors probably will not work. Because transgene expression is transient, the treatments will need to be repeated periodically. However, given that secondary immune responses are very rapid and efficient, subsequent treatments with the same viral vector probably will be ineffective. Human gene therapy is performed under strict guidelines developed by the National Institutes of Health (NIH). Each proposed gene-therapy procedure is scrutinized by review committees at both the local (institution or medical center) and national (NIH) levels. Several requirements must be fulfilled before a gene-therapy procedure will be approved:

- 1. The gene must be cloned and well characterized; that is, it must be available in pure form.
- **2.** An effective method must be available for delivering the gene into the desired tissue(s) or cells.
- **3.** The risks of gene therapy to the patient must have been carefully evaluated and shown to be minimal.
- 4. The disease must not be treatable by other strategies.
- **5.** Data must be available from preliminary experiments with animal models or human cells and must indicate that the proposed gene therapy should be effective.

A gene-therapy proposal will not be approved by the local and national review committees until they are convinced that all of the above conditions have been fulfilled. Moreover, with the unfortunate death in September 1999 of Jesse Gelsinger, an 18-year-old with ornithine transcarbamylase deficiency, due to a severe immune reaction to the adenovirus vector used in his experimental gene therapy, the review committees are being especially cautious in their evaluation of gene-therapy proposals.

The first use of gene therapy in humans occurred in 1990, when a four-yearold girl with **adenosine deaminase-deficient severe combined immunodeficiency disease** (ADA⁻ SCID) received her first transgene treatment. SCID is a rare autosomal disease of the immune system. Individuals with SCID have essentially no immune system, so that even minor infections are often fatal. In the absence of adenosine deaminase (ADA), toxic levels of the phosphorylated form of its substrate, deoxyadenosine, accumulate in T lymphocytes (white blood cells essential to an immune response) and kill them. T lymphocytes stimulate cells called B lymphocytes to develop into antibody-producing plasma cells. Thus, in the absence of T lymphocytes, no immune response is possible, and newborns with ADA⁻ SCID seldom live more than a few years.

After her gene therapy in 1990, the girl's transgenic T lymphocytes did synthesize adenosine deaminase for a while, but not long-term. Fortunately, enzyme therapy has subsequently proven successful in treating ADA⁻ SCID. Injections of bovine adenosine deaminase stabilized with *polyethylene glycol* (PEG, the key component in antifreeze) are now used to treat ADA⁻ SCID. The four-year-old pioneer of gene therapy is now a healthy and active young woman with a special interest in music. She is also a strong advocate of gene therapy.

To avoid the limitations resulting from the short lifespan of white blood cells, the bone marrow stem cells that give rise to white blood cells could be used to treat immune disorders such as ADA⁻ SCID. The modified stem cells should continually produce T lymphocytes with the *ADA* transgene and could provide a permanent or long-term treatment of the disease. Indeed, stem-cell gene therapy was first used to treat two infants with ADA⁻ SCID in 1993, and this procedure has become the method of choice. Unfortunately, ADA synthesis was still short-term when the transgene was present in stem cells.

During the year 2000, British and French physicians performed what at the time appeared to be the first successful somatic-cell gene-therapy treatment of individuals with a fatal inherited disease. They treated boys with a type of SCID similar to the ADA⁻ SCID previously discussed but caused by mutations in a gene on the X chromosome. This X-linked SCID results from the loss or inactivation of the γ subunit of the interleukin-2 receptor. Interleukin-2 is a signaling molecule required for the development of cells of the immune system. However, the γ polypeptide of the interleukin-2 receptor is also a component of several other lymphocyte-specific growth factors. Collectively, they stimulate the development of B and T lymphocytes—cells

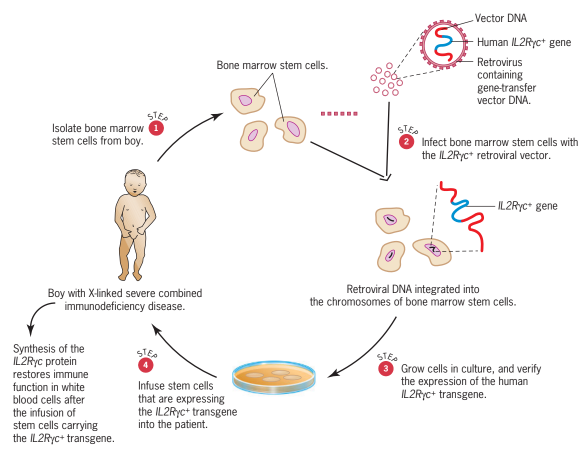


FIGURE 16.7 Treatment of X-linked severe combined immunodeficiency disease (IL2R γ c⁻ SCID) by somatic-cell gene therapy. This form of X-linked SCID results from the loss or lack of activity of the γ polypeptide of the interleukin-2 receptor (the γ polypeptide is also a component of other interleukins). Gene therapy is performed by isolating bone marrow stem cells from the patient, introducing a wild-type copy of the *IL2R* γ c⁺ gene into these cells with a retroviral vector, verifying the expression of the transgene in cultured cells, and infusing the transformed stem cells back into the patient.

required for the production of antibody-producing plasma cells and killer T cells, respectively. In the absence of the γ polypeptide, an individual has no functional immune system and seldom survives for more than a few years.

Like the individuals with ADA⁻ SCID, boys with X-linked SCID seemed to be good candidates for treatment by somatic-cell gene therapy. Thus, the gene encoding the γ subunit of the human interleukin-2 receptor was cloned, inserted into a retroviral vector, introduced into hematopoietic stem cells (precursors to cells of the circulatory system) isolated from patients with X-linked SCID, and checked for gene expression while the cells were still growing in culture medium. After verifying expression of the gene (designated *IL2R* γc for *interleukin-2 receptor* γ *common*), the stem cells were transfused back into the SCID patients from whom they had been isolated (**Figure 16.7**). During the next two years, 14 boys with X-linked SCID were treated. In all 14 cases, gene therapy cured the immunodeficiency, resulting in normal T-cell levels within a few months after treatment. Thus, for two years, everything indicated that the gene therapy had been a major success. Then one of the boys developed acute T-cell leukemia, a cancer of the white blood cells. Later, the same T-cell leukemia was detected in three more of the gene-therapy patients. Clearly, something had gone wrong.

One advantage of retroviral vectors is that they insert themselves into the chromosomes of host cells and, therefore, are transmitted to progeny cells during cell Identification of the LMO2 oncogene by its association with a translocation between chromosomes 7 and 11 in individuals with T-cell acute lymphoblastic leukemia (T-cell ALL).

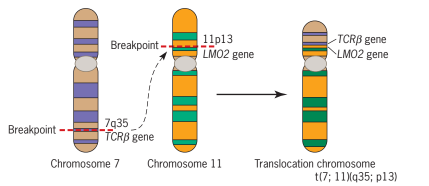


FIGURE 16.8 The *LMO2* gene (*LIM-only* gene 2) encodes a small protein that functions as a bridge joining different transcription factors. It was identified in studies of individuals with T-cell acute lymphoblastic leukemia (T-cell ALL, a cancer affecting white blood cells). In these patients, a translocation had occurred between chromosomes 7 and 11. This translocation moved the *TCR* β (*T-cell* receptor β subunit) gene on chromosome 7 next to the *LMO2* gene on chromosome 11 and resulted in the overexpression of *LMO2*. When overexpressed, *LMO2* behaves as an oncogene (cancer-causing gene; see Chapter 21) in a pathway leading to T-cell leukemia.

division. However, like transposable elements, they can cause mutation by inserting themselves into genes of host cells (see Figure 13.13). In addition, some retroviral DNAs upregulate the expression of genes close to their sites of integration, and the vector (derived from components of the Moloney murine leukemia virus) used to introduce the *IL2Ryc* gene into X-linked SCID patients was of this type.

When the location of the viral DNA carrying the $IL2R\gamma c$ gene was determined in the first two boys to develop leukemia, the vector was found in the same gene in both cases. The retroviral DNA had integrated into a gene that was known to be associated with T-cell acute lymphoblastic leukemia (T-cell ALL) in individuals carrying a unique translocation chromosome. The translocation fused the $TCR\beta$ (*T*-cell receptor β subunit) gene on chromosome 7 with the 5' region of the *LMO2* (LIM-only) gene on chromosome 11 (**Figure 16.8**). LMO2 encodes a protein that is essential for the formation of certain transcription factor complexes. The expression of *LMO2* is normally downregulated during the development of T cells. When it is overexpressed in T cells, it stimulates cell division. As such, LMO2 is classified as a proto-oncogene, a gene that can become a cancer-causing oncogene by mutation or altered expression (see Chapter 21). Indeed, LMO2 is overexpressed in the T cells of individuals with acute leukemia resulting from the translocation shown in Figure 16.8. It is also overexpressed in the boys with X-linked SCID who underwent gene therapy and subsequently developed leukemia or leukemia-like symptoms.

Scientists have known that the retroviral vectors used in gene therapy might cause mutations by integrating within genes. However, the risk was thought to be small. If a vector integrated at random into the human genome (3×10^9 nucleotide pairs), the chance that the vector would insert into a specific gene would be about 1 in a million. However, retroviral vectors are known to insert preferentially into expressed genes. Given that there are about 20,500 genes in the human genome, even if all insertions were into genes, the random insertion of vectors into genes would hit a given gene with a probability of about 1 in 20,500. Obviously, with 2 out of 15 insertions occurring within the *LMO2* gene, insertions are not occurring at random. Instead, this particular vector exhibits a strong tendency to insert into or near the *LMO2* gene.

Clearly, we still have a lot to learn before gene therapy can be used as an effective treatment of inherited human disorders. We need safer vectors, and we need to learn how to regulate the expression of the genes in these vectors. How long will it take to develop effective and safe gene-therapy protocols? We do not have an answer to that question; however, we can predict that there will be a time when gene therapy is used routinely and safely in the treatment of inherited human diseases.

Two recent applications of gene therapy have provided encouraging results. One involves the treatment of children with a rare form of congenital blindness—Leber's congenital amaurosis type II, which was discussed in the opening section of this chapter. The other involves the treatment of Canavan disease, an autosomal recessive neurodegenerative disorder. Individuals with Canavan disease lack an enzyme that breaks down the N-acetylaspartate produced in neurons. When the gene encoding the enzyme was introduced into brain cells, the missing enzyme was synthesized and neurological functions were improved. So far, both of these gene-therapy treatments appear to have been successful.

All past and current somatic-cell gene-therapy protocols are gene-addition procedures; they simply add functional copies of the gene that is defective in the patient to the genomes of recipient cells. They do not replace the defective gene with a functional gene. In fact, the introduced genes are inserted at random or nearly random sites in the chromosomes of the host cells. The ideal gene-therapy protocol would replace the defective gene with a functional gene. Gene replacements would be mediated by homologous recombination and would place the introduced gene at its normal location in the host genome. In humans, gene replacements are usually referred to as targeted gene transfers. Oliver Smithies and coworkers first used homologous recombination to target DNA sequences to the β -globin locus of human tissue-culture cells in 1985. However, the frequency of the targeted gene transfer was very low (about 10⁻⁵). Since then, Smithies, Mario Capecchi, and others have developed improved gene-targeting vectors and selection strategies. As a result, more efficient targeted gene replacements are possible, and cells with the desired gene replacement can be identified more easily. In the future, targeted gene replacements will probably become the method of choice for somatic-cell gene therapy treatment of human diseases.

- Gene therapy involves the addition of a normal (wild-type) copy of a gene to the genome of an individual who carries defective copies of the gene.
- Although somatic-cell gene therapy effectively restored immunological function in boys with X-linked severe combined immunodeficiency disease, four of the boys subsequently developed leukemia or leukemia-like disorders.
- Somatic-cell gene therapy holds promise for the treatment of many inherited human diseases; however, the results to date have been disappointing.

KEY POINTS

DNA Profiling

Fingerprints have played a central role in human identity cases for decades. Indeed, fingerprints have often provided the key evidence that places a suspect at a crime DNA profiles—recorded patterns of DNA polymorphisms— v provide strong evidence of an individual's identity or nonidentity.

scene. The use of fingerprints in forensic cases is based on the premise that no two individuals will have identical prints. Similarly, no two individuals, except for identical twins, will have genomes with the same nucleotide sequences. The human genome contains 3×10^9 nucleotide pairs; each site is occupied by one of the four base pairs in DNA. Moreover, the human genome contains DNA polymorphisms of many different types, polymorphisms that can provide invaluable evidence in cases of uncertain identity.

■ FIGURE 16.9 Ground zero after the collapse of the Twin Towers of the World Trade Center on September 11, 2001. The bodies of some of the nearly 3000 people killed in the collapse could be identified only by comparing their DNA sequences with those of close relatives, a process called DNA profiling.



Recorded patterns of DNA polymorphisms—DNA profiles (originally called DNA prints)—are now used routinely to identify and/or distinguish individuals. The use of DNA sequence data in personal identity cases is called DNA profiling (formerly DNA fingerprinting); it is a valuable tool in cases of uncertain identity, such as paternity,

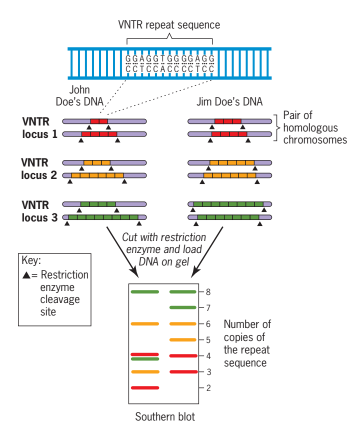


FIGURE 16.10 Simplified diagram of the use of variable number tandem repeats (VNTRs) and Southern blots to prepare DNA profiles.

rape, murder, and the identification of mutilated bodies after explosions, crashes, or other tragedies. DNA profiling was used extensively to identify bodies and body parts recovered in the debris after the collapse of the Twin Towers of the World Trade Center in New York City on September 11, 2001 (**■** Figure 16.9).

Two types of DNA polymorphisms have proven to be especially useful in DNA profiling. Variable number tandem repeats (VNTRs, also called minisatellites) are composed of repeated sequences 10 to 80 nucleotide pairs long, and short tandem repeats (STRs; also called microsatellites) are composed of repeated sequences 2 to 10 nucleotide pairs long. These sequences exhibit highly variable copy number, making them ideal for use in DNA profiling.

For many years, most DNA profiles contained specific banding patterns on Southern blots of genomic DNA cleaved with a specific restriction enzyme and hybridized to appropriate DNA probes (**■** Figure 16.10). Today, most DNA profiles are electropherograms produced by using PCR primers tagged with fluorescent dyes to amplify the genomic DNA segments of interest, capillary gel electrophoresis to separate the PCR products, and lasers and photocells (fluorescence detectors) to record the sizes of the fluorescent PCR products (**■** Figure 16.11). The separation and detection steps are performed using the automated DNA sequencing machines discussed in Chapter 14.

In 1997, the Federal Bureau of Investigation (FBI) adopted a panel of 13 STR loci to be used as the standard database in criminal investigations. Collectively, these 13 STR loci make up the *Combined DNA Index System* (CODIS) that is widely used in DNA profiling. These loci are located on 12 different chromosomes (Table 16.1). By selecting PCR primers that yield products of distinct sizes, three or more STR loci can be amplified with primer-pairs labeled with the same fluorescent

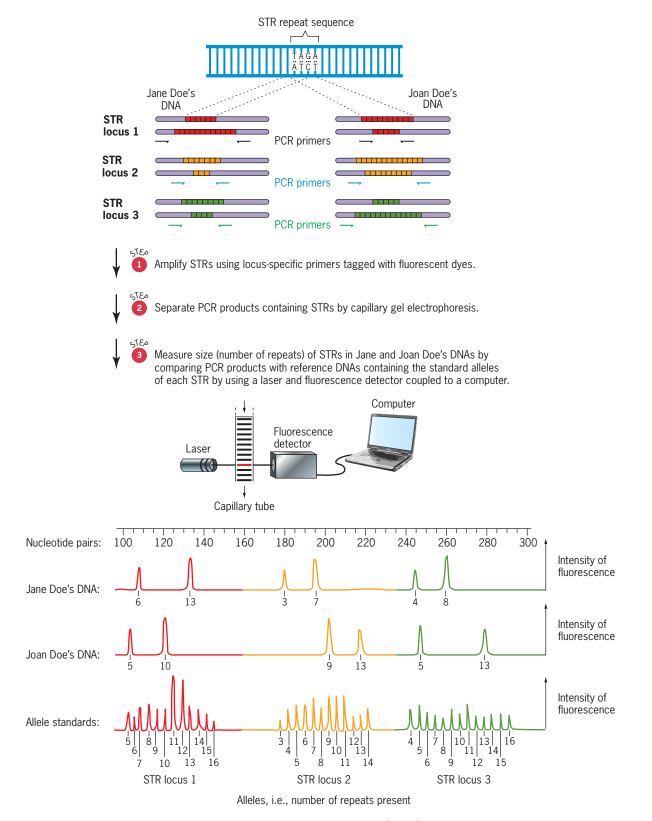
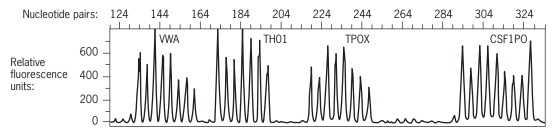
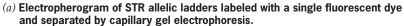


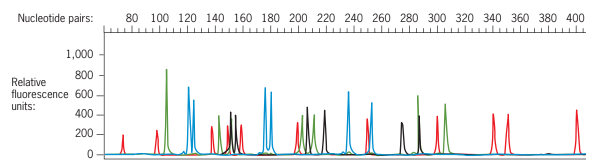
FIGURE 16.11 Diagram illustrating the use of short tandem repeats (STRs), PCR performed with fluorescently tagged primers, capillary gel electrophoresis, and fluorescence detectors to prepare DNA profiles. The sizes of the PCR products are shown in nucleotide pairs above the DNA profiles.

TABLE 16.1 The 13 STR Loci in the Core CODIS Panel					
Locus Chromosome Repeat Motif Number of Alleles Observer					
LUCUS	Chromosome	Repeat Moth	Number of Atteles observed		
1. TPOX	2	GAAT	15		
2. D3S1358	3	[TCTG][TCTA]	25		
3. FGA	4	CTTT	80		
4. D5S818	5	AGAT	15		
5. CSF1P0	5	TAGA	20		
6. D7S820	7	GATA	30		
7. D8S1179	8	[TCTA][TCTG]	15		
8. TH01	11	TCAT	20		
9. VWA	12	[TCTG][TCTA]	29		
10. D13S317	13	TATC	17		
11. D16S539	16	GATA	19		
12. D18S51	18	AGAA	51		
13. D21S11	21	[TCTA][TCTG]	89		

dye and separated by gel electrophoresis (**■** Figure 16.12*a*) and up to nine STR loci can be amplified using three PCR primer-pairs labeled with distinct fluorescent dyes and separated in a single capillary gel electrophoresis tube (**■** Figure 16.12*b*). The separation of families of STR alleles in one to three PCR amplifications and one or two gel electrophoresis separations is called multiplex STR analysis. Several companies have developed fluorescent dye-labeled multiplex PCR primers that allow characterization







(b) Electropherogram of STR alleles in genomic DNA using three pairs of PCR primers each labeled with a different fluorescent dye (shown as blue, green, and black peaks). The red peaks represent DNA size standards. In this multiplex STR analysis, nine STR loci are characterized simultaneously.

FIGURE 16.12 Electropherograms of (*a*) multiplex STR ladders labeled with a single fluorescent dye and separated by capillary gel electrophoresis and (*b*) multiplex analysis of nine STR loci performed using three pairs of PCR primers labeled with three different fluorescent dyes. The red peaks represent added DNA size markers.

of the alleles of all 13 standard STR loci in just two PCR amplifications and gel electrophoresis separations.

The power and utility of DNA profiles in personal identity cases are obvious to anyone familiar with molecular genetics and the techniques utilized in the production of the profiles. Nevertheless, numerous disputes have arisen over the use of DNA profiles in forensic cases over the years. Most of these controversies were related to the competency of the research laboratories involved, the probability of human error in producing profiles, and the methods for calculating the probability that two individuals have identical DNA profiles.

To make accurate estimates of the likelihood of identical profiles, researchers must have reliable information about the frequency of the polymorphisms in the population in question. For example, if inbreeding (matings between related individuals) is common in the population, the probability of identical DNA profiles will increase. Thus, accurate estimates of the probability that two individuals will have matching profiles require reliable information about the frequencies of the polymorphisms in the relevant population. Data obtained from one population should never be extrapolated to another population because different polymorphism frequencies may be present in different populations. For this reason, forensic scientists have collected extensive data on the frequencies of the CODIS STR alleles in populations throughout the world, and these data are used as references in forensic cases using DNA profiles.

DNA profiling provides a powerful forensic tool if used properly. Profiles can be prepared from minute amounts of blood, semen, hair bulbs, or other cells. The DNA is extracted from these cells, amplified by PCR, and the STRs are characterized by PCR using fluorescent primers, capillary gel electrophoresis, and fluorescence detectors/recorders (see Figure 16.11). Although DNA profiles are applicable in all cases of questionable identity, they have proven especially useful in paternity and forensic cases.

PATERNITY TESTS

In the past, cases of uncertain paternity often have been decided by comparing the blood types of the child, the mother, and possible fathers. Blood-type data can be used to prove that men with particular blood types could not have fathered the child. Unfortunately, these blood-type comparisons contribute little toward a positive identification of the father. In contrast, DNA profiles not only exclude misidentified fathers, but also come close to providing a positive

identification of the true father. DNA samples are obtained from cells of the child, the mother, and possible fathers, and DNA profiles are prepared as described in Figures 16.10 or 16.11. When the profiles are compared, all the bands in the child's DNA profile should be present in the combined DNA profiles of the parents. For each pair of homologous chromosomes, the child will have received one from each parent. Thus, approximately half of the DNA bands or peaks in the child's DNA profiles will result from DNA sequences inherited from the mother, and the other half from DNA sequences inherited from the father.

Figure 16.13 shows the DNA profiles of a child, the mother, and two men suspected of being the child's father. In this case, the DNA profiles indicate that the second father candidate is probably the child's biological father. The accuracy of DNA profiles in identifying child–parent relationships increases with the number of polymorphic loci used in the analysis. If all 13 CODIS STR loci are analyzed, the results are usually very accurate. Test your understanding of the use of DNA profiling in paternity cases by working Solve It: How Can DNA Profiles Be Used to Establish Identity?

FORENSIC APPLICATIONS

DNA profiles were first used as evidence in a criminal case in 1988. In 1987, a Florida judge denied the prosecutor's request to present statistical interpretations of DNA

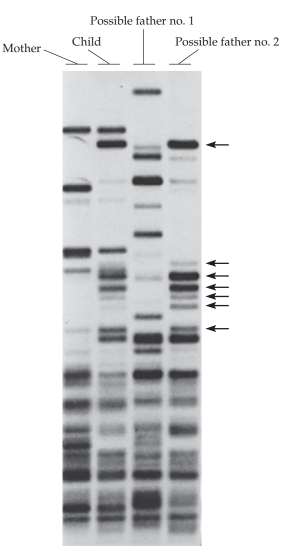


FIGURE 16.13 DNA profiles of a mother, her child, and two men, each of whom claimed to be the child's father. Arrows mark bands that identify male no. 2 as the biological father.

Solve It!

How Can DNA Profiles Be Used to Establish Identity?

A tragic airplane crash killed 17 people everyone on board. The plane burst into flames on impact leaving the bodies burned beyond recognition. Two 10-yearold boys were on the flight, one traveling with his parents and the other on his way home from visiting his grandparents. How can DNA profiles be used to distinguish the bodies of the two boys, so that the surviving parents can bury their own son?

To see the solution to this problem, visit the Student Companion site.

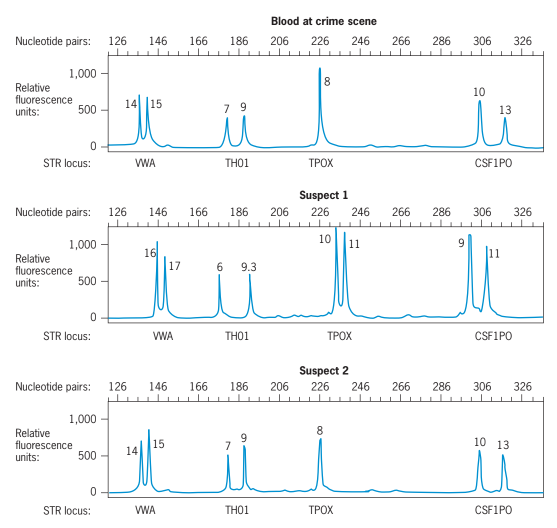


FIGURE 16.14 DNA profiles at four STR loci prepared from DNA isolated from a bloodstain at the site of a crime and from blood obtained from two individuals suspected of committing the crime. In actual forensic cases, the DNA profiles of all 13 CODIS STR loci would be compared.

evidence against an accused rapist. After a mistrial, the suspect was released. Three months later, he was again in court, accused of another rape. This time the judge allowed the prosecutor to present a statistical analysis of the data based on appropriate population surveys. The analysis showed that the DNA profiles prepared from semen recovered from the victim had a probability of about one in 10 billion of matching the DNA profile of the suspect purely by chance. This time the suspect was convicted. There can be no question about the value of DNA profiles in forensic cases of this type when good tissue or cell samples are obtained from the scene of the crime. If performed carefully by trained scientists and interpreted conservatively using valid population-based data on the frequencies of the polymorphisms involved, DNA profiles can provide a much-needed and powerful tool in the ongoing fight against crime.

Figure 16.14 illustrates the type of STR profiles used in forensic cases. For the sake of simplicity, the DNA profiles are shown for only 4 of the standard 13 CODIS STR loci. In practice, the profiles of all 13 loci would be compared. The DNA profile prepared from the bloodstain at the crime scene matches the DNA profile from suspect 2, but not the profile from suspect 1. Of course, these matching DNA profiles by themselves do not prove that suspect 1 committed the crime, but, if combined with additional DNA profiles and supporting evidence, they provide strong evidence that suspect 1 was at the scene of the crime. Perhaps more importantly, these profiles clearly show that the blood cells in the stain were not from suspect 1. Thus, DNA profiles have proven invaluable in reducing the frequency of wrongful convictions,

and in several cases they have exonerated prisoners in jail for crimes they did not commit.

By comparing STR profiles at all 13 CODIS loci, perhaps supplemented with mitochondrial DNA evidence, the possibility that DNA profiles from two individuals will match just by chance can virtually be eliminated. Indeed, the chance that two unrelated Caucasians in a randomly mating population will have identical DNA profiles at all 13 CODIS loci is approximately one in 5.75 trillion. Clearly, DNA profiling is a powerful tool in personal identity cases.

- DNA profiles detect and record polymorphisms in the genomes of individuals.
- DNA profiles provide strong evidence of individual identity, evidence that is extremely valuable in paternity and forensic cases.

KEY POINTS

Production of Eukaryotic Proteins in Bacteria

For decades, microorganisms have been used to produce important products for humans. We are all aware of the impact of antibiotics on human health; fewer of us are aware of their economic importance. The wholesale market value of antibiotics in the United States is over \$2 billion annually.

Human insulin, human growth hormone, and other valuable eukaryotic proteins can be produced economically in genetically engineered bacteria.

Microbes also play important roles in the production of many other materials, for example, antifungal drugs, amino acids, and vitamins. Today, because of genetic engineering, bacteria are being used in the production of important eukaryotic proteins such as human insulin, human growth hormone, and the entire family of human interferons. In addition, genetically engineered microbes are being used to synthesize valuable enzymes and other organic molecules and to provide metabolic machinery for the detoxification of pollutants and the conversion of biomass to combustible compounds.

HUMAN GROWTH HORMONE

In 1982, human insulin became the first commercial success of the new recombinant DNA technologies in the field of pharmaceuticals. Since then, several other human proteins with medicinal value have been synthesized in bacteria. Some of the first human proteins to be produced in microorganisms were blood-clotting factor VIII (lacking in individuals with one type of hemophilia), plasminogen activator (a protein that disperses blood clots), and human growth hormone (a protein deficient in certain types of dwarfism). As an example, let's examine the synthesis of human growth hormone (hGH) in *E. coli*. hGH, which is required for normal growth, is a single polypeptide chain 191 amino acids in length. In contrast to insulin, porcine and bovine pituitary growth hormones do not work in humans. Only growth hormones from humans or from closely related primates will function in humans. Thus, prior to 1985, the major source of growth hormone suitable for treatment of humans was from human cadavers.

To obtain expression in *E. coli*, the hGH coding sequence must be placed under the control of *E. coli* regulatory elements. Therefore, the hGH coding sequence was joined to the promoter and ribosome-binding sequences of the *E. coli lac* operon (a set of genes encoding proteins required for growth on the sugar lactose; see Chapter 18). To accomplish this, a *Hae*III cleavage site in the nucleotide-pair triplet specifying codon 24 of hGH was used to fuse a synthetic DNA sequence encoding amino acids 1–23 to a partial cDNA sequence encoding amino acids 24–191. This unit was then inserted into a plasmid carrying the *lac* regulatory signals and introduced into *E. coli* by transformation. The structure of the first plasmid used to produce hGH in *E. coli* is shown in **E Figure 16.15**.

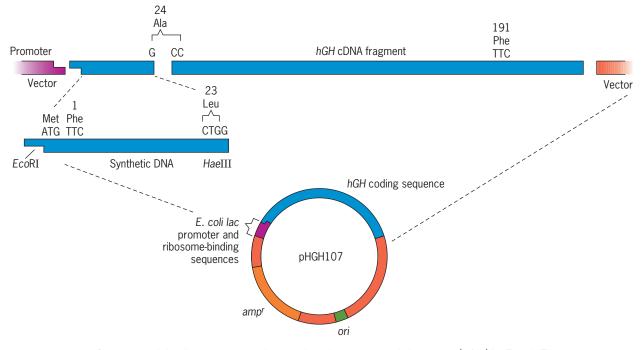


FIGURE 16.15 Structure of the first vector used to produce human growth hormone (hGH) in *E. coli*. The *amp*^r gene provides resistance to ampicillin; *ori* is the plasmid's origin of replication. The amino acids are numbered one through 191 beginning at the amino terminus.

The hGH produced in E. coli in these first experiments contained methionine at the amino terminus (the methionine specified by the ATG initiator codon). Native hGH has an amino-terminal phenylalanine: a methionine is initially present but is then enzymatically removed. E. coli also removes many amino-terminal methionine residues posttranslationally. However, the excision of the terminal methionine is sequence-dependent, and E. coli cells do not excise the amino-terminal methionine residue from hGH. Nevertheless, the hGH synthesized in *E. coli* was found to be fully active in humans despite the presence of the extra amino acid. More recently, a DNA sequence encoding a signal peptide (the amino acid sequence required for transport of proteins across membranes) has been added to an HGH gene construct similar to the one shown in Figure 16.15. With the signal sequence added, hGH is both secreted and correctly processed; that is, the methionine residue is removed with the rest of the signal peptide during the transport of the primary translation product across the membrane. This product is identical to native hGH. In 1985, hGH became the second genetically engineered pharmaceutical to be approved for use in humans by the U.S. Food and Drug Administration. Human insulin produced in E. coli had been approved for use by diabetics in 1982.

PROTEINS WITH INDUSTRIAL APPLICATIONS

Some enzymes with important industrial applications have been manufactured for many years by using microorganisms to carry out their synthesis. For example, proteases have been produced from *Bacillus licheniformis* and other bacteria. These proteases have been employed extensively as cleaning aids in detergents and in smaller amounts as meat tenderizers and as digestive aids in animal feeds. Amylases have been widely used to break down complex carbohydrates such as starch to glucose. The glucose is then converted to fructose with the enzyme glucose isomerase, and this fructose is used as a food sweetener. The amylases and glucose isomerase are all manufactured by microbiological processes.

The protein rennin is used in making cheeses. Prior to the advent of genetic engineering, rennin was extracted from the fourth stomach of cattle. Genetically engineered bacteria are now used for the commercial production of rennin. These examples are all proteins that have had important industrial applications for some time. In the future, we can expect many additional enzymes to be manufactured and used in industrial applications because of the ease of producing these proteins by means of recombinant microorganisms (or by transgenic plants and animals; see the next section).

- Valuable proteins that could be isolated from eukaryotes only in small amounts and at great expense can now be produced in large quantities in genetically engineered bacteria.
- Proteins such as human insulin and human growth hormone are valuable pharmaceuticals used to treat diabetes and pituitary dwarfism, respectively.

KEY POINTS

Transgenic Animals and Plants

Although a complete discussion of the methods used to produce transgenic animals and plants is beyond the scope of this book, let's examine a couple of the commonly used procedures, and some of the initial applications of recombinant DNA technologies in animal and plant breeding. Synthetic, modified, or other foreign genes can be introduced into animals and plants, and the resulting transgenic organisms can be used to study the functions of the genes, for example, by insertional mutagenesis, to produce novel products, or to serve as animal models for studies of inherited human diseases.

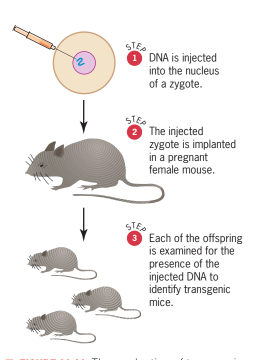
TRANSGENIC ANIMALS:

Many different animals have been modified by the introduction of foreign DNA. The mouse, however, has been studied more than any other vertebrate, and we will restrict our discussion of the techniques used to produce transgenic animals to those used with mice. There are two general methods of introducing transgenes into mouse chromosomes. One relies on the injection of DNA into fertilized eggs or embryos and the other involves the transfection of embryonic stem cells growing in culture.

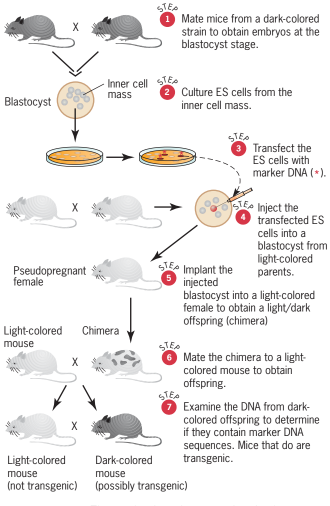
The first transgenic mice were produced by microinjection of DNA into fertilized eggs. Indeed, this procedure has been used almost exclusively to produce transgenic pigs, sheep, cattle, and other domestic animals. Prior to the microinjection of DNA, the eggs are surgically removed from the female parent and are fertilized *in vitro*. The DNA is then microinjected into the male pronucleus (the haploid nucleus contributed by the sperm, prior to nuclear fusion) of the fertilized egg through a very fine-tipped glass needle (**■** Figure 16.16). Usually, several hundred to several thousand copies of the gene of interest are injected into each egg, and multiple integrations often occur. Surprisingly, when multiple copies do integrate into the genome, they usually do so as tandem, head-to-tail arrays at a single chromosomal site. The integration of injected DNA molecules appears to occur at random sites in the genome.

Because the DNA is injected into the fertilized egg, integration of the injected DNA molecules usually occurs early during embryonic development. As a result, some germ-line cells may carry the transgene. As would be expected, the animals that develop from the injected eggs—called the G_0 generation—are almost always genetic mosaics, with some somatic cells carrying the transgene and others not carrying it. The initial (G_0) transgenic animals must be mated and G_1 progeny produced to obtain animals in which all cells carry the transgene. In most of the cases where their inheritance has been studied, the transgenes were transmitted to the progeny in a stable fashion.

The other procedure that is now widely used to produce transgenic mice relies on the injection or transfection of DNA into large populations of cultured cells that



■ **FIGURE 16.16** The production of transgenic mice by injecting DNA into eggs and implanting them into females to complete their development.



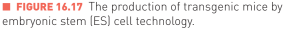




FIGURE 16.18 The transgenic mouse on the left, which carries a chimeric human growth hormone gene, is about twice the size of the control mouse on the right.

were derived from very young mouse embryos (**■** Figure 16.17). These **embryonic stem cells** (or **ES cells**) come from the inner cell mass, a group of cells found in the blastula stage of mouse embryos. Such cells can be cultured *in vitro*, transfected or injected with DNA, and then introduced into other developing mouse embryos. By chance, some of the introduced ES cells may contribute to the formation of adult tissues, so that when the mouse is born, it may consist of a mixture of two types of cells, its own and those derived from the cultured (and potentially transfected) ES cells. Such mice are called **chimeras**. If the ES cells happened to contribute to the chimera's germ line, the introduced foreign DNA has a chance of being transmitted to the next generation. Breeding a chimeric mouse may therefore establish a transgenic strain.

Transgenic mice are produced routinely in laboratories throughout the world, with thousands of transgenic strains having been created. They provide valuable tools for the study of gene expression in mammals and an excellent model system with which to test various gene-transfer vectors and methodologies for possible use in humans. In most cases, the transgenes show normal patterns of inheritance, indicating that they have been integrated into the host genome. We discuss one important application of this technology in the section Knockout Mutations in the Mouse later in this chapter.

One of the first experiments with transgenic mice showed that growth rate could be increased when rat, bovine, or human growth hormone genes were expressed in the mice (**■** Figure 16.18). This prompted animal breeders to ask whether the introduction of either (1) extra copies of the homologous (same-species) growth hormone gene or (2) copies of heterologous growth hormone genes from related species might result in domestic animals with enhanced growth rates. Thus, animal scientists introduced growth hormone transgenes into pigs, fish, and chickens with the goal of enhancing growth rate.

Another potentially important use of transgenic animals is for the production and secretion of valuable proteins in milk. Many native human proteins contain carbohydrate or lipid side groups that are added posttranslationally. Bacteria do not contain the enzymes that catalyze the addition of these moieties to nascent proteins. In such cases, recombinant bacteria cannot be used to synthesize the final product; they will synthesize the polypeptide only in its unmodified form. For this reason, some researchers have been exploring alternative methods for producing valuable human proteins, especially glycoproteins and lipoproteins. Indeed, mouse and hamster cells growing in culture are now commonly used for the production of human proteins with medicinal applications.

TRANSGENIC PLANTS: THE TI PLASMID OF AGROBACTERIUM TUMEFACIENS

Plant breeders have modified plants genetically for decades. Today, however, plant breeders can directly modify the DNA of plants, and they can quickly add genes from other species to plant genomes by recombinant DNA techniques. Indeed, transgenic plants can be produced by several different procedures. One widely used procedure, called **microprojectile bombardment**, involves shooting DNA-coated tungsten or gold particles into plant cells. Another procedure, called **electroporation**, uses a short burst of electricity to get the DNA into cells. However, the most widely used method of generating transgenic plants, at least in dicots, is *Agrobacterium tumefaciens*-mediated transformation. *A. tumefaciens* is a soil bacterium that has evolved a natural genetic engineering system; it contains a segment of DNA that is transferred from the bacterium to plant cells.

An important feature of plant cells is their **totipotency**—that is, the ability of a single cell to produce all the differentiated cells of the mature plant. Many differentiated plant cells are able to dedifferentiate to the embryonic state and subsequently to redifferentiate to new cell types. Thus, there is no separation of germ-line cells from somatic cells as in higher animals. This totipotency of plant cells is a major advantage for genetic engineering because it permits the regeneration of entire plants from individual modified somatic cells.

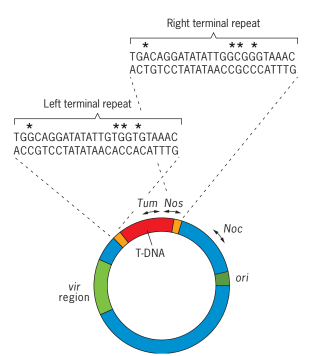
A. tumefaciens is the causative agent of crown gall disease of dicotyledonous plants. The name refers to the galls or tumors that often form at the crown (junction between the root and the stem) of infected plants. Because the crown of the plant is usually located at the soil surface, it is here that a plant is most likely to be wounded (for example, from a soil abrasion as it blows in a strong wind) and infected by a soil bacterium such as A. tumefaciens. After the infection of a wound site by A. tumefaciens, two key events occur: (1) the plant cells begin to proliferate and form tumors, and (2) they begin to synthesize an arginine derivative called an opine. The opine synthesized is usually either nopaline or octopine depending on the strain of A. tumefaciens. These opines are catabolized and used as energy sources by the infecting bacteria. A. tume*faciens* strains that induce the synthesis of nopaline can grow on nopaline, but not on octopine, and vice versa. Clearly, an interesting interrelationship has evolved between A. tumefaciens strains and their plant hosts. A. tumefaciens is able to divert the metabolic resources of the host plant to the synthesis of opines, which are of no apparent benefit to the plant but which provide sustenance to the bacterium.

The ability of *A. tumefaciens* to induce crown galls in plants is controlled by genetic information carried on a large (about 200,000 nucleotide pairs) plasmid called the **Ti plasmid** for its *t*umor-*i*nducing capacity. Two components of the Ti plasmid, the **T-DNA** and the *vir* **region**, are essential for the transformation of plant cells. During the transformation process, the T-DNA (for *T*ransferred *DNA*) is excised from the Ti plasmid, transferred to a plant cell, and integrated (covalently inserted) into the DNA of the plant cell. The available data indicate that integration of the T-DNA occurs at random chromosomal sites; moreover, in some cases, multiple T-DNA integration events occur in the same cell. In nopaline-type Ti plasmids that we will discuss, the T-DNA is a 23,000-nucleotide-pair segment that carries 13 known genes.

The structure of a typical nopaline Ti plasmid is shown in Figure 16.19. Some of the genes on the T-DNA segment of the Ti plasmid encode enzymes that catalyze the synthesis of phytohormones (the auxin indoleacetic acid and the cytokinin isopentenyl adenosine). These phytohormones are responsible for the tumorous growth of cells in crown galls. The T-DNA region is bordered by 25-nucleotide-pair imperfect repeats, one of which must be present in *cis* for T-DNA excision and transfer. The deletion of the right border sequence completely blocks the transfer of T-DNA to plant cells.

The vir (for virulence) region of the Ti plasmid contains the genes required for the T-DNA transfer process. These genes encode the DNA processing enzymes required for excision, transfer, and integration of the T-DNA segment during the transformation process. The vir genes can supply the functions needed for T-DNA transfer when located either cis or trans to the T-DNA. They are expressed at very low levels in *A. tumefaciens* cells growing in soil. However, exposure of the bacteria to wounded plant cells or exudates from plant cells induces enhanced levels of expression of the vir genes. This induction process is very slow for bacteria, taking 10 to 15 hours to reach maximum levels of expression. Phenolic compounds such as acetosyringone act as inducers of the vir genes, and transformation rates can often be increased by adding these inducers to plant cells inoculated with Agrobacterium. The transformation of plant cells by the Ti plasmid of *A. tumefaciens* occurs as illustrated in **Figure 16.20**.

Once it had been established that the T-DNA region of the Ti plasmid of *A. tumefaciens* is transferred to plant cells and becomes integrated in plant chromosomes, the potential use of *Agrobacterium* in plant genetic engineering was obvious. Foreign genes could be inserted into the T-DNA and then transferred to the plant



■ FIGURE 16.19 Structure of the nopaline Ti plasmid pTi C58, showing selected components. The Ti plasmid is 210 kb in size. Symbols used are: *ori*, origin of replication; *Tum*, genes responsible for tumor formation; *Nos*, genes involved in nopaline biosynthesis; *Noc*, genes involved in the catabolism of nopaline; *vir*, virulence genes required for T-DNA transfer. The nucleotide-pair sequences of the left and right terminal repeats are shown at the top; the asterisks mark the four base pairs that differ in the two border sequences.

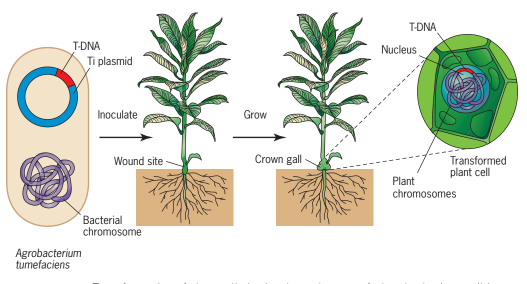


FIGURE 16.20 Transformation of plant cells by *Agrobacterium tumefaciens* harboring a wild-type Ti plasmid. Plant cells in the tumor contain the T-DNA segment of the Ti plasmid integrated into chromosomal DNA.

with the rest of the T-DNA. This procedure works very well given modifications to the Ti plasmid such as the deletion of the genes responsible for tumor formation, the addition of a selectable marker, and the addition of appropriate regulatory elements.

The *kan^r* gene from the *E. coli* transposon Tn5 has been used extensively as a selectable marker in plants; it encodes an enzyme called neomycin phosphotransferase type II (NPTII). NPTII is one of several prokaryotic enzymes that detoxify the kanamycin family of aminoglycoside antibiotics by phosphorylating them. Because the promoter sequences and transcription-termination signals are different in bacteria and plants, the native Tn5 *kan^r* gene cannot be used in plants. Instead, the NPTII coding sequence must be provided with a plant promoter (5' to the coding sequence) and plant termination and polyadenylation signals (3' to the coding sequence). Such constructions with prokaryotic coding sequences flanked by eukaryotic regulatory sequences are called **chimeric selectable marker genes**.

Regulatory sequences from several different plant genes have been used to construct chimeric marker genes. One widely used chimeric selectable marker gene contains the cauliflower mosaic virus (CaMV) 35S (transcript size) promoter, the NPTII coding sequence, and the Ti nopaline synthase (*nos*) termination sequence; this chimeric gene is usually symbolized 35S/NPTII/*nos*. The Ti vectors used to transfer genes into plants have the tumor-inducing genes of the plasmid replaced with a chimeric selectable marker gene such as 35S/NPTII/*nos*. A large number of sophisticated Ti plasmid gene-transfer vectors are now used routinely to transfer genes into plants.

The powerful new tools that permit plant and animal breeders to produce transgenic plants and animals with relative ease have a vast array of applications. In Chapter 1, we discussed the production of corn borer-resistant corn. The most widely used transgenes are those that produce herbicide resistance in agronomic crops. With the development of these and other genetically modified plants and animals have come questions about their safety. Indeed, the safety of genetically modified (GM) crops and other foods is a major concern in some countries.

KEY POINTS

- DNA sequences of interest can now be introduced into most plant and animal species.
 - The resulting transgenic organisms provide valuable resources for studies of gene function and biological processes.
 - The Ti plasmid of Agrobacterium tumefaciens is an important tool for transferring genes into plants.

Reverse Genetics: Dissecting Biological Processes by Inhibiting Gene Expression

The explosion of new information in biology during the twentieth century resulted in part because of the application of genetic approaches to the dissection of biological processes (see Chapter 13). The classical genetic approach was to identify organisms Reverse genetic approaches make use of known nucleotide sequences to devise procedures for inhibiting the expression of specific genes.

with abnormal phenotypes and to characterize the mutant genes responsible for these phenotypes. Comparative molecular studies were then performed on mutant and wild-type organisms to determine the effects of the mutations. These studies identified genes encoding products that were involved in the biological processes under investigation. In some cases, the results of these studies allowed biologists to determine the precise sequence of events or pathway by which a process occurs. The complete pathway of morphogenesis in bacteriophage T4 (see Figure 13.20) provided early documentation of the power of the mutational dissection approach.

During the last couple of decades, the nucleotide sequences of entire genomes have become available. Today, we often know the nucleotide sequence of a gene before we know its function. This knowledge has led to new approaches to the genetic dissection of biological processes, approaches collectively called **reverse genetics**. Reverse genetic approaches use the nucleotide sequences of genes to devise procedures for either isolating null mutations in them or shutting off their expression. The function of a specific gene often can be deduced by studying organisms lacking any functional product of the gene. In the remaining sections of this chapter, we examine three important reverse genetic approaches: foreign DNA insertions producing "knockout" mutations in mice, T-DNA and transposon insertions in plants, and RNA interference.

KNOCKOUT MUTATIONS IN THE MOUSE

We discussed the procedures used to generate transgenic mice in an earlier section of this chapter (see Figures 16.16 and 16.17). Normally, the transgenes are inserted into the genome at random sites. However, if the injected or transfected DNA contains a sequence homologous to a sequence in the mouse genome, it will sometimes be inserted into that sequence by homologous recombination. The insertion of this foreign DNA into a gene will disrupt or "knock out" the function of the gene just like the insertion of a transposable genetic element (see Figure 13.13). Indeed, this approach has been used to generate knockout mutations in hundreds of mouse genes.

The first step in the production of mice carrying a knockout mutation in a gene of interest is to construct a gene-targeting vector, a vector with the potential to undergo homologous recombination with one of the chromosomal copies of the gene and, in so doing, insert foreign DNA into the gene and disrupt its function. A gene (neor) that confers resistance to the antibiotic neomycin is inserted into a cloned copy of the gene of interest, splitting it into two parts and making it nonfunctional (**Figure 16.21**, step 1). The presence of the *neo^r* gene in the vector will allow neomycin to be used to eliminate cells not carrying an integrated copy of the gene-targeting vector or the *neo^r* gene. The segments of the gene retained on either side of the inserted *neo^r* gene provide sites of homology for recombination with chromosomal copies of the gene. The thymidine kinase gene (tk^{HSV}) from herpes simplex virus is inserted into the cloning vector (Figure 16.21, step 2) for subsequent use in eliminating transgenic mouse cells resulting from the random integration of the vector. The thymidine kinase from herpes simplex virus (HSV) phosphorylates the drug gancyclovir, and when this phosphorylated nucleotide-analog is incorporated into DNA, it kills the host cell. In the absence of the HSV thymidine kinase, gancyclovir is harmless to the host cell.

The next step is to transfect embryonic stem (ES) cells (from dark-colored mice) growing in culture with linear copies of the gene-targeting vector (Figure 16.21,

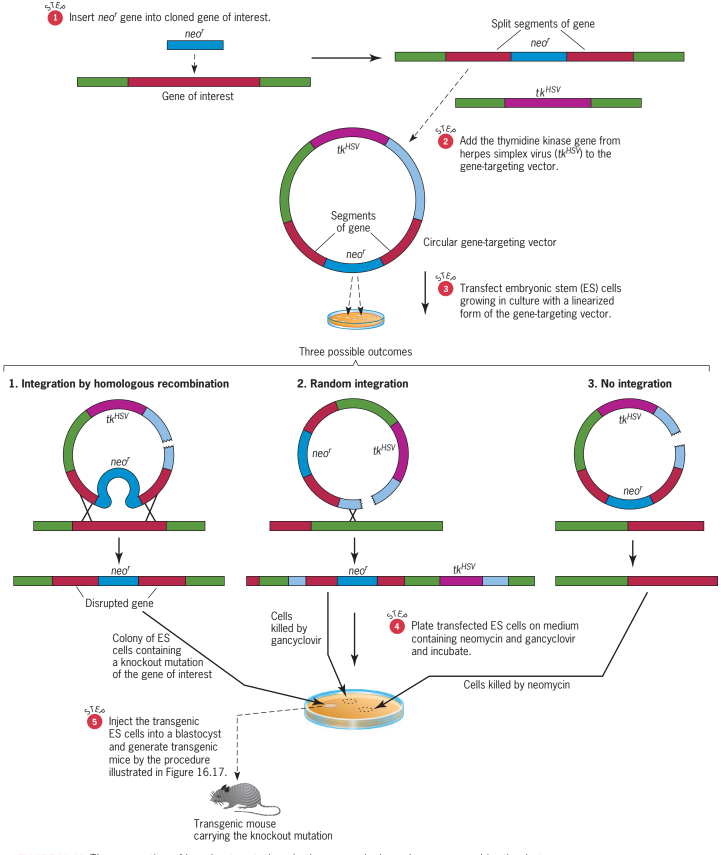


FIGURE 16.21 The generation of knockout mutations in the mouse by homologous recombination between gene-targeting vectors and chromosomal genes in transfected embryonic stem (ES) cells. The procedure used to produce transgenic mice from transgenic ES cells growing in culture is illustrated in Figure 16.17. The *neo^r* gene confers mouse cells with resistance to the antibiotic neomycin, and the *tk^{HSV}* gene makes them sensitive to the nucleotide-analog gancyclovir. See text for additional details.

step 3) and subsequently plate them on medium containing neomycin and gancyclovir (Figure 16.21, step 4). Three different events will occur in the transfected ES cells. (1) Homologous recombination may occur between the split sequences of the gene in the vector and a chromosomal copy of the gene inserting the *neo^r* gene into the chromosomal gene and disrupting its function. When this event occurs, the tk^{HSV} gene will not be inserted into the chromosome. As a result, these cells will be resistant to neomycin, but not sensitive to gancyclovir. (2) The gene-targeting vector may integrate at random into the host chromosome. When this occurs, both the *neo^r* gene and the tk^{HSV} gene will be present in the chromosome. These cells will be resistant to neomycin, but killed by gancyclovir. (3) There may be no recombination between the gene-targeting vector and the chromosome and, thus, no integration of any kind. In this case, the cells will be killed by neomycin. Thus, only the ES cells with the knockout mutation produced by the insertion of the *neo^r* gene into the gene of interest on the chromosome will be able to grow on medium containing both neomycin and gancyclovir.

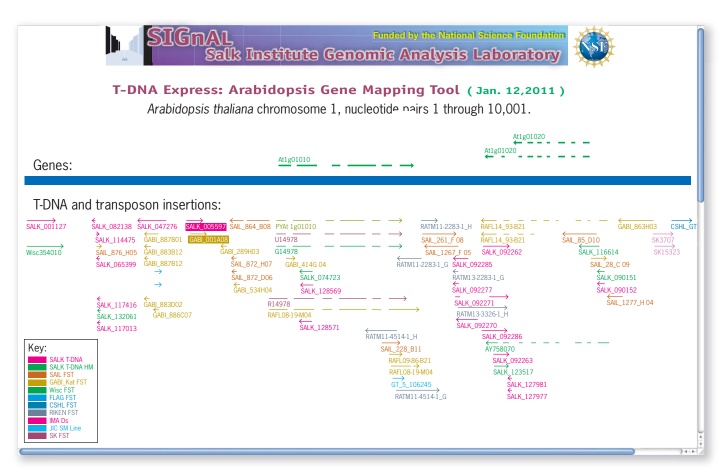
The selected ES cells containing the knockout mutation are injected into blastocysts from light-colored parents, and the blastocysts are implanted into light-colored females (see Figure 16.17). Some of the offspring will be chimeric with patches of light and dark fur. The chimeric offspring are mated with light-colored mice, and any dark-colored progeny produced by this mating are examined for the presence of the knockout mutation. In the last step, male and female offspring that carry the knockout mutation are crossed to produce progeny that are homozygous for the mutation. Depending on the function of the gene, the homozygous progeny may have normal or abnormal phenotypes. Indeed, if the product of the gene is essential early during development, homozygosity for the knockout mutation will be lethal during embryonic development. In other cases, for example, when there are related genes with overlapping or identical functions, mice that are homozygous for the knockout mutation may have wild-type phenotypes, and PCR or Southern blots will have to be performed to verify the presence of the knockout mutation.

Knockout mice have been used to study a wide range of processes in mammals including development, physiology, neurobiology, and immunology. Knockout mice have provided model systems for studies of numerous inherited human disorders from sickle-cell anemia to heart disease to many different types of cancer.

Because of the value of knockout mice for studies of processes related to human health, the National Institutes of Health initiated the Knockout Mouse Project in 2006 with the goal of producing knockout mutations in as many mouse genes as possible. This project has subsequently been expanded to the North American Conditional Mouse Mutagenesis Project and is working together with the European Conditional Mouse Mutagenesis Project to produce at least one knockout mutation in each of the over 20,000 genes in the mouse genome. All of the knockout strains produced by this collaborative effort are being made available to researchers throughout the world.

T-DNA AND TRANSPOSON INSERTIONS

In a preceding section of this chapter, we discussed how the T-DNA segment of the Ti plasmid of *Agrobacterium tumefaciens* is transferred into plant cells and inserted into the chromosomes of the plant (see Figure 16.20). When the T-DNA inserts into a gene, it disrupts the function of the gene. Transposons are genetic elements that have the ability to move from one location in the genome to another location (Chapter 17). Like the T-DNA of the Ti plasmid, a transposon will disrupt the function of a gene into which it inserts (see Figure 13.13). Thus, T-DNAs and transposons provide powerful tools for reverse genetic analysis. In both cases, the genetic element is used to perform insertional mutagenesis—the induction of null mutations (often called "knockout" mutations) by the insertion of foreign DNAs into genes. Insertional mutagenesis is basically the same whether performed with the Ti plasmid or a transposon. Thus, we will illustrate the use of insertional mutagenesis for reverse genetics by discussing the utilization of T-DNA insertions to dissect gene function in the plant *Arabidopsis thaliana*.



■ FIGURE 16.22 Map of T-DNA and transposon insertions in the 10-kb region at the tip of chromosome 1 in *Arabidopsis*. The positions of the *f*lanking sequence *t*ags (FSTs) are shown as arrows below the chromosome (dark blue box). The data shown are from the SIGnAL (Salk *I*nstitute *G*enomic *A*nalysis *L*aboratory) web site, http://signal.salk.edu/cgi-bin/tdnaexpress. The two genes (At1g01010 and At1g01020) in this region of chromosome 1 have unknown functions. The T-DNA and transposon insertion lines are from the Salk Institute (Salk T-DNA), the Syngenta *Arabidopsis* Insertion *Library* (SAIL), the German collection (GABI-Kat), the University of Wisconsin collection (Wisc), the French collection (FLAG), the *Cold Spring Harbor Laboratory* (CSHL) collection, the Riken BioResource Center in Japan (RIKEN), the Institute of Molecular Agrobiology (IMA) in Singapore, the John Innes Centre (JIC), and the Saskatoon (SK) collection.

When T-DNA is transferred from *A. tumefaciens* to plant cells, it integrates into essentially all components of the genome; that is, T-DNAs are found scattered along each of the five chromosomes of *Arabidopsis*. Therefore, if a large enough population of transformed *Arabidopsis* plants is examined, it should be possible to identify T-DNA insertions in all of the approximately 26,000 genes of this species.

Indeed, hundreds of thousands of T-DNA insertions have been mapped throughout the *Arabidopsis* genome, and seed stocks containing these insertions are available upon request from the *Arabidopsis B*iological *R*esource *C*enter (ABRC) at Ohio State University. In addition, seeds of T-DNA and transposon insertion lines characterized at the *V*ersailles *G*enomic *R*esource *C*enter (VGRC) in France, the *N*ottingham *Arabidopsis* Stock *C*entre (NASC) in Germany, and the Riken BioResource Center in Japan are also available to the *Arabidopsis* research community. Researchers at the Salk Institute in La Jolla, California, have integrated their map of T-DNA insertions with the maps of T-DNA and transposon insertions characterized by other research groups. Their sequence-based map of these insertions is available at http://signal.salk. edu/cgi-bin/tdnaexpress; an abbreviated version of their map of the tip of chromosome 1 is shown in **Figure 16.22**. Therefore, if someone is interested in the function of a particular *Arabidopsis* gene, she or he can search the Salk web site for T-DNA and transposon insertions in that gene; once the insertions have been identified, seeds carrying the desired insertion mutations can be ordered online. These large collections of insertional mutations have proven to be invaluable resources for studies of gene function in this model plant.

RNA INTERFERENCE

Although its effects were first observed in petunias a few years earlier, the discovery of the third reverse genetics approach—**RNA** interference **(RNAi)**—is usually credited to the work of Andrew Fire, Craig Mello, and colleagues, published in 1998. Indeed, Fire and Mello shared the 2006 Noble Prize in Physiology or Medicine in recognition of this work. When they injected double-stranded RNA (dsRNA) into *C. elegans*, it "interfered with" (or shut off) the expression of genes containing the same nucleotide sequence. During the last decade, RNAi has moved to the cutting edge in molecular biology. We now know that double-stranded RNA (dsRNA) plays important roles in preventing viral infections, in combating the expansion of populations of transposable genetic elements, and in regulating gene expression (see Chapter 19). Indeed, RNAi is not only at the cutting edge of molecular biology, but it has great potential for use in the fight against human diseases. In this chapter, however, we will focus on the use of RNAi as a tool for reverse genetics, a tool with which to study gene function and dissect biological processes.

RNAi is used extensively to silence genes-to turn down or turn off their expression—in C. elegans, D. melanogaster, and many plants. It has potential uses in all species, including humans. RNAi can be carried out in several different ways. The common feature in all RNAi experiments is the presence of dsRNA carrying at least a portion of the nucleotide sequence of the gene that one wishes to silence in the organism or cells under investigation. Two different approaches are frequently used to achieve this goal. In one approach, the dsRNA is synthesized in vitro and microinjected into the organism (**Figure 16.23a**). In the second approach, a geneexpression cassette is constructed that carries two copies of at least a portion o the gene of interest in inverse orientations and is introduced into the organism by transformation or transfection (**Figure 16.23b**). When the introduced transgene is transcribed, it produces an RNA molecule that is self-complementary and forms a partially double-stranded stem-and-loop, or "hairpin" structure. In both cases, the dsRNAs stimulate RNA-induced gene silencing. The dsRNAs are ultimately bound by the RNA-induced silencing complex (RISC) and the corresponding mRNAs synthesized from endogenous genes are either degraded or translationally repressed (see Chapter 19 for details).

RNAi is quite easy to perform in *C. elegans*; these little worms can be microinjected with the dsRNA, soaked in media containing the dsRNA, or fed bacteria synthesizing the dsRNA of interest. All three procedures lead to effective gene silencing in *C. elegans*.

The sequence of 99 percent of the genome of *C. elegans* was published in December 1998. Within two years, collaborative research groups in Great Britain, Germany, Switzerland, and Canada had used RNAi to systematically silence more than 90 percent of the 2769 predicted genes on chromosome I and more than 96 percent of the 2300 predicted genes on chromosome III of *C. elegans*. These studies provided new information about the functions of over 400 genes. Clearly, RNAi is a powerful tool with which to dissect biological processes. RNAi makes use of natural pathways involved in the regulation of gene expression. There are hundreds of genes in plant and animal genomes that encode microRNAs, which form dsRNAs *in vivo*. At present, we know the regulatory functions of only a few of these microRNAs (see Chapter 19); however, the functions of the rest of the microRNAs are the subject of many ongoing investigations.

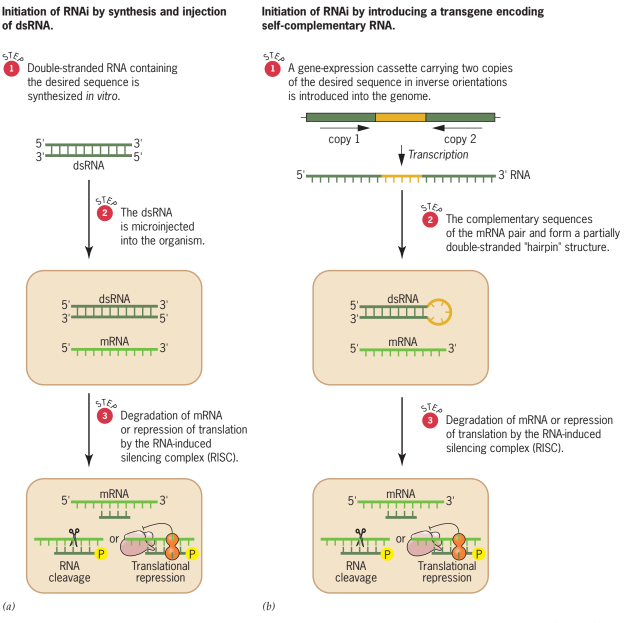
Can RNAi be used to inhibit the reproduction of viruses such as the human immunodeficiency virus (HIV) or to downregulate the expression of oncogenes

Solve It!

How Might RNA Interference Be Used to Treat Burkitt's Lymphoma?

Burkitt's lymphoma is a white blood cell cancer that occurs when a translocation moves the *c-myc* oncogene (cancercausing gene) on chromosome 8 close to one of the three immunoglobin (antibody chain) gene clusters on chromosomes 2, 14, and 22 (see Chapter 21). The resulting juxtaposition of *c-myc* next to the cluster of the highly expressed antibody genes causes its overexpression, which, in turn, leads to uncontrolled cell division, that is, cancer. How might RNA interference be used to suppress this cancer? Design an experimental approach using RNA interference to treat Burkitt's lymphoma. Explain the rationale behind your proposal and how to evaluate its potential effectiveness.

To see the solution to this problem, visit the Student Companion site.



■ FIGURE 16.23 Two procedures for initiating RNAi with double-stranded RNA (dsRNA). (a) A dsRNA molecule containing a portion of the nucleotide sequence of the gene to be silenced is synthesized *in vitro* and injected into the organism. (b) A gene-expression cassette containing two copies of a segment of the gene in inverse orientations is constructed and introduced into the organism under investigation. The self-complementary RNA transcript forms a partially double-stranded RNA hairpin. In both cases, the dsRNA initiates silencing of the targeted gene via the RNA-induced silencing complex (RISC) pathway, which results in the degradation of the targeted mRNA or repression of its translation (see Chapter 19 for details).

(cancer-causing genes)? We don't know the answer to these questions. However, we do know that the business world is excited about the potential therapeutic applications of RNAi. Not only are the big pharmaceutical firms investing heavily in RNAi technology, but a plethora of start-up companies have been formed specifically to exploit RNAi for commercial goals. Whether or not the RNAi technologies will live up to expectations remains to be seen. To test your understanding of RNAi, try Solve It: How Might RNA Interference Be Used to Treat Burkitt's Lymphoma?

- Reverse genetic approaches use known nucleotide sequences to devise procedures for isolating null mutations of genes or inhibiting gene expression.
- Knockout mutations of genes in the mouse can be produced by inserting foreign DNAs into chromosomal genes by homologous recombination.
- T-DNA or transposon insertions provide a source of null mutations of genes.
- RNA interference—blocking gene expression with double-stranded RNA—can be used to dissect biological processes by inhibiting the functions of specific genes.

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. How were restriction fragment-length polymorphisms (RFLPs) used in the search for the mutant gene that causes Huntington's disease (HD)?
- **Answer:** The HD research teams screened members of two large families for linkage between RFLPs and the *HTT* (*huntingtin*) gene. They found an RFLP on chromosome 4 that was tightly linked to the *HTT* gene (4 percent recombination).
- 2. Once tight linkage had been established between the *HTT* gene and the RFLP on chromosome 4, what was the research teams' next step in their search for the mutant *HTT* gene?
- **Answer:** They next prepared a detailed restriction map of this region (spanning 500 kb) of chromosome 4 (see Figure 16.1).
- **3.** How did the research teams identify candidate genes within the mapped region of chromosome 4?
- **Answer:** They used cDNA clones to identify the coding segments or exons of genes in the region and to screen genomic libraries for clones overlapping the exons. The sequences of the cDNAs and genomic DNAs were then compared to deduce the exon-intron structures of genes in the mapped region.
- 4. How did the HD research teams determine which of the candidate genes was the *HTT* gene?
- **Answer:** They sequenced the candidate genes of individuals with HD and nonaffected members of their families and

looked for structural abnormalities in the genes of affected individuals. Their results showed that one gene, now called the *huntingtin* (*HTT*) gene, contains a trinucleotide repeat, (CAG)_n, which was present in 11 to 34 copies in nonaffected individuals and in 42 to over 100 copies in affected individuals. They identified this expanded trinucleotide repeat in the *huntingtin* alleles of affected members of 72 different families, leaving little doubt that *huntingtin* is the gene responsible for HD.

- 5. Of what value is knowledge of the nucleotide sequence of the *buntingtin* gene to genetic counselors?
- **Answer:** Knowing the nucleotide sequence of the *huntingtin* gene has provided counselors with a simple and accurate diagnostic test for the presence of mutant alleles of the gene. Oligonucleotide primers to sequences flanking the trinucleotide repeat region of the gene can be used to amplify this segment of the gene, and the number of trinucleotide repeats can be determined by polyacrylamide gel electrophoresis (see Figure 16.2). As a result, individuals at risk of transmitting the mutant gene can be tested for its presence before starting a family. If the mutant gene is present in one of the parents, fetal cells or even a single cell from an eight-cell pre-implantation embryo can be tested for its presence. Thus, genetic counselors are able to provide families at risk for the disorder with accurate information regarding the presence of the gene in individuals planning families, in fetal cells, and even in eight-cell embryos.

Testing Your Knowledge

Integrate Different Concepts and Techniques

1. Spinocerebellar ataxia (type 1) is a progressive neurological disease with onset typically occurring between ages 30 and 50. The neurodegeneration results from the selective loss of specific neurons. Although it is not understood why selective neuronal death occurs, it is known that the disease is caused by the expansion of a CAG trinucleotide repeat, with normal alleles containing about 28 copies and mutant alleles harboring 43 to 81 copies of the trinucleotide. Given the nucleotide sequences on either side of the repeat region, how would you test for the presence of the expanded trinucleotide repeat region responsible for type 1 spinocerebellar ataxia?

Answer: The DNA test for spinocerebellar ataxia (type 1) would be similar to the test for the *huntingtin* allele described in Figure 16.2. You would first synthesize PCR primers

KEY POINTS

corresponding to DNA sequences on either side of the CAG repeat region. These primers would be used to amplify the desired CAG repeat region from genomic DNA of the individual being tested by PCR. Then, the sizes of the trinucleotide repeat regions would be determined by measuring the sizes of the PCR products by gel electrophoresis. Any gene with fewer than 30 copies of the CAG repeat would be considered a normal allele, whereas the presence of a gene with 40 or more copies of the trinucleotide would be diagnostic of the mutant alleles that cause spinocerebellar ataxia.

2. Assume that you have just performed the DNA test for spinocerebellar ataxia on a 25-year-old woman whose mother died from the disease. The results came back positive for the ataxia mutation. The woman and her husband long for their own biological children, but do not want to risk

transmitting the defective gene to any of these children. What are their options?

Answer: Their options will depend on their religious and moral convictions. One possibility involves the use of amniocentesis or chorionic biopsy to obtain fetal cells early in pregnancy, performing the DNA test for the expanded trinucleotide region responsible for spinocerebellar ataxia on the fetal cells, and allowing the pregnancy to continue only if the defective gene is not present. Another possibility is the use of *in vitro* fertilization. The ataxia DNA test is then performed on a cell from an eight-cell pre-embryo, and the pre-embryo is implanted only if the test for the defective ataxia gene is negative. A third option may become available in the future, namely, an effective method of treating the disease prior to the onset of neurodegeneration, perhaps by gene-replacement therapy.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **16.1** What are CpG islands? Of what value are CpG islands in positional cloning of human genes?
- **16.2** Why is the mutant gene that causes Huntington's disease called *huntingtin?* Why might this gene be renamed in the future?
- **16.3** How was the nucleotide sequence of the *CF* gene used to obtain information about the structure and function of its gene product?
- **16.4** How might the characterization of the *CF* gene and its product lead to the treatment of cystic fibrosis by somatic-cell gene therapy? What obstacles must be overcome before cystic fibrosis can be treated successfully by gene therapy?
- 16.5 Myotonic dystrophy (MD), occurring in about 1 of 8000 individuals, is the most common form of muscular dystrophy in adults. The disease, which is characterized by progressive muscle degeneration, is caused by a dominant mutant gene that contains an expanded CAG repeat region. Wild-type alleles of the *MD* gene contain 5 to 30 copies of the trinucleotide. Mutant *MD* alleles contain 50 to over 2000 copies of the CAG repeat. The complete nucleotide sequence of the *MD* gene is available. Design a diagnostic test for the mutant gene responsible for myotonic dystrophy that can be carried out using genomic DNA from newborns, fetal cells obtained by amniocentesis, and single cells from eight-cell pre-embryos produced by *in vitro* fertilization.
- **16.6** In humans, the absence of an enzyme called purine nucleoside phosphorylase (PNP) results in a severe T-cell immunodeficiency similar to that of severe combined immunodeficiency disease (SCID). PNP deficiency exhibits an autosomal recessive pattern of inheritance,

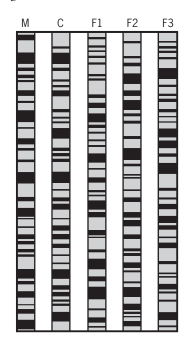
and the gene encoding human PNP has been cloned and sequenced. Would PNP deficiency be a good candidate for treatment by gene therapy? Design a procedure for treatment of PNP deficiency by somatic-cell gene therapy.

- **16.7** Human proteins can now be produced in bacteria such as *E. coli.* However, one cannot simply introduce a human gene into *E. coli* and expect it to be expressed. What steps must be taken to construct an *E. coli* strain that will produce a mammalian protein such as human growth hormone?
- **16.8** You have constructed a synthetic gene that encodes an enzyme that degrades the herbicide glyphosate. You wish to introduce your synthetic gene into *Arabidopsis* plants and test the transgenic plants for resistance to glyphosate. How could you produce a transgenic *Arabidopsis* plant harboring your synthetic gene by *A. tumefaciens*-mediated transformation?
- **16.9** A human STR locus contains a tandem repeat $(TAGA)_n$, where *n* may be between 5 and 15. How many alleles of this locus would you expect to find in the human population?
- 16.10 A group of bodies are found buried in a forest. The police suspect that they may include the missing Jones family (two parents and two children). They extract DNA from bones and examine the DNA profiles of STR loci *A* and *B*, which are known to contain tandem repeats of variable length. They also analyze the DNA profiles of two other men. The results are shown in the following table where the numbers indicate the number of copies of the tandem repeat in a particular allele; for example, male 1 has one allele with 8 and another allele with 9 copies of a tandem repeat in locus *A*.

Locus A	Locus B
8/9	5/7
6/8	5/5
7/10	7/7
8/8	3/5
7/8	5/7
8/8	3/7
	8/9 6/8 7/10 8/8 7/8

Could the woman have been the mother of both children? Why or why not? Which man, if any, could have been the father of child 1?

- **16.11** DNA profiles have played central roles in many rape and murder trials. What is a DNA profile? What roles do DNA profiles play in these forensic cases? In some cases, geneticists have been concerned that DNA profile data were being used improperly. What were some of their concerns, and how can these concerns be properly addressed?
- **16.12 (co)** The DNA profiles shown in this problem were prepared using genomic DNA from blood cells obtained from a woman, her daughter, and three men who all claim to be the girl's father.



Based on the DNA profiles, what can be determined about paternity in this case?

16.13 Most forensic experts agree that profiles of DNA from blood samples obtained at crime scenes and on personal items can provide convincing evidence for murder convictions. However, the defense attorneys sometimes argue successfully that sloppiness in handling blood samples results in contamination of the samples. What problems would contamination of blood samples present in the

interpretation of DNA profiles? Would you expect such errors to lead to the conviction of an innocent person or the acquittal of a guilty person?

- **16.14** The Ti plasmid contains a region referred to as T-DNA. Why is this region called T-DNA, and what is its significance?
- **16.15** The generation of transgenic plants using *A. tumefaciens*mediated transformation often results in multiple sites of insertion. These sites frequently vary in the level of transgene expression. What approaches could you use to determine whether or not transgenic plants carry more than one transgene and, if so, where the transgenes are inserted into chromosomes?
- **16.16** "Disarmed" retroviral vectors can be used to introduce genes into higher animals, including humans. What advantages do retroviral vectors have over other kinds of gene-transfer vectors? What disadvantages?
- **16.17** Transgenic mice are now routinely produced and studied in research laboratories throughout the world. How are transgenic mice produced? What kinds of information can be obtained from studies performed on transgenic mice? Does this information have any importance to the practice of medicine? If so, what?
- **16.18** Two men claim to be the father of baby Joyce Doe. Joyce's mother had her CODIS STR DNA profile analyzed and was homozygous for allele 8 at the TPOX locus (allele 8 contains eight repeats of the GAAT sequence at this polymorphic locus). Baby Joyce is heterozygous for alleles 8 and 11 at this locus. In an attempt to resolve the disputed paternity, the two men were tested for their STR DNA profiles at the TPOX locus on chromosome 2. Putative father 1 was heterozygous for alleles 8 and 11 at this locus. Can these results resolve this case of disputed paternity? If so, who is the biological father? If not, why not?
- **16.19** Many valuable human proteins contain carbohydrate or lipid components that are added posttranslationally. Bacteria do not contain the enzymes needed to add these components to primary translation products. How might these proteins be produced using transgenic animals?
- **16.20** Richard Meagher and coworkers have cloned a family of 10 genes that encode actins (a major component of the cytoskeleton) in *Arabidopsis thaliana*. The 10 actin gene products are similar, often differing by just a few amino acids. Thus, the coding sequences of the 10 genes are also very similar, so that the coding region of one gene will cross-hybridize with the coding regions of the other nine genes. In contrast, the noncoding regions of the 10 genes are quite divergent. Meagher has hypothesized that the 10 actin genes exhibit quite different temporal and spatial patterns of expression. You have been hired by Meagher to test this hypothesis. Design experiments that will allow you to determine the temporal and spatial pattern of expression of each of the 10 actin genes in *Arabidopsis*.

- **16.21** The first transgenic mice resulted from microinjecting fertilized eggs with vector DNA similar to that diagrammed in Figure 16.15 except that it contained a promoter for the mammalian metallothionein gene linked to the *HGH* gene. The resulting transgenic mice showed elevated levels of HGH in tissues of organs other than the pituitary gland, for example, in heart, lung, and liver, and the pituitary gland underwent atrophy. How might the production of HGH in transgenic animals be better regulated, with expression restricted to the pituitary gland?
- **16.22** How do the reverse genetic approaches used to dissect biological processes differ from classical genetic approaches?
- **16.23** How can RNAi gene silencing be used to determine the function of genes?
- **16.24** How do insertional mutagenesis approaches differ from other reverse genetic approaches?
- **16.25** Insertional mutagenesis is a powerful tool in both plants and animals. However, when performing large-scale insertional mutagenesis, what major advantage do plants have over animals?
- **16.26** We discussed the unfortunate effects of insertional mutagenesis in the four boys who developed leukemia after treatment of X-linked severe combined immunodeficiency disease by gene therapy. How might this consequence of gene therapy be avoided in the future? Do you believe that the use of somatic-cell gene therapy to treat human diseases can ever be made 100 percent risk free? Why? Why not?

16.27 One strand of a gene in *Arabidopsis thaliana* has the following nucleotide sequence:

The function of this gene is still uncertain. (a) How might insertional mutagenesis be used to investigate its function? (b) Design an experiment using RNA interference to probe the function(s) of the gene.

16.28 Let's check the Salk Institute's Genome Analysis Laboratory web site (http://signal.salk.edu/cgi-bin/tdnaexpress) to see if any of their T-DNA lines have insertions in the gene shown in the previous question. At the SIGnAL web site, scroll down to "Blast" and paste or type the sequence in the box. The resulting map will show the location of mapped T-DNA insertions relative to the location of the gene (green rectangle at the top). The blue arrows at the top right will let you focus on just the short region containing the gene or relatively long regions of chromosome 4 of *Arabidopsis*. Are there any T-DNA insertions in the gene in question? near the gene?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

Muscular dystrophy is a group of human disorders that involve progressive muscle weakness and loss of muscle cells.

- **1.** How many different types of inherited muscular dystrophy have been characterized in humans to date?
- **2.** What are the chromosomal locations of the defective genes responsible for the different forms of muscular dystrophy?
- **3.** Duchenne and Becker muscular dystrophy both result from mutations in a gene on the X chromosome that encodes a protein called dystrophin. How do these two types of muscular dystrophy differ?
- **4.** The dystrophin (*DMD*) gene has been cloned and sequenced. What are the unique features of this gene and the protein that it encodes? What obstacles do they present for the treatment of Duchenne and Becker muscular dystrophy by gene therapy?

5. Are gene tests for Duchenne muscular dystrophy available? How are they performed?

Hint: At the NCBI web site, click on OMIM (Online Mendelian Inheritance in Man) and search using "muscular dystrophy" as the query. From the resulting list of muscular dystrophies, note the chromosomal locations of the genes responsible for the various forms of the disorder to estimate the number of different genes involved. For information on Duchenne and Becker muscular dystrophy, click on #310200, "Muscular Dystrophy, Duchenne Type," and #300376, "Muscular Dystrophy, Becker Type." For information on gene tests, click on "Gene Tests" and follow the links provided to the different types of tests available.

Transposable Genetic Elements



CHAPTER OUTLINE

- Transposable Elements: An Overview
- Transposable Elements in Bacteria
- Cut-and-Paste Transposons in Eukaryotes
- Retroviruses and Retrotransposons
- Transposable Elements in Humans
- The Genetic and Evolutionary Significance of Transposable Elements

Maize: A Staple Crop with a Cultural Heritage

Maize is one of the world's most important crop plants. The cultivation of maize began at least 5000 years ago in Central America. By the time Christopher Columbus arrived in the New World, maize cultivation had spread north to Canada and south to Argentina. The native peoples of North and South America developed many different varieties of maize, each adapted to particular conditions. They developed varieties that had colorful kernels—red, blue, yellow, white, and purple—and associated each color with a special aesthetic or religious value. To the peoples of the American Southwest, for example, blue maize is considered sacred, and each of the four cardinal directions of the compass is represented by a particular maize color. Some groups consider kernels with stripes and spots to be signs of strength and vigor.

The colorful patterns that we see on maize ears also have an important scientific significance. Modern research has shown that the stripes and spots on maize kernels are the result of a genetic phenomenon called *transposition*. Within the maize genome indeed, within the genomes of most organisms—geneticists have found DNA sequences that can move from one position to another. These *transposable elements*—or, more simply, *transposons*—



Color variation among kernels of maize. Studies of the genetic basis of this variation led to the discovery of transposable elements.

constitute an appreciable fraction of the genome. In maize, for example, they account for 85 percent of all the DNA. When transposable elements move from one location to another, they may break chromosomes or mutate genes. Thus, these elements have a profound genetic significance.

Transposable Elements: An Overview

Transposable elements—transposons—are found in the genomes of many kinds of organisms; they are structurally and functionally diverse. Many different kinds of transposable elements have been identified in an assortment of organisms, including bacteria, fungi, protists, plants, and animals. These elements are prominent components of genomes—for example, more than 40 percent of the human genome—and they clearly

have roles in shaping the structure of chromosomes and in modulating the expression of genes. In this chapter we explore the structural and behavioral diversity of different types of transposable elements, and we investigate their genetic and evolutionary significance.

Although each kind of transposable element has its own special characteristics, most can be classified into one of three categories based on how they transpose (Table 17.1). In the first category, transposition is accomplished by excising an element from its position in a chromosome and inserting it into another position. The excision and insertion events are catalyzed by an enzyme called the transposase, which is usually encoded by the element itself. Geneticists refer to this mechanism as *cut-and-paste transposition* because the element is physically cut out of one site in a chromosome and pasted into a new site, which may even be on a different chromosome. We will refer to the elements in this category as **cut-and-paste transposons**.

In the second category, transposition is accomplished through a process that involves replication of the transposable element's DNA. A transposase encoded by the element mediates an interaction between the element and a potential insertion site. During this interaction, the element is replicated, and one copy of it is inserted at the new site; one copy also remains at the original site. Because there is a net gain of one copy of the element, geneticists refer to this mechanism as *replicative transposition*. We will refer to the elements in the category as **replicative transposons**.

In the third category, transposition is accomplished through a process that involves the insertion of copies of an element that were synthesized from the element's RNA. An enzyme called reverse transcriptase uses the element's RNA as a

TABLE 17.1

Categorization of Transposable Elements by Transposition Mechanism

Category	Examples	Host Organism
I. Cut-and-paste transposons	IS elements (e.g., IS <i>50</i>) Composite transposons	Bacteria
	(e.g., Tn <i>5</i>)	Bacteria
	<i>Ac/Ds</i> elements	Maize
	P elements hobo elements	Drosophila Drosophila
	piggyBac	<i>Drosophila</i> moth
	Sleeping Beauty	salmon
II. Replicative transposons	Tn3 elements	Bacteria
 III. Retrotransposons A. Retroviruslike elements (also called long terminal repeat, or LTR, retrotransposons) 	Ty1 copia gypsy	Yeast Drosophila Drosophila
B. Retroposons	<i>F</i> , <i>G</i> , and / elements Telomeric retroposons LINEs (e.g., <i>L1</i>) SINEs (e.g., <i>Alu</i>)	Drosophila Drosophila Humans Humans

template to synthesize DNA molecules, which are then inserted into new chromosomal sites. Because this mechanism reverses the usual direction in which genetic information flows in cells—that is, it flows from RNA to DNA instead of from DNA to RNA—geneticists refer to it as *retrotransposition*. We will refer to the elements in this category as **retrotransposons**. Some of the elements that transpose in this way are related to a special group of viruses that utilize reverse transcriptase—the retroviruses; consequently, they are called *retroviruslike elements*. Other elements that engage in retrotransposition are simply called *retroposons*.

We will encounter many different transposable elements in this chapter, each with its own peculiar story. Table 17.1 categorizes these elements according to their transposition mechanisms. The cut-and-paste transposons are found in both prokaryotes and eukaryotes. The replicative transposons are found only in prokaryotes, and the retrotransposons are found only in eukaryotes.

- A cut-and-paste transposon is excised from one genomic position and inserted into another by an enzyme, the transposase, which is usually encoded by the transposon itself.
- A replicative transposon is copied during the process of transposition.
- A retrotransposon produces RNA molecules that are reverse-transcribed into DNA molecules; these DNA molecules are subsequently inserted into new genomic positions.

Transposable Elements in Bacteria

Although transposable elements were originally discovered in eukaryotes, bacterial transposons were the first to be studied at the molecular level. There are three main types: the insertion sequences, or IS elements, the composite transposons, and the

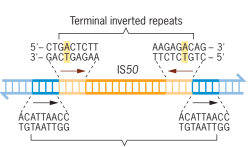
Tn3-like elements. These three types of transposons differ in size and structure. The IS elements are the simplest, containing only genes that encode proteins involved in transposition. The composite transposons and Tn3-like elements are more complex, containing some genes that encode products unrelated to the transposition process.

IS ELEMENTS

The simplest bacterial transposons are the insertion sequences, or IS elements, so named because they can insert at many different sites in bacterial chromosomes and plasmids. IS elements were first detected in certain lac^- mutations of *E. coli*. These mutations had the unusual property of reverting to wild-type at a high rate. Molecular analyses revealed that these unstable mutations possessed extra DNA in or near the *lac* genes. When DNA from the wild-type revertants of these mutations was compared with that from the mutations themselves, it was found that the extra DNA had been lost. Thus, these genetically unstable mutations were caused by DNA sequences that had inserted into *E. coli* genes, and reversion to wild-type was caused by excision of these sequences. Similar insertion sequences have been found in many other bacterial species.

IS elements are compactly organized. Typically, they consist of fewer than 2500 nucleotide pairs and contain only genes whose products are involved in promoting or regulating transposition. Many distinct types of IS elements have been identified. The smallest, IS1, is 768 nucleotide pairs long. Each type of IS element is demarcated by short identical, or nearly identical, sequences at its ends (**■** Figure 17.1). Because these terminal sequences are always in inverted orientation with respect to each other, they are called terminal inverted repeats. Their lengths range from 9 to 40 nucleotide pairs. Terminal inverted repeats are characteristic of most—but not all—types

Bacterial transposons move within and between chromosomes and plasmids.



Target site duplication

■ FIGURE 17.1 Structure of an inserted IS50 element showing its terminal inverted repeats and target site duplication. The terminal inverted repeats are imperfect because the fourth nucleotide pair (highlighted) from each end is different.

KEY POINTS

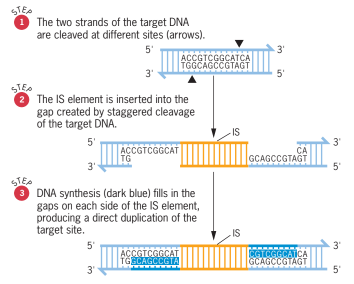


FIGURE 17.2 Production of target site duplications by the insertion of an IS element.

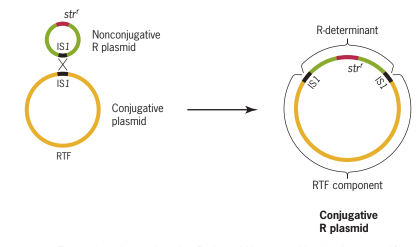
of transposons. When nucleotides in these repeats are mutated, the transposon usually loses its ability to move. These mutations therefore demonstrate that terminal inverted repeats play an important role in the transposition process.

IS elements usually encode a protein, the **transposase**, that is needed for transposition. The transposase binds at or near the ends of the element and then cuts both strands of the DNA. This cleavage excises the element from the chromosome or plasmid, so that it can be inserted at a new position in the same or a different DNA molecule. IS elements are therefore cut-and-paste transposons. When IS elements insert into chromosomes or plasmids, they create a duplication of part of the DNA sequence at the site of the insertion. One copy of the duplication is located on each side of the element. These short (2 to 13 nucleotide pairs), directly repeated sequences, called **target site duplications**, arise from staggered cleavage of the double-stranded DNA molecule (**Transposition**).

A bacterial chromosome may contain several copies of a particular type of IS element. For example, 6 to 10 copies of IS1 are found in the *E. coli* chromosome. Plasmids may also contain IS elements. The F plasmid, for example, typically has at least two different IS

elements, IS2 and IS3. When a particular IS element resides in two different DNA molecules, it creates the opportunity for homologous recombination between them. For instance, an IS element in the F plasmid may pair and recombine with the same kind of IS element in the *E. coli* chromosome. Both the *E. coli* chromosome and the F plasmid are circular DNA molecules. When an IS element mediates recombination between these molecules, the smaller plasmid is integrated into the larger chromosome, creating a single circular molecule. Such integration events produce Hfr strains capable of transferring their chromosomes during conjugation. These strains vary in the integration site of the F plasmid because the IS elements that mediate recombination occupy different chromosomal positions in different *E. coli* strains—a result of their ability to transpose.

IS elements may also mediate recombination between two different plasmids. For example, consider the situation diagrammed in **Figure 17.3**, where a plasmid that carries a gene for resistance to the antibiotic streptomycin (*str^r*) recombines with a plasmid that can be transferred between cells during conjugation (a conjugative plasmid). The recombination event is mediated by IS1 elements present in both plasmids, and it creates a large plasmid that has both the *str^r* gene and the capability to





be transferred during conjugation. Such plasmids have a medical significance because they allow the antibiotic resistance gene to spread horizontally between individuals in a bacterial population. Eventually, all or nearly all the bacterial cells acquire the resistance gene, and the antibiotic is no longer useful as a treatment for whatever infections the cells may cause.

Plasmids that transfer genes for antibiotic resistance between cells are called **conjugative R plasmids**. These plasmids have two components: the *resistance transfer factor*, or *RTF*, which contains the genes needed for conjugative transfer between cells, and the *R-determinant*, which contains the gene or genes for antibiotic resistance. Conjugative R plasmids can be transferred rapidly between cells in a bacterial population, even between quite dissimilar cell types—for example, between a coccus and a bacillus. Thus, once they have evolved in a part of the microbial kingdom, they can spread to other parts with relative ease.

Some conjugative R plasmids carry several different antibiotic resistance genes. These plasmids are formed by the successive integration of resistance genes through IS-mediated recombination events. The evolution of multiple drug resistance has occurred in several species pathogenic to humans, including strains of *Staphylococcus, Enterococcus, Neisseria, Shigella*, and *Salmonella*. Today many bacterial infections causing diseases such as dysentery, tuberculosis, and gonorrhea are difficult to treat because the pathogen has acquired resistance to several different antibiotics. To explore the evolution of these multi-drug-resistance plasmids, work through Solve It: Accumulating Drug-Resistance Genes.

COMPOSITE TRANSPOSONS

Composite transposons are created when two IS elements insert near each other. The region between the two IS elements can then be transposed when the elements act jointly. In effect, the two IS elements "capture" a DNA sequence that is otherwise immobile and endow it with the ability to move. **Figure 17.4** gives three examples of composite transposons, each denoted by the symbol Tn. In Tn9, the flanking IS elements are in the same orientation with respect to each other, whereas in Tn5 and Tn10, the orientation is inverted. The region between the IS elements in each of these transposons contains genes that have nothing to do with transposition. In fact, in all three transposons, the genes between the flanking IS elements confer resistance to antibiotics—a feature with obvious medical significance. Composite

transposons, like the IS elements that are part of them, create target site duplications when they insert into DNA.

Sometimes the flanking IS elements in a composite transposon are not quite identical. For instance, in Tn5, the element on the right, called IS50R, is capable of producing a transposase to stimulate transposition, but the element on the left, called IS50L, is not. This difference is due to a change in a single nucleotide pair that prevents IS50L from encoding the active transposase.

THE Tn3 ELEMENT

Bacteria contain other large transposons that do not have IS elements at each of their ends. Instead, these transposons terminate in simple inverted repeats 38 to 40 nucleotide pairs long; however, like the

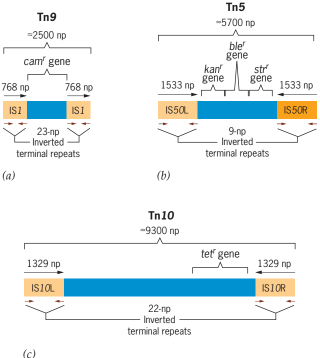
■ FIGURE 17.4 Genetic organization of composite transposons. The orientation and length (in nucleotide pairs, np) of the constituent sequences are indicated. (a) Tn9 consists of two IS1 elements flanking a gene for chloramphenicol resistance. (b) Tn5 consists of two IS50 elements flanking genes for kanamycin, bleomycin, and streptomycin resistance. (c) Tn10 consists of two IS10 elements flanking a gene for tetracycline resistance.

Solve It!

Accumulating Drug-Resistance Genes

An *E. coli* cell has a conjugative R plasmid that carries the gene for streptomycin resistance (*str'*) flanked by IS1 elements. Another *E. coli* cell has a nonconjugative plasmid that carries a gene for tetracycline resistance (*tet'*), as well as one copy of IS1. Outline how a conjugative R plasmid that carries both the *str'* and *tet'* genes might evolve.

To see the solution to this problem, visit the Student Companion site.



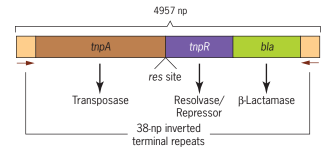


FIGURE 17.5 Genetic organization of Tn*3*. Lengths of DNA sequences are given in nucleotide pairs (np).

cut-and-paste transposons, they create target site duplications when they insert into DNA. The element known as Tn3 is the prime example of this type of transposon.

The genetic organization of Tn3 is shown in **Figure 17.5**. There are three genes, *tnpA*, *tnpR*, and *bla*, encoding, respectively, a transposase, a resolvase/repressor, and an enzyme called beta lactamase. The beta lactamase confers resistance to the antibiotic ampicillin, and the other two proteins play important roles in transposition.

Tn3 is a replicative transposon that moves in a two-stage process (**Figure 17.6**). In the first stage, the transposase mediates the fusion of two circular molecules—for instance, two plasmids, one carrying Tn3 (the donor plasmid) and the other not carrying it (the recipient plasmid). The resulting structure is called a **cointegrate**. During the formation of the cointegrate,

Tn3 is replicated, and one copy is inserted at each point where the two plasmids have fused; within the cointegrate, these two copies of Tn3 are oriented in the same direction. In the second stage of transposition, the *tnpR*-encoded resolvase mediates a site-specific recombination event between the two Tn3 copies. This event occurs at a sequence in Tn3 called *res*, the *resolution site*, and when it is completed, the cointegrate is resolved into its two constituent plasmids, each with a copy of Tn3.

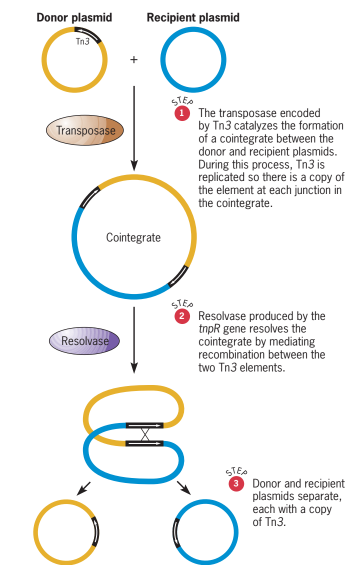


FIGURE 17.6 Transposition of Tn3 via the formation of a cointegrate.

The tnpR gene product of Tn3 has yet another function—to repress the synthesis of both the transposase and resolvase proteins. Repression occurs because the *res* site is located between the tnpA and tnpR genes. By binding to this site, the tnpR protein interferes with the transcription of both genes, leaving their products in chronic short supply. As a result, the Tn3 element tends to remain immobile.

- Insertion sequences (IS elements) are cut-and-paste transposons that reside in bacterial chromosomes and plasmids.
- IS elements can mediate recombination between different DNA molecules.
- Conjugative plasmids can move genes for antibiotic resistance from one bacterial cell to another.
- Composite transposons consist of two IS elements flanking a region that contains one or more genes for antibiotic resistance.
- Tn3 is a replicative transposon that transposes by temporarily fusing DNA molecules into a cointegrate; when the cointegrate is resolved, each of the constituent DNA molecules emerges with a copy of Tn3.
- Bacterial transposons are demarcated by terminal inverted repeats; when they insert into a DNA molecule, they create a duplication of sequences at the insertion site (a target site duplication).

KEY POINTS

Cut-and-Paste Transposons in Eukaryotes

Geneticists have found many different types of transposons in eukaryotes. These elements vary in size, structure, and behavior. Some are abundant in the genome, others rare. In the following sections, we discuss a few of the eukaryotic transposons that move by a cut-and-paste mechanism. All these elements have inverted repeats at their termini and create target site duplications when they insert into DNA molecules. Some encode a transposase that catalyzes the movement of the element from one position to another.

Ac AND Ds ELEMENTS IN MAIZE

The *Ac* and *Ds* elements in maize were discovered by the American scientist Barbara McClintock. Through genetic analysis, McClintock showed that the activities of these elements are responsible for the striping and spotting of maize kernels. Many years later, Nina Federoff, Joachim Messing, Peter Starlinger, Heinz Saedler, Susan Wessler, and their colleagues isolated the elements and determined their molecular structure.

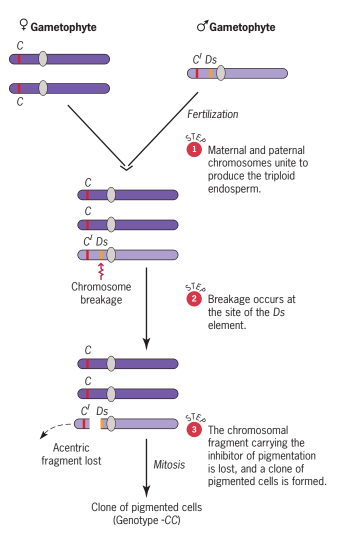
McClintock discovered the *Ac* and *Ds* elements by studying chromosome breakage. She used genetic markers that controlled the color of maize kernels to detect the breakage events. When a particular marker was lost, McClintock inferred that the chromosome segment on which it was located had also been lost, an indication that a breakage event had occurred. The loss of a marker was detected by a change in the color of the aleurone, the outermost layer of the triploid endosperm of maize kernels.

In one set of experiments, the genetic marker that McClintock followed was an allele of the *C* locus on the short arm of chromosome 9. Because this allele, C^{l} , is a dominant inhibitor of aleurone coloration, any kernel possessing it is colorless. McClintock fertilized *CC* ears with pollen from $C^{l}C^{l}$ tassels, producing kernels in which the endosperm was $C^{l}CC$. (The triploid endosperm receives two alleles from the female parent and one from the male parent; see Chapter 2.) Although McClintock found that most of these kernels were colorless, as expected, some showed patches of brownish-purple pigment (**■** Figure 17.7). McClintock guessed that in such

Transposable elements were discovered by analyzing genetic instabilities in maize; genetic analyses have also revealed transposable elements in *Drosophila*.



■ **FIGURE 17.7** Maize kernel (top view) showing loss of the *C*['] allele for the inhibition of pigmentation in the aleurone. The brownish purple patches are *-CC*, whereas the yellow patches are *C'CC*.



■ **FIGURE 17.8** Chromosome breakage caused by the transposable element *Ds* in maize. The allele *C* on the short arm of chromosome 9 produces normal pigmentation in the aleurone; the allele *C'* inhibits this pigmentation.

mosaics, the inhibitory C^{i} allele had been lost sometime during endosperm development, leading to a clone of tissue that was able to make pigment. The genotype in such a clone would be *-CC*, where the dash indicates the missing C^{i} allele.

The mechanism that McClintock proposed to explain the loss of the C^{I} allele is diagrammed in **Figure 17.8**. A break at the site labeled by the arrow detaches a segment of the chromosome from its centromere, creating an acentric fragment. Such a fragment tends to be lost during cell division; thus, all the descendants of this cell will lack part of the paternally derived chromosome. Because the lost fragment carries the C^{I} allele, none of the cells in this clone is inhibited from forming pigment. If any of them produces a part of the aleurone, a patch of purple tissue will appear, creating a mosaic kernel similar to the one shown in Figure 17.7.

McClintock found that the breakage responsible for these mosaic kernels occurred at a particular site on chromosome 9. She named the factor that produced these breaks Ds, for Dissociation. However, by itself, this factor was unable to induce chromosome breakage. In fact, McClintock found that Ds had to be stimulated by another factor, called Ac, for Activator. The Ac factor was present in some maize stocks but absent in others. When different stocks were crossed, Ac could be combined with Ds to create the condition that led to chromosome breakage.

This two-factor *Ac/Ds* system provided an explanation for the genetic instability that McClintock had observed on chromosome 9. Additional experiments demonstrated that this was only one of many instabilities present in the maize genome. McClintock found other instances of breakage at different sites on chromosome 9 and also on other chromosomes. Because breakage at these sites depended on activation by *Ac*, she concluded that *Ds* factors were also involved. To explain all these observations, McClintock proposed that *Ds* could exist at many different sites in the genome and that it could move from one site to another.

This explanation has been borne out by subsequent analyses. The Ac and Ds elements belong to a family of transposons. These elements are structurally related to each other and can insert at many different sites on the chromosomes. Multiple copies of the Ac and Ds elements are often present in the maize genome. Through genetic analysis, McClintock demonstrated that both Ac and Ds can move. When one of these elements

inserts in or near a gene, McClintock found that the gene's function is altered—sometimes completely abolished. Thus, *Ac* and *Ds* can induce mutations by inserting into genes. To emphasize this effect on gene expression, McClintock called the *Ac* and *Ds* transposons **controlling elements**.

DNA sequencing has shown that Ac elements consist of 4563 nucleotide pairs bounded by inverted repeats that are 11 nucleotide pairs long (**■** Figure 17.9*a*); these terminal inverted repeats are essential for transposition. Each Ac element is also flanked by direct repeats 8 nucleotide pairs long. Because the direct repeats are created at the time the element is inserted into the chromosome, they are target site duplications, not integral parts of the element.

Unlike Ac, Ds elements are structurally heterogeneous. They all possess the same inverted terminal repeats as Ac elements, demonstrating that they belong to the same transposon family, but their internal sequences vary. Some Ds elements appear to have been derived from Ac elements by the loss of internal sequences (**■** Figure 17.9*b*). The deletions in these elements may have been caused by incomplete DNA synthesis during replication or transposition. Other Ds elements contain non-Ac DNA between their inverted terminal repeats (**■** Figure 17.9*c*). These unusual members of the Ac/Ds family are called *aberrant* Ds elements. A third class of Ds element is inserted into another but in an inverted orientation. These so-called *double* Ds elements appear to have been responsible for the chromosome breakage that McClintock observed in her experiments.

The activities of the Ac/Ds elements—excision and transposition, and all their genetic correlates, including mutation and chromosome breakage—are caused by a transposase encoded by the Ac elements. The Ac transposase interacts with sequences at or near the ends of Ac and Ds elements, catalyzing their movement. Deletions or mutations in the gene that encodes the transposase abolish this catalytic function. Thus Ds elements, which have such lesions, cannot activate themselves. However, they can be activated if a transposaseproducing Ac element is present somewhere in the genome. The transposase made by this element can diffuse through the nucleus, bind to a Ds element, and activate it. The Ac transposase is, therefore, a *trans*-acting protein.

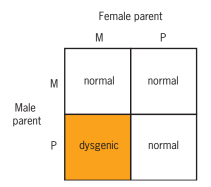
Transposons related to the *Ac/Ds* elements have been found in other species, including animals. Perhaps the best-studied of these elements is one called *hobo*, whimsically named for its ability to transpose. The *hobo* element is found in some species of *Drosophila*. To explore other genetic effects of the *Ac/Ds* elements, work through Problem-Solving Skills: Analyzing Transposon Activity in Maize.

P ELEMENTS AND HYBRID DYSGENESIS IN DROSOPHILA

Some of the most extensive research on transposable elements has focused on the P elements of *Drosophila melanogaster*. These transposons were identified through the cooperation of geneticists working in several different laboratories. In 1977 Margaret and James Kidwell, working in Rhode Island, and John Sved, working in Australia, discovered that crosses between certain strains of *Drosophila* produce hybrids with an assortment of aberrant traits, including frequent mutation, chromosome breakage, and sterility. The term **hybrid dysgenesis**, derived from Greek roots meaning "a deterioration in quality," was used to denote this syndrome of abnormalities.

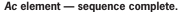
Kidwell and her colleagues found that they could classify *Drosophila* strains into two main types based on whether or not they produce dysgenic hybrids in testcrosses. The two types of strains are denoted M and P. Only

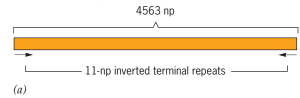
crosses between M and P strains produce dysgenic hybrids, and they do so only if the male in the cross is from the P strain. Crosses between two different P strains, or between two different M strains, produce hybrids that are normal. We can summarize the phenotypes of the hybrid offspring from these different crosses in a simple table:



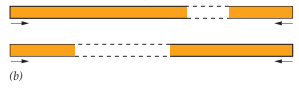
The parents of the different strains therefore contribute *maternally* or *paternally* to the formation of dysgenic hybrids—hence, their designations as M and P.

To Kidwell and her colleagues, these findings suggested that the chromosomes of P strains carry genetic factors that are activated when they enter eggs made by M females, and that once activated, these factors induce mutations and chromosome breakage. Inspired by this work, William Engels, a graduate student at the University of Wisconsin, began to study mutations induced in dysgenic hybrids. In 1979 Engels





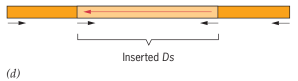
Ds elements — internal sequences missing.



Aberrant Ds element — internal sequences unrelated to Ac.







■ **FIGURE 17.9** Structural organization of the members of the *Ac/Ds* family of transposable elements in maize. The terminal inverted repeats (short arrows underneath) and DNA sequence lengths (in nucleotide pairs, np) are indicated.

PROBLEM-SOLVING SKILLS

Analyzing Transposon Activity in Maize

THE PROBLEM

In maize, the wild-type allele of the *C* gene is needed for dark coloration of the aleurone in kernels. Without this allele, the aluerone is pale yellow. c^{Ds} is a recessive mutation caused by the insertion of a *Ds* element into the 5' untranslated region of the *C* gene—that is, into the region between the transcription start site and the first codon in the polypeptide coding sequence. Inbred strains of maize that are homozygous for this mutation produce pale yellow kernels, just like inbred strains that are homozygous for a deletion of the *C* gene (c^{Δ}). A maize breeder crosses an inbred $c^{Ds}c^{Ds}$ strain as female parent to an inbred $c^{\Delta}c^{\Delta}$ strain as male parent. Among the kernels in the F₁, he sees many that have patches of brownish purple tissue on an otherwise pale yellow aleurone. (a) Explain the F₁ phenotype. (b) Would you expect this phenotype if the *Ds* element were inserted somewhere in the coding sequence of the *C* gene?

FACTS AND CONCEPTS

- 1. *Ds*, the nonautonomous member of the *Ac/Ds* transposon family, moves only in the presence of *Ac*, the autonomous member.
- **2.** The 5' untranslated region of a gene does not contain codons for amino acids in the polypeptide specified by the gene.
- **3.** A transposon insertion into a gene may interfere with the gene's expression.
- **4.** Excision of a transposon usually leaves at least a portion of the target site duplication that was created when the transposon inserted.

ANALYSIS AND SOLUTION

a. To explain the F_1 phenotype, we note that the expression of the c^{Ds} allele is disrupted by a Ds insertion into the 5' untranslated

region of the *C* gene. If this *Ds* element were to be excised, the gene's expression might be restored. When the maize breeder crossed the two inbred strains, he unwittingly crossed a strain with a *Ds* insertion in the *C* gene to a strain that carried a cryptic *Ac* element. The triploid aleurone in the F_1 kernels must have been $c^{Ds}c^{Ds}c^4$ (*Ac*). The two copies of the c^{Ds} allele were derived from the female parent, and the single copy of the c^4 deletion allele and the single copy of *Ac* were derived from the male parent. In this hybrid genotype, *Ac* can activate the *Ds* element, causing it to excise from the *C* gene. Because the element was inserted into noncoding DNA, its excision is expected to restore *C* gene expression. Therefore, if cells in which such excisions occur give rise to aleurone tissue, that tissue will be brownish purple in an otherwise pale yellow kernel.

b. Ds excisions are seldom precise. Usually, several nucleotides in the gene's sequence around the Ds insertion site are either duplicated or deleted when the Ds element excises. For instance, the Ds element often leaves the target site duplication that it generated when it inserted into the gene—a kind of transposon footprint. These extra nucleotides are not likely to disrupt gene expression if they are located in the gene's 5' untranslated region, which does not contain any coding information. However, if they are located in the gene's coding region, they are likely to cause serious problems. They could alter the length or composition of the polypeptide encoded by the gene. Thus, excising a Ds element from the coding sequence of the C gene is not likely to restore that gene's function. With such a Ds insertion, we would not expect to see patches of brownish purple tissue in the F₁ kernels.

For further discussion visit the Student Companion site.

found a particular mutation that reverted to wild-type at a high rate. This instability, which is reminiscent of the behavior of IS-induced mutations in *E. coli*, strongly suggested that a transposable element was involved.

The discovery by Michael Simmons and Johng Lim of dysgenesis-induced mutations in the *white* gene allowed the transposon hypothesis to be tested. In 1980, Simmons and Lim, working in Minnesota and Wisconsin, respectively, sent the newly discovered *white* mutations to Paul Bingham, a geneticist in North Carolina. Bingham and his collaborator, Gerald Rubin, a geneticist in Maryland, had just finished isolating DNA from the *white* gene. Using this DNA as a probe, Bingham and Rubin were able to isolate DNA from the mutant *white* alleles and compare it to the wild-type *white* DNA. In each mutation, they found that a small element had been inserted into the coding region of the *white* gene. Additional experiments demonstrated that these elements are present in multiple copies and at different locations in the genomes of P strains; however, they are completely absent from the genomes of M strains. Geneticists therefore began calling these P strain–specific transposons *P* elements.

DNA sequence analysis has shown that P elements vary in size. The largest elements are 2907 nucleotide pairs long, including terminal inverted repeats of 31 nucleotide pairs. These *complete* P elements carry a gene that encodes a transposase. When the P transposase cleaves DNA near the ends of a complete P element, it can

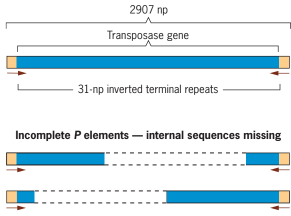


move that element to a new location in the genome. *Incomplete* P elements (**Figure 17.10**) lack the ability to produce the transposase because some of their internal sequences are deleted; however, they do possess the terminal and subterminal sequences recognized by the transposase. Consequently, these elements can be mobilized if a transposase-producing complete element is present somewhere in the genome.

In dysgenic hybrids, P elements transpose only in the cells of the germ line. This restriction is due to the inability of the somatic cells to remove one of the introns from the P element's pre-mRNA. When translated, this incompletely spliced RNA produces a polypeptide that does not have the transposase's ability to catalyze P element movement. As a result, the somatic cells are spared from the ravages of P element activity. Hybrid dysgenesis is, therefore, a strictly germ-line phenomenon.

Drosophila's germ-line cells also have ways of minimizing the damage that *P* elements can cause. The most effective mechanism involves small RNA molecules that are derived from the *P* elements themselves. These RNAs form complexes with a special group of proteins, whimsically called the Piwi proteins; hence, they are designated as piwi-interacting or piRNAs. Females from P strains produce these piRNAs and transmit them to their offspring through the cytoplasm of their eggs. Once in the offspring, the piRNAs repress *P* element activity in the germ line and prevent hybrid dysgenesis from occurring. Maternal transmission of the repressing piRNAs therefore explains why the offspring of crosses between P females and M males, as well as the offspring of crosses between P females and P males, are not dysgenic. On the Cutting Edge: Small RNAs Repress *P* Element Activity highlights some of the recent discoveries about this mechanism of transposon regulation.

Complete P element — all sequences present



■ **FIGURE 17.10** Structure of *P* elements in *Drosophila* showing orientations and lengths (in nucleotide pairs, np) of DNA sequences.

ON THE CUTTING EDGE

SMALL RNAs REPRESS P ELEMENT ACTIVITY

he discovery that *P* elements cause hybrid dysgenesis raised two basic questions. Why doesn't dysgenesis occur in the somatic tissues, and why doesn't it occur in flies whose mothers have come from a P strain? Geneticists answered the first question in 1986 when they learned that the *P* element's transposase is not produced in somatic cells. Without transposase to catalyze movement, *P* elements remain inactive in the somatic cells. The second question was answered more recently when geneticists learned that flies from P strains make small RNAs that interfere with *P* element activity.

Genetic analyses have shown that repression of hybrid dysgenesis in the germ line is correlated with the presence of a *P* element in one site in the *Drosophila* genome—the left telomere of the X chromosome. Many flies in natural populations carry this kind of *P* element, and they repress dysgenesis handsomely. An X-linked telomeric *P* element therefore appears to have an almost magical power to control all the other *P* elements in the genome, no matter where they are located. However, an X-linked telomeric *P* element can only exert its power if it is inherited maternally. A telomeric *P* element that is inherited paternally loses its ability to prevent hybrid dysgenesis.

Why is a maternally inherited telomeric *P* element so special? It turns out that the element is inserted in a site in the genome that produces piRNAs. Many different loci produce piRNAs, but the left telomere of the X chromosome is an unusually strong source. When a *P* element is inserted into this locus, it generates *P*-specific piRNAs—that is, piRNAs consisting of *P* element sequences. These RNAs are 23–29 nucleotides long, and they may be either sense or antisense in sequence. Furthermore, these piRNAs are transmitted maternally through the cytoplasm of *Drosophila* eggs. Thus, a female that carries a telomeric *P* element can produce *P*-specific piRNAs and transmit them to her offspring.

What is the significance of this maternal endowment? In the offspring, piRNAs with antisense sequences clearly pose a threat to the expression of the *P* element's transposase gene. In the germ line, this gene is transcribed into a pre-mRNA, which is then spliced to form an mRNA encoding the P transposase. But if antisense piRNAs base-pair with the mRNA, translation of the mRNA will be blocked. Even worse, the piRNA-mRNA duplex molecules may induce the cell's machinery to destroy the mRNA. In either case, the *P* element transposase will not be made, and without it, none of the *P* elements in the genome can move. Thus, maternally inherited piRNAs generated from the telomeric *P* element will prevent hybrid dysgenesis from occurring.

These discoveries have raised many new questions. How does the telomeric locus produce piRNAs with both sense and antisense sequences? How are the piRNAs transmitted through the egg cytoplasm? Do the Piwi proteins play a role? What is the state of the piRNA locus in males? Is it active or inactive, and if inactive, can it be reactivated if the locus is transmitted to a female? Are *P*-specific piRNAs generated in somatic cells? Do piRNAs with different specificities regulate the movement of other kinds of transposable elements in *Drosophila*? Does the piRNA mechanism operate in other organisms? KEY POINTS

- The maize transposable element Ds, discovered because of its ability to break chromosomes, is activated by another transposable element, Ac, which encodes a transposase.
 - Transposable P elements are responsible for hybrid dysgenesis, a syndrome of germ-line abnormalities that occurs in the offspring of crosses between P and M strains of Drosophila.
 - Within the germ line, P element activity is regulated by small RNAs (piRNAs) derived from the P elements themselves.

Retroviruses and Retrotransposons

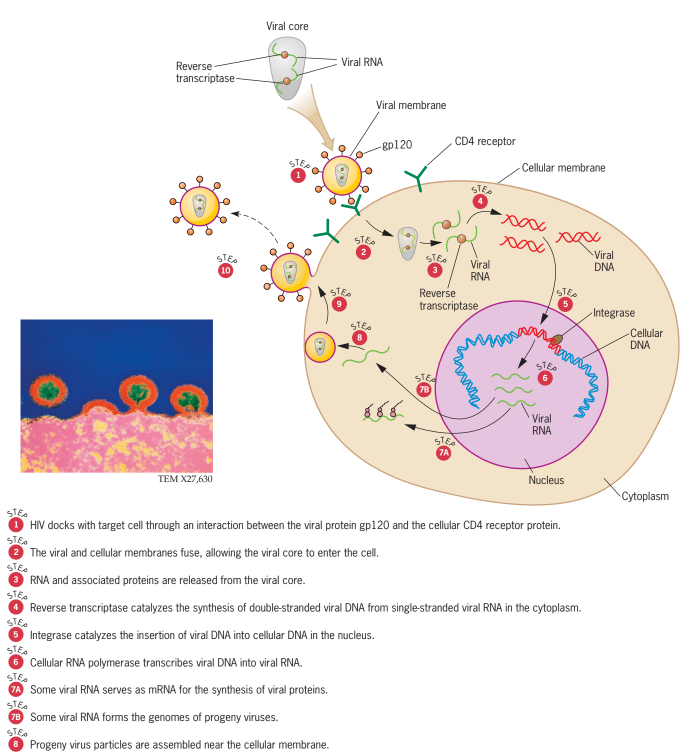
Retroviruses and related transposable elements utilize the enzyme reverse transcriptase to copy RNA into DNA. The DNA copies are subsequently inserted at different positions in genomic DNA. In addition to cut-and-paste transposons such as Ac and P, eukaryotic genomes contain transposable elements whose movement depends on the reverse transcription of RNA into DNA. This reversal in the flow of genetic information has led geneticists to call these elements **retrotransposons**, from a Latin prefix meaning "backward." Reverse transcription also plays a crucial role in the life cycles of some viruses. The genomes of

these viruses are composed of single-stranded RNA. When one of these viruses infects a cell, its RNA is copied into double-stranded DNA. Because the genetic information moves from RNA to DNA, these viruses are called **retroviruses**. We will begin our investigation of retrotransposons with a discussion of the retroviruses. Later, we will delve into the two main classes of retrotransposons.

RETROVIRUSES

The retroviruses were discovered by studying the causes of certain types of tumors in chickens, cats, and mice. In each case, an RNA virus was implicated in the production of the tumor. An important advance in understanding the life cycles of these viruses came in 1970 when David Baltimore, Howard Temin, and Satoshi Mizutani discovered an RNA-dependent DNA polymerase—that is, a **reverse transcriptase**, which allows these viruses to copy RNA into DNA. This discovery initiated research on the process of reverse transcription and provided a glimpse into what might be called the "retro-world"—that vast collection of DNA sequences derived from reverse transcription. We now know that reverse transcription is responsible for populating genomes with many kinds of DNA sequences, including, of course, the retroviruses. The discovery of reverse transcriptase therefore opened a view onto a component of genomes that had previously been unexplored.

Many different types of retroviruses have been isolated and identified. However, the epitome is the human immunodeficiency virus (HIV), which causes acquired immune deficiency syndrome, or AIDS, a disease that now affects tens of millions of people. AIDS was first detected in the last quarter of the twentieth century. It is a serious disease of the immune system. As it progresses, a person loses the ability to fight off infections by an assortment of pathogens, including organisms that are normally benign. Without treatment, infected individuals succumb to these infections, and eventually they die. AIDS is transmitted from one individual to another through bodily fluids such as blood or semen that have been contaminated with HIV. The initial symptoms of the disease are flulike. Infected individuals experience aches, fever, and fatigue. After a few weeks, these symptoms abate and health is seemingly restored. This asymptomatic state may last several years. However, the virus continues to multiply and spread through the body, targeting specialized cells that play important roles in the immune system. Eventually, these cells are so depleted by the killing action of the virus that the immune system fails and opportunistic pathogens assert themselves. Many types of illnesses, such as pneumonia, may ensue. AIDS is a major cause of death among subpopulations in many countries-for example, among intravenous drug users and sex industry workers-and in sub-Saharan Africa, it is a major cause of death in the population at large.



- STE
 Progeny virus particles are extruded from the cell by budding.
- SIED
- 10 Progeny virus particles are free to infect other cells.
- **FIGURE 17.11** The HIV life cycle. The inset shows virus particles budding from a cell.

Because of its lethality and pandemic status, HIV/AIDS has been the focus of an enormous amount of research. One outcome of this effort has been a detailed understanding of HIV's life cycle (**■** Figure 17.11). The spherical virus enters a host cell by interacting with specific receptor proteins, called CD4 receptors, which are located on the cell's surface. This interaction is mediated by a glycoprotein (a protein to which sugars have been attached) called gp120, which is embedded in the lipid

membrane that surrounds the viral particle. Once gp120 has "docked" with the CD4 receptor, the viral and cellular membranes fuse and the viral particle is admitted to the cell. Inside the cell, the lipid membrane and the protein coat that surround the virus particle are removed, and materials within the virus's core are released into the cell's cytoplasm. This core contains two identical single-stranded RNA molecules—the virus's genome—and a small number of proteins that facilitate replication of the genome, including two molecules of the viral reverse transcriptase, one bound to each strand of viral RNA.

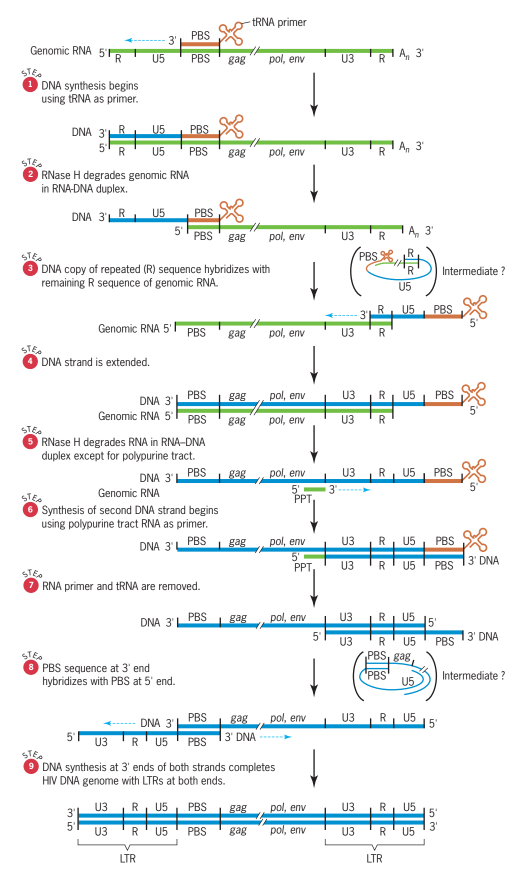
HIV's reverse transcriptase—and other reverse transcriptases as well—converts single-stranded RNA into double-stranded DNA. The resulting double-stranded DNA molecules are then inserted at random positions in the chromosomes of the infected cell, in effect populating that cell's genome with many copies of the viral genome. These copies can then be transcribed by the cell's ordinary RNA polymerases to produce a large amount of viral RNA, which serves to direct the synthesis of viral proteins and also provides genomic RNA for the assembly of new viral particles. These particles are extruded from the cell by a process of budding through the cell's membrane. The extruded particles may then infect other cells by interacting with the CD4 receptors on their surfaces. In this way, HIV's genetic material is replicated and disseminated through a population of susceptible immune cells.

The HIV genome, slightly more than 10 kilobases long, contains several genes. Three of these genes, denoted *gag*, *pol*, and *env*, are found in all other retroviruses. The *gag* gene encodes proteins of the viral particle; the *pol* gene encodes the reverse transcriptase and another enzyme called integrase, which catalyzes the insertion of the DNA form of the HIV genome into the chromosomes of a host cell; and the *env* gene encodes the glycoproteins that are embedded in the virus's lipid envelope.

Let's now take a closer look at replication of the HIV genome (**Figure 17.12**). This process, catalyzed by reverse transcriptase, begins with the synthesis of a single DNA strand complementary to the single-stranded RNA of the viral genome. It is primed by a tRNA that is complementary to a sequence called PBS (*primer binding site*) situated to the left of center in the HIV RNA (step 1 in Figure 17.12). This tRNA is packaged already bound to the PBS in the HIV core. After reverse transcriptase catalyzes the synthesis of the 3' end of the viral DNA, ribonuclease H (RNase H) degrades the genomic RNA in the RNA-DNA duplex (step 2). This degradation leaves the repeated (R) sequence of the nascent DNA strand free to hybridize with the R sequence at the 3' end of the HIV RNA. The net result is that the R region of the nascent DNA strand "jumps" from the 5' end of the HIV RNA to the 3' end of the HIV RNA (step 3). Reverse transcriptase next extends the DNA copy by using the 5' region of the HIV RNA as template (step 4).

In step 5, RNaseH degrades all the RNA in the RNA-DNA duplex except a small region, the polypurine tract (PPT), which is composed mostly of the purines adenine and guanine. This polypurine tract is used to prime second-strand DNA synthesis of part of the HIV genome (step 6). After the tRNA and the genomic RNA present in the RNA-DNA duplexes are removed (step 7), a second DNA "jump" occurs during which the PBS at the 5' end of the second DNA strand hybridizes with the complementary PBS at the 5' end of the first DNA strand (step 8). The 3'-hydroxyl termini of the two DNA strands are then used to prime DNA synthesis to complete the synthesis of double-stranded HIV DNA (step 9). Note that the conversion of the viral RNA to viral DNA produces signature sequences at both ends of the DNA molecule. These sequences, called **long terminal repeats (LTRs),** are required for integration of the viral genome into the DNA of the host cell.

Integration (**Figure 17.13**) of the viral DNA is catalyzed by the enzyme integrase, which has endonuclease activity. Integrase first produces recessed 3' ends in the HIV DNA by making single-stranded cuts near the ends of both LTRs (step 1). These recessed ends are next used for integrase-catalyzed attacks on phosphodiester bonds in a target sequence in the DNA of the host cell. This process results in the formation of new phosphodiester linkages between the 3' ends of the HIV DNA and 5' phosphates in the host DNA (step 2). In the final stage of integration, DNA repair enzymes of the host cell fill in the single-strand gaps to produce an HIV DNA genome covalently



■ **FIGURE 17.12** Conversion of HIV genomic RNA into double-stranded DNA. R, repeated sequence; U5, *u*nique sequence near 5' terminus; U3, *u*nique sequence near 3' terminus; PBS, *p*rimer *b*inding site; A_n poly(A) tail; *gag*, *pol*, and *env*, sequences encoding HIV proteins; PPT, polypurine tract rich in adenine and guanine; LTR, long terminal repeat. The dashed arrows indicate the direction in which DNA synthesis will occur at each step in the process.

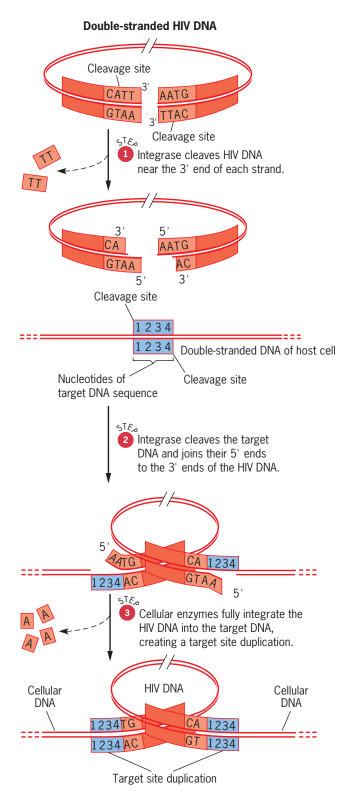


FIGURE 17.13 Integration of the HIV double-stranded DNA into the chromosomal DNA of the host cell.

inserted into the chromosomal DNA of the host cell (step 3). Notice that the target sequence at the site of integration is duplicated in the process. The integrated HIV genome thereafter becomes a permanent part of the host cell genome, replicating just like any other segment of the host DNA.

Integrated retroviruses of many different types are present in vertebrate genomes, including our own. Because these retroviruses are replicated along with the rest of the DNA, they are transmitted to daughter cells during division, and if they are integrated in germ-line cells, they are also passed on to the next generation through the gametes. Geneticists call the heritable DNA sequences that are derived from the reverse transcription and integration of viral genomes *endogenous retroviruses*. For the most part, these sequences have lost their ability to produce infectious viral particles; they are, therefore, innocuous remnants of ancient viral infections. HIV is not an endogenous retrovirus, but if it should lose its lethal potential and be transmitted in integrated form through the germ line, it could become one.

We now turn our attention to two classes of retrotransposons: the retroviruslike elements, which resemble the integrated forms of retroviruses, and the retroposons, which are DNA copies of polyadenylated RNA.

RETROVIRUSLIKE ELEMENTS

Retroviruslike elements are found in many different eukaryotes, including yeast, plants, and animals. Despite differences in size and nucleotide sequence, they all have the same basic structure: a central coding region flanked by long terminal repeats, or LTRs, which are oriented in the same direction. The repeated sequences are typically a few hundred nucleotide pairs long. Each LTR is, in turn, usually bounded by short, inverted repeats like those found in other types of transposons. Because of their characteristic LTRs, the retroviruslike elements are sometimes called *LTR retrotransposons*.

The coding region of a retroviruslike element contains a small number of genes, usually only two. These genes are homologous to the *gag* and *pol* genes found in retroviruses; *gag* encodes a structural protein of the virus capsule, and *pol* encodes a reverse transcriptase/integrase protein. The retroviruses have a third gene, *env*, which encodes a protein component of the virus envelope. In the retroviruslike elements, the gag and pol proteins play important roles in the transposition process.

One of the best-studied retroviruslike elements is the Ty1 transposon from the yeast *Saccharomyces cerevisiae*. This element is about 5.9 kilobase pairs long; its LTRs are about 340 base pairs long, and it creates a 5-bp target site duplication upon insertion into a chromosome. Most yeast strains have about 35 copies of the Ty1 element in their genome. Ty1 elements have only two genes, *TyA* and *TyB*, which are homologous to the *gag* and *pol* genes of the retroviruses. Biochemical studies have shown that the products of these two genes can form viruslike particles in the cytoplasm of yeast cells. The transposition of Ty1 elements involves reverse transcription of RNA (**Figure 17.14**). After the RNA is synthesized from Ty1 DNA, a reverse transcriptase encoded by

the TyB gene uses it as a template to make double-stranded DNA, probably in the viruslike particles. Then the newly synthesized DNA is transported to the nucleus and inserted somewhere in the genome, creating a new Ty1 element.

Retroviruslike elements have also been found in *Drosophila*. One of the first that was identified is called *copia*, so named because it produces copious amounts of RNA. The *copia* element is structurally similar to the Ty1 element of yeast. The *gypsy* element, another *Drosophila* retrotransposon, is larger than the *copia* element because

it contains a gene similar to the *env* gene of retroviruses. Both the *copia* and *gypsy* elements form viruslike particles inside *Drosophila* cells; however, only the particles that contain *gypsy* RNA can move across cell membranes, possibly because they also contain *gypsy*'s *env* gene product. The *gypsy* element therefore appears to be a genuine retrovirus. Many other families of retroviruslike transposons have been found in *Drosophila*, but their activities are poorly understood.

RETROPOSONS

The **retroposons**, or non-LTR retrotransposons, are a large and widely distributed class of retrotransposons, including the F, G, and I elements of *Drosophila* and several types of elements in mammals. These elements move through an RNA molecule that is reverse transcribed into DNA, usually by a protein encoded by the elements themselves. Although they create a target site duplication when they insert into a chromosome, they do not have inverted or direct repeats as integral parts of their termini. Instead, they are distinguished by a homogeneous sequence of A:T base pairs at one end. This sequence is derived from reverse transcription of the poly(A) tail that is added near the 3' end of the retroposon RNA during its maturation. Integrated retroposons therefore exhibit a vestige of their origin as reverse transcripts of polyadenylated RNAs.

In *Drosophila*, special retroposons are found at the ends (telomeres) of chromosomes, where they perform the critical function of replenishing DNA that is lost by incomplete chromosome replication. With

each round of DNA replication, a chromosome becomes shorter. Shortening takes place because the DNA polymerase can only move in one direction, adding nucleotides to the 3' end of a primer (Chapter 10). Usually, the primer is RNA, and when it is removed, a single-stranded region is left at the end of the DNA duplex. In the next round of replication, the deficient strand produces a duplex that is shorter than the original. As this process continues, cycle after cycle, the chromosome loses material from its end.

To counterbalance this loss, *Drosophila* has evolved a curious mechanism involving at least two different retroposons, one called *HeT-A* and another called *TART* (for *t*elomere-*a*ssociated *r*etro*t*ransposon). Mary Lou Pardue, Robert Levis, Harald Biessmann, James Mason, and their colleagues have shown that these two elements transpose preferentially to the ends of chromosomes, extending them by several kilobases. Eventually, the transposed sequences are lost by incomplete DNA replication, but then a new transposition occurs to restore them. The *HeT-A* and *TART* retroposons therefore perform the important function of regenerating lost chromosome ends.

- Retrovirus genomes are composed of single-stranded RNA comprising at least three genes: gag (coding for structural proteins of the viral particle), pol (coding for a reverse transcriptase/ integrase protein), and env (coding for a protein embedded in the virus's lipid envelope).
- The human retrovirus HIV infects cells of the immune system and causes the life-threatening disease AIDS.
- *Retroviruslike elements possess genes homologous to gag and pol, but not to env.*
- Retroviruslike elements and the DNA forms of retroviruses inserted in cellular chromosomes are demarcated by long terminal repeat (LTR) sequences.
- Retroposons lack LTRs; however, at one end they have a sequence of A:T base pairs derived from the reverse transcription of a poly(A) tail attached to the retroposon's RNA.
- The retroposons HeT-A and TART are components of the ends of Drosophila chromosomes.

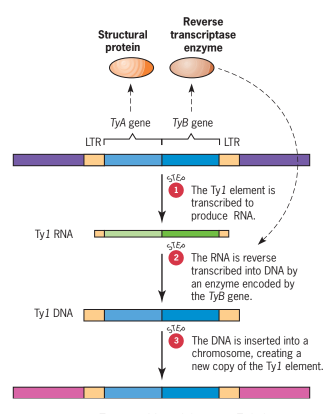


FIGURE 17.14 Transposition of the yeast Ty1 element.

KEY POINTS

Transposable Elements in Humans

The human genome is populated by a diverse array of transposable elements that collectively account for 44 percent of all human DNA. With the sequencing of the human genome, it is now possible to assess the significance of transposable elements in our own species. At least 44 percent of human DNA is derived from transposable elements, including retroviruslike elements (8 percent of the sequenced genome), retroposons (33 percent), and several families of elements that transpose by a cut-and-paste mechanism (3 percent).

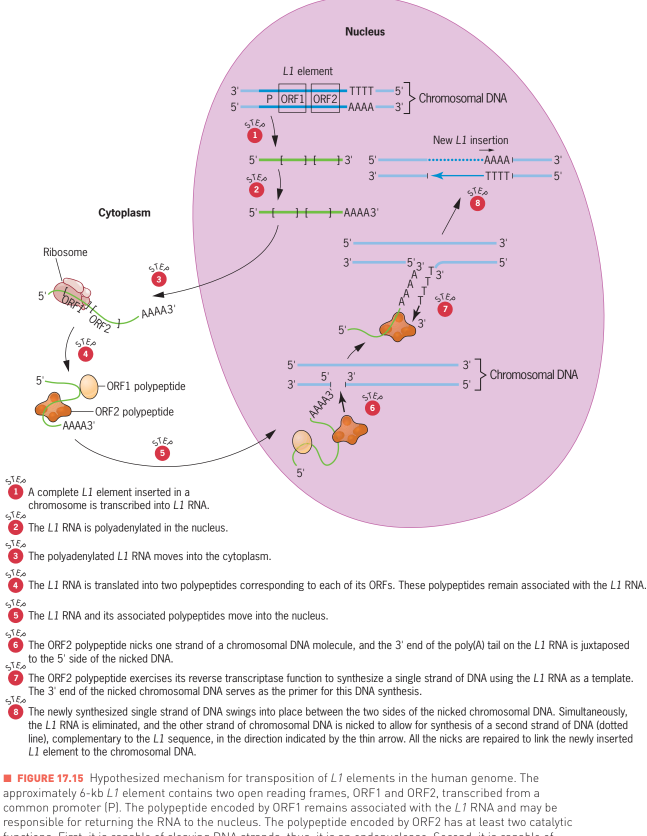
The principal transposable element is a retroposon called L1. This element belongs to a class of sequences known as the **long interspersed nuclear elements**, or **LINEs**. Complete L1 elements are about 6 kb long, they have an internal promoter that is recognized by RNA polymerase II, and they have two open reading frames: ORF1, which encodes a nucleic acid-binding protein, and ORF2, which encodes a protein with endonuclease and reverse transcriptase activities. The human genome contains between 3000 and 5000 complete L1 elements. In addition, it contains more than 500,000 L1 elements that are truncated at their 5' ends; these incomplete L1 elements are transpositionally inactive. Each L1 element in the genome, whether complete or incomplete, is usually flanked by a short target site duplication.

L1 transposition involves the transcription of a complete L1 element into RNA and the reverse transcription of this RNA into DNA (**Figure 17.15**). Both processes take place in the nucleus. However, before the L1 RNA is reverse transcribed, it journeys to the cytoplasm where it is translated into polypeptides that apparently remain associated with it when it returns to the nucleus. The polypeptide encoded by ORF2 possesses an endonuclease function that catalyzes cleavage of one strand of the DNA duplex at a prospective insertion site in a chromosome. The exposed 3' end of this cleaved DNA strand then serves as a primer for DNA synthesis using the L1 RNA as a template and the reverse transcriptase activity provided by the ORF2 polypeptide. In this way, an L1 DNA sequence is synthesized at the point in the chromosome where the ORF2 polypeptide has introduced a single-strand nick. The newly synthesized L1 DNA is subsequently made double-stranded by further DNA synthesis, and the double-stranded product is then covalently integrated into the chromosome to create a new L1 element. Sometimes the 5' region of the L1 RNA is not copied into DNA. When this happens, the resulting L1 insertion will lack 5' sequences, including the promoter, and will be unable to generate RNA through ordinary transcription. Thus, these incomplete L1 elements will be transpositionally inactive.

Transposed copies of certain complete L1 elements have been discovered through analysis of individuals with genetic diseases such as hemophilia and muscular dystrophy. The rarity of these cases suggests that the frequency of L1 transposition in humans is low. Two other types of LINE sequences, L2 (315,000 copies) and L3(37,000 copies), are found in the human genome; however, neither of these elements is transpositionally active.

The short interspersed nuclear elements, or SINES, are the second most abundant class of transposable elements in the human genome. These elements are typically less than 400 base pairs long and do not encode proteins. Like all retroposons, they have a sequence of A:T base pairs at one end. SINEs transpose through a process that involves reverse transcription of an RNA that has been transcribed from an internal promoter. Although the details of the transposition process are not well understood, it seems that the reverse transcriptase needed for the synthesis of DNA from the SINE RNA is furnished by a LINE-type element. Thus, the SINEs depend on the LINEs to multiply and insert within the genome. In this sense, they can be considered as retroposons that are parasites on the functionally autonomous and authentic retroposons such as *L1*. The human genome contains three families of SINEs, the *Alu*, *MIR*, and *Ther2/MIR3* elements. However, only the *Alu* elements—named for an enzyme that recognizes a specific nucleotide sequence within them—are transpositionally active.

The human genome possesses more than 400,000 sequences that are derived from retroviruslike elements. Most of these sequences are solitary LTRs. Although



functions. First, it is capable of cleaving DNA strands; thus, it is an endonuclease. Second, it is capable of synthesizing DNA from an RNA template; thus, it is a reverse transcriptase. The size of the new L1 insertion will depend on how far the reverse transcriptase travels along the L1 RNA template. If it fails to reach the 5' end, the insertion will be incomplete. Incomplete insertions usually do not have functional promoters and therefore cannot produce L1 RNA for future transpositions.

more than 100 different families of retroviruslike elements have been identified in human DNA, only a few appear to have been transpositionally active in recent evolutionary history. Like the inactive LINEs and SINEs, nearly all of the human retroviruslike sequences are genetic fossils left over from a time when they were actively transposing.

Cut-and-paste transposons are a small component of the human genome. DNA sequencing has identified two elements that are distantly related to the Ac/Ds elements of maize, as well as a few other types of elements. All the available evidence indicates that these types of transposons have been transpositionally inactive for many millions of years.

- **KEY POINTS** The human genome contains four basic types of transposable elements: LINEs, SINEs, retroviruslike elements, and cut-and-paste transposons.
 - The L1 LINE and the Alu SINE are transpositionally active; other human transposons appear to be inactive.

The Genetic and Evolutionary Significance of Transposable Elements

Transposable elements are used as tools by geneticists. In nature, they play a role in genome evolution.

TRANSPOSONS AS MUTAGENS

Spontaneous mutations are often the result of transposable element activity. In Drosophila, for example, many of the spontaneous mutant alleles of the *white* gene are due to transposon insertions. In fact, the

very first mutant allele of white, w¹, discovered by T. H. Morgan, resulted from a transposon insertion. These observations suggest that transposons are nature's intrinsic mutagens. As they wander through the genome, they mutate genes and break chromosomes.

Geneticists have exploited the mutagenic potential of transposons to disrupt genes. Transposon mutagenesis was pioneered in the 1970s and 1980s using the Pelements of Drosophila. Crosses between males from P strains and females from M strains produce dysgenic hybrids in which the P elements inherited from the father become highly active. As these elements transpose in the germ-line cells of the hybrid offspring, they cause mutations that can be recovered by crossing the hybrids appropriately. A researcher might, for example, use H. J. Muller's ClB technique (Chapter 13) to recover *P*-induced recessive lethal mutations on the X chromosome. By following this general strategy, geneticists have obtained P element insertions in a large fraction of all the genes in the *Drosophila* genome.

Other types of transposons have been used to induce mutations in the genomes of nematodes, fish, mice, and various plants. Mutagenesis with transposons has an advantage over traditional methods of inducing mutations because a gene that has been mutated by the insertion of a transposable element is "tagged" with a known DNA sequence. The transposon tag can subsequently be used to isolate the gene from a large, heterogeneous mixture of DNA by using a probe derived from a cloned version of the transposon. Mutagenesis by transposon tagging is therefore a standard genetic technique today.

GENETIC TRANSFORMATION WITH TRANSPOSONS

Some bacterial transposons—for example, the composite transposons and Tn3—carry genes whose products are unrelated to tranposition. This observation suggests that transposons might be used to move different kinds of genes around in a genome-in

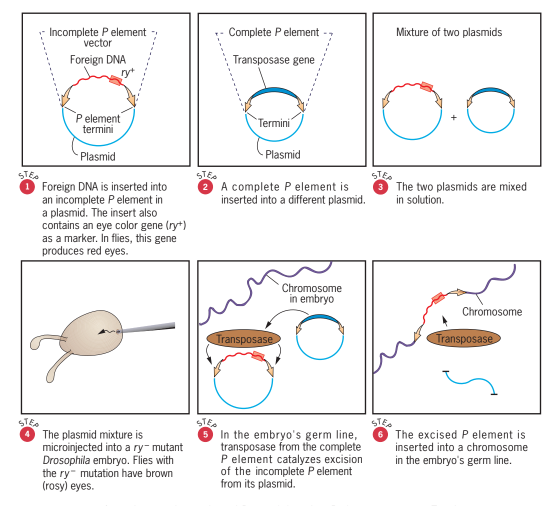


FIGURE 17.16 Genetic transformation of *Drosophila* using *P* element vectors. Foreign DNA inserted between *P* element termini is integrated into the genome through the action of a transposase encoded by the complete *P* element. Flies with this DNA in their genomes can be propagated in laboratory cultures.

effect, the genes become transposon cargo. It might also be possible to use transposons to move genes between organisms—that is, to transform one organism with DNA obtained from another organism.

These ideas inspired Gerald Rubin and Allan Spradling to see if a transposon could carry a cloned gene into an organism. As a test case, they chose one of the many genes that control eye color in *Drosophila*. This gene, called *rosy* (symbol ry), encodes the enzyme xanthine dehydrogenase. Flies lacking this enzyme-that is, homozygous ry mutants—have brown eyes, whereas flies homozygous for the wildtype allele ry^+ have red eyes. Rubin and Spradling used recombinant DNA techniques to insert the $r\gamma^+$ gene into an incomplete P element that had been cloned in a bacterial plasmid (**Figure 17.16**). Let's denote this recombinant element as $P(ry^+)$. In another plasmid, they cloned a complete P element capable of encoding the P element's transposase. Rubin and Spradling then injected a mixture of the two plasmids into *Drosophila* embryos that were homozygous for a mutant ry allele. They hoped that the transposase produced by the complete P element would catalyze the incomplete element to jump from its plasmid into the chromosomes of the germ-line cells and carry the ry^+ gene along as cargo. When the injected animals matured, Rubin and Spradling mated them to ry mutant flies. Among the offspring, they found many that had red eyes. Subsequent molecular analysis demonstrated that these red-eyed flies carried the $P(r\gamma^+)$ element. In effect, Rubin and Spradling had corrected the mutant eye color by inserting a copy of the wild-type rosy gene into the fly genome—that is, they had genetically transformed mutant flies with DNA from wild-type flies. A Milestone in Genetics: Transformation of *Drosophila* with P elements on the Student Companion site provides more details about this important achievement.

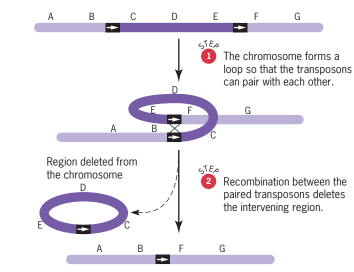
The technique that Rubin and Spradling developed is now routinely used to transform *Drosophila* with cloned DNA. An incomplete P element serves as the *transformation vector*; and a complete P element serves as the source of the transposase that is needed to insert the vector into the chromosomes of an injected embryo. The term *vector* comes from the Latin word for "carrier." It is used in this context because the incomplete P element *carries* a fragment of DNA into the genome. Practically any DNA sequence can be placed into the vector and ultimately inserted into the animal.

Unfortunately, *P* elements are not effective as transformation vectors in other species. However, geneticists have identified several transposons that can be used in their place. For example, the *piggyBac* transposon from a moth can serve as a transformation vector in many different species, and the *Sleeping Beauty* transposon from salmon works well in vertebrates, including humans, where it is being developed as a possible agent for gene therapy.

TRANSPOSONS AND GENOME ORGANIZATION

Some genomic regions are especially rich in transposon sequences. In *Drosophila*, for example, transposons are concentrated in the centric heterochromatin and in the heterochromatin abutting the euchromatin of each chromosome arm. However, many of these transposons have mutated to the point where they cannot be mobilized; genetically, they are the equivalent of "dead." Heterochromatin therefore seems to be a kind of graveyard filled with degenerate transposable elements.

Some evidence, especially from cytological studies of *Drosophila* by Johng Lim, suggests that transposable elements play a role in the evolution of chromosome structure. Several *Drosophila* transposons have been implicated in the formation of chromosome rearrangements, and a few seem to rearrange chromosomes at high frequencies. One possible mechanism is crossing over between homologous transposons located at different positions in a chromosome. If two transposons in the same orientation pair and cross over, the segment between them will be deleted (**■** Figure 17.17). You can explore the consequence of crossing over between two transposons in opposite orientations in a chromosome by working through Solve It: Transposon-Mediated Chromosome Rearrangements.



Chromosome with region C D E deleted

FIGURE 17.17 Formation of a deletion by intrachromosomal recombination between two transposons in the same orientation.

Solve It!

Transposon-Mediated Chromosome Rearrangements

Suppose a chromosome carries two copies of a transposon in opposite orientations. The order of the genes on the chromosome is $A \ B \ C \ D \ E \ F \ G$, and one transposon is located between genes B and C and the other is located between genes E and F. If the two transposons pair and then a crossover occurs, what will the order of the genes be in the resulting chromosome? Does your answer depend on whether the transposons are facing each other or pointing away from each other?

To see the solution to this problem, visit the Student Companion site.

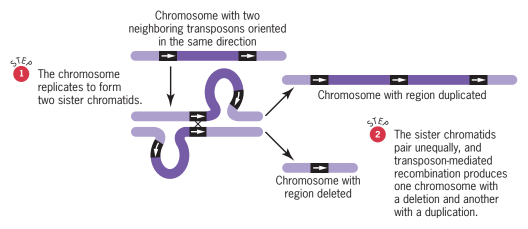


FIGURE 17.18 Origin of duplications and deletions by transposon-mediated unequal crossing over between sister chromatids.

Crossing over can also occur between transposons located in different chromosomes. In \blacksquare Figure 17.18 we consider a case where the crossover involves two sister chromatids. Each chromatid carries two neighboring transposons oriented in the same direction. The transposon on the left in one chromatid has paired with the transposon on the right in the other chromatid. A crossover between these paired transposons yields two structurally altered chromatids, one lacking the segment between the two transposons, the other with an extra copy of this segment. Crossing over between neighboring transposons can therefore duplicate or delete chromosome segments—that is, it can expand or contract a region of the genome.

- Transposons are used in genetic research to induce mutations.
- Transposons are used as vectors to move DNA within and between genomes.
- Crossing over between paired transposons can create chromosome rearrangements.

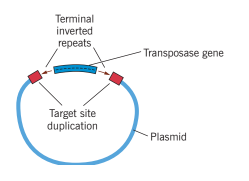
KEY POINTS

Basic Exercises

Illustrate Basic Genetic Analysis

1. Sketch a bacterial *IS* element inserted in a circular plasmid. Indicate the positions of (a) the transposase gene, (b) the terminal inverted repeats, and (c) the target site duplication.

Answer:



- 2. What factor must be present in maize to mobilize a *Ds* element inserted in a chromosome arm?
- **Answer:** A *Ds* element is mobilized when the transposase encoded by an *Ac* element acts on it. An *Ac* element must therefore be present somewhere in the maize genome.
- 3. A geneticist has two strains of *Drosophila*. One, a longstanding laboratory stock with white eyes, is devoid of *P* elements; the other, recently derived from wild-type flies collected in a fruit market, has *P* elements in its genome. Which of the following crosses would be expected to produce dysgenic hybrid offspring: (a) white females × wildtype males, (b) white males × wild-type females, (c) white females × white males, (d) wild-type females × wild-type males?

- **Answer:** (a) white females \times wild-type males. The white females lack *P* elements in their genomes, and they also lack the ability to make and transmit the piRNAs that could repress *P* elements in the germ line of the offspring. The wild-type males have *P* elements in their genomes, and they also have the capacity to produce repressing piRNAs. However, piRNAs cannot be transmitted to the offspring through the sperm. Thus, when the wild-type males are crossed to the white females, the offspring inherit *P* elements from their fathers and they do not inherit an ability to repress these elements from their mothers. This combination of factors allows the paternally inherited *P* elements to become active in the germline tissues of the offspring, and hybrid dysgenesis ensues.
- **4.** What are the similarities and differences among retroviruses, retroviruslike elements, and retroposons?
- **Answer:** All three types of retroelements use reverse transcription to insert DNA copies of their RNA into new sites in the cell's genome. Furthermore, the enzyme (reverse transcriptase) that catalyzes reverse transcription is encoded by each type of element. For retroviruses and retroviruslike elements, reverse transcription of the RNA occurs in the cytoplasm, whereas for retroposons, it occurs in the nucleus. Retroviruses and retroviruslike elements encode

another protein that functions in the assembly of virus or viruslike particles in the cytoplasm. Retroposons encode a different protein that appears to bind to the retroposon RNA and convey it into the nucleus. Retroviral RNA is packaged into viral particles, which can exit from the cell. This exiting capability requires a protein encoded by the *env* gene in the viral genome. Because neither retroviruslike elements nor retroposons carry an *env* gene, their RNA cannot be packaged for exit from the cell. Retroviruses are infectious; retroviruslike elements and retroposons are not.

- 5. What transposable element is most abundant in the human genome?
- **Answer:** The LINE known as *L1* is the most abundant human transposon. It accounts for about 17 percent of all human DNA.
- 6. How could two transposons in the same family cause deletion of DNA between them on a chromosome?
- **Answer:** The two transposons would have to be in the same orientation. Pairing between the transposons followed by recombination would excise the chromosomal material between them. See Figure 17.17.

Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. A copy of the wild-type *white* gene (w⁺) from *Drosophila* was inserted in the middle of an incomplete P element contained within a plasmid. The plasmid was mixed with another plasmid that contained a complete P element, and the mixture was carefully injected into *Drosophila* embryos homozygous for a null mutation (w⁻) of the *white* gene. The adults that developed from these injected embryos all had white eyes, but when they were mated to uninjected white flies, some of their progeny had red eyes. Explain the origin of these red-eyed progeny.
- **Answer:** The complete *P* element in one of the plasmids would produce the P transposase, the enzyme that catalyzes Pelement transposition, in the germ lines of the injected embryos. The incomplete P element in the other plasmid would be a target for this transposase. If this incomplete Pelement were mobilized by the transposase to jump from its plasmid into the chromosomes of the injected embryo, the fly that developed from this embryo would carry a copy of the wild-type white gene in its germ line. (P element movement is limited to the germ line; therefore, the incomplete P element would not jump into the chromosomes of the somatic cells, such as those that eventually form the eye.) Such a genetically transformed fly would, in effect, have the germ-line genotype w^{-}/w^{-} ; $P(w^{+})$ or w^{-}/Y ; $P(w^{+})$, where $P(w^+)$ denotes the incomplete P element that contains the w^+ gene. This element could be inserted on any

of the chromosomes. If the transformed fly were mated to an uninjected white fly, some of its offspring would inherit the $P(w^+)$ insertion, which, because it carries a wild-type *white* gene, would cause red eyes to develop. The red-eyed progeny are therefore the result of genetic transformation of a mutant white fly by the w^+ gene within the incomplete P element.

- 2. The *Alu* element is one of the SINEs in the human genome. Each *Alu* retroposon is about 300 base pairs long—not long enough to encode a reverse transcriptase that could catalyze the conversion of *Alu* RNA into *Alu* DNA during the process of retrotransposition. In spite of this deficiency, the *Alu* elements have accumulated to such an extent that they constitute 11 percent of human DNA—over 1 million copies. How might this dramatic expansion of *Alu* elements have occurred during the evolutionary history of the human lineage without an *Alu*-encoded reverse transcriptase?
- Answer: The *Alu* elements may have "borrowed" the services of a reverse transcriptase encoded by a different retroposon such as the *L1* element, which is large enough to encode a reverse transcriptase and at least one other polypeptide. If *L1*-encoded reverse transcriptase, or the reverse transcriptase encoded by some other retrotransposon—perhaps another LINE—can bind to *Alu* RNA, then it is conceivable

that the reverse transcriptase could use the *Alu* RNA to synthesize *Alu* DNA, which could subsequently be integrated into chromosomal DNA. Repetition of this process over evolutionary time could explain the accumulation of so many copies of the *Alu* element in the human genome.

- **3.** What techniques could be used to demonstrate that a mutation in a man with hemophilia is due to the insertion of an *Alu* element into the coding sequence of the X-linked gene for factor VIII, which is one of the proteins needed for efficient blood clotting in humans?
- **Answer:** A molecular geneticist would have several ways of showing that the mutant gene for hemophilia is due to an *Alu* insertion in the gene's coding sequence. One technique is genomic Southern blotting. Genomic DNA from the hemophiliac could be digested with different restriction endonucleases, size-fractionated by gel electrophoresis, and blotted to a DNA-binding membrane. The bound DNA fragments could then be hybridized with labeled DNA probes made from a cloned *factor VIII* gene. By analyzing the sizes of the DNA fragments that hybridize

with the probes, it should be possible to construct a restriction map of the mutant gene and compare it to a map of a nonmutant gene. This comparison should show the presence of an insertion in the mutant gene. It might also reveal the identity of the inserted sequence. (Alu elements are cleaved by a particular restriction endonuclease, Alu I, which could be one of the enzymes used in the analysis.) A simpler technique is to amplify portions of the coding sequence of the *factor VIII* gene by using the polymerase chain reaction (PCR). Pairs of primers positioned appropriately down the length of the coding sequence could be used in a series of amplification reactions, each of which would be seeded with template DNA from the hemophiliac. Each pair of primers would be expected to amplify a segment of the factor VIII gene. The sizes of the PCR products could then be determined by gel electrophoresis. An *Alu* insertion in a particular segment of the gene would increase the size of that segment by about 300 base pairs. The putative Alu insertion could be identified definitively by sequencing the DNA of the larger-than-normal PCR product.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

17.1 Which of the following pairs of DNA sequences could qualify as the terminal repeats of a bacterial IS element? Explain.

(a) 5'-GAATCCGCA-3' and 5'-ACGCCTAAG-3'
(b) 5'-GAATCCGCA-3' and 5'-CTTAGGCGT-3'
(c) 5'-GAATCCGCA-3' and 5'-GAATCCGCA-3'
(d) 5'-GAATCCGCA-3' and 5'-TGCGGATTC-3'

17.2 Which of the following pairs of DNA sequences could qualify as target site duplications at the point of an IS50 insertion? Explain.

(a) 5'-AATTCGCGT-3' and 5'-AATTCGCGT-3'
(b) 5'-AATTCGCGT-3' and 5'-TGCGCTTAA-3'
(c) 5'-AATTCGCGT-3' and 5'-TTAAGCGCA-3'
(d) 5'-AATTCGCGT-3' and 5'-ACGCGAATT-3'

- **17.3** One strain of *E. coli* is resistant to the antibiotic streptomycin, and another strain is resistant to the antibiotic ampicillin. The two strains were cultured together and then plated on selective medium containing streptomycin and ampicillin. Several colonies appeared, indicating that cells had acquired resistance to both antibiotics. Suggest a mechanism to explain the acquisition of double resistance.
- 17.4 What distinguishes IS and Tn3 elements in bacteria?
- **17.5** The circular order of genes on the *E. coli* chromosome is $^*A B C D E F G H^*$, with the * indicating that the ends of the chromosome are attached to each other. Two copies of an IS element are located in this chromosome, one

between genes C and D, and the other between genes Dand E. A single copy of this element is also present in the F plasmid. Two Hfr strains were obtained by selecting for integration of the F plasmid into the chromosome. During conjugation, one strain transfers the chromosomal genes in the order D E F G H A B C, whereas the other transfers them in the order D C B A H G F E. Explain the origin of these two Hfr strains. Why do they transfer genes in different orders? Does the order of transfer reveal anything about the orientation of the IS elements in the *E. coli* chromosome?

- 17.6 The composite transposon Tn5 consists of two IS50 elements, one on either side of a group of three genes for antibiotic resistance. The entire unit IS50L kan^r ble^r str^r IS50R can transpose to a new location in the *E. coli* chromosome. However, of the two IS50 elements in this transposon, only IS50R produces the catalytically active transposase. Would you expect IS50R to be able to be excised from the Tn5 composite transposon and inserted elsewhere in the chromosome? Would you expect IS50L to be able to do this?
- 17.7 By chance, an IS1 element has inserted near an IS2 element in the *E. coli* chromosome. The gene between them, *sug*⁺, confers the ability to metabolize certain sugars. Will the unit IS1 *sug*⁺ IS2 behave as a composite transposon? Explain.
- **17.8** A researcher has found a new Tn5 element with the structure IS50L *str^r ble^r kan^r* IS50L. What is the most likely origin of this element?

- **17.9** Would a Tn3 element with a frameshift mutation early in the *tnpA* gene be able to form a cointegrate? Would a Tn3 element with a frameshift mutation early in the *tnpR* gene be able to form a cointegrate?
- **17.10** What enzymes are necessary for replicative transposition of Tn *3*? What are their respective functions?
- 17.11 What is the medical significance of bacterial transposons?
- **17.12** Describe the structure of the *Ac* transposon in maize. In what ways do the *Ds* transposons differ structurally and functionally from the *Ac* transposon?
- **17.13** In homozygous condition, a deletion mutation of the c locus, c^n , produces colorless (white) kernels in maize; the dominant wild-type allele, C, causes the kernels to be purple. A newly identified recessive mutation of the c locus, c^m , has the same phenotype as the deletion mutation (white kernels), but when $c^m c^m$ and $c^n c^n$ plants are crossed, they produce white kernels with purple stripes. If it is known that the $c^n c^n$ plants harbor Ac elements, what is the most likely explanation for the c^m mutation?
- 17.14 In maize, the O2 gene, located on chromosome 7, controls the texture of the endosperm, and the C gene, located on chromosome 9, controls its color. The gene on chromosome 7 has two alleles, a recessive, o2, which causes the endosperm to be soft, and a dominant, O2, which causes it to be hard. The gene on chromosome 9 also has two alleles, a recessive, c, which allows the endosperm to be colored, and a dominant, C^{I} , which inhibits coloration. In one homozygous C^{I} strain, a Ds element is inserted on chromosome 9 between the C gene and the centromere. This element can be activated by introducing an Ac element by appropriate crosses. Activation of Ds causes the C¹ allele to be lost by chromosome breakage. In C¹/c/c kernels, such loss produces patches of colored tissue in an otherwise colorless background. A geneticist crosses a strain with the genotype o2/o2; C^{I} Ds/ C^{I} Ds to a strain with the genotype O2/o2; c/c. The latter strain also carries an Ac element somewhere in the genome. Among the offspring, only those with hard endosperm show patches of colored tissue. What does this tell you about the location of the Ac element in the O2/o2; c/c strain?
- **17.15** In maize, the recessive allele *bz* (*bronze*) produces a lighter color in the aleurone than does the dominant allele, *Bz*. Ears on a homozygous *bz/bz* plant were fertilized by pollen from a homozygous *Bz/Bz* plant. The resulting cobs contained kernels that were uniformly dark except for a few on which light spots occurred. Suggest an explanation.
- 17.16 The X-linked *singed* locus is one of several in *Drosophila* that controls the formation of bristles on the adult cuticle. Males that are hemizygous for a mutant *singed* allele have bent, twisted bristles that are often much reduced in size. Several *P* element insertion mutations of the *singed* locus have been characterized, and some have been shown to revert to the wild-type allele by excision of the inserted

element. What conditions must be present to allow such reversions to occur?

- 17.17 Dysgenic hybrids in *Drosophila* have elevated mutation rates as a result of *P* element transposition. How could you take advantage of this situation to obtain *P* element insertion mutations on the X chromosome?
- 17.18 If DNA from a *P* element insertion mutation of the *Drosophila white* gene and DNA from a wild-type *white* gene were purified, denatured, mixed with each other, renatured, and then viewed with an electron microscope, what would the hybrid DNA molecules look like?
- 17.19 When complete P elements are injected into embryos from an M strain, they transpose into the chromosomes of the germ line, and progeny reared from these embryos can be used to establish new P strains. However, when complete P elements are injected into embryos from insects that lack these elements, such as mosquitoes, they do not transpose into the chromosomes of the germ line. What does this failure to insert in the chromosomes of other insects indicate about the nature of P element transposition?
- **17.20** (a) What are retroviruslike elements? (b) Give examples of retroviruslike elements in yeast and *Drosophila*. (c) Describe how retroviruslike elements transpose. (d) After a retroviruslike element has been inserted into a chromosome, is it ever expected to be excised?
- **17.21** Sometimes solitary copies of the LTR of Ty*1* elements are found in yeast chromosomes. How might these solitary LTRs originate?
- **17.22** Would you ever expect the genes in a retrotransposon to possess introns? Explain.
- 17.23 Suggest a method to determine whether the *TART* retroposon is situated at the telomeres of each of the chromosomes in the *Drosophila* genome.
- 17.24 **(i)** It has been proposed that the *hobo* transposable elements in *Drosophila* mediate intrachromosomal recombination—that is, two *hobo* elements on the same chromosome pair and recombine with each other. What would such a recombination event produce if the *hobo* elements were oriented in the same direction on the chromosome? What if they were oriented in opposite directions?
- **17.25** What evidence suggests that some transposable elements are not simply genetic parasites?
- **17.26** Approximately half of all spontaneous mutations in *Drosophila* are caused by transposable element insertions. In human beings, however, the accumulated evidence suggests that the vast majority of spontaneous mutations are *not* caused by transposon insertions. Propose a hypothesis to explain this difference.
- 17.27 Z. Ivics, Z. Izsvák, and P. B. Hackett have "resurrected" a nonmobile transposable element isolated from the DNA of salmon. These researchers altered 12 codons within the coding sequence of the transposase gene of

the salmon element to restore the catalytic function of its transposase. The altered element, called *Sleeping Beauty*, is being tested as an agent for the genetic transformation of vertebrates such as mice and zebra fish (and possibly humans). Suppose that you have a bacterial plasmid containing the gene for green fluorescent protein (gfp) inserted between the ends of a *Sleeping Beauty* element. How would you go about obtaining mice or zebra fish that express the gfp gene?

17.28 The human genome contains about 5000 "processed pseudogenes," which are derived from the insertion of DNA copies of mRNA molecules derived from many different genes. Predict the structure of these pseudogenes. Would each type of processed pseudogene be expected to found a new family of retrotransposons within the human genome? Would the copy number of each type of processed pseudogene be expected to increase significantly over evolutionary time, as the copy number of the *Alu* family has? Explain your answers.

Genomics on the Web at http://www.ncbi.nlm.nih.gov

Here is a path to the sequence of a complete *P* element inserted in genomic DNA of *Drosophila melanogaster*:

Genomic biology \rightarrow Insects \rightarrow Drosophila melanogaster \rightarrow Resources: Flybase \rightarrow Files \rightarrow Transposons (Dmel) \rightarrow P-element \rightarrow Sequence Accession \rightarrow X06779.

- 1. Click on "repeat region (direct repeat)" to find the target site duplication created when this *P* element inserted into the genome. Note the length of the duplication and its sequence.
- **2.** Click on "repeat region (P element)" to find the 2907 basepair sequence of the *P* element itself; the first and last 31 base pairs are the terminal inverted repeats. Note the sequence.
- **3.** Copy the first line of nucleotides in the entire sequence (genomic DNA plus inserted *P* element) and use the BLAST function under the Tools tab on the Flybase web site to locate this sequence in the *D. melanogaster* genome (be sure to delete the spaces between segments of 10 nucleotides when you carry out your search). What chromosome is the insertion on, and what gene is it near? What phenotype is associated with mutations in this gene? Would the *P* element insertion be expected to cause a mutant phenotype?

18

Regulation of Gene Expression in Prokaryotes

CHAPTER OUTLINE

- Constitutive, Inducible, and Repressible Gene Expression
- Positive and Negative Control of Gene Expression
- Operons: Coordinately Regulated Units of Gene Expression
- The Lactose Operon in *E. coli*: Induction and Catabolite Repression
- The Tryptophan Operon in E. coli: Repression and Attenuation
- Translational Control of Gene Expression
- Posttranslational Regulatory Mechanisms

D'Hérelle's Dream of Treating Dysentery in Humans by Phage Therapy

In 1910, the French-Canadian microbiologist Felix d'Hérelle was in Mexico investigating a bacterial disease that was killing entire populations of locusts. The infected locusts developed severe diarrhea, excreting almost pure suspensions of bacilli prior to death. When he studied the bacteria in the feces of the locusts. d'Hérelle observed circular clear spots in the bacterial cultures grown on agar. However, when he examined the material in the clear spots microscopically, he could not see anything. In 1915, d'Hérelle returned to the Pasteur Institute in Paris, where he studied an epidemic of bacterial dysentery that was raging through army units stationed in France. He once again observed clear spots in lawns of bacteria. In addition, he demonstrated that whatever was killing the *Shigella*—a bacterium that causes dysentery in humans—could pass through a porcelain filter that retained all known bacteria. In 1917, d'Hérelle published his results and named the submicroscopic bacteriocidal agents bacteriophages (from the Greek for "bacteria-devouring"). About the same time, an English medical bacteriologist, Frederick W. Twort, discovered a similar submicroscopic agent that killed micrococci. Unfortunately, Twort's research was soon interrupted by a call to serve in the Royal Army Medical Corps in World War I.

Meanwhile, d'Hérelle continued to study the submicroscopic agents that killed *Shigella*. He provided the following account of one of his experiments: ".... in a flash I had understood: what caused my clear spots was in fact an invisible microbe, a filtrable virus, but a virus parasitic on bacteria.... 'If this is true, the same thing has probably occurred during the night in the sick man, who yesterday was in serious condition. In his intestine, as in my test tube, the dysentery bacilli will have dissolved away under the action of their parasite. He should now be cured.' I dashed to the hospital. In fact, during the night, his condition had greatly improved and convalescence was beginning" (d'Hérelle, F. 1949. The Bacteriophage. *Science News* 14:44–59).

Indeed, d'Hérelle became obsessed with his belief that human diseases caused by bacteria could be treated, perhaps even eradicated, by bacteriophage therapy. Unfortunately, it was soon demonstrated that this simple form of bacteriophage therapy is not effective in treating bacterial infections because, too frequently, the bacteria mutate to phage-resistant forms. Nevertheless, d'Hérelle's work set the stage for research that would eventually produce a whole new field—microbial genetics—and yield insights into the mechanisms by which gene expression is regulated. In this chapter,

we examine some of these mechanisms.



olorized electron micrograph of bacteriophage *lambda*.

Levels at which gene expression is regulated in prokaryotes

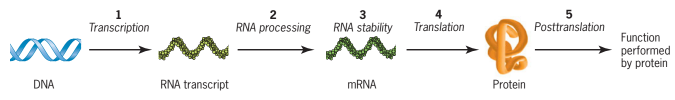


FIGURE 18.1 An abbreviated pathway of gene expression, showing five important levels of regulation in prokaryotes.

Microorganisms exhibit remarkable capacities to adapt to diverse environmental conditions. This adaptability depends in part on their ability to turn on and turn off the expression of specific sets of genes in response to changes in the environment. The expression of particular genes is turned on when the products of these genes are needed for growth. Their expression is turned off when the gene products are no longer needed. The synthesis of gene transcripts and translation products requires the expenditure of considerable energy. By turning off the expression of genes when their products are not needed, an organism can save energy and can utilize the conserved energy to synthesize products that maximize growth rate. What, then, are the mechanisms by which microorganisms regulate gene expression in response to changes in the environment?

Gene expression in prokaryotes is regulated at several different levels: transcription, mRNA processing, mRNA turnover, translation, and posttranslation (**■** Figure 18.1). However, the regulatory mechanisms with the largest effects on phenotype act at the level of transcription.

Based on what is known about the regulation of transcription, the various regulatory mechanisms seem to fit into two general categories:

- 1. Mechanisms that involve the rapid turn-on and turn-off of gene expression in response to environmental changes. Regulatory mechanisms of this type are important in microorganisms because of the frequent exposure of these organisms to sudden changes in environment. They provide microorganisms with considerable "plasticity," an ability to adjust their metabolic processes rapidly in order to achieve maximal growth and reproduction under a wide range of environmental conditions.
- 2. Mechanisms referred to as preprogrammed circuits or cascades of gene expression. In these cases, some event triggers the expression of one set of genes. The product(s) of one or more of these genes functions by turning off the transcription of the first set of genes or turning on the transcription of a second set of genes. Then, one or more of the products of the second set acts by turning on a third set, and so on. In these cases, the sequential expression of genes is genetically preprogrammed, and the genes cannot usually be turned on out of sequence. Such preprogrammed sequences of gene expression are well documented in prokaryotes and the viruses that attack them. For example, when a lytic bacteriophage infects a bacterium, the viral genes are expressed in a predetermined sequence, and this sequence is directly correlated with the temporal sequence of gene-product involvement in the reproduction and morphogenesis of the virus. In most of the known examples of preprogrammed gene expression, the circuitry is cyclical. For example, during viral infections, some event associated with the packaging of the viral DNA or RNA in the protein coat resets the genetic program so that the proper sequence of gene expression occurs once again when a progeny virus infects a new host cell.

Constitutive, Inducible, and Repressible Gene Expression

Genes that specify cellular components that perform housekeeping functions—for example, the ribosomal RNAs and proteins involved in protein synthesis—are expressed constitutively. Other genes often are expressed only when their products are required for growth.

Certain gene products—such as tRNA molecules, rRNA molecules, ribosomal proteins, RNA polymerase subunits, and enzymes catalyzing metabolic processes that are frequently referred to as cellular "housekeeping" functions—are essential components of almost all living cells. Genes that specify products of this type are continually being expressed in most cells. Such genes are said to be expressed constitutively and are referred to as **constitutive genes**.

Other gene products are needed for cell growth only under certain environmental conditions. Constitutive synthesis of such gene products would be wasteful, using energy that could otherwise be utilized for more rapid growth. The evolution of regulatory mechanisms that provide for the synthesis of such gene products only when

> and where they are needed would clearly endow the organisms that possess these regulatory mechanisms with a selective advantage over organisms that lack them. This undoubtedly explains why currently existing organisms, including bacteria and viruses, exhibit highly efficient mechanisms for the control of gene expression.

> Escherichia coli and most other bacteria are capable of growth using any one of several carbohydrates-for example, glucose, sucrose, galactose, arabinose, and lactose-as an energy source. If glucose is present in the environment, it will be preferentially metabolized by E. coli cells. However, in the absence of glucose, E. coli cells can grow very well on other carbohydrates. Cells growing in medium containing the sugar lactose, for example, as the sole carbon source synthesize two enzymes, β-galactosidase and β -galactoside permease, which are uniquely required for the catabolism of lactose. β-Galactoside permease pumps lactose into the cell, where β -galactosidase cleaves it into glucose and galactose. Neither of these enzymes is of any use to *E. coli* cells if no lactose is available to them. The synthesis of these two enzymes requires considerable energy (in the form of ATP and GTP; see Chapters 11 and 12). Thus, E. coli cells have evolved a regulatory mechanism by which the synthesis of these lactose-catabolizing enzymes is turned on in the presence of lactose and turned off in its absence.

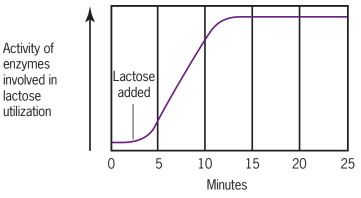
> In natural environments (intestinal tracts and sewers), *E. coli* cells probably encounter an absence of glucose and the presence of lactose relatively infrequently. Therefore, the *E. coli* genes encoding the enzymes involved in lactose utilization are probably turned off most of the time. If cells growing on a carbohydrate other than lactose are transferred to medium containing lactose as the only carbon source, they quickly begin to synthesize the enzymes required for lactose utilization (**Theorem 18.2a**). This process of turning on the expression of genes in response to a substance in the environment is called **induction**. Genes whose expression is regulated in this manner are called **induction ducible genes**; their products, if enzymes, are called **inducible enzymes**.

Enzymes that are involved in **catabolic** (degradative) **pathways**, such as in lactose, galactose, or arabinose utilization, are characteristically inducible. As we discuss later in this chapter, induction occurs at the level of transcription. Induction alters the rate of enzyme

synthesis, not the activity of existing enzyme molecules. Induction should not be confused with enzyme activation, which occurs when the binding of a small molecule to an enzyme increases the activity of the enzyme, but does not affect its rate of synthesis.

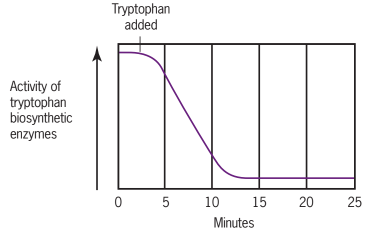
Bacteria can synthesize most of the organic molecules required for growth, such as amino acids, purines, pyrimidines, and vitamins. For example, the *E. coli* genome

Induction of enzyme synthesis



(a)

Repression of enzyme synthesis



(b)

■ FIGURE 18.2 (a) Induction of the synthesis of enzymes required for the utilization of lactose as an energy source and (b) repression of the synthesis of the enzymes required for the biosynthesis of tryptophan, both in *E. coli*. Note that low levels of enzyme synthesis occur whether the metabolites are present or absent. contains five genes encoding enzymes that catalyze steps in the biosynthesis of tryptophan. These five genes must be expressed in *E. coli* cells growing in an environment devoid of tryptophan in order to provide adequate amounts of this amino acid for ongoing protein synthesis.

When *E. coli* cells are present in an environment containing enough tryptophan to support optimal growth, the continued synthesis of the tryptophan biosynthetic enzymes would be a waste of energy. Thus, a regulatory mechanism has evolved in *E. coli* that turns off the synthesis of the tryptophan biosynthetic enzymes when external tryptophan is available (**Tigure 18.2b**). A gene whose expression has been turned off in this way is said to be "repressed"; the process is called **repression**. When the expression of this gene is turned on, it is said to be "derepressed"; such a response is called **derepression**.

Enzymes that are components of **anabolic** (biosynthetic) **pathways** often are repressible. Repression, like induction, occurs at the level of transcription. Repression should not be confused with feedback inhibition, which occurs when the product of a biosynthetic pathway binds to and inhibits the activity of the first enzyme in the pathway, but does not affect the synthesis of the enzyme.

- In prokaryotes, genes that specify housekeeping functions such as rRNAs, tRNAs, and ribosomal proteins are expressed constitutively. Other genes usually are expressed only when their products are needed.
- Genes that encode enzymes involved in catabolic pathways often are expressed only in the presence of the substrates of the enzymes; their expression is inducible.
- Genes that encode enzymes involved in anabolic pathways usually are turned off in the presence of the end product of the pathway; their expression is repressible.
- Although gene expression can be regulated at many levels, transcriptional regulation is the most common.

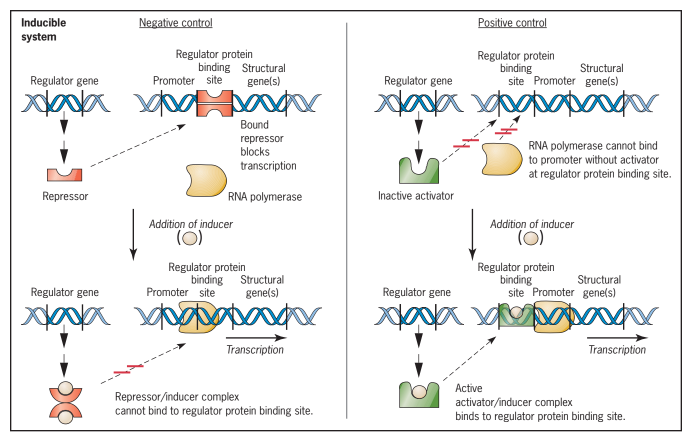
Positive and Negative Control of Gene Expression

The regulation of gene expression—induction, or turning genes on, and repression, or turning genes off—can be accomplished by both positive control mechanisms and negative control mechanisms. Both mechanisms involve the participation of **regulator genes**—genes encoding products that regulate the expression of other genes. In **positive control mecha**- In some cases, the product of a regulatory gene is required to initiate the expression of one or more genes. In other cases, the product of a regulatory gene is required to turn off the expression of one or more genes.

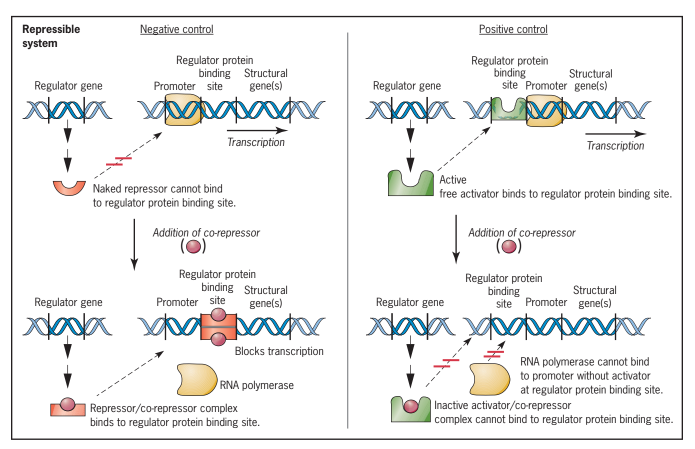
nisms, the product of the regulator gene is required to turn on the expression of one or more structural genes (genes specifying the amino acid sequences of enzymes or structural proteins), whereas in **negative control mechanisms**, the product of the regulator gene is necessary to shut off the expression of structural genes. Positive and negative regulation are illustrated for both inducible and repressible systems in **Figure 18.3**.

Recall that a gene is expressed when RNA polymerase binds to its promoter and synthesizes an RNA transcript that contains the coding region of the gene (Chapter 11). The product of the regulator gene acts by binding to a site called the regulator protein-binding site (*RPBS*) adjacent to the promoter of the structural gene(s). When the product of the regulator gene is bound at the *RPBS*, transcription of the structural gene(s) is turned on in a positive control system (Figure 18.3, right) or turned off in a negative control system (Figure 18.3, left). The regulator gene products are called **activators**—because they activate gene expression—in positive control systems, and **repressors**—because they repress gene expression—in negative control systems. Whether or not a regulator protein can bind to the *RPBS* depends on the presence or absence of **effector molecules** in the cell. The effectors are usually small molecules such as amino acids, sugars, and similar metabolites. The effector molecules involved in repression of gene expression are called **inducers**; those involved in repression of gene expression are called **co-repressors**.

KEY POINTS



(a)



(b)

FIGURE 18.3 Negative and positive control of inducible (*a*) and repressible (*b*) gene expression. The regulator gene product is required to turn on gene expression in positive control systems and to turn off gene expression in negative control systems.

The effector molecules (inducers and co-repressors) bind to regulator gene products (activators and repressors) and cause changes in the three-dimensional structures of these proteins. Conformational changes in protein structure resulting from the binding of small molecules are called **allosteric transitions**. Conformational changes in proteins frequently result in alterations in their activity. In the case of activators and repressors, allosteric transitions caused by the binding of effector molecules usually alter their ability to bind to regulator protein-binding sites adjacent to the promoters of the structural genes they control.

In a negative, inducible control mechanism (Figure 18.3*a*, left), the free repressor binds to the *RPBS* and prevents the transcription of the structural gene(s) in the absence of inducer. When inducer is present, it is bound by the repressor, and the repressor/inducer complex cannot bind to the *RPBS*. With no repressor bound to the *RPBS*, RNA polymerase binds to the promoter and transcribes the structural gene(s). In a positive, inducible control mechanism (Figure 18.3*a*, right), the activator cannot bind to the *RPBS* unless inducer is present, and RNA polymerase cannot transcribe the structural gene(s) unless the activator/inducer complex is bound to the *RPBS*. Thus, transcription of the structural genes is turned on only in the presence of inducer.

In a negative, repressible regulatory mechanism (Figure 18.3*b*, left), transcription of the structural gene(s) occurs in the absence of the co-repressor, but not in its presence. When the repressor/co-repressor complex is bound to the *RPBS*, it prevents RNA polymerase from transcribing the structural genes. In the absence of co-repressor, free repressor cannot bind to the *RPBS*; thus, RNA polymerase can bind to the promoter and transcribe the structural genes. In a positive, repressible control mechanism (Figure 18.3*b*, right), the product of the regulator gene, the activator, must be bound to the *RPBS* in order for RNA polymerase to bind to the promoter and transcribe the structural gene(s). When co-repressor is present, it forms a complex with the activator protein, and this activator/co-repressor complex is unable to bind to the *RPBS*; consequently, RNA polymerase cannot bind to the promoter and transcribe the structural gene(s).

In order to understand the details of these four mechanisms of regulation, focus on the key differences between them. (1) The regulator gene product, the activator, participates in turning on gene expression in a positive control mechanism, whereas the regulator gene product, the repressor, is involved in turning off gene expression in a negative control mechanism. (2) With both positive and negative control mechanisms, whether gene expression is inducible or repressible depends on whether the free regulator protein or the regulator protein/effector molecule complex binds to the regulator protein-binding site (*RPBS*).

- Gene expression is controlled by both positive and negative regulatory mechanisms.
- In positive control mechanisms, the product of a regulator gene, an activator, is required to turn on the expression of the structural gene(s).
- In negative control mechanisms, the product of a regulator gene, a repressor, is necessary to turn off the expression of the structural gene(s).
- Activators and repressors regulate gene expression by binding to sites adjacent to the promoters of structural genes.
- Whether or not the regulator proteins can bind to their binding sites depends on the presence or absence of small effector molecules that form complexes with the regulator proteins.
- The effector molecules are called inducers in inducible systems and co-repressors in repressible systems.

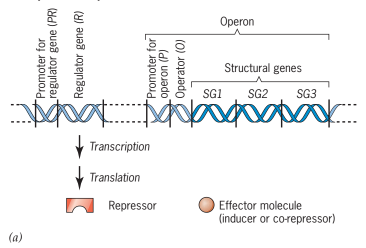
Operons: Coordinately Regulated Units of Gene Expression

The **operon model**, a negative control mechanism, was developed in 1961 by François Jacob and Jacques Monod to explain the regulation of genes required for lactose utilization in *E. coli*. We discuss some of the experimental results

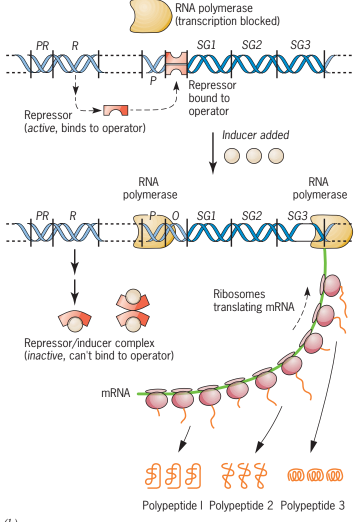
In prokaryotes, genes with related functions often are present in coordinately regulated genetic units called operons.

KEY POINTS





The operon: induction



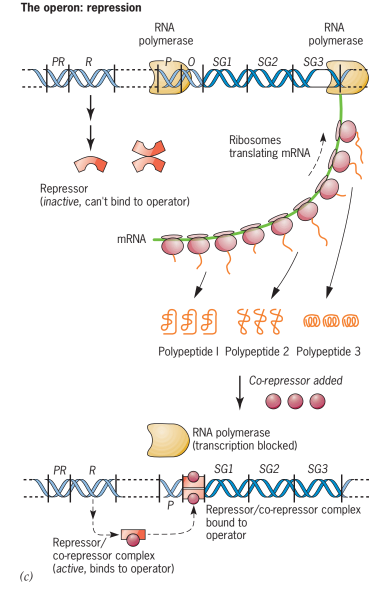


FIGURE 18.4 Regulation of gene expression by the operon mechanism. (a) Components of an operon: one or more structural genes (three, SG1, SG2, and SG3, are shown) and the adjoining operator (0) and promoter (P) sequences. One operator and one promoter are shown; however, some operons have multiple operators and promoters. The transcription of the regulator gene (R) is initiated by RNA polymerase, which binds to its promoter (PR). When repressor is bound to the operator, it sterically prevents RNA polymerase from initiating transcription of the structural genes. The difference between an inducible operon (b) and a repressible operon (c) is that free repressor binds to the operator(s) of an inducible operon, whereas the repressor/effector molecule complex binds to the operator(s) of a repressible operon. Thus, an inducible operon is turned off in the absence of the effector (inducer) molecule, and a repressible operon is turned on in the absence of the effector (co-repressor) molecule.

that led to the development of this model in A Milestone in Genetics: Jacob, Monod, and the Operon Model on the Student Companion site. Jacob and Monod proposed that the transcription of a set of contiguous structural genes is regulated by two controlling elements (**Figure 18.4a**). One of the elements, the repressor gene, encodes a repressor, which (under the appropriate conditions) binds to the second element, the **operator**. The operator is always contiguous with the structural genes whose expression it regulates. Some operons—including the lactose operon discussed in the next section—contain multiple operators; however, for now, we will consider only a single operator to keep the mechanism as simple as possible.

Transcription is initiated at promoters located just upstream (5') from the coding regions of structural genes. When repressor is bound to the operator, it sterically prevents RNA polymerase from transcribing the structural genes in the operon. Operator regions are contiguous with promoter regions; sometimes operators and promoters even overlap, sharing a short DNA sequence. Operator regions are often located between the promoters and the structural genes that they regulate. The complete contiguous unit, including the structural genes, the operator, and the promoter, is called an **operon** (Figure 18.4*a*).

Whether the repressor will bind to the operator and turn off the transcription of the structural genes in an operon is determined by the presence or absence of effector molecules as discussed in the preceding section. Inducible operons and repressible operons can be distinguished from one another by determining whether the naked repressor or the repressor/effector molecule complex is active in binding to the operator.

- 1. In the case of an inducible operon, the free repressor binds to the operator, turning off transcription (Figure 18.4*b*).
- **2.** In the case of a repressible operon, the situation is reversed. The free repressor cannot bind to the operator. Only the repressor/effector molecule (co-repressor) complex is active in binding to the operator (Figure 18.4*c*).

Except for this difference in the operator-binding behavior of the free repressor and the repressor/effector molecule complex, inducible and repressible operons are identical.

A single mRNA transcript carries the coding information of an entire operon. Thus, the mRNAs of operons consisting of more than one structural gene are multigenic. For example, the tryptophan operon mRNA of *E. coli* contains the coding sequences of five different genes. Because they are co-transcribed, all structural genes in an operon are coordinately expressed.

Although the molar quantities of the different gene products need not be the same (because of different efficiencies of initiation of translation), the relative amounts of the different polypeptides specified by genes in an operon usually remain the same, regardless of the state of induction or repression. In some cases, the differential use of transcription termination signals can alter the amounts of gene products synthesized.

- In bacteria, genes with related functions frequently occur in coordinately regulated units called operons.
- Each operon contains a set of contiguous structural genes, a promoter (the binding site for RNA polymerase), and an operator (the binding site for a regulatory protein called a repressor).
- When a repressor is bound to the operator, RNA polymerase cannot transcribe the structural genes in the operon. When the operator is free of repressor, RNA polymerase can transcribe the operon.

The Lactose Operon in *E. coli:* Induction and Catabolite Repression

Jacob and Monod proposed the operon model based on their studies of the lactose (*lac*) operon in *E. coli* (see A Milestone in Genetics: Jacob, Monod, and the Operon Model on the Student Companion site). The *lac* operon contains a promoter (*P*), three operators The structural genes in the *lac* operon are transcribed only when lactose is present and glucose is absent.

KEY POINTS

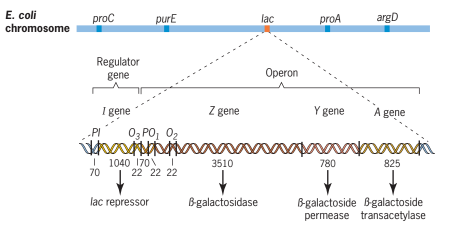
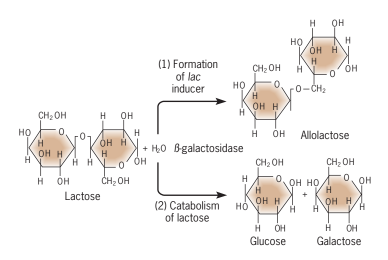


FIGURE 18.5 The *lac* operon of *E. coli*. The *lac* operon consists of three structural genes, *Z*, *Y*, and *A*, plus the promoter (*P*) and three operators $(O_1, O_2, and O_3)$. The regulator gene (*I*) is contiguous with the operon in the case of *lac* and has its own promoter (*PI*). The numbers below the various genetic elements indicate their sizes in nucleotide pairs.

 $(O_1, O_2, \text{ and } O_3)$, and three structural genes, *lacZ*, *lacY*, and *lacA*, encoding the enzymes β -galactosidase, β -galactoside permease, and β -galactoside transacetylase, respectively (**Figure 18.5**). β -galactoside permease "pumps" lactose into the cell, where β -galactosidase cleaves it into glucose and galactose (**Figure 18.6**). The biological role of the transacetylase is unknown.

In Jacob and Monod's model, the *lac* operon contained a single operator (now designated O_1). However, two additional operators (O_2 and O_3) were subsequently discovered. Initially, O_2 and O_3 were thought to play very minor roles. Then, Benno Müller-Hill and coworkers demonstrated that the deletion of both "minor" operators had a large effect on the level of transcription of the operon. More recent studies have shown that efficient repression of the *lac* operon requires the major operator (O_1) and at least one of the minor operators (O_2 or O_3) and maximum repression requires all three operators. Nevertheless, we will first discuss Jacob and Monod's model of the *lac* operon, which involved only one operator, now designated O_1 . Then, we will extend the model and examine the functions of all three operators in the section entitled Protein–DNA Interactions That Control Transcription of the *lac* Operon.



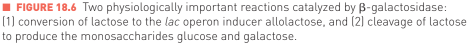


TABLE 18.1

Phenotypic Effects of Mutations in the Repressor Gene	and the	• Operator (<i>0</i>) Re	gion of the <i>lac</i> Operon

	β -Galactosid	lase Activityª	β -Galactoside Permease Activity ^a		
Genotype	With Lactose	Without Lactose	With Lactose	Without Lactose	Deduction
$I^+P^+O^+Z^+Y^+$	100 units	1 unit	100 units	1 unit	Wild-type is inducible
<i>I+P+0+Z+Y+/</i> F' <i>I+P+0+<mark>Z-Y-</mark></i>	100 units	1 unit	100 units	1 unit	Z^+ is dominant to Z^-
					Y ⁺ is dominant to <mark>Y⁻</mark>
I+P+O+Z+Y+/F'I+P+O+Z+Y+	200 units	2 units	200 units	2 units	Activity depends on gene dosage
$I^{-}P^{+}O^{+}Z^{+}Y^{+}$	100 units	100 units	100 units	100 units	<i>lacl</i> -mutants are constitutive
I+P+0+Z+Y+/F'I-P+0+Z+Y+	200 units	2 units	200 units	2 units	/+ is dominant to <mark>/-</mark>
$I^+P^+ O^c Z^+Y^+$	100 units	100 units	100 units	100 units	lacO ^c mutants are constitutive
<i>I+P+<mark>0</mark>^cZ+<mark>Y-</mark>/F'I+P+0+<mark>Z-</mark>Y+</i>	100 units	100 units	100 units	1 unit	O ^c and O ⁺ are <i>cis</i> -acting regulators

^aActivity levels in wild-type bacteria have been set at 100 units for both β -galactosidase (the product of gene Z) and β -galactoside permease (the product of gene Y). The A gene and its product β -galactoside transacetylase are not shown for the sake of brevity.

INDUCTION

The *lac* operon is a negatively controlled inducible operon; the *lacZ*, *lacY*, and *lacA* genes are expressed only in the presence of lactose. The *lac* regulator gene, designated the *I* gene, encodes a repressor that is 360 amino acids long. However, the active form of the *lac* repressor is a tetramer containing four copies of the *I* gene product. In the absence of inducer, the repressor binds to the *lac* operators, which in turn prevents RNA polymerase from catalyzing the transcription of the three structural genes (see Figure 18.4*b*). (Note: only the original operator (O_1) discovered by Jacob and Monod is shown in Figures 18.4, 18.7, and 18.8.) A few molecules of the *lacZ*, *lacY*, and *lacA* gene products are synthesized in the uninduced state, providing a low background level of enzyme activity. This background activity is essential for induction of the *lac* operon because the inducer of the operon, allolactose, is derived from lactose in a reaction catalyzed by β -galactosidase (Figure 18.6). Once formed, allolactose is bound by the repressor, causing the release of the repressor from the operator. In this way, allolactose induces the transcription of the *lacZ*, *lacY*, and *lacA* structural genes (see Figure 18.4*b*).

The *lacI* gene, *lac* operator O_1 , and the *lac* promoter were all initially identified genetically by the isolation of mutant strains that exhibited altered expression of the *lac* operon genes. Mutations in the *I* gene and the operator frequently result in constitutive synthesis of the *lac* gene products. These mutations are designated I^- and O^c , respectively. The I^- and O^c constitutive mutations can be distinguished not only by map position, but also by their behavior in partial diploids in which they are located in *cis* and *trans* configurations relative to mutations in *lac* structural genes (**Table 18.1**). Recall that partial diploids can be constructed using fertility (F) factors that carry chromosomal genes—F' factors (Chapter 8). F' factors that carry the *lac* operon have been used to study the interactions between the various components of the operon.

Like monoploid wild-type $(I^+P^+O^+Z^+Y^+A^+)$ cells, partial diploids (also called "merozygotes") of genotype F' $I^+P^+O^+Z^+A^+/I^+P^+O^+Z^-Y^-A^-$ or of genotype F' $I^+P^+O^+Z^-Y^-A^-/I^+P^+O^+Z^+Y^+A^+$ are inducible for the utilization of lactose as a carbon source. The wild-type alleles $(Z^+, Y^+, \text{ and } A^+)$ of the three structural genes are dominant to their mutant alleles $(Z^-, Y^-, \text{ and } A^-)$. This dominance is expected because the wild-type alleles produce functional enzymes, whereas the mutant alleles produce no enzymes or defective (inactive) enzymes. Partial diploids of genotype $I^+P^+O^+Z^+Y^+A^+/I^-P^+O^+Z^+Y^+A^+$ (I^+/I^-) are also inducible for the synthesis of the three enzymes specified by the *lac* operon. Thus, I^+ is dominant to I^- as expected, because I^+ encodes a functional repressor molecule and its I^- allele specifies an inactive repressor. The dominance of I^+ over I^- also indicates that the repressor is diffusible, because

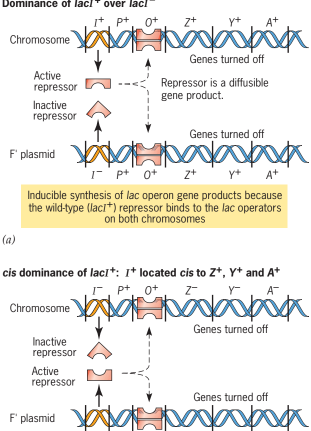
Solve It!

Constitutive Mutations in the *E. coli lac* Operon

You have isolated two mutants of E. coli K12 that synthesize β-galactosidase, β-galactoside permease, and β-galactoside transacetylase constitutively, that is, whether or not lactose is present in the medium. You next introduce an F' that carries wild-type copies of the lacl gene, the lac promoter, and the three lac operators, but contains a deletion of the distal segment of *lacZ* and all of *lacY* and *lacA*, into each of your constitutive mutants. The resulting partial diploid containing constitutive mutant 1 continues to synthesize the three lactose catabolic enzymes constitutively, whereas the partial diploid containing constitutive mutant 2 exhibits inducible synthesis of the three enzymes. Explain the difference between mutants 1 and 2.

To see the solution to this problem, visit the Student Companion site.

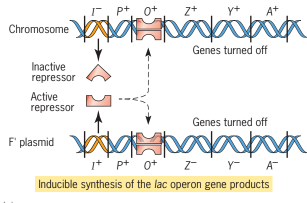
Dominance of lac1⁺ over lac1⁻







Inducible synthesis of the lac operon gene products



(c)

FIGURE 18.7 Studies of *E. coli* partial diploids have shown that the *lacl*⁺ gene is dominant to lacl- alleles (a) and controls lac operators located either cis (b) or trans (c) to itself. These effects demonstrate that the lacl gene product is diffusible. Although the functional form of the *lac* repressor is a tetramer, the two molecules at the back of the tetramer are not shown for the sake of simplicity.

the repressor produced by the *lacI*⁺ allele on one chromosome can turn off the *lac* structural genes on both operons in the cell (**Figure 18.7***a*).

Like wild-type cells, partial diploids of genotype F' $I^+P^+O^+Z^+Y^+A^+/$ $I^{-}P^{+}O^{+}Z^{-}Y^{-}A^{-}$ or genotype F' $I^{+}P^{+}O^{+}Z^{-}Y^{-}A^{-}/I^{-}P^{+}O^{+}Z^{+}Y^{+}A^{+}$ are inducible for β -galactosidase, β -galactoside permease, and β -galactoside transacetylase. The inducibility of these genotypes demonstrates that the *lac* repressor (I^+ gene product) controls the expression of structural genes located either *cis* (**I** Figure 18.7*b*) or *trans* (**I** Figure 18.7*c*) to the $lacI^+$ allele.

The operator constitutive (O^{c}) mutations act only in *cis*; that is, O^{c} mutations affect the expression of only those structural genes located on the same chromosome. The *cis*-acting nature of O mutations is logical given the function of the operator. O mutations should not act in trans if the operator is the binding site for the repressor; as such, the operator does not encode any product, diffusible or otherwise. A regulator gene should act in trans only if it specifies a diffusible product. Therefore, a partial diploid of genotype F' $I^+P^+O^c Z^-Y^-A^-/I^+P^+O^+Z^+Y^+A^+$ is inducible for the three enzymes specified by the structural genes of the *lac* operon (Table 18.2, ■ Figure 18.8a), whereas a partial diploid of genotype F' $I^+P^+O^c Z^+Y^+A^+/I^+P^+O^+Z^-Y^-A^-$ synthesizes these enzymes constitutively (Table 18.2, Figure 18.8b). Once you are confident that you understand how the components of the operon interact to regulate the transcription of the lac structural genes, try Solve It: Constitutive Mutations in the E. coli lac Operon and see Problem-Solving Skills: Testing Your Understanding of the lac Operon.

Some of the I gene mutations, those designated I^{-d} , are dominant to the wild-type allele (I^+) . This dominance results from the inability of heteromultimers (proteins composed of two or more different forms of a polypeptide; recall that the *lac* repressor functions as a tetramer) that contain both wild-type and mutant polypeptides to bind to the operator. Other I gene mutations, those designated I^s (s for superrepressed), cause the lac operon to be uninducible. In strains carrying these I' mutations, the lac structural genes can usually be induced to some degree with high concentrations of inducer, but they are not induced at normal concentrations of inducer. When studied *in vitro*, the mutant I^s polypeptides form tetramers that bind to *lac* operator DNA. However, they either do not bind inducer or exhibit a low affinity for inducer. Thus, the I^s mutations alter the inducer binding site of the *lac* repressor.

Promoter mutations do not change the inducibility of the lac operon. Instead, they modify the levels of gene expression in the induced and uninduced state by changing the frequency of initiation of lac operon transcription—that is, the efficiency of RNA polymerase binding.

The *lac* promoter actually contains two separate components: (1) the RNA polymerase binding site and (2) a binding site for another protein called catabolite activator protein (abbreviated CAP) that prevents the lac operon from being induced in the presence of glucose. This second control circuit, which we consider next, assures the preferential utilization of glucose as an energy source when it is available.

CATABOLITE REPRESSION

The presence of glucose has long been known to prevent the induction of the lac operon, as well as other operons controlling enzymes involved in carbohydrate catabolism. This phenomenon, called **catabolite repression** (or the *glucose effect*), assures that glucose is metabolized when present, in preference to other, less efficient, energy sources.

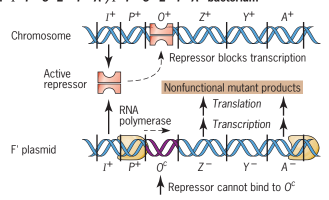
The catabolite repression of the *lac* operon and several other operons is mediated by a regulatory protein called CAP (for catabolite activator protein) and a small effector molecule called cyclic AMP (adenosine-3', 5'-monophosphate; abbreviated cAMP)

	β -Galactosidase Activity ^a		β -Galactoside Permease Activity ^a		
Genotype	With Lactose	Without Lactose	With Lactose	Without Lactose	Deduction
$I^{+}P^{+}O^{+}Z^{+}Y^{+}$	100 units	1 unit	100 units	1 unit	Wild-type is inducible
I+P+O+ Z+Y+/F' I-P+O+ <mark>Z-Y-</mark>	100 units	1 unit	100 units	1 unit	
$I^{-}P^{+}O^{+}Z^{+}Y^{+}/F'$ $I^{+}P^{+}O^{+}Z^{-}Y^{-}$	100 units	1 unit	100 units	1 unit	I ⁺ acts both <i>cis</i> and <i>trans</i>
I+P+O+Z+Y+/F'I+P+O-Z-Y-	100 units	1 unit	100 units	1 unit	0 ⁺ acts only in <i>cis</i>
$I^+P^+O^+Z^-Y^-/F'I^+P^+O^cZ^+Y^+$	100 units	100 units	100 units	100 units	O ^c acts only in <i>cis</i>

Inducible synthesis of the lac operon gene products in an F' $I^+ P^+ O^c Z^- Y^- A^- / I^+ P^+ O^+ Z^+ Y^+ A^+$ bacterium

TARI F 18 2

(a)



Inducer absent; no functional *lac* operon gene products are synthesized

Constitutive synthesis of the lac operon gene products in an F' $I^+ P^+ O^c Z^+ Y^+ A^+ / I^+ P^+ O^+ Z^- Y^- A^-$ bacterium

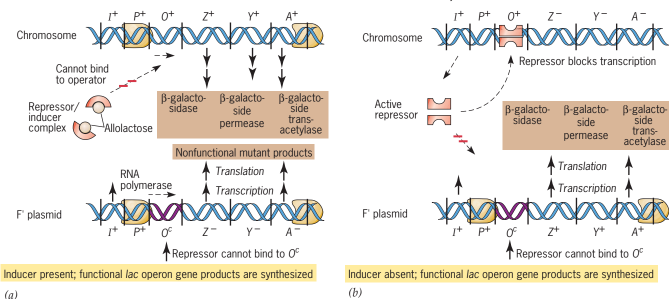


FIGURE 18.8 Studies of *E. coli* partial diploids have shown that the operator acts only in the *cis* configuration. The synthesis of functional β -galactosidase, β -galactoside permease, and β -galactoside transacetylase is (a) inducible in a partial diploid of genotype F' $I^+P^+O^cZ^-Y^-A^-/I^+P^+O^+Z^+Y^+A^+$ and (b) constitutive in a partial diploid of genotype F' $I^+P^+O^cZ^+Y^+A^+/I^+P^+O^+Z^-Y^-A^-$. These results demonstrate that the operator (0) is *cis*-acting; that is, it regulates only those structural genes located on the same chromosome.

PROBLEM-SOLVING SKILLS

Testing Your Understanding of the lac Operon

THE PROBLEM

The following table gives the relative activities of the enzymes β -galactosidase and β -galactoside permease in cells with different genotypes at the *lac* locus in *E. coli*. The induced level of activity of each enzyme in wild-type *E. coli* cells that do **not** carry an F' was arbitrarily set at 100 units, and all other enzyme levels were measured relative to the levels observed in these wild-type cells. Based on the data given in the table for genotypes 1 through 4, fill in the levels of activity that would be expected for genotype 5 in the spaces (parentheses) provided.

	β -Galactosidase		β-Galactoside Permease		
Genotype	- inducer	+inducer	— inducer	+ inducer	
1. <i>I</i> + <i>O</i> + <i>Z</i> + <i>Y</i> +	0.2	100	0.2	100	
2. <i>I-O+Z+Y</i> +	100	100	100	100	
3. $I^+ O^c Z^+ Y^+$	75	100	75	100	
4. <i>I-0+Z+Y-</i> / F' <i>I-0+Z+Y+</i>	200	200	100	100	
5. I-0°Z-Y+ / F' I+0+Z+Y+	[]	[]	[]	[]	

FACTS AND CONCEPTS

- 1. The *lacZ* and *lacY* genes encode the enzymes β -galactosidase and β -galactoside permease, respectively. β -galactoside permease transports lactose into cells where β -galactosidase cleaves it into glucose and galactose. The *lacZ*⁺ and *lacY*⁺ alleles of these genes encode functional enzymes, whereas the *lacZ*⁻ and *lacY*⁻ alleles encode nonfunctional gene products.
- 2. In wild-type *E. coli* cells, the *lacZ*⁺ and *lacY*⁺ genes are transcribed only in the presence of lactose. Their transcription is repressed (turned off) in the absence of lactose when β -galactosidase and β -galactoside permease have nothing to catabolize or transport. Their transcription is induced when lactose is added to the medium in which the cells are growing (see Figure 18.4*b*).
- **3.** Constitutive mutants of *E. coli* synthesize β -galactosidase and β -galactoside permease continually whether or not lactose is present. These constitutive muations are of two types and map at two distinct sites in and near the *lac* operon on the *E. coli* chromosome. Some of the constitutive mutations—*lacl⁻* mutations—map in the gene that encodes the *lac* repressor; others—*lacO^c* mutations—map in the operator region—the site where the *lac* repressor binds.
- 4. The lac repressor (lacl⁺ gene product) binds to the lac operator (0) and prevents RNA polymerase from binding to the lac promoter and transcribing the genes in the lac operon (see Figure 18.4). The lacl⁻ mutant alleles encode inactive repressors that cannot bind to the lac operator. Allele lacl⁺ is dominant to lacl⁻.
- **5.** The *lac* repressor is a diffusible protein; thus, *lacl*⁺ regulates the expression of *lac* operon genes located both *cis* (on the same chromosome) and *trans* (on a different chromosome) to it. Regulatory elements of this type are said to be *cis* and *trans* acting.

6. The wild-type operator (0⁺) contains a nucleotide sequence that functions as a binding site for the *lac* repressor. Operator-constitutive (0^c) mutants contain an operator with an altered nucleotide sequence (often a deletion) to which the *lac* repressor either no longer binds or binds inefficiently. Thus, the constitutive level of enzyme synthesis will depend on whether the repressor binds to the mutant operator weakly or not at all. Because *lacO*⁺ and *lacO*^c operators only regulate the expression of *lac* genes on the same chromosome, they are called *cis*-acting regulators.

7. The amount of β -galactosidase and β -galactoside permease synthesized in a cell depends on the number of functional copies of the *lacZ*⁺ and *lacY*⁺ genes in the cell.

ANALYSIS AND SOLUTION

- **1.** The data given for genotype 1 $(I^+O^+Z^+Y^+ = \text{wild-type})$ show that these cells synthesize 0.2 unit of each enzyme in the absence of lactose and 100 units in the presence of lactose.
- **2.** The data for genotype 2 $(I^-O^+Z^+Y^+)^+$ = repressor-constitutive mutant) show that in the absence of a functional repressor cells synthesize 100 units of each enzyme whether lactose is present or absent.
- **3.** The operator-constitutive mutant (genotype 3, $l^+O^cZ^+Y^+$) in this question makes 75 units of each enzyme in the absence of lactose and 100 units in the presence of lactose. Although enzyme synthesis is constitutive, there is some binding of the *lac* repressor to the *lac* operator in the absence of lactose. When lactose is present, that binding no longer occurs, and synthesis of the *lac* enzymes increases to the fully induced level (100 units).
- 4. The data presented for genotype 4 (the partial diploid I⁻O⁺Z⁺Y⁻/ F' I⁻O⁺Z⁺Y⁺) shows the effect of gene dosage. Cells make twice as much enzyme when two copies of a wild-type gene are present as when only one is present.
- **5.** Genotype 5 $(I^-O^c Z^- Y^+/F' I^+O^+Z^+Y^+)$ is a partial diploid with two copies of the *lac* operon. It has two copies of Y^+ , but only one copy of Z^+ . It has an I^+ allele on the F', so functional repressor will be present in the cells. Transcription of chromosomal genes will be controlled by O^c , whereas transcription of genes on the F' will be controlled by O^+ . All of the β -galactosidase will be produced by the Z^+ allele on the F'; there is a Z^- mutation on the chromosome. The F' contains a wild-type *lac* operon, so 0.2 unit of β -galactosidase will be synthesized in the absence of lactose, and 100 units will be synthesized in the presence of lactose. In the case of β -galactoside permease, the contributions of both copies of the Y^+ gene must be considered and combined to calculate the total amount of the enzyme per cell. In the absence of lactose, 75 units will be produced from the chromosomal copy of the Y^+ gene and 0.2 unit from the copy on the F', for a total of 75.2 units. In the presence of lactose, 100 units will be made from each copy of the Y^+ gene, for a total of 200 units.

For further discussion visit the Student Companion site.



(**Figure 18.9**). Because CAP binds cAMP when this mononucleotide is present at sufficient concentrations, it is sometimes called the cyclic AMP receptor protein.

The *lac* promoter contains two separate binding sites, one for RNA polymerase and one for the CAP/cAMP complex (**Figure 18.10**). The CAP/cAMP complex must be present at its binding site in the *lac* promoter in order for the operon to be induced normally. The CAP/cAMP complex thus exerts positive control over the transcription of the *lac* operon. It has an effect exactly opposite to that of repressor binding to an operator. Although the precise mechanism by which CAP/cAMP stimulates RNA polymerase binding to the promoter is still uncertain, its positive control of *lac* operon transcription is firmly established by the results of both *in vivo* and *in vitro* experiments. CAP functions as a dimer; thus, like the *lac* repressor, it is multimeric in its functional state.

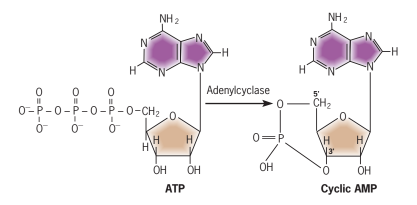


FIGURE 18.9 The adenylcyclase-catalyzed synthesis of cyclic AMP (cAMP) from ATP.

Only the CAP/cAMP complex binds to the *lac* promoter; in the absence of cAMP, CAP does not bind. Thus, cAMP acts as the effector molecule, determining the effect of CAP on *lac* operon transcription. The intracellular cAMP concentration is sensitive to the presence or absence of glucose. High concentrations of glucose cause sharp decreases in the intracellular concentration of cAMP. Glucose prevents the activation of adenylcyclase, the enzyme that catalyzes the formation of cAMP from ATP. Thus, the presence of glucose results in a decrease in the intracellular concentration of cAMP. In the presence of a low concentration of cAMP, CAP cannot bind to the *lac* operon promoter. In turn, RNA polymerase cannot bind efficiently to the *lac* operon transcription never exceeds 2 percent of the induced rate observed in the absence of glucose. By similar mechanisms, CAP and cAMP keep the arabinose (*ara*) and galactose (*gal*) operons of *E. coli* from being induced in the presence of glucose.

PROTEIN-DNA INTERACTIONS THAT CONTROL TRANSCRIPTION OF THE *lac* OPERON

The nucleotide-pair sequence of the *lac* operon regulatory region is shown in Figure 18.10. Comparative nucleotide-sequence studies of mutant and wild-type

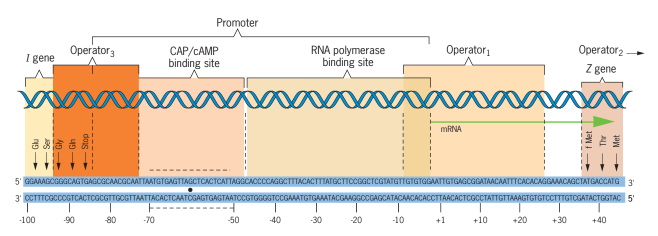
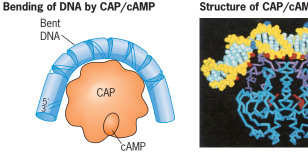


FIGURE 18.10 Organization of the promoter-operator region of the *lac* operon. The promoter consists of two components: (1) the site that binds the CAP/cAMP complex and (2) the RNA polymerase binding site. The adjacent segments of the *lacl* (repressor) and *lacZ* (β -galactosidase) structural genes and the *lac* operators O_1 and O_3 are also shown. Operator O_2 is located downstream (centered at position +412) in the *lacZ* gene. The horizontal line labeled mRNA shows the position at which transcription of the operon begins (the 5' end of the *lac* mRNA). The numbers at the bottom give distances in nucleotide pairs from the site of transcript initiation (position +1). The dot between the two nucleotide strands indicates the center of symmetry of an imperfect palindrome.





■ FIGURE 18.11 The interaction of CAP/cAMP with its binding site in the *lac* promoter. (a) When CAP/cAMP, a positive regulator, binds to the lac promoter, it produces a bend of over 90° in the DNA. (b) Structure of the complex formed by CAP/cAMP and a synthetic 30-bp DNA molecule containing the CAP/cAMP binding site based on X-ray studies.

Structure of CAP/cAMP/DNA complex



promoters and operators, in addition to in vitro CAP/ cAMP, RNA polymerase, and repressor binding studies and X-ray crystallographic data, have provided important information about the sequence-specific protein-nucleic acid interactions that regulate the transcription of the lac operon.

One key interaction involves the binding of RNA polymerase to its binding site in the *lac* promoter (see Chapter 11). Another important interaction is the binding of CAP/cAMP to its binding site in the lac promoter (discussed in the preceding section). A third is the binding of the lac repressor to the lac operators.

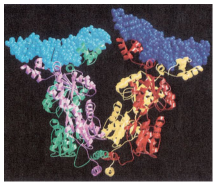
Let's first examine the binding of CAP/cAMP to its binding site in the *lac* promoter. CAP/cAMP controls catabolite

repression; the binding of CAP/cAMP to the promoter is required for efficient induction of the lac operon. How does the binding of CAP/cAMP stimulate transcription of the lac structural genes? RNA polymerase cannot bind efficiently to its binding site in the lac promoter unless CAP/cAMP is already bound. When CAP/cAMP binds to DNA, it bends the DNA (**Figure 18.11***a*). X-ray studies show that the DNA is bent as it is wrapped on the surface of the CAP/cAMP complex (**Figure 18.11b**). Recall that the CAP/cAMP and RNA polymerase binding sites are adjacent to one another in the lac promoter (see Figure 18.10). Presumably, the bending of the DNA by CAP/ cAMP promotes a more open site for RNA polymerase and thus enhanced binding and transcription of the structural genes. However, there is also evidence for contact between RNA polymerase and CAP/cAMP, so the complete picture may be more complex than just the bending of the DNA.

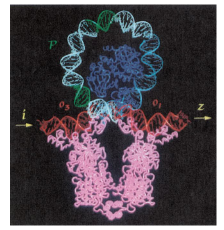
Next, let's examine the binding of the *lac* repressor to the *lac* operators, which prevents RNA polymerase from transcribing the structural genes in the operon. Recall that the *lac* operon is controlled by three operators: the primary operator— O_1 —and two secondary operators— O_2 and O_3 (see Figures 18.5 and 18.10). O_1 is the original operator identified by Jacob and Monod; it is located between the promoter and the Z gene. O_1 is located downstream from O_1 within the Z gene, and O_3 is located upstream of the promoter. Maximum repression requires all three operators; however, strong repression occurs as long as O_1 and either O_2 or O_3 are present. Why are two of the operators required for efficient repression? To answer this question, we need to look at the sequence-specific binding of the repressor to the operators.

The active form of the *lac* repressor is a tetramer containing four copies of the product of the *lacI* gene. X-ray studies of the structures formed by the *lac* repressor and 21-bp-synthetic binding sites showed that each tetrameric repressor binds two operator sequences simultaneously (**Figure 18.12a**). In effect, the tetramer consists

Binding of lac repressor to two synthetic operator DNAs



Structure of the lac repressor/ O_1 - O_3 operator DNAs/CAP/cAMP complex



■ FIGURE 18.12 The interaction of *lac* repressor with its binding sites in the *lac* operators. (a) Binding of the tetrameric lac repressor to two 21-bp DNAs containing repressor recognition sequences. (b) Montage structure of the 93-bp loop formed when tetrameric repressor is bound to *lac* operators O_1 and O_3 . CAP/cAMP (blue) is shown inside the loop associated with its binding site in the *lac* promoter.

of two dimers, each with a sequence-specific binding site. One of the dimers binds to O_1 , and the other binds to either O_2 or O_3 . In so doing, the repressor bends the DNA forming either a hairpin (O_1 and O_2) or a loop (O_1 and O_3). The proposed structure of the O_1 - O_3 -repressor complex is shown in **Figure 18.12b**. Note the presence of CAP/cAMP within the DNA loop formed when *lac* repressor is bound to both O_1 and O_3 (Figure 18.12b).

Similar DNA loops are known to be formed by the binding of protein activators and repressors of other operons in *E. coli* and other bacteria. Regulatory proteins have the ability to bind to DNA in a sequence-specific manner, to alter the structure of the DNA, and to stimulate or repress the transcription of structural genes in the vicinity. A complete understanding of the regulation of gene expression will require detailed knowledge of these important interactions.

- The E. coli lac operon is a negative inducible and catabolite repressible system; the three structural genes in the lac operon are transcribed at high levels only in the presence of lactose and the absence of glucose.
- In the absence of lactose, the lac repressor binds to the lac operators and prevents RNA
 polymerase from initiating transcription of the operon.
- Catabolite repression keeps operons such as lac encoding enzymes involved in carbohydrate catabolism from being induced in the presence of glucose, the preferred energy source.
- The binding of the CAP/cAMP complex to its binding site in the lac promoter bends the DNA and makes it more accessible to RNA polymerase.
- The lac repressor binds to two operators—either O₁ and O₂ or O₁ and O₃—simultaneously and bends the DNA into a hairpin or a loop, respectively.

KEY POINTS

The Tryptophan Operon in *E. coli:* Repression and Attenuation

The *trp* operon of *E. coli* controls the synthesis of the enzymes that catalyze the biosynthesis of the amino acid tryptophan. The functions of the five structural genes and the adjacent regulatory sequences of the *trp* operon have been analyzed in detail by Charles Yanofsky and colleagues. The five structural genes encode enzymes that convert chorismic acid to tryptophan. The expression of the *trp* operon is regulated at two levels: repression, which controls the initiation of transcription,

The structural genes in the tryptophan operon are transcribed only when tryptophan is absent or present in low concentrations. The expression of the genes in the *trp* operon is regulated by repression of transcriptional initiation and by attenuation (premature termination) of transcription when tryptophan is prevalent in the environment.

and attenuation, which governs the frequency of premature transcript termination. We will discuss these regulatory mechanisms in the following two sections.

REPRESSION

The *trp* operon of *E. coli* is a negative repressible operon. The organization of the *trp* operon and the pathway of biosynthesis of tryptophan are shown in **Figure 18.13**. The *trpR* gene, which encodes the *trp* repressor, is not closely linked to the *trp* operon. The operator (*O*) region of the *trp* operon lies within the primary promoter (P_1) region. There is also a weak promoter (P_2) at the operator-distal end of the *trpD* gene. The P_2 promoter increases the basal level of transcription of the *trpC*, *trpB*, and *trpA* genes. Two transcription termination sequences (*t* and *t'*) are located downstream from *trpA*. The *trpL* region specifies a 162-nucleotide-long mRNA leader sequence.

The regulation of transcription of the *trp* operon is diagrammed in Figure 18.4*c*. In the absence of tryptophan (the co-repressor), RNA polymerase binds

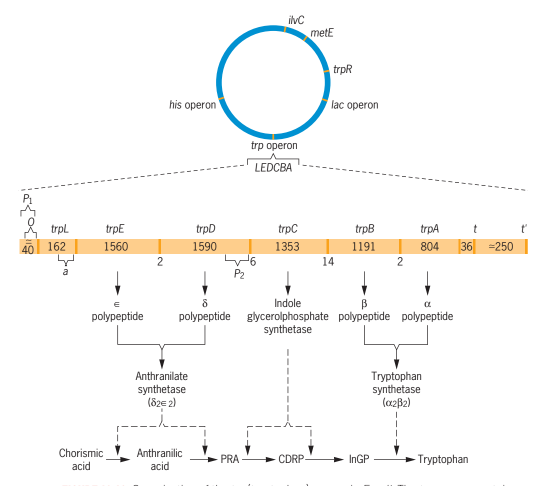


FIGURE 18.13 Organization of the *trp* (tryptophan) operon in *E. coli*. The *trp* operon contains five structural genes that encode enzymes involved in the biosynthesis of tryptophan, as shown at the bottom, and the *trpL* regulatory region. The length of each gene or region is given in nucleotide pairs; the intergenic distances are shown below the gene sequence. Key: PRA, phosphoribosyl anthranilate; CDRP, carboxyphenylamino-deoxyribulose phosphate; InGP, indole-glycerol phosphate.

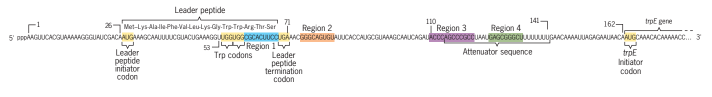
to the promoter region and transcribes the structural genes of the operon. In the presence of tryptophan, the co-repressor/repressor complex binds to the operator region and prevents RNA polymerase from initiating transcription of the genes in the operon.

The rate of transcription of the *trp* operon in the derepressed state (absence of tryptophan) is 70 times the rate that occurs in the repressed state (presence of tryptophan). In *trpR* mutants, which lack a functional repressor, the rate of synthesis of the tryptophan biosynthetic enzymes is still reduced about tenfold by the addition of tryptophan to the medium. This additional reduction in *trp* operon expression is caused by attenuation, which is discussed next.

ATTENUATION

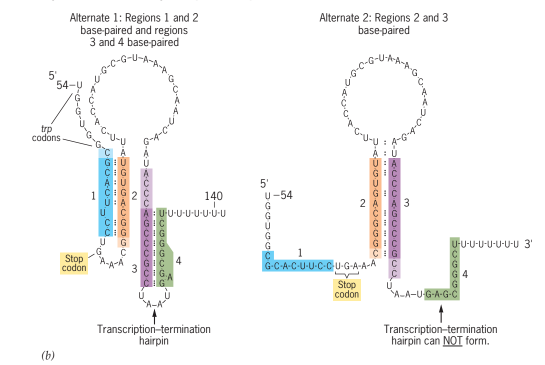
Deletions that remove part of the trpL region (Figure 18.13) result in increased rates of expression of the trp operon. However, these deletions have no effect on the repressibility of the trp operon; that is, repression and derepression occur just as in $trpL^+$ strains. These results indicate that the synthesis of the tryptophan biosynthetic enzymes is regulated at a second level by a mechanism that is independent of repression/derepression and requires nucleotide sequences present in the trpL region of the trp operon.





(a)

Alternate secondary structures formed by the trpL transcript



■ **FIGURE 18.14** Sequences in the leader region of the *trp* mRNA responsible for attenuation. (a) The *trpL* sequence, highlighting the sequence encoding the leader peptide, the two tandem tryptophan codons responsible for the control of attenuation by tryptophan, and the four regions (shaded) that form the stem-and-loop or hairpin structures shown in (b). (b) Alternate secondary structures formed by the *trpL* mRNA—either (1) region 1 will pair with region 2 and region 3 with region 4, forming a transcription–termination hairpin, or (2) region 2 will base-pair with region 3, preventing region 3 from pairing with region 4. The concentration of tryptophan in the cell determines which of these structures will form during the transcription of the *trp* operon.

This second level of regulation of the trp operon is called **attenuation**, and the sequence within trpL that controls this phenomenon is called the **attenuator** (**Figure 18.14a**). Attenuation occurs by control of the termination of transcription at a site near the end of the mRNA leader sequence. This "premature" termination of trp operon transcription occurs only in the presence of tryptophan-charged tRNA^{Trp}. When this premature termination or attenuation occurs, a truncated (140 nucleotides) trp transcript is produced.

The attenuator region has a nucleotide-pair sequence essentially identical to the *transcription-termination signals* found at the ends of most bacterial operons. These termination signals contain a G:C-rich palindrome followed by several A:T base pairs. Transcription of these termination signals yields a nascent RNA with the potential to form a hydrogen-bonded hairpin structure followed by several uracils.

Solve It!

Regulation of the Histidine Operon of Salmonella typhimurium

The amino acid histidine is synthesized from 5-phosphoribosyl 1-pyrophosphate and ATP via a series of 10 reactions catalyzed by enzymes encoded by eight contiquous genes in the histidine operon of Salmonella typhimurium. The his operon is transcribed as a unit yielding a multigenic mRNA. The operon is expressed at high levels when histidine concentrations are low, but at low levels when histidine levels are high. The nucleotide sequence of the nontemplate strand of the 5' untranslated leader region of the his operon is shown in the following sequence, along with the predicted amino acid sequences (using the single-letter code) of a leader peptide specified by the small ORF and the first five amino acids of the *hisG* product. Also, six regions capable of forming basepaired stem-and-loop (hairpin) structures are designated 1–6.



mechanism by which the expression of the *his* operon might be regulated.

To see the solution to this problem, visit the Student Companion site. When a nascent transcript forms this hairpin structure, it causes a conformational change in the associated RNA polymerase, resulting in termination of transcription within the following, more weakly hydrogen-bonded $(A:U)_n$ region of DNA-RNA base-pairing.

The nucleotide sequence of the attenuator therefore explains its ability to terminate *trp* operon transcription prematurely. But how can this be regulated by the presence or absence of tryptophan?

First, recall that transcription and translation are coupled in prokaryotes; that is, ribosomes begin translating mRNAs while they are still being synthesized. Thus, events that occur during translation may also affect transcription.

Second, note that the 162-nucleotide-long leader sequence of the trp operon mRNA contains sequences that can base-pair to form alternate stem-and-loop or hairpin structures (**Figure 18.14b**). The four leader regions that can base-pair to form these structures are: (1) nucleotides 60–68, (2) nucleotides 75–83, (3) nucleotides 110–121, and (4) nucleotides 126–134. The actual lengths of these regions involved in base-pairing vary depending on which regions pair. The nucleotide sequences of these four regions are such that region 1 can base-pair with region 2, region 2 can pair with region 3, and region 3 can pair with region 4. Region 2 can base-pair with either region 1 or region 3, but, obviously, it can pair with only one of these regions at any given time. Thus, there are two possible secondary structures for the trpleader sequence: (1) region 1 paired with region 2 and region 3 paired with region 4 or (2) region 2 paired with region 3, leaving regions 1 and 4 unpaired. The pairing of regions 3 and 4 produces the previously mentioned transcription-termination hairpin. If region 3 is base-paired with region 2, it cannot pair with region 4, and the transcription-termination hairpin cannot form. As you have probably guessed by now, the presence or absence of tryptophan determines which of these alternative structures will form.

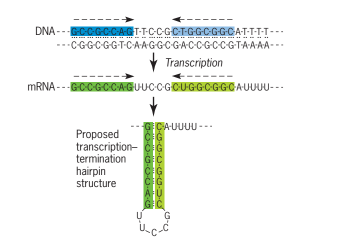
Third, note that the leader sequence contains an AUG translation-initiation codon, followed by 13 codons for amino acids, followed in turn by a UGA translation-termination codon (Figure 18.14*a*). In addition, the *trp* leader sequence contains an efficient ribosome-binding site located in the appropriate position for the initiation of translation at the leader AUG initiation codon. All the available evidence indicates that a 14-amino-acid "leader peptide" is synthesized as diagrammed in Figure 18.14*a*.

The normal *trp* operon transcription-termination hairpin is shown in **Figure 18.15***a*, and the proposed mechanism of attenuation of *trp* operon transcription is diagrammed in **Figure 18.15***b* and *c*. The leader peptide contains two contiguous tryptophan residues. The two Trp codons are positioned such that in low concentrations of tryptophan (and thus low concentrations of Trp-tRNA^{Trp}), the ribosome will stall before it encounters the base-paired structure formed by leader regions 2 and 3 (Figure 18.15*b*). Because the pairing of regions 2 and 3 precludes the formation of the transcription-termination hairpin by the base-pairing of regions 3 and 4, transcription will continue past the attenuator into the *trpE* gene in the absence of tryptophan.

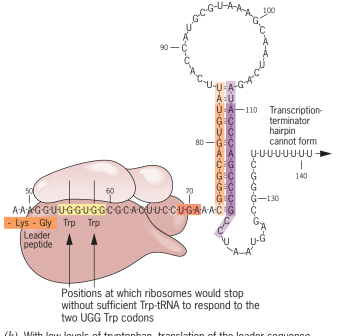
In the presence of sufficient tryptophan, the ribosome can translate past the Trp codons to the leader-peptide termination codon. In the process, it will disrupt the base-pairing between leader regions 2 and 3. This disruption leaves region 3 free to pair with region 4, forming the transcription–termination hairpin (Figure 18.15*c*). Thus, in the presence of sufficient tryptophan, transcription frequently (about 90 percent of the time) terminates at the attenuator, reducing the amount of mRNA for the *trp* structural genes.

The transcription of the *trp* operon can be regulated over a range of almost 700-fold by the combined effects of repression (up to 70-fold) and attenuation (up to 10-fold).

Regulation of transcription by attenuation is not unique to the *trp* operon. Five other operons (*thr*; *ilv*, *leu*, *phe*, and *his*) are known to be regulated by attenuation. The *his* operon, which for many years was thought to be repressible, is now believed to be regulated entirely by attenuation. Although minor details vary from operon to operon, the main features of attenuation are the same for all six operons. Try Solve It: Regulation



(a) Structure of *trp* operon transcription–termination sequence *t* and formation of the transcription–termination hairpin.

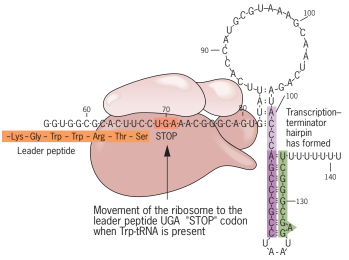


(b) With low levels of tryptophan, translation of the leader sequence stalls at one of the Trp codons. This stalling allows leader regions 2 and 3 to pair, which prevents region 3 from pairing with region 4 to form the transcription-termination hairpin. Thus transcription proceeds through the entire *trp* operon. **FIGURE 18.15** Control of the *trp* operon by attenuation. (*a*) The transcriptiontermination signal in *E. coli* contains a region of dyad symmetry (arrows) that results in mRNA sequences that can form hairpin structures. (*b*) In low concentrations of tryptophan, transcription proceeds past the attenuator sequence through the entire *trp* operon. (*c*) In the presence of sufficient tryptophan, transcription frequently terminates at the attenuator sequence.

of the Histidine Operon of *Salmonella typhimurium* to test your understanding of attenuation. In addition, see On the Cutting Edge: The Lysine Riboswitch for a discussion of a related regulatory mechanism.

- The E. coli trp operon is a negative repressible system; transcription of the five structural genes in the trp operon is repressed in the presence of significant concentrations of tryptophan.
- Operons such as trp that encode enzymes involved in amino acid biosynthetic pathways often are controlled by a second regulatory mechanism called attenuation.
- Attenuation occurs by the premature termination of transcription at a site in the mRNA leader sequence (the sequence 5' to the coding region) when tryptophan is prevalent in the environment in which the bacteria are growing.

KEY POINTS



(c) In the presence of sufficient tryptophan, translation proceeds past the Trp codons to the termination codon and disrupts the basepairing between leader regions 2 and 3. This process leaves region 3 free to pair with region 4 to form the transcription-termination hairpin, which stops transcription at the attenuator sequence.

ON THE CUTTING EDGE

THE LYSINE RIBOSWITCH

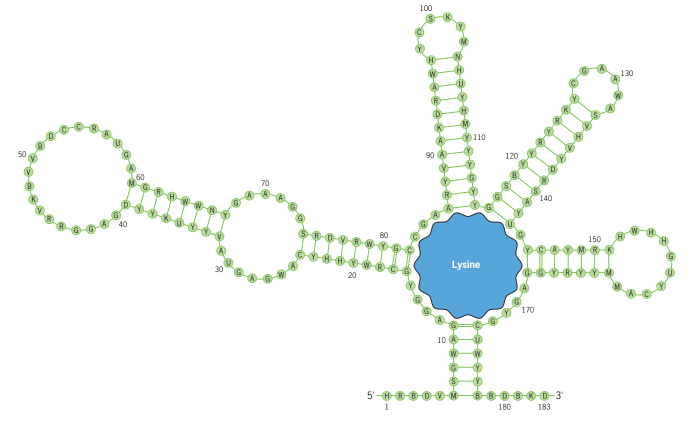
nzymes have long been known to undergo changes in conformation and activity after binding small molecules. During the last decade, RNA molecules have been shown to bind metabolites and undergo similar changes in conformation. Indeed, the metabolite-binding domains of many bacterial mRNAs play central roles in regulating gene expression. Together, the metabolite-binding domains of these RNAs and the domains that undergo changes in conformation are called **riboswitches.** They regulate gene expression by changing conformation after binding specific metabolites. The changes in conformation can activate or terminate either transcription or translation.

Riboswitches commonly terminate transcription by forming transcription-termination hairpins similar to the one responsible for attenuation in the *trp* operon in *E. coli* (see Figure 18.15*c*). Alternatively, they can block translation by sequestering the Shine-Dalgarno sequence (ribosome-binding site) within a hydrogen-bonded hairpin, so that ribosomes cannot bind to the mRNA. Most of the riboswitches characterized to date occur in bacteria; however, riboswitches also have been identified in archaea, fungi, and plants. In fungi and plants, riboswitches sometimes alter mRNA splicing and 3' processing.



Riboswitches contain two essential components: (1) an **aptamer domain**, a folded region that has the ability to bind a specific metabolite, and (2) an **expression domain**, which can fold into two distinct structures, one facilitating gene expression and the other blocking gene expression. Both domains are usually present in mRNAs upstream from the translation start codon.

Let's examine one particular riboswitch, the lysine riboswitch, which regulates the biosynthesis of lysine as well as its transport into cells. Bacteria synthesize lysine from aspartate in a sequence of enzyme-catalyzed reactions. In *E. coli*, the *lysC* gene encodes an aspartokinase that catalyzes the first step in the biosynthesis of lysine, and mutations in E. coli that result in the constitutive synthesis of lysine map within the leader region of the lysC gene. Now, it turns out that this region of the *lysC* mRNA is highly conserved and folds into a structure with five helical regions surrounding a lysinebinding pocket. A very similar lysine riboswitch was found in the 5'-untranslated region of the *B. subtilis lysC* mRNA. The conserved sequences in the *E. coli* and *B. subtilis* riboswitches were then used to search for similar sequences in other bacterial genomes. The results were clear-cut; lysine riboswitches are highly conserved and widely distributed in the bacterial kingdom. **Figure 1** shows the predicted structure of the aptamer (lysine-binding) domain of the



■ FIGURE 1 Structure of the lysine-binding domain of the lysine riboswitch. The structure shows the conserved stem-and-loop regions surrounding the lysine-binding pocket (blue). The sequence shown is derived from a comparative genomics analysis of a large number of lysine riboswitches from many species. The five stem-and-loop structures are highly conserved, as are the nucleotides that make up the lysine-binding pocket. Gs, As, Cs, and Us represent invariant nucleotides. The other symbols are: R = either purine, G or A; Y = either pyrimidine, C or U; W = either A or U; S = either G or C; M = either A or C; K = either G or U; H = A, U or C; B = G, C or U; D = A, G or U; V = A, C or U; and N = A, G, C or U.

lysine riboswitch based on a comparison of 71 lysine riboswitches from 37 different bacterial species.

In the absence of lysine, the expression domain of the lysine riboswitch forms a hydrogen-bonded hairpin region called the antiterminator upstream from the coding region of the *lysC* gene (**Figure 2a**) and other regulated genes. The downstream sequence present in this antiterminator hairpin overlaps the upstream sequence in the transcription-terminator hairpin that forms when lysine is present. Thus, the presence of the antiterminator hairpin precludes the formation of the terminator hairpin and facilitates the ongoing transcription of *lysC* and other regulated genes in the absence of lysine. If lysine is added to the medium in which bacteria are growing, it is bound by the aptamer (see Figure 1) and triggers a conformational change in the expression domain of the lysine riboswitch (**Figure 2b**). This conformational change sequesters the upstream sequence of the antiterminator hairpin into a basal hydrogen-bonded hairpin that is part of the lysine-binding pocket of the aptamer. As a result, the downstream sequence of the antiterminator is free to become part of the transcription-terminator hairpin and terminate transcription upstream of the *lysC* coding region (Figure 2b). As a result, bacteria shut off the biosynthesis of lysine when synthesis is no longer needed and use the conserved energy to enhance other metabolic processes.

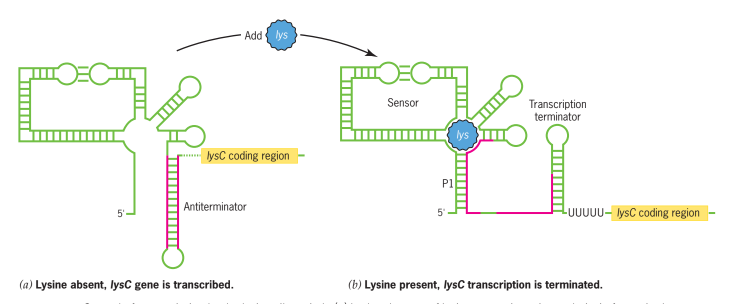


FIGURE 2 Control of transcription by the lysine riboswitch. (a) In the absence of lysine, an antiterminator hairpin forms in the expression domain upstream from the start of translation of the regulated gene (*lysC* in the diagram) and precludes the formation of the transcription-terminator hairpin. As a result, transcription of the gene is turned on. (b) When present, lysine is bound by the aptamer, and the upstream antiterminator sequence is sequestered in a basal hairpin that is part of the lysine-binding pocket. As a result, the downstream antiterminator sequence is free to participate in the formation of a transcription-termination hairpin upstream from the coding region of the regulated gene (*lysC* is shown). Therefore, transcription is turned off.

Translational Control of Gene Expression

Although gene expression in prokaryotes is regulated predominantly at the level of transcription, fine-tuning often occurs at the level of translation. In prokaryotes, mRNA molecules are frequently multigenic, carrying the coding sequences of several genes. For example, The regulation of gene expression is often fine-tuned by modulating either the frequency of initiation of translation or the rate of polypeptide chain elongation.

the *E. coli lac* operon mRNA harbors nucleotide sequences encoding β -galactosidase, β -galactoside permease, and β -galactoside transacetylase. Thus, the three genes encoding these proteins must be turned on and turned off together at the transcription level because the genes are co-transcribed. Nevertheless, the three gene products are not synthesized in equal amounts. An *E. coli* cell that is growing on rich medium with lactose as the sole carbon source contains about 3000 molecules of β -galactosidase, 1500 molecules of β -galactoside permease, and 600 molecules of β -galactoside transacetylase. Clearly, the different molar quantities of these proteins per cell must be controlled posttranscriptionally.

Remember that transcription, translation, and mRNA degradation are coupled in prokaryotes; an mRNA molecule usually is involved in all three processes at any given time. Thus, gene products may be produced in different amounts from the same transcript by several mechanisms.

- **1.** *Unequal efficiencies of translational initiation* are known to occur at the ATG start codons of different genes.
- **2.** Altered efficiencies of ribosome movement through intergenic regions of a transcript are quite common. Decreased translation rates often result from hairpins or other forms of secondary structure that impede ribosome migration along the mRNA molecule.
- 3. Differential rates of degradation of specific regions of mRNA molecules also occur.

KEY POINT

Regulatory fine-tuning frequently occurs at the level of translation by modulation of the rate
of polypeptide chain initiation or chain elongation.

Posttranslational Regulatory Mechanisms

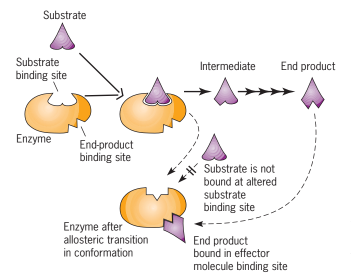
Feedback inhibition occurs when the product of a biosynthetic pathway inhibits the activity of the first enzyme in the pathway, rapidly shutting off the synthesis of the product. Earlier in this chapter, we discussed the mechanism by which the transcription of bacterial genes encoding enzymes in a biosynthetic pathway is repressed when the product of the pathway is present in the medium in which the cells are growing. A second, and more rapid, regulatory fine-tuning of metabolism often occurs at the level of enzyme activity. The presence of a sufficient concentration of the end product of a biosyn-

thetic pathway frequently results in the inhibition of the first enzyme in the pathway (**■** Figure 18.16). This phenomenon is called **feedback** inhibition or **end-product** inhibition. Feedback inhibition results in an almost instantaneous arrest of the synthesis of the end product when it is added to the medium.

The tryptophan biosynthetic pathway in *E. coli* provides a good illustration of feedback inhibition. The end product—tryptophan—is bound by the first enzyme in the pathway—anthranilate synthetase (see Figure 18.13)—and completely arrests its activity, stopping the synthesis of tryptophan almost immediately.

Feedback inhibition-sensitive enzymes contain an end-product binding site (or sites) in addition to the substrate binding site (or sites). In the case of multimeric enzymes, the *end-product* or *regulatory binding site* often is on a subunit (polypeptide) different from that of the substrate site. Upon binding the end-product, such enzymes undergo allosteric transitions that reduce their affinity for their substrates. Proteins that undergo such conformational changes are referred to as allosteric proteins. Many, perhaps most, enzymes undergo allosteric transitions of some kind.

Allosteric transitions also appear to be responsible for enzyme activation, which often occurs when an enzyme binds one or more of its substrates or some other small molecule. Some enzymes exhibit a broad spectrum of activation and inhibition by many different effector molecules. An example is the enzyme glutamine synthetase, which catalyzes the final step in the biosynthesis of the amino acid glutamine. Glutamine synthetase is a complex multimeric enzyme in both prokaryotes and eukaryotes. The glutamine synthetase of *E. coli* has been shown to respond, either by activation or inhibition, to 16 different metabolites, presumably through allosteric transitions.



■ FIGURE 18.16 Feedback inhibition of gene-product activity. The end product of a biosynthetic pathway often binds to and arrests the activity of the first enzyme in the pathway, quickly blocking the synthesis of the end product.

- Feedback inhibition occurs when the product of a biosynthetic pathway inhibits the activity of the first enzyme in the pathway, rapidly arresting the biosynthesis of the product.
- Enzyme activation occurs when a substrate or other effector molecule enhances the activity of an enzyme, increasing the rate of synthesis of the product of the biosynthetic pathway.

Basic Exercises

Illustrate Basic Genetic Analysis

- 1. How can positive and negative regulatory mechanisms be distinguished?
- **Answer:** Mutations in regulator genes that yield nonfunctional products will have very different effects in positive and negative control systems. In positive control circuits, such mutations will make it impossible to turn on the expression of the regulated genes, whereas in negative control circuits, these mutations will make it impossible to turn off the expression of the regulated genes.
- 2. How can inducible and repressible operons be distinguished?
- **Answer:** In the absence of the effector molecule, inducible operons will be turned off, whereas repressible operons will be turned on.
- **3.** How can *cis* and *trans*-acting regulatory elements be distinguished?
- Answer: They can be distinguished by constructing partial diploids in which the regulatory elements are positioned (1) *cis* to the regulated genes and (2) *trans* to the regulated genes. A *cis*-acting element will only influence the expression of the genes when present in the *cis* configuration, whereas a *trans*-acting element will exert its effect in either the *cis* or *trans* configuration (compare Figures 18.7 and 18.8).
- 4. What is attenuation, and how does it work?
- **Answer:** Attenuation is a mechanism for regulating gene expression by the premature termination of transcription in the leader region of a transcript. In the case of the tryptophan (*trp*) operon of *E. coli*, for example, the presence or absence of the end product, tryptophan, determines whether or

not attenuation occurs. The leader region of the mRNA has sequences that can base-pair to form alternative hairpin structures, one of which is a typical transcriptiontermination signal. Whether or not this hairpin forms depends on the translation of a leader peptide containing two tryptophan residues. When low levels of tryptophan are present, translation stops at the Trp codons, which prevents the formation of the transcription-termination hairpin (see Figure 18.15b). When sufficient tryptophan is present, translation proceeds past the Trp codons to the translation-termination codon, disrupting the first hairpin. This, in turn, allows the transcription-termination hairpin to form and attenuation (termination of transcription at the attenuator) to occur (see Figure 18.15c). Attenuation decreases the synthesis of the tryptophan biosynthetic enzymes tenfold. Attenuation is possible in prokaryotes because transcription and translation are coupled, so events occurring during translation can affect transcription.

- 5. When histidine is added to the medium in which *E. coli* cells are growing, its synthesis stops very quickly, long before the synthesis of the histidine biosynthetic enzymes stops. How can this be explained?
- **Answer:** In addition to turning off the *synthesis* of the histidine biosynthetic enzymes, histidine also inhibits the *activity* of the first enzyme—N'-5'-phosphoribosyl-ATP transferase—in the histidine biosynthetic pathway by a process called feedback inhibition. The enzyme contains a histidine-binding site, and when it binds histidine, it undergoes a change in conformation that inhibits its activity (see Figure 18.16). Thus, feedback inhibition results in an almost instantaneous shutoff of histidine synthesis.

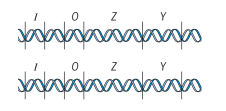
Testing Your Knowledge

Integrate Different Concepts and Techniques

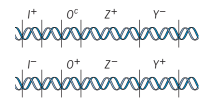
1. The operon model for the regulation of enzyme synthesis concerned in lactose utilization by *E. coli* includes a regulator gene (*I*), an operator region (*O*), a structural gene (*Z*) for the enzyme β -galactosidase, and another structural gene (*Y*) for β -galactoside permease. β -Galactoside permease transports lactose into the bacterium, where β -galactosidase cleaves it into galactose and glucose. Mutations in the *lac* operon have the following effects: *Z*⁻ and *Y*⁻

mutant strains are unable to make functional β-galactosidase and β-galactoside permease, respectively, whereas I^- and O^c mutant strains synthesize the *lac* operon gene products constitutively. The following figure shows a partially diploid strain of *E. coli* that carries two copies of the *lac* operon. On the diagram, fill in a genotype that will result in the constitutive synthesis of β-galactosidase and the inducible synthesis of β-galactoside permease by this partial diploid.

KEY POINTS



Answer: Several different genotypes will produce β -galactosidase constitutively and β -galactoside permease inducibly. They must meet two key requirements: (1) the cell must contain at least one copy of the I^+ gene, which encodes the repressor, and (2) the Z^+ gene and an O^e mutation must be on the same chromosome because the operator acts only in *cis*; that is, it only affects the expression of genes on the same chromosome. In contrast, the cell can be either homozygous or heterozygous for the I^+ gene, and, if heterozygous, I^+ may be on either chromosome because I^+ is dominant to I^- and I^+ acts in both the *cis* and *trans* arrangement. One possible genotype is given in the following diagram.



How many other genotypes can you devise that will synthesize β -galactosidase constitutively and β -galactoside permease inducibly?

- 2. Wild-type *E. coli* cells have been growing exponentially in culture medium containing very low concentrations of tryptophan for 20 minutes when someone adds a large amount of tryptophan to the culture medium. What physiological changes will occur in these cells after the addition of tryptophan?
- **Answer:** (a) The first thing that will happen is that tryptophan will be bound by the first enzyme-anthranilate synthetase-in the tryptophan biosynthetic pathway, inhibiting the activity of the enzyme and arresting the synthesis of tryptophan almost immediately. This regulatory mechanism is called feedback inhibition (see Figure 18.16). (b) The second thing that will happen is that the high concentration of this amino acid will decrease the rates of synthesis of the tryptophan biosynthetic enzymes by the premature termination-attenuation-of transcription of the genes in the tryptophan operon (see Figures 18.14 and 18.15). (c) The third thing that will happen is that the high concentration of tryptophan will lead to the repression of transcription of the trp operon, further decreasing the rates of synthesis of the tryptophan biosynthetic enzymes (see Figure 18.4c). Working in concert, feedback inhibition, attenuation, and repression/derepression quickly and rather precisely adjust the rates of synthesis of metabolites such as tryptophan in bacteria in response to changes in environmental conditions.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **18.1** How can inducible and repressible enzymes of microorganisms be distinguished?
- **18.2** Distinguish between (a) repression and (b) feedback inhibition caused by the end product of a biosynthetic pathway. How do these two regulatory phenomena complement each other to provide for the efficient regulation of metabolism?
- **18.3** In the lactose operon of *E. coli*, what is the function of each of the following genes or sites: (a) regulator, (b) operator, (c) promoter, (d) structural gene *Z*, and (e) structural gene *Y*?
- 18.4 What would be the result of inactivation by mutation of the following genes or sites in the *E. coli* lactose operon:(a) regulator, (b) operator, (c) promoter, (d) structural gene *Z*, and (e) structural gene *Y*?
- **18.5** Groups of alleles associated with the lactose operon are as follows (in order of dominance for each allelic series): repressor, I^s (superrepressor), I^+ (inducible), and I^- (constitutive); operator, O^c (constitutive, cis^- dominant)

and O^+ (inducible, *cis*-dominant); structural, Z^+ and Y^+ . (a) Which of the following genotypes will produce β -galactosidase and β -galactoside permease if lactose is present: (1) $I^+O^+Z^+Y^+$, (2) $I^-O^cZ^+Y^+$, (3) $I^sO^cZ^+Y^+$, (4) $I^sO^+Z^+Y^+$, and (5) $I^-O^+Z^+Y^+$? (b) Which of the above genotypes will produce β -galactosidase and β -galactoside permease if lactose is absent? Why?

- **18.6** Sume that you have discovered a new strain of *E. coli* that has a mutation in the *lac* operator region that causes the wild-type repressor protein to bind irreversibly to the operator. You have named this operator mutant O^{sb} for "superbinding" operator. (a) What phenotype would a partial diploid of genotype $I^+O^{sb}Z^-Y^+/I^+O^+Z^+Y^$ have with respect to the synthesis of the enzymes β -galactosidase and β -galactoside permease? (b) Does your new O^{sb} mutation exhibit *cis* or *trans* dominance in its effects on the regulation of the *lac* operon?
- **18.7** Why is the *O^c* mutation in the *E. coli lac* operon epistatic to the *I^s* mutation?

- **18.8 ••** For each of the following partial diploids indicate whether enzyme synthesis is constitutive or inducible (see Problem 18.5 for dominance relationships):
 - (a) $I^+O^+Z^+Y^+/I^+O^+Z^+Y^+$, (b) $I^+O^+Z^+Y^+/I^+O^cZ^+Y^+$, (c) $I^+O^cZ^+Y^+/I^+O^cZ^+Y^+$, (d) $I^+O^+Z^+Y^+/I^-O^+Z^+Y^+$, (e) $I^-O^+Z^+Y^+/I^-O^+Z^+Y^+$. Why?
- **18.9** Write the partial diploid genotype for a strain that will (a) produce β -galactosidase constitutively and permease inducibly and (b) produce β -galactosidase constitutively but not permease either constitutively or inducibly, even though a Y^+ gene is known to be present.
- 18.10 As a genetics historian, you are repeating some of the classic experiments conducted by Jacob and Monod with the lactose operon in E. coli. You use an F' plasmid to construct several E. coli strains that are partially diploid for the *lac* operon. You construct strains with the following genotypes: (1) $I^+O^cZ^+Y^-/I^+O^+Z^-Y^+$, (2) $I^+ O^c Z^- Y^+ / I^+ O^+ Z^+ Y^-$, (3) $I^- O^+ Z^+ Y^+ / I^+ O^+ Z^- Y^-$ (4) $I^{s}O^{+}Z^{-}Y^{-}/I^{+}O^{+}Z^{+}Y^{+}$, and (5) $I^{+}O^{c}Z^{+}Y^{+}/I^{s}O^{+}Z^{-}Y^{+}$. (a) Which of these strains will produce functional β-galactosidase in both the presence and absence of lactose? (b) Which of these strains will exhibit constitutive synthesis of functional β -galactoside permease? (c) Which of these strains will express both gene Z and gene Y constitutively and will produce functional products (β-galactosidase and β-galactoside permease) of both genes? (d) Which of these strains will show cis dominance of lac operon regulatory elements? (e) Which of these strains will exhibit trans dominance of lac operon regulatory elements?
- 18.11 Constitutive mutations produce elevated enzyme levels at all times; they may be of two types: O^c or I⁻. Assume that all other DNA present is wild-type. Outline how the two constitutive mutants can be distinguished with respect to (a) map position, (b) regulation of enzyme levels in O^c/O⁺ versus I⁻/I⁺ partial diploids, and (c) the position of the structural genes affected by an O^c mutation versus the genes affected by an I⁻ mutation in a partial diploid.
- **18.12** How could the tryptophan operon in *E. coli* have developed and been maintained by evolution?
- **18.13** Of what biological significance is the phenomenon of catabolite repression?
- **18.14** How might the concentration of glucose in the medium in which an *E. coli* cell is growing regulate the intracellular level of cyclic AMP?
- **18.15** Is the CAP–cAMP effect on the transcription of the *lac* operon an example of positive or negative regulation? Why?

- **18.16** Would it be possible to isolate *E. coli* mutants in which the transcription of the *lac* operon is not sensitive to catabolite repression? If so, in what genes might the mutations be located?
- **18.17** Using examples, distinguish between negative regulatory mechanisms and positive regulatory mechanisms.
- **18.18** The following table gives the relative activities of the enzymes β -galactosidase and β -galactoside permease in cells with different genotypes at the *lac* locus in *E. coli*. The level of activity of each enzyme in wild-type *E. coli* not carrying F's was arbitrarily set at 100; all other values are relative to the observed levels of activity in these wild-type bacteria. Based on the data given in the table for genotypes 1 through 4, fill in the levels of enzyme activity that would be expected for the fifth genotype.

Genotype	β-Galactoside		β-Galactosidase Permease	
	-Inducer	+Inducer	-Inducer	+Inducer
1. $I^+O^+Z^+Y^+$	0.1	100	0.1	100
2. $I^- O^+ Z^+ Y^+$	100	100	100	100
3. $I^+ O^c Z^+ Y^+$	25	100	25	100
4. $I^-O^+Z^+Y^-/F'$ $I^-O^+Z^+Y^-$	200	200	100	100
5. $I^-O^cZ^-Y^+/F'$ $I^+O^+Z^+Y^+$				

- **18.19** The rate of transcription of the *trp* operon in *E. coli* is controlled by both (1) repression/derepression and (2) attenuation. By what mechanisms do these two regulatory processes modulate *trp* operon transcript levels?
- **18.20** What effect will deletion of the *trpL* region of the *trp* operon have on the rates of synthesis of the enzymes encoded by the five genes in the *trp* operon in *E. coli* cells growing in the presence of tryptophan?
- **18.21** By what mechanism does the presence of tryptophan in the medium in which *E. coli* cells are growing result in premature termination or attenuation of transcription of the *trp* operon?
- **18.22** Suppose that you used site-specific mutagenesis to modify the *trpL* sequence such that the two UGG Trp codons at positions 54–56 and 57–60 (see Figure 18.14) in the mRNA leader sequence were changed to GGG Gly codons. Will attenuation of the *trp* operon still be regulated by the presence or absence of tryptophan in the medium in which the *E. coli* cells are growing?
- **18.23** What do *trp* attenuation and the lysine riboswitch have in common?
- **18.24** Would attenuation of the type that regulates the level of *trp* transcripts in *E. coli* be likely to occur in eukaryotic organisms?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

The *E. coli* catabolite activator protein (CAP) plays an important regulatory role by preventing the induction of the *lac* operon in the presence of high concentrations of glucose, which is a more efficient energy source than lactose. High concentrations of glucose prevent the activation of the enzyme adenylcyclase, which catayzes the synthesis of cyclic AMP (cAMP) from ATP. CAP must form a complex with cAMP in order to bind to the *lac* promoter and, in turn, stimulate the binding of RNA polymerase.

Without CAP-cAMP bound to the promoter, transcription of the *lac* operon never exceeds 2 percent of the level observed in the absence of glucose. The CAP-cAMP complex has the same effect on the *gal* operon, the *ara* operon, and several other operons. It serves as a global regulator of catabolic pathways in bacteria. This phenomenon—catabolite repression or the "glucose effect"—involves specific interactions between DNA-binding domains of the CAP-cAMP complex and nucleotide sequences in bacterial promoters.

- **1.** What kinds of interactions are involved in the binding of CAP-cAMP to DNA?
- 2. What is the three-dimensional structure of CAP-cAMP?
- **3.** What are the three-dimensional structures of CAP-cAMP-DNA complexes?
- **4.** Does the binding of CAP-cAMP have any effect on DNA structure?
- **5.** Does CAP share any three-dimensional structural domains with other DNA-binding proteins?

Hint: At the NCBI web site, click on "Molecular databases," scroll down and click on "Structure (MMDB = *M*olecular *M*odeling *D*atabase)," and search using "CAP-cAMP" as a query. Click on "103T," "Crystal Structures of CAP-DNA Complexes," "1G6N, 2.1 Angstrom Structure of CAP-cAMP," and others, to view three-dimensional models of these molecular interactions.

Regulation of Gene Expression in Eukaryotes



CHAPTER OUTLINE

- Ways of Regulating Eukaryotic Gene Expression: An Overview
- Induction of Transcriptional Activity by Environmental and Biological Factors
- Molecular Control of Transcription in Eukaryotes
- Posttranscriptional Regulation of Gene Expression by RNA Interference
- Gene Expression and Chromatin Organization
- Activation and Inactivation of Whole Chromosomes

parasite, a type of trypanosome, is now called *Trypanosoma brucei* in Bruce's honor. Humans can also be infected with tsetse-borne trypanosomes, whereupon they develop the debilitating illness known as African sleeping sickness.

African Trypanosomes:

A Wardrobe of Molecular Disguises

Near the end of the nineteenth century, David Bruce, a surgeon

in the British Medical Service, summarized his observations and experiments on a disease of wild and domesticated animals

in southern Africa. The disease, called nagana from a Zulu

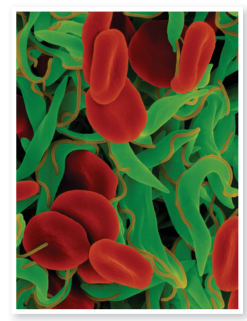
word meaning "loss of spirit," is characterized by fever,

swelling, lethargy, and emaciation. Bruce recognized that nagana is transmitted by the tsetse, a type of biting fly common

in the open spaces of the African scrub plain. Furthermore, his

examination of diseased animals led him to conclude that the causative agent is a flagellated, unicellular protozoan that is injected into the animal's blood when the tsetse bites. This blood

In both animals and humans, trypanosome infections last a long time. This is remarkable because, in the blood, trypanosomes are subjected to repeated attacks by the immune system. With each immune attack, most of the trypanosomes are destroyed; however, a few always survive to repopulate the blood and maintain the infection. The key to this resurgence is the trypanosome's ability to change the protein that coats its surface. Each trypanosome is covered with about 10 million molecules of a single glycoprotein. When the immune system recognizes this protein coat, the infecting trypanosome is in trouble; immune cells will trap and destroy it. However, before all the trypanosomes in the animal are completely wiped out, a few manage to change their



Trypanosomes among red blood cells.

surface glycoprotein to one that is not immediately recognized by the immune system. These altered trypanosomes escape destruction and proliferate. Eventually, the immune system will learn to recognize them too, but in the meantime another group of altered trypanosomes arises to keep the infection going. The seemingly endless supply of molecular disguises available to trypanosomes is due to a large array of genes that encode the variant surface glycoproteins (VSGs) coating these organisms. At any one time, only one of these genes is expressed; all the others are silent. However, during the course of an infection the identity of the expressed gene changes. With each change, the trypanosomes acquire a new surface protein and manage to stay one step ahead of the animal's immune defenses. Thus, the infection is maintained for weeks or even months until, through exhaustion, the animal dies.

Ways of Regulating Eukaryotic Gene Expression: An Overview

Eukaryotic gene expression can be regulated at the transcriptional, processing, or translational levels.

DIMENSIONS OF EUKARYOTIC GENE REGULATION

The story of how trypanosomes evade attacks by the immune system is a story about gene regulation. Different *vsg* genes are expressed at different times—that is, the *vsg* genes are temporally regulated. Among eukaryotes, especially multicellular organisms like ourselves, genes are also regulated in a spatial dimension. Multicellular organisms contain many different cell types organized into tissues and organs. A particular gene might be expressed in blood cells, but never in nerve cells. Another gene might have just the opposite expression profile. The regulation that creates such differences in gene expression underlies the anatomical and physiological complexity of multicellular eukaryotes.

As in prokaryotes, the expression of genes in eukaryotes involves the transcription of DNA into RNA and the subsequent translation of that RNA into polypeptides. However, prior to translation, most eukaryotic RNA is "processed." During processing, the RNA is capped at its 5' end, polyadenylated at its 3' end, and altered internally by losing its noncoding intron sequences (see Chapter 11). Prokaryotic RNAs typically do not undergo these terminal and internal modifications.

Gene expression is more complicated in eukaryotes than it is in prokaryotes because eukaryotic cells are compartmentalized by an elaborate system of membranes. This compartmentalization subdivides the cells into separate organelles, the most conspicuous one being the nucleus; eukaryotic cells also possess mitochondria, chloroplasts (if they are plant cells), and an endoplasmic reticulum. Each of these organelles performs a different function. The nucleus stores the genetic material, the mitochondria and chloroplasts recruit energy, and the reticulum transports materials within the cell.

The subdivision of eukaryotic cells into organelles physically separates the events of gene expression. The primary event, transcription of DNA into RNA, occurs in the nucleus. RNA transcripts are also modified in the nucleus by capping, polyadenylation, and the removal of introns. The resulting messenger RNAs are then exported to the cytoplasm where they become associated with ribosomes, many of which are located on the membranes of the endoplasmic reticulum. Once associated with ribosomes, these mRNAs are translated into polypeptides. This physical separation of the events of gene expression makes it possible for regulation to occur in different places (**■ Figure 19.1**). Regulation can occur in the nucleus at either the DNA or RNA level, or in the cytoplasm at either the RNA or polypeptide level.



In prokaryotes, gene expression is regulated mainly by controlling the transcription of DNA into RNA. A gene that is not transcribed is simply not expressed. Transcription occurs in prokaryotes when negative regulatory molecules such as the *lac* repressor protein have been removed from the vicinity of a gene and positive regulatory molecules such as the catabolite activator protein (CAP)/cyclic AMP complex have bound to it (Chapter 18). These protein–DNA interactions control whether or not a gene is accessible to RNA polymerase. Furthermore, the mechanisms that have evolved to control transcription in these organisms respond quickly to environmental changes. As we discussed in Chapter 18, this hair-trigger control is an efficient strategy for prokaryotic survival.

The control of transcription is more complex in eukaryotes than it is in prokaryotes. One reason is that genes are sequestered in the nucleus. Before environmental signals can have any effect on the level of transcription, they must be transmitted from

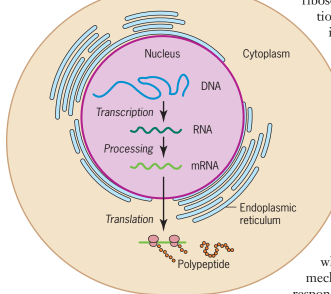


FIGURE 19.1 Eukaryotic gene expression showing the stages at which expression can be regulated: transcription, processing, and translation.

the cell surface, where they are usually received, through the cytoplasm and the nuclear membrane, and onto the chromosomes. Eukaryotic cells therefore need fairly elaborate internal signaling systems to control the transcription of DNA. Another complicating factor is that many eukaryotes are multicellular. Environmental cues may have to pass through layers of cells in order to have an impact on the transcription of genes in a particular tissue. Intercellular communication is therefore an important aspect of eukaryotic transcriptional regulation.

As in prokaryotes, eukaryotic transcriptional regulation is mediated by protein–DNA interactions. Positive and negative regulator proteins bind to specific regions of the DNA and stimulate or inhibit transcription. As a group, these proteins are called **transcription factors**. Many different types have been identified, and most seem to have characteristic domains that allow them to interact with DNA. The structure of these proteins, and the nature of their interactions with DNA, will be discussed in a later section.

ALTERNATE SPLICING OF RNA

Most eukaryotic genes possess introns, noncoding regions that interrupt the sequence that specifies the amino acids of a polypeptide. Each intron must be removed from the RNA transcript of a gene in order for the coding sequence to be expressed properly. As we discussed in Chapter 11, this process involves the precise joining of the coding sequences, or exons, into a messenger RNA. The formation of the mRNA is mediated by tiny nuclear organelles called spliceosomes.

Genes with multiple introns present a curious problem to the RNA splicing machinery. These introns can be removed separately or in combination, depending on how the splicing machinery interacts with the RNA. If two successive introns are removed together, the exon between them will also be removed. Thus, the splicing machinery has the opportunity to modify the coding sequence of an RNA by deleting some of its exons. This phenomenon of splicing an RNA transcript in different ways is apparently a way of economizing on genetic information. Instead of duplicating genes, or pieces of genes, the *alternate splicing* of transcripts makes it possible for a single gene to encode different polypeptides.

One example of alternate splicing occurs during the expression of the gene for troponin T, a protein found in the skeletal muscles of vertebrates; the size of this protein ranges from about 150 to 250 amino acids. In the rat, the troponin T gene is more than 16 kb long and contains 18 different exons (**Figure 19.2**). Transcripts of this gene are spliced in different ways to create a large array of mRNAs. When these are translated, many different troponin T polypeptides are produced. All these polypeptides share amino acids from exons 1–3, 9–15, and 18. However, the regions encoded by exons 4–8 may be present or absent, depending on the splicing pattern, and apparently in any combination. Additional variation is provided by the presence or absence of regions encoded by exons 16 and 17; if 16 is present, 17 is not, and vice versa. These different forms of troponin T presumably function in slightly different ways within the muscles, contributing to the variability of muscle cell action. To appreciate the variation that can be generated by alternate splicing of RNA, work through Solve It: Counting mRNAs.

CYTOPLASMIC CONTROL OF MESSENGER RNA STABILITY

Messenger RNAs are exported from the nucleus to the cytoplasm where they serve as templates for polypeptide synthesis. Once in the cytoplasm, a particular mRNA can be translated by several ribosomes that move along it in sequential fashion. This translational assembly line continues until the mRNA is degraded. Messenger RNA

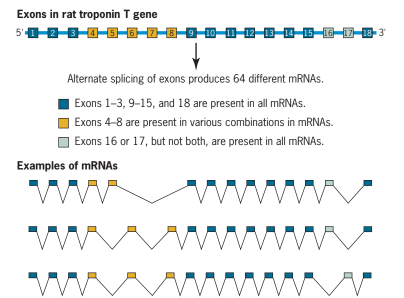


FIGURE 19.2 Alternate splicing of transcripts from the rat troponin T gene. Only 3 of the possible 64 different mRNAs are shown.

Solve It!

Counting mRNAs

The primary transcript of a gene with one intron is spliced to produce a single kind of mRNA. With two introns, the transcript can be alternately spliced; each of the introns can be removed separately, or the two introns can be removed together along with the exon between them. Thus, two different mRNAs can be generated from the transcript of such a gene. How many different mRNAs can be generated by alternate splicing of transcripts from genes with three or four introns? Assume that the first and last exons will be present in all the mRNAs, but that the internal exons may be present or absent, depending on the splicing pattern. What is the general formula for the number of mRNAs generated by alternate splicing of a transcript from a gene with *n* introns?

To see the solution to this problem, visit the Student Companion site.

degradation is therefore another control point in the overall process of gene expression. Long-lived mRNAs can support multiple rounds of polypeptide synthesis, whereas short-lived mRNAs cannot.

An mRNA that is rapidly degraded must be replenished by additional transcription; otherwise, the polypeptide it encodes will cease to be synthesized. This cessation of polypeptide synthesis may, of course, be part of a developmental program. Once the polypeptide has had its effect, it may no longer be needed; in fact, its continued synthesis may be harmful. In such cases, rapid degradation of the mRNA would be a reasonable way of preventing undesired polypeptide synthesis.

Messenger RNA longevity can be influenced by several factors. Poly(A) tails seem to stabilize mRNAs. The sequence of the 3' untranslated region (3' UTR) preceding a poly(A) tail also seems to affect mRNA stability. Several short-lived mRNAs have the sequence AUUUA repeated several times in their 3' untranslated regions. When this sequence is artificially transferred to the 3' untranslated region of more stable mRNAs, they, too, become unstable. Chemical factors, such as hormones, may also affect mRNA stability. In the toad *Xenopus laevis*, the *vitellogenin* gene is transcriptionally activated by the steroid hormone estrogen. However, in addition to inducing transcription of this gene, estrogen also increases the longevity of its mRNA.

Recent research has revealed that the stability of mRNAs and the translation of mRNAs into polypeptides are also regulated by small, noncoding RNA molecules called small interfering RNAs (siRNAs) or microRNAs (miRNAs). These regulatory RNA molecules, which are between 21 and 28 nucleotides long, are produced from larger, double-stranded RNAs in a wide variety of eukaryotic organisms, including fungi, plants, and animals. Short interfering and microRNAs base-pair with sequences in specific mRNAs; once paired, they either cause the mRNA to be cleaved and subsequently degraded, or they prevent the mRNA from being translated into a polypeptide. In plants, these small RNA molecules provide a critical defense against infection by RNA viruses, and in both plants and animals they regulate the expression of genes involved in maturation and development. We will discuss them in more detail later in this chapter.

- **KEY POINTS**
 - Proteins called transcription factors interact with DNA to control the transcription of eukaryotic genes.
 - Eukaryotic gene transcripts may be alternately spliced to produce messenger RNAs that encode distinct, but related, polypeptides.
 - The stability of eukaryotic messenger RNAs can influence the level of polypeptide synthesis.

Induction of Transcriptional Activity by Environmental and Biological Factors

Eukaryotic gene expression can be induced by environmental factors such as heat and by signaling molecules such as hormones and growth factors.

In their study of the *lactose* operon in *E. coli*, Jacob and Monod discovered that the genes for lactose metabolism were specifically transcribed when lactose was given to the cells. Thus, they demonstrated that lactose was an **inducer** of gene transcription. Following in the footsteps of Jacob and Monod, many researchers have attempted to identify specific inducers of eukaryotic gene transcription. Although these efforts

have met with considerable success, the overall extent to which eukaryotic genes are induced by environmental and nutritional factors seems to be less than it is in prokaryotes. Here we will consider two examples of inducible gene expression in eukaryotes.

TEMPERATURE: THE HEAT-SHOCK GENES

When organisms are subjected to the stress of high temperature, they respond by synthesizing a group of proteins that help to stabilize the internal cellular environment. These *heat-shock proteins*, found in both prokaryotes and eukaryotes, are among the most conserved polypeptides known. Comparisons of the amino acid sequences of heat-shock proteins from organisms as diverse as *E. coli* and *Drosophila* show that they are 40 to 50 percent identical—a remarkable finding considering the length of evolutionary time separating these organisms.

The expression of the heat-shock proteins is regulated at the transcriptional level; that is, heat stress specifically induces the transcription of the genes encoding these proteins (**Figure 19.3**). In *Drosophila*, for example, one of the heat-shock proteins called HSP70 (for *heat-shock protein*, molecular weight 70 kilodaltons) is encoded by a family of genes located in two nearby clusters on one of the autosomes. Altogether, there are five to six copies of these *hsp70* genes in the two clusters. When the temperature exceeds 33°C, as it does on hot summer days, each of the genes is transcribed into RNA, which is then processed and translated to produce HSP70 polypeptides. This heat-induced transcription of the *hsp70* genes is mediated by a polypeptide called the heat-shock transcription factor, or HSTF, which is present in the nuclei of *Drosophila* cells. When *Drosophila* are heat stressed, the HSTF is chemically altered by phosphorylation. In this altered state, it binds specifically to nucleotide sequences upstream of the *hsp70* genes and makes the genes more accessible to RNA polymerase II, the enzyme that transcribes most protein-encoding genes. The transcription of the

hsp70 genes is then vigorously stimulated. The sequences to which the phosphorylated

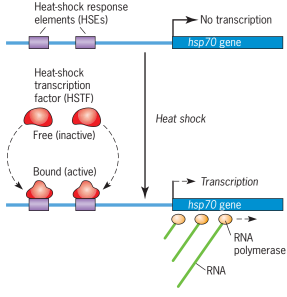
SIGNAL MOLECULES: GENES THAT RESPOND TO HORMONES

HSTF binds are called *heat-shock response elements* (HSEs).

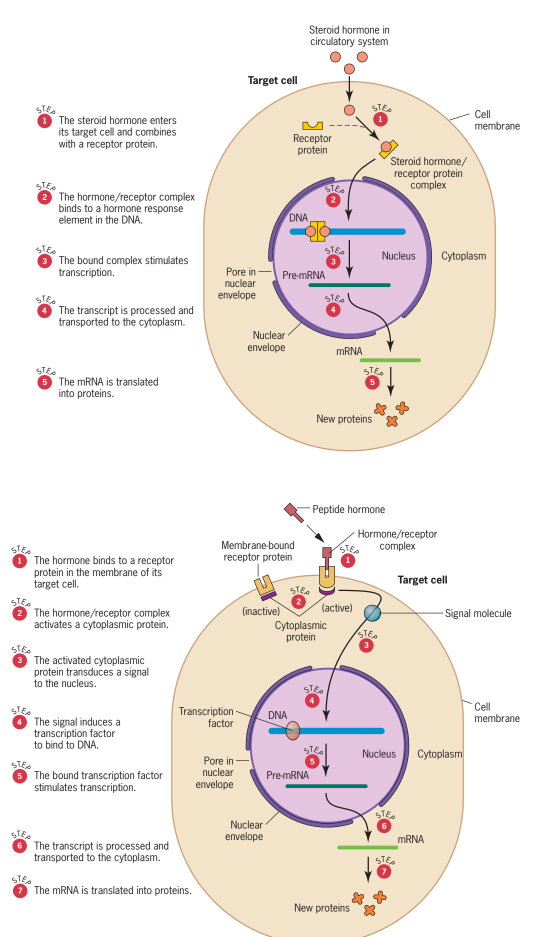
In multicellular eukaryotes, one type of cell can signal another by secreting a **hormone**. Hormones circulate through the body, make contact with their target cells, and then initiate a series of events that regulate the expression of particular genes. In animals there are two general classes of hormones. The first class, the *steroid hormones*, are small, lipid-soluble molecules derived from cholesterol. Because of their lipid nature, they have little or no trouble passing through cell membranes. Examples are estrogen and progesterone, which play important roles in female reproductive cycles; testosterone, a hormone of male differentiation and behavior; the glucocorticoids, which are involved in regulating blood sugar levels; and ecdysone, a hormone that controls developmental events in insects. Once these hormones have entered a cell, they interact with cytoplasmic or nuclear proteins called *hormone receptors*. The receptor/ hormone complex that is formed then interacts with the DNA where it acts as a transcription factor to regulate the expression of certain genes (**■** Figure 19.4).

The second class of hormones, the *peptide hormones*, are linear chains of amino acids. Like all other polypeptides, these molecules are encoded by genes. Examples are insulin, which regulates blood sugar levels, somatotropin, which is a growth hormone, and prolactin, which targets tissues in the breasts of female mammals. Because peptide hormones are typically too large to pass freely through cell membranes, the signals they convey must be transmitted to the interior of cells by *membrane-bound receptor proteins* (**■** Figure 19.5). When a peptide hormone interacts with its receptor, it causes a conformational change in the receptor that eventually leads to changes in other proteins inside the cell. Through a cascade of such changes, the hormonal signal is transmitted through the cytoplasm of the cell and into the nucleus, where it ultimately has the effect of regulating the expression of specific genes. This process of transmitting the hormonal signal through the cell and into the nucleus is called signal transduction.

Hormone-induced gene expression is mediated by specific sequences in the DNA. These sequences, called *hormone response elements* (*HREs*), are analogous to



■ FIGURE 19.3 Induction of transcription from the *Drosophila hsp70* gene by heat shock. The HSEs are located between 40 and 90 base pairs upstream of the transcription initiation site (bent arrow). ■ FIGURE 19.4 Regulation of gene expression by steroid hormones. The hormone interacts with a receptor inside its target cell. In this example the receptor is in the cytoplasm; other steroid hormone receptors are located in the nucleus. The steroid/ hormone receptor complex moves into the nucleus where it activates the transcription of particular genes.



■ FIGURE 19.5 Regulation of gene expression by peptide hormones. The hormone (an extracellular signal) interacts with a receptor in the membrane of its target cell. The resulting hormone/receptor complex activates a cytoplasmic protein that triggers a cascade of intracellular changes. These changes transmit the signal into the nucleus, where a transcription factor stimulates the expression of particular genes. the heat-shock response elements discussed earlier. They are situated near the genes they regulate and serve to bind specific proteins, which then act as transcription factors. With steroid hormones such as estrogen, the HREs are bound by the hormone/ receptor complex, which then stimulates transcription. The vigor of this transcriptional response depends on the number of HREs present. When there are multiple response elements, hormone/receptor complexes bind cooperatively with each other, significantly increasing the rate of transcription; that is, a gene with two response elements is transcribed more than twice as vigorously as a gene with only one. With peptide hormones, the receptor usually remains in the cell membrane, even after it has formed a complex with the hormone. The hormonal signal is therefore conveyed to the nucleus by other proteins, some of which bind to sequences near the genes that are regulated by the hormone. These proteins then act as transcription factors to control the expression of the genes.

Transcriptional activity can be induced by many other kinds of proteins that are not hormones in the classical sense—that is, not produced by a particular gland or organ. These include a variety of secreted, circulating molecules such as nerve growth factor, epidermal growth factor, and platelet-derived growth factor, and other noncirculating molecules associated with cell surfaces or with the matrix between cells. Although each of these proteins has its own peculiarities, the general mechanism whereby they induce transcription resembles that of the peptide hormones. An interaction between the signaling protein and a membrane-bound receptor initiates a chain of events inside the cell that ultimately results in specific transcription factors binding to particular genes, which are then transcribed.

- Transcription of the hsp70 genes in response to increased temperature is mediated by a heat-shock transcription factor.
- Steroid hormones and their receptor proteins form complexes that act as transcription factors to regulate the expression of specific genes.
- Peptide hormones interact with membrane-bound receptor proteins to activate a signaling system that regulates the expression of specific genes.

KEY POINTS

Molecular Control of Transcription in Eukaryotes

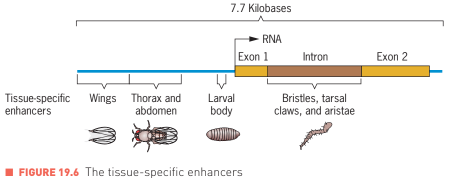
Much of the current research on eukaryotic gene expression focuses on the factors that control transcription. This heavy emphasis on transcriptional control is partly due to the development of experimental techniques that have permitted this aspect of The transcription of eukaryotic genes is regulated by interactions between proteins and DNA sequences within or near the genes.

gene regulation to be analyzed in great detail. However, it is also due to the appeal of ideas that emerged from the study of prokaryotic genes. In both prokaryotes and eukaryotes, transcription is the primary event in gene expression; it is therefore the most fundamental level at which gene expression can be controlled.

DNA SEQUENCES INVOLVED IN THE CONTROL OF TRANSCRIPTION

Transcription is initiated in the promoter of a gene, the region recognized by the RNA polymerase. However, as we discussed in Chapter 11, the accurate initiation of transcription from eukaryotic gene promoters requires several accessory proteins, or *basal transcription factors*. Each of these proteins binds to a sequence within the promoter to facilitate the proper alignment of the RNA polymerase on the template strand of the DNA.





of the *Drosophila yellow* gene.

The transcription of eukaryotic genes is also controlled by a variety of *special transcription factors*, such as those involved in the regulation of the heatand hormone-inducible genes we have discussed. These factors bind to response elements, or, more generally, to sequences called **enhancers** located in the vicinity of a gene. The special transcription factors that bind to these enhancers may interact with the basal transcription factors and the RNA polymerase, which bind to the promoter of a gene. The interactions that take place among the special transcription factors, the basal transcription factors, and the RNA polymerase regulate the transcriptional activity of a gene.

Enhancers exhibit three fairly general properties: (1) they act over relatively large distances—up to several thousand base pairs from their regulated gene(s); (2) their influence on gene expression is independent of orientation—they function equally well in either the normal or inverted orientation within the DNA; and (3) their effects are independent of position—they can be located upstream, downstream, or within an intron of a gene and still have profound effects on the gene's expression. These three characteristics distinguish enhancers from promoters, which are typically located immediately upstream of the gene and which function only in one orientation.

Enhancers can be relatively large, up to several hundred base pairs long. They sometimes contain repeated sequences that have partial regulatory activity by themselves. Most enhancers function in a tissue-specific manner; that is, they stimulate transcription only in certain tissues. In other tissues they are simply ignored. A clear example of this tissue specificity comes from the study of the yellow gene in Drosophila (**Figure 19.6**). This gene is responsible for pigmentation in many parts of the body in the wings, legs, thorax, and abdomen. Wild-type flies show a dark brownish-black pigment in all these structures, whereas mutant flies show a lighter yellowish-brown pigment. However, in some mutants, there is a mosaic pattern of pigmentation, brownish-black in some tissues and yellowish-brown in others. These mosaic patterns are due to mutations that alter the transcription of the *yellow* gene in some tissues but not in others. Pamela Geyer and Victor Corces have shown that the yellow gene is regulated by several enhancers, some of which are located within an intron, and that each enhancer activates transcription in a different tissue. If, for example, the enhancer for expression in the wing is mutated, the bristles on the wings are vellowish-brown instead of brownish-black. The battery of enhancers associated with the yellow gene allows its expression to be controlled in a tissue-specific way. To see another way of studying enhancers, work through Problem-Solving Skills: Defining the Sequences Required for a Gene's Expression.

How do enhancers influence the transcription of genes? The results of many studies indicate that the proteins that bind to enhancers influence the activity of the proteins that bind to promoters, including the basal transcription factors and the RNA polymerase. The two types of proteins are brought into physical contact by a multimeric complex consisting of at least 20 different proteins. This *mediator complex* appears to bend the DNA in such a way that the proteins bound to an enhancer are juxtaposed to those bound at the promoter. In this way, then, proteins bound to the enhancer exert control over transcription, which is initiated at the promoter.

PROTEINS INVOLVED IN THE CONTROL OF TRANSCRIPTION: TRANSCRIPTION FACTORS

Research over the last three decades has identified a large number of eukaryotic proteins that stimulate transcription. Many of these proteins appear to have at least two important chemical domains: a DNA-binding domain and a transcriptional activation domain. These domains may occupy separate parts of the molecule, or they

PROBLEM-SOLVING SKILLS

Defining the Sequences Required for a Gene's Expression

THE PROBLEM

The tubulins are important proteins of the cytoskeleton in eukaryotes. In Arabidopsis thaliana the tubulin encoded by the TUA1 gene is expressed primarily in pollen. To determine the sequences responsible for this tissue-specific expression, 533 base pairs of DNA upstream of the TUA1 transcription start site plus the first 56 base pairs of the 5' untranslated region of the TUA1 gene were fused to the coding sequence of the β -glucuronidase (GUS) gene from *E. coli.* β -glucuronidase catalyzes the conversion of a colorless substance called X-gluc into a dark blue pigment. Thus, the appearance of blue pigment in X-gluc-treated material is an indication that the GUS gene is being expressed. When this assay was applied to Arabidopsis plants that had been genetically transformed with the GUS gene fused behind the upstream sequences of the TUA1 gene, the pollen turned dark blue; all the other tissues remained colorless. The entire experiment was then repeated using progressively shorter segments from the TUA1 upstream sequences to drive expression of the *GUS* gene. From the results shown in **Figure 1**, what part of the upstream region is required for expression of the TUA1 gene?

FACTS AND CONCEPTS

- 1. The region upstream of a gene's transcription start site contains the gene's promoter.
- 2. This region may also contain enhancers that regulate the gene's expression in a spatially or temporally specific way.
- 3. The 5' untranslated region of a gene lies between the transcription start site and the translation start site.
- **4.** *E. coli* genes such as *GUS* can be expressed in eukaryotes such as Arabidopsis if they are fused to eukaryotic promoters.

ANALYSIS AND SOLUTION

In this series of experiments, GUS is a "reporter" that tells us if the upstream sequences from the TUA1 gene are able to drive gene expression. All but the smallest of the upstream sequences can function as a successful driver. Thus, there must be a sequence between base pairs -97 and -39 in the upstream sequence of TUA1 that is

in pollen 5' UTR GUS TUAI Upstream sequences + -533 +1 56 + -380 + -332 -271 -217 -97 -39

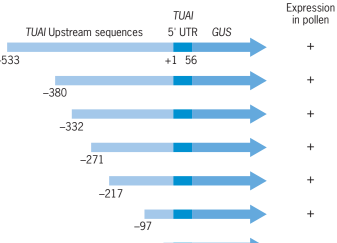
FIGURE 1 Expression of *TUA1/GUS* transgenes in *Arabidopsis* pollen. Progressively shorter segments from the upstream region of the *TUA1* gene and a short sequence from the 5' untranslated region (UTR) of this gene have been fused to the coding sequences of the GUS gene from E. coli. +1 is the transcription start site of the TUA1 gene. Nucleotides to the left of this site are indicated by negative numbers. GUS activity in transgenic pollen is indicated by a plus sign; no GUS activity is indicated by a minus sign. For further details see Carpenter, J., S. E. Ploense, D. P. Snustad, and C. D. Silflow. 1992. Preferential expression of an α -tubulin gene of Arabidopsis in pollen. The Plant Cell 4: 557–571.

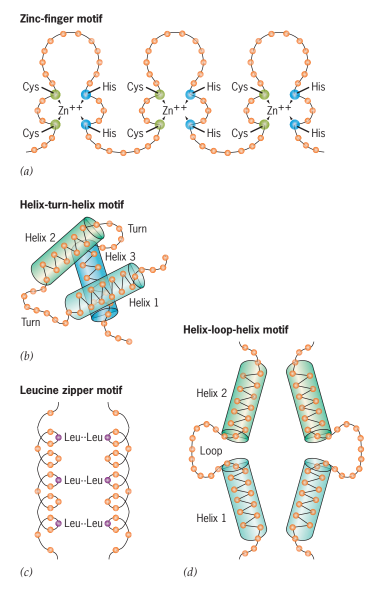
critical for the gene's expression. Without this sequence, the TUA1 gene cannot be expressed. Furthermore, this sequence is sufficient to drive TUA1 expression in mature pollen. Thus, it functions as an enhancer controlling the tissue-specific expression of the TUA1 gene.

For further discussion visit the Student Companion site.

may be overlapping. In the GAL4 transcription factor from yeast, for example, the DNA-binding domain is situated near the amino terminus of the polypeptide. Two transcriptional activation domains are present in this polypeptide, one more or less in the middle and one near the carboxy terminus. In the steroid hormone receptor proteins, which are transcription factors in animals, the DNA-binding domain is centrally located and seems to overlap a transcriptional activation domain that extends toward the amino terminus. Steroid hormone receptors also have a third domain that specifically binds the steroid hormone.

Transcriptional activation appears to involve physical interactions between proteins. A transcription factor that has bound to an enhancer may make contact with one or more proteins at other enhancers, or it may interact directly with proteins that have bound in the promoter region. Through these contacts and interactions, the transcriptional activation domain of the factor may then induce conformational changes in the assembled proteins, paving the way for the RNA polymerase to initiate transcription.





■ FIGURE 19.7 Structural motifs within different types of transcription factors. (a) Zinc-finger motifs in the mammalian transcription factor SP1. (b) Helix-turn-helix motif in a homeodomain transcription factor. (c) A leucine zipper motif that allows two polypeptides to dimerize and then bind to DNA. (d) A helix-loop-helix motif that allows two polypeptides to dimerize and then bind to DNA.

Many eukaryotic transcription factors have characteristic structural motifs that result from associations between amino acids within their polypeptide chains. One of these motifs is the *zinc finger*; a short peptide loop that forms when two cysteines in one part of the polypeptide and two histidines in another part nearby jointly bind a zinc ion; the peptide segment between the two pairs of amino acids then juts out from the main body of the protein as a kind of finger (**Figure 19.7a**). Mutational analysis has demonstrated that these fingers play important roles in DNA binding.

A second motif in many transcription factors is the *helix-turnhelix*, a stretch of three short helices of amino acids separated from each other by turns (**Figure 19.7***b*). Genetic and biochemical analyses have shown that the helical segment closest to the carboxy terminus is required for DNA binding; the other helices seem to be involved in the formation of protein dimers. In many transcription factors, the helix-turn-helix motif coincides with a highly conserved region of approximately 60 amino acids called the *homeodomain*, so named because it occurs in proteins encoded by the homeotic genes of Drosophila. Classical analyses have demonstrated that mutations in these genes alter the developmental fates of groups of cells (Chapter 20). Thus, for example, mutations in the Antennapedia gene can cause antennae to develop as legs. This bizarre phenotype is an example of a homeotic transformationthe substitution of one body part for another during the developmental process. Molecular analyses of the homeotic genes in Drosophila have demonstrated that each encodes a protein with a homeodomain and that these proteins can bind to DNA. The homeodomain proteins stimulate the transcription of particular genes in a spatially and temporally specific manner during development. Homeodomain proteins have also been identified in other organisms, including humans, where they may play important roles as transcription factors.

A third structural motif found in transcription factors is the *leucine zipper*; a stretch of amino acids with a leucine at every seventh position (**Figure 19.7***c*). Polypeptides with this feature can form dimers by interactions between the leucines in each of their zipper regions. Usually, the zipper sequence is adjacent to a positively charged stretch of amino acids. When two zippers interact, these charged regions splay out in opposite directions, forming a surface that can bind to negatively charged DNA.

A fourth structural motif found in some transcription factors is the *helix-loop-helix*, a stretch of two helical regions of amino acids separated by a nonhelical loop (**Figure 19.7***d*). The helical regions permit dimerization between two polypeptides.

Sometimes the helix-loop-helix motif is adjacent to a stretch of basic (positively charged) amino acids, so that when dimerization occurs, these amino acids can bind to negatively charged DNA. Proteins with this feature are denoted *basic HLH*, or *bHLH*, proteins.

Transcription factors with dimerization motifs such as the leucine zipper or the helix-loop-helix could, in principle, combine with polypeptides like themselves to form homodimers, or they could combine with different polypeptides to form heterodimers. This second possibility suggests a way in which complex patterns of gene expression can be achieved. The transcription of a gene in a particular tissue might depend on activation by a heterodimer, which could form only if its constituent polypeptides were synthesized in that tissue. Moreover, these two polypeptides would have to be present in the correct amounts to favor the formation of the heterodimer over the corresponding homodimers. Subtle modulations in gene expression might therefore be achieved by shifting the concentrations of the two components of a heterodimer.

- Enhancers act in an orientation-independent manner over considerable distances to regulate transcription from a gene's promoter.
- Transcription factors recognize and bind to specific DNA sequences within enhancers.
- Transcription factors possess characteristic structural motifs such as the zinc finger, the helix-turn-helix, the leucine zipper, and the helix-loop-helix.

KEY POINTS

Posttranscriptional Regulation of Gene Expression by RNA Interference

Although a great deal of eukaryotic gene regulation occurs at the transcriptional level, recent research has demonstrated that posttranscriptional mechanisms also play important roles in regulating the expression of eukaryotic genes. Some of these mechanisms involve small, noncoding RNAs. By base-pairing with Short noncoding RNAs may regulate the expression of eukaryotic genes by interacting with the messenger RNAs produced by these genes.

target sequences in messenger RNA molecules, these small RNAs interfere with gene expression. Hence, this type of posttranscriptional gene regulation is called **RNA inter-ference**, often abbreviated as **RNAi**. Most types of eukaryotic organisms are capable of RNAi. Among the model genetic organisms, this phenomenon has been well studied in the nematode *Caenorhabditis elegans*, in *Drosophila*, and in *Arabidopsis*. It also exists in mammals, including humans. As we will see, the widespread capacity of eukaryotic organisms to regulate gene expression by RNAi has allowed geneticists to analyze the functions of genes in organisms that are not amenable to standard genetic approaches.

RNAi PATHWAYS

The phenomenon of RNA interference, which is summarized in **Figure 19.8**, involves small RNA molecules called short interfering RNAs (siRNAs) or microRNAs (miRNAs). These molecules, 21 to 28 base pairs long, are produced from larger, double-stranded RNA molecules by the enzymatic action of proteins that are double-stranded RNA-specific endonucleases. Because these endonucleases "dice" large RNA into small pieces, they are called *Dicer* enzymes. The nematode *Caenorhabditis elegans* produces a single kind of Dicer enzyme; *Drosophila* produces two different Dicer enzymes; and *Arabidopsis* produces at least three. In *C. elegans* and *Drosophila*, these enzymes act in the cytoplasm; in *Arabidopsis*, they probably act in the nucleus. The siRNAs and miRNAs produced by Dicer activity are base-paired throughout their lengths except at their 3' ends, where two nucleotides are unpaired.

In the cytoplasm, siRNAs and miRNAs become incorporated into ribonucleoprotein particles. The double-stranded siRNA or miRNA in these particles is unwound, and one of its strands is preferentially eliminated. The surviving single strand of RNA is then able to interact with specific messenger RNA molecules. This interaction is mediated by base-pairing between the single strand of RNA in the RNA–protein complex and a complementary sequence in the messenger RNA molecule. Because this interaction prevents the expression of the gene that produced the mRNA, the RNA–protein particle is called an **RNA-Induced Silencing Complex (RISC)**.

RISCs from different organisms vary in size and composition. However, they all contain at least one molecule from the whimsically named Argonaute family of proteins. The function of these proteins is not fully understood. Whenever the base-pairing between the RNA within the RISC and the target sequence in the mRNA is perfect or nearly so, the RISC cleaves the target mRNA in the middle of the basepaired region. The cleaved mRNA is then degraded. After cleavage, the RISC may associate with another molecule of mRNA and induce its cleavage. Because a RISC

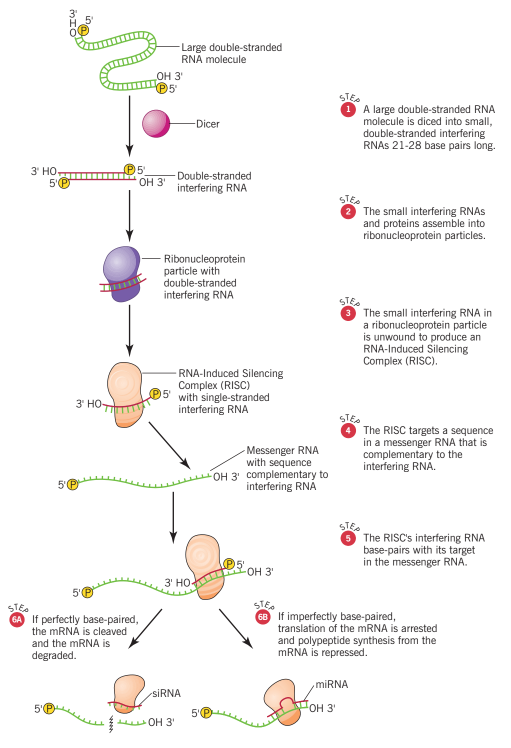


FIGURE 19.8 Summary of events involved in RNA interference pathways.

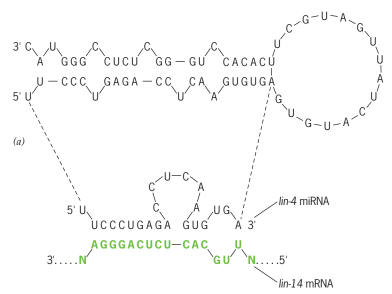
may be used repeatedly without losing its ability to target and cleave mRNA, it behaves as a catalyst. RISC-associated RNAs that result in mRNA cleavage are usually termed **short interfering RNAs**. Whenever the RNA within the RISC pairs imperfectly with its target sequence, the mRNA is usually not cleaved; instead, translation of the mRNA is inhibited. RISC-associated RNAs that have this effect are usually termed **microRNAs**. In animals, the sequences targeted by RISCs are found in the 3' untranslated regions of mRNA molecules, and often these sequences are present several times within the 3' untranslated region (UTR). In plants, the sequences targeted by RISCs are usually located within the coding region of the mRNA, or within the mRNA's 5' UTR.

SOURCES OF SHORT INTERFERING RNAs AND MicroRNAs

Some of the small RNA molecules that induce RNAi are derived from the transcripts of microRNA genes. These genes, usually denoted by the symbol *mir*, are found in the genomes of many kinds of eukaryotes; about 100 mir genes are present in the C. elegans and Drosophila genomes, and about 250 are present in vertebrate genomes. Initially, a few of these genes were identified through analysis of mutations that altered the regulation of other genes. When the *mir* genes defined by these mutations were analyzed at the molecular level, they were found to have little or no proteincoding potential. Instead, they possessed a peculiar structure. Each of them contained a short stretch of nucleotides repeated in opposite orientations around a short intervening segment of DNA. When transcribed, this inverted repeat structure generates an RNA that can fold back on itself to form a short double-stranded stem at the base of a single-stranded loop (**Figure 19.9a**). An enzyme called Drosha recognizes this stem-loop region and excises it from the primary transcript of the *mir* gene. The liberated stem-loop is then exported to the cytoplasm where it is cleaved by Dicer to form an miRNA. In *C. elegans*, where this process was discovered, Dicer removes the loop and trims the stem to a length of 22 nucleotides on each of its strands. After maturing in a RISC, the miRNA—now single-stranded—can target a sequence in the mRNA produced by another gene. **Figure 19.9** shows base-pairing between the miRNA from the C. elegans mir gene lin-4 and one of this miRNA's targets in the 3' UTR of the mRNA from a protein-coding gene, *lin-14*. Through this base-pairing, the *lin-4* miRNA represses translation of the *lin-14* mRNA.

Since the discovery of these mutationally defined *mir* genes, many other *mir* genes have been found by using computer programs to screen the genomic DNA sequences of *C. elegans*, *Drosophila*, and other model organisms for the characteristic inverted repeat structure. Many of the candidate *mir* genes identified by this computer-based genomic approach have been verified by detecting miRNAs derived from these genes in cell extracts. Genes whose mRNAs contain sequences targeted by miRNAs are also being identified by a combination of computer-based analysis and *in vivo* experimentation. Many of these genes encode transcription factors or other developmentally significant proteins.

Some of the RNAs that induce RNAi are derived from the transcription of other elements in the genome such as transposons and transgenes, and they are also derived from RNA viruses. The ways in which these types of interfering RNAs are formed are not fully



■ **FIGURE 19.9** Regulation of gene expression by RNA interference. (a) Stem-loop structure of a transcript from the *C. elegans* microRNA gene *lin-4*. (b) Base-pairing between the microRNA derived from the *lin-4* transcript and a sequence in the 3' untranslated region of the *lin-14* messenger RNA. understood. Some aspect of the transposon, transgene, or viral RNA marks it as unusual. In plants and nematodes, these unusual RNAs can be copied into complementary RNA molecules by enzymes known as RNA-dependent RNA polymerases (RdRPs). If the complementary RNA strand remains base-paired with the template from which it was made, the resulting double-stranded RNA molecule can be diced into siRNAs by Dicer-type enzymes; then, the siRNAs produced by Dicer can enter an RNAi pathway and target the RNA population that originally gave rise to them. In this fashion, potentially troublesome RNAs derived from transposons, transgenes, or viruses can be targeted for repression or degradation. This application of RNAi may represent its most primitive function—to protect organisms against viral infections and runaway transposition. By contrast, the intricate miRNA-based systems for gene regulation evident in organisms like *C. elegans* seem to represent highly evolved applications of RNAi.

Researchers have discovered that RNAi can also be induced by double-stranded RNA that has been prepared in vitro by transcription from cloned genes or gene segments (see A Milestone in Genetics: The Discovery of RNA Interference on the Student Companion site). The DNA is transcribed in both directions by inserting it between promoters in opposite orientations in a suitable cloning vector or by inserting inverted copies of the DNA downstream of a single promoter (see Chapter 16). Double-stranded RNA molecules derived from the transcripts of such clones can be introduced into cultured cells; they can also be injected into living organisms. Once inside cells, the double-stranded RNA enters an RNAi pathway. It is diced into siRNA molecules, which are then incorporated into RNA-protein complexes and targeted to mRNAs containing complementary sequences. The targeted mRNAs are usually degraded. Thus, treating cells or organisms with a particular type of double-stranded RNA has the effect of knocking out or knocking down the expression of the gene that corresponds to that RNA. It is therefore equivalent to inducing an amorphic or hypomorphic mutation in the gene. Using this approach, geneticists have been able to study the consequences of ablating or attenuating the expression of particular genes in a wide variety of organisms, including some in which genetic analysis is difficult, slow, or impossible. Thus, RNAi is now being used to analyze the function of genes in fish, rodents, and humans, as well as in simpler model organisms such as C. elegans, Drosophila, and Arabidopsis. To see one application of this technology, work through Solve It: Using RNAi in Cell Research.

- **KEY POINTS** Short interfering RNAs and microRNAs are produced from larger double-stranded precursors by the action of Dicer-type endonucleases.
 - In RNA-Induced Silencing Complexes (RISCs), siRNAs and miRNAs become single stranded so they can target complementary sequences in messenger RNA molecules.
 - Messenger RNA that has been targeted by siRNA is cleaved, and mRNA that has been targeted by miRNA is prevented from serving as a template for polypeptide synthesis.
 - Hundreds of genes for miRNAs are present in eukaryotic genomes.
 - Transposons and transgenes may stimulate the synthesis of siRNAs.
 - RNA interference is used as a research tool to knock out or knock down the expression of genes in cells and whole organisms.

Gene Expression and Chromatin Organization

Various aspects of chromatin organization influence the transcription of genes.

S. Eukaryotic chromosomes are composed of about equal parts of DNA and protein. Collectively, we refer to this material as **chromatin**. The chemical characteristics of chromatin vary along the length of a chromosome. In some regions, for example, the histones, which constitute the bulk of the protein in chromatin, are acetylated, and in other regions, some of the nucleotides in the DNA are methylated. These chemical modifications can influence the transcriptional activity of genes. Other aspects of chromatin organization—for instance, the presence of "packaging" proteins—play roles in gene regulation. In this section, we consider how the composition and organization of chromatin affects gene expression.

EUCHROMATIN AND HETEROCHROMATIN

Variation in the density of chromatin within the nuclei of cells leads to differential staining of sections of chromosomes. The deeply staining material is called **hetero-chromatin**, and its lightly staining counterpart is called **euchromatin**. What, if any, is the functional significance of these different types of chromatin?

A combination of genetic and molecular analyses has shown that the vast majority of eukaryotic genes are located in euchromatin. Moreover, when euchromatic genes are artificially transposed to a heterochromatic environment, they tend to function abnormally, and, in some cases, not to function at all. This impaired ability to function can create a mixture of normal and mutant characteristics in the same individual, a condition referred to as position-effect variegation. This term is used because the variability in the phenotype is caused by changing the position of the euchromatic gene, specifically by relocating it to the heterochromatin. Many examples of position-effect variegation have been discovered in Drosophila, usually in association with inversions or translocations that move a euchromatic gene into the heterochromatin. The *white mottled* allele is a good example. In this case, a wild-type allele of the *white* gene has been relocated by an inversion, with one break near the euchromatic white locus and the other in the basal heterochromatin of the X chromosome. This rearrangement interferes with the normal expression of the *white* gene and causes a mottled-eye phenotype (**Figure 19.10**). Apparently, the euchromatic white gene cannot function well in a heterochromatic environment. This and other examples have led to the view that heterochromatin represses gene function, perhaps because it is condensed into a form that is not accessible to the transcriptional machinery.

The behavior of the *white* gene in flies with this rearranged X chromosome indicates that gene expression can be influenced by conditions that do not alter the nucleotide sequence of the gene. Moreover, because the *white* gene is expressed in some patches of the eye, but not in others, we know that once these conditions are established, they are inherited clonally as the eye's cells divide. Because these conditions are superimposed on the basic structure of the *white* gene, we say that they are **epigenetic**. The Greek prefix "epi" means "above," and here it is used to convey the idea that a heritable state other than the actual sequence of the gene regulates the gene's expression. In this case, the heritable epigenetic state involves some aspect of chromatin organization near the repositioned *white* gene. In the sections that follow, we will encounter other examples of epigenetic regulation of gene expression.

MOLECULAR ORGANIZATION OF TRANSCRIPTIONALLY ACTIVE DNA

What is the molecular organization of transcriptionally active DNA? Is this DNA more "open" than nontranscribed DNA? These questions have been answered by measuring the sensitivity of DNA in chromatin to the action of pancreatic deoxyribonuclease I (DNase I), an enzyme that cleaves DNA molecules and degrades them into their constituent nucleotides. In 1976, Mark Groudine and Harold Weintraub demonstrated that transcriptionally active DNA is more sensitive to DNase I than nontranscribed DNA. Groudine and Weintraub extracted chromatin from chicken red blood cells and partially digested it with DNase I. Then they probed the residual chromatin material for sequences of two genes, β -globin, which is actively transcribed in red blood cells, and

Solve It!

Using RNAi in Cell Research

A researcher is studying the formation of centrosomes inside cultured human cells. These small organelles play an important role in orchestrating cell division. Centrosomes can be visualized by staining cells appropriately. The researcher hypothesizes that two proteins, γ -tubulin and CEP135, are needed for centrosome formation. The genes for these two proteins have been cloned from the human genome, and their sequences have been analyzed. Outline how the technique of RNA interference could be used to test the researcher's hypothesis. Explain what materials would be needed and how you would ascertain if the synthesis of γ -tublulin and CEP135 was blocked or reduced in cultured cells. How would you ascertain if centrosome formation was impaired? What controls would you include in these experiments?

To see the solution to this problem, visit the Student Companion site.

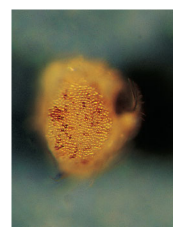


FIGURE 19.10 The variegated eye color phenotype of *Drosophila* that carry the *white mottled* allele in a rearranged X chromosome.

ovalbumin, which is not. They found that over 50 percent of the β -globin DNA had been digested by the DNase I enzyme, compared with only 10 percent of the ovalbumin DNA. These results strongly implied that the actively transcribed gene was more "open" to nuclease attack. Subsequent research has shown that the nuclease sensitivity of transcriptionally active genes depends on at least two small nonhistone proteins, HMG14 and HMG17 (HMG for *b*igh *m*obility group, because they have high mobility during gel electrophoresis). When these proteins are removed from active chromatin, nuclease sensitivity is lost; when they are added again, it is restored.

The treatment of isolated chromatin with a very low concentration of DNase I causes the DNA to be cleaved at a few specific sites, appropriately called *DNase I hypersensitive sites*. Some of these sites have been shown to lie upstream of transcriptionally active genes, in either promoter or enhancer regions. The functional significance of these hypersensitive sites is still unclear, but some evidence suggests that they may mark regions in which the DNA is locally unwound, perhaps because transcription has begun.

In the case of the human genes for β -globin, several DNase I hypersensitive sites are located in a 15-kb-long *locus control region* (*LCR*) upstream of the genes themselves (**■** Figure 19.11). The human β -globin genes reside in a cluster spanning 28 kilobases on chromosome 11. Each of the genes in the cluster is a duplicate of an ancestral β -globin gene. Over evolutionary time, the individual genes in the cluster have diverged from one another by random mutation so that today, each one of them encodes a slightly different polypeptide. In one of the genes, a nonsense mutation has abolished the ability to make a polypeptide. Such noncoding genes are called pseudogenes, and they are usually denoted by the Greek letter psi (Ψ)—thus, the (Ψ) β gene in this cluster.

The human β -globin genes are spatially and temporally regulated. In fact, a remarkable feature of this gene cluster is that its members are expressed at different times during development. The ϵ gene is expressed in the embryo, the two γ genes are expressed in the fetus, and the δ and β genes are expressed in infants and adults. This sequential activation of genes from one side to the other in the cluster is apparently related to the need to produce slightly different kinds of hemoglobin during the course of human development. Embryo, fetus, and infant have different oxygen requirements, different circulatory systems, and different physical environments. The temporal switching in β -globin gene expression is apparently an adaptation to this changing array of conditions.

The LCR of the β -globin gene cluster contains binding sites for transcription factors that preactivate the individual genes for transcription. Preactivation is detected by an increase in the sensitivity of the DNA within the LCR to digestion with low concentrations of DNase I. Transcription of the β -globin genes appears to require this preactivation and is stimulated by transcription factors that bind to specific enhancers in the β -globin gene complex. However, the tissue and temporal specificity of β -globin gene expression depends on sequences embedded in the LCR. Studies with transgenic mice indicate that the LCR is not simply a large collection of enhancers that exert control over the various β -globin genes. The LCR must be situated upstream of the β -globin genes and in its natural orientation in order to control gene expression properly. That is, it functions in an orientation-dependent manner.

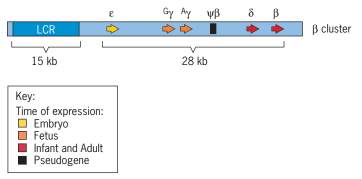


FIGURE 19.11 The β-globin gene cluster on human chromosome 11.

Enhancers typically function in an orientation-independent manner and in different positions relative to a gene's promoter. The LCR has one other feature that distinguishes it from simple enhancers: it can control β -globin gene expression when the entire gene cluster (LCR plus β -globin genes) is inserted in a different chromosomal position. Enhancers, by contrast, often fail to function when they and their associated genes are transposed to a different chromosomal location. Thus, the LCR seems to insulate the β -globin genes from the influence of the chromatin around them.

CHROMATIN REMODELING

Experiments that assess the sensitivity of DNA to digestion with DNase I have established that transcribed DNA is more accessible to

nuclease attack than nontranscribed DNA. Is transcribed DNA packaged in nucleosomes? If it is, what structural changes occur in the nucleosomes during transcription? Are the nucleosomes "opened" and "closed" as the RNA polymerase passes along the DNA template? Efforts to answer these questions have involved a combination of genetic and biochemical approaches that have demonstrated that transcribed DNA is indeed packaged into nucleosomes. However, in transcribed DNA, the nucleosomes are altered by multiprotein complexes that ultimately facilitate the action of the RNA polymerase. This alteration of nucleosomes in preparation for transcription is called **chromatin remodeling**.

Two general types of chromatin-remodeling complexes have been identified. One type is composed of enzymes that transfer acetyl groups to the amino acid lysine at specific positions in the histones of the nucleosomes. As a class, these enzymes are called *histone acetyl transferases (HATs)*. Numerous studies have shown that acetylation of histones is correlated with increased gene expression, perhaps because the addition of the acetyl groups loosens the association between the DNA and the histone octamers in the nucleosomes. *Kinases*—enzymes that transfer phosphate groups to molecules—may also play a role along with these chromatin-remodeling complexes. It is known, for example, that acetylation of lysine-14 in histone H4 is often preceded by phosphorylation of serine-10 in that molecule. Together, these two modifications of histone H4 seem to "open" the chromatin for increased transcriptional activity.

Another type of chromatin-remodeling complex disrupts nucleosome structure in the vicinity of a gene's promoter. The most intensively studied of these complexes is the SWI/SNF complex found in baker's yeast. This complex is named for the two types of mutations (*sw*itching-*i*nhibited and *s*ucrose *n*onfermenter) that led to the discovery of its constituent proteins. Related complexes have been found in the cells of other organisms, including humans. The SWI/SNF complex consists of at least eight proteins. It regulates transcription by sliding histone octamers along the associated DNA in nucleosomes; it can also transfer these octamers to other locations on a DNA molecule. The nucleosome shifting catalyzed by the SWI/SNF complex apparently gives transcription factors access to the DNA. These factors then stimulate a gene's expression.

We have discussed chromatin remodeling from the point of view of gene activation. However, active chromatin can also be remodeled into inactive chromatin. This reverse remodeling seems to involve two biochemical modifications to the histones in nucleosomes: deacetylation, catalyzed by the *bistone deacetylases (HDACs)*, and methylation, catalyzed by the *bistone methyl transferases (HMTs)*. As discussed in the next section, some of the nucleotides in the DNA may also be methylated by a group of enzymes called the *DNA methyl transferases (DNMTs)*. Chromatin that has been subjected to these modifications tends to be transcriptionally silent.

DNA METHYLATION

The chemical modification of nucleotides also appears to be important for the regulation of genes in some eukaryotes, especially mammals. Of the approximately 3 billion base pairs in a typical mammalian genome, about 40 percent are G:C base pairs, and about 2 to 7 percent of these are modified by the addition of a methyl group to the cytosine (**■** Figure 19.12). Most of the methylated cytosines are found in base-pair doublets with the structure

5' mCpG 3'

3' GpCm 5'

where mC denotes methylcytosine and the p between C and G denotes the phosphodiester bond between adjacent nucleotides in each DNA strand. This structure is often simply abbreviated by giving the composition of one strand, thus, mCpG. Methylated CpG dinucleotides can be detected by digesting DNA with restriction enzymes that are sensitive to chemical modifications of their recognition sites. For example, the enzyme *Hpa*II recognizes and cleaves the sequence CCGG; however, when the

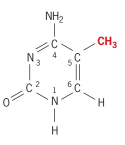


FIGURE 19.12 The structure of 5-methylcytosine.

second cytosine in this sequence is methylated, *Hpa*II cannot cleave the sequence. Thus, methylated and unmethylated DNAs give different patterns of restriction fragments when they are digested with this enzyme.

CpG dinucleotides occur less often than expected in mammalian genomes, probably because they have been mutated into TpG dinucleotides over the course of evolution. Moreover, the distribution of CpG dinucleotides is uneven, with numerous short segments of DNA having a much higher density of CpG dinucleotides than other regions of the genome. These CpG-rich segments, usually about 1 to 2 kb long, are called **CpG islands**. In the human genome, there are about 30,000 such islands, most being situated near transcription start sites. Molecular analysis has demonstrated that the cytosines in these islands are rarely, if ever, methylated, and that this un- or undermethylated state is conducive to transcription. Thus, DNA in the vicinity of a CpG island is hypersensitive to digestion with DNase I, and its nucleosomes are usually somewhat different than nucleosomes elsewhere in the genome—typically, there is less histone H1, and some of the core histones are acetylated.

Where methylated DNA is found, it is associated with transcriptional repression. This is most dramatically seen in female mammals where the inactive X chromosome is extensively methylated. Regions of the mammalian genome that contain repetitive sequences, including those regions that are rich in transposable elements, are also methylated, perhaps as a way of protecting the organism against the deleterious effects of transposon expression and movement. The mechanisms that cause methylated DNA to be transcriptionally silent are not thoroughly understood; however, at least two proteins that repress transcription are known to bind to methylated DNA, and one of these, denoted MeCP2, has been shown to cause changes in chromatin structure. Thus, it is possible that methylated CpG dinucleotides bind specific proteins and that these proteins form a complex that prevents the transcription of neighboring genes.

The methylated state is transmitted clonally through cell division. When a DNA sequence is methylated, both strands of the sequence acquire methyl groups. After the DNA is replicated, each daughter duplex will have one methylated parental DNA sequence and one unmethylated sequence. DNA methyl transferases, the enzymes that attach methyl groups to DNA, can recognize this asymmetry and add a methyl group to the unmethylated sequence. Thus, the fully methylated state is reestablished in the daughter DNA duplexes. In this way, the methylation pattern can be transmitted more or less faithfully through every round of DNA replication—that is, through every cell division. In this sense, DNA methylation is an epigenetic modification of chromatin. Histone acetylation is also considered to be an epigenetic modification, although it is not yet clear how the acetylation pattern is transmitted through cell division. On the Cutting Edge: The Epigenetics of Twins discusses the potential significance of these modifications in humans.

IMPRINTING

DNA methylation in mammals is also responsible for unusual cases in which the expression of a gene is controlled by its parental origin. For example, in mice, the Igf2 gene, which encodes an *i*nsulin-like growth *f*actor, is expressed when it is inherited from the father but not from the mother. By contrast, a gene known as H19 is expressed when it is inherited from the mother but not from the father. Whenever the expression of a gene is conditioned by its parental origin, geneticists say that the gene has been **imprinted**—a term intended to convey the idea that the gene has been marked in some way so that it "remembers" which parent it came from.

Recent molecular analysis has demonstrated that the mark that conditions the expression of a gene is methylation of one or more CpG dinucleotides in the gene's vicinity. These methylated dinucleotides are initially formed in the parental germ line (**Figure 19.13**). Thus, for example, the *Igf*2 gene is methylated in the female germ line but not in the male germ line. At fertilization, a methylated, maternally contributed *Igf*2 gene is combined with an unmethylated, paternally contributed *Igf*2 gene. During embryogenesis, the methylated and unmethylated states are preserved each

ON THE CUTTING EDGE

THE EPIGENETICS OF TWINS

any human twins look and act alike, so much so that we have a hard time telling them apart. But the parents of "identical" twins know that each twin is distinct, and their distinctiveness becomes more apparent with age. One twin may become confident while the other becomes shy. One may become an athlete, the other an artist. Later in life, though they still look alike, one twin may succumb to a chronic illness such as diabetes while the other does not, and in old age, one may develop Alzheimer's disease while the other does not. These differences arouse our curiosity because we know that these types of twins began life with exactly the same genotype. The fertilized egg had split to form two embryos, each of which then developed into a separate person. To emphasize their origin from a single fertilized egg, we say that such twins are **monozygotic**.

In 2005, an international research team explored the possibility that genetically identical twins might be epigentically different.¹ They studied 40 pairs of monozygotic twins from Spain. These twins ranged in age from 3 years to 74 years and varied in the extent to which they shared life's experiences. The researchers examined two types of epigenetic modifications in the chromatin of white blood cells taken from the twins: DNA methylation and histone acetylation.

Most of the twin pairs showed amazingly similar epigenetic profiles. However, in 35 percent of the pairs, there were notable differences in the overall levels of DNA methylation and histone acetylation. These differences were more prevalent among the older twin pairs, and in pairs who spent less of their lives together or who had different health histories. A closer look at the differences in DNA



methylation showed that about half of them were associated with retrotransposons in the genomes of the twins; the other half were associated with known or suspected genes. Cytogenetic mapping demonstrated that the differences were distributed throughout the genome. They localized to the telomeres of the chromosomes and to certain gene-rich regions such as the long and short arms of chromosome 1, the short arm of chromosome 3, and the long arm of chromosome 8. When RNA levels were assayed, the DNA sequences that were hypermethylated were either silent or underexpressed. Thus, the epigenetic differences between the twins seemed to have a functional significance.

This study—the first of its kind—demonstrated that twins with the same genotype can have different "epigenotypes," and it suggested that some of the phenotypic differences between twins might be due to epigenetic differences, which could, in turn, be due to the twins' different life histories. Thus, this study implies that over time, a person's experiences—diet, social and physical activities, medical treatments, exposure to different environments, and so on—might have a role in shaping the "epigenome," which may then influence how the underlying genome is expressed.

In the fall of 2010, another international team was formed to study epigenetic differences in twins. This "Epitwin" project is being led by scientists in the United Kingdom and China, and it will search for epigenetic modifications that influence susceptibility to various conditions and diseases such as obesity, diabetes, osteoporosis, and longevity. Five thousand twins will be analyzed. This largescale study therefore has the potential to reveal how epigenetically regulated gene expression affects the etiology of complex traits.

¹Fraga, M. F., et al., 2005. Epigenetic differences arise during the lifetime of monozygotic twins. *Proc. Natl. Acad. Sci. USA* 102: 10604–10609.

time the genes replicate. Because a methylated gene is silent, only the paternally contributed Igf2 gene is expressed in the developing animal. Exactly the opposite happens with the H19 gene, which is methylated in the male germ line but not in the female germ line. More than 20 different imprinted genes have been identified in mice and humans. For each, the methylation imprint is established in the parental germ line. However, a methylated gene that was inherited from one sex can be unmethylated when it passes through an offspring of the opposite sex. Thus, the methylation imprints are reset each generation, depending on the sex of the animal. The fact that some genes are methylated in one sex but not in the other implies that sex-specific factors control the methylation machinery.

- Heterochromatin is associated with the repression of transcription.
- Position-effect variegation is an example of the epigenetic regulation of gene expression.
- Transcription occurs preferentially in loosely organized chromatin.
- Transcriptionally active DNA tends to be more sensitive to digestion with DNase I.
- During transcriptional activation, chromatin is remodeled by multiprotein complexes.
- Methylation of DNA is associated with gene silencing in mammals.
- The expression of a gene that is imprinted is conditioned by the gene's parental origin.

KEY POINTS

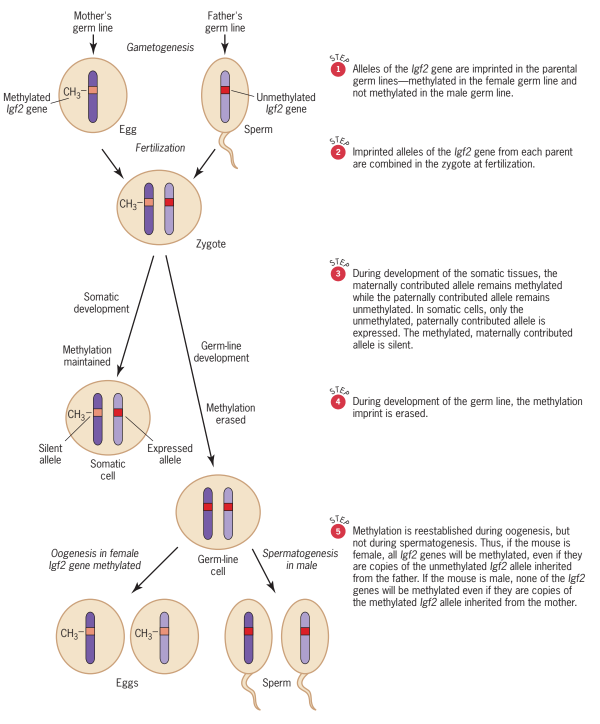


FIGURE 19.13 Methylation and imprinting of the *lgf*2 gene in mice. The gene is methylated in females but not in males.

Activation and Inactivation of Whole Chromosomes

Mammals, flies, and worms have distinct ways of compensating for different dosages of X chromosomes in males and females. Organisms with an XX/XY or XX/XO sex-determination system face the problem of equalizing the activity of X-linked genes in the two sexes. In mammals, this problem is solved by randomly inactivating one of the two X chromosomes in females; each female therefore has the same number of transcriptionally

active X-linked genes as a male. In *Drosophila*, neither of the two X chromosomes in a female is inactivated; instead, the genes on the single X chromosome in a male are

transcribed more vigorously to bring their output in line with that of the genes on the two X chromosomes in a female. Still another solution to the problem of unequal numbers of X-linked genes has been found in the nematode *Caenorhabditis elegans*. In this organism, XX individuals are hermaphrodites (they function as both male and female), and XO individuals are males. X-linked transcriptional activity is equalized in these two genotypes by partial repression of the genes on both of the X chromosomes in the hermaphrodites. Therefore, mammals, flies, and worms have solved the problem of X-linked gene dosage in different ways (**■** Figure 19.14). In mammals, one of the X chromosomes in females is inactivated; in *Drosophila*, the single X chromosome in males is hyperactivated; and in *C. elegans*, both of the X chromosomes in hermaphrodites are hypoactivated.

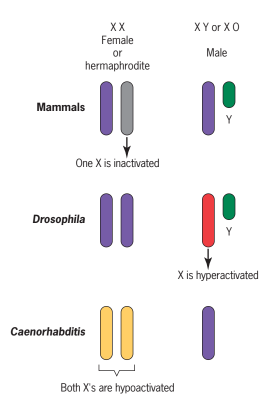
These three different mechanisms of **dosage compensation**—*inactivation*, *hyperactivation*, and *hypoactivation*—have an important feature in common: many different genes are coordinately regulated because they are on the same chromosome. This chromosomewide regulation is superimposed on all other regulatory mechanisms involved in the spatial and temporal expression of these genes. What might be responsible for such a global regulatory system? For decades, geneticists have been trying to elucidate the molecular basis of dosage compensation. The working hypothesis has been that some factor or factors bind specifically to the X chromosome and alter its transcriptional activities. Recent discoveries indicate that this idea is correct.

INACTIVATION OF X CHROMOSOMES IN MAMMALS

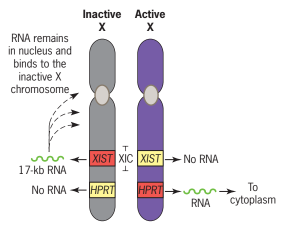
In mammals, X chromosome inactivation begins at a particular site called the X *inactivation center* (XIC) and then spreads in opposite directions toward the ends of the chromosome. Curiously, not all genes on an inactivated X chromosome are transcriptionally silent. One that remains active is called XIST (for X *i*nactive specific *t*ranscript); this gene is located within the XIC (**Figure 19.15**). In human beings the XIST gene encodes a 17-kb transcript devoid of any significant open reading frames. It therefore seems unlikely that the XIST gene codes for a protein. Instead, the RNA itself is probably the functional product of the XIST gene. Though polyadenylated, this RNA is restricted to the nucleus and is specifically localized to inactivated X chromosomes; it does not appear to be associated with active X chromosomes in either males or females.

In mice, where fairly detailed experimental analysis has been possible, researchers have found that the homologue of the human XIST gene is transcribed during the early stages of embryonic development at a low level from both of the X chromosomes that are present in females. The transcripts from each of a female mouse's Xist genes are unstable and remain closely associated with their respective genes. As development proceeds, the transcripts from one of the genes stabilize and eventually envelop the entire X chromosome on which that gene is located; the transcripts from the other Xist gene disintegrate, and further transcription from that gene is repressed by methylation of nucleotides in the gene's promoter. Thus, in the female mouse, one X chromosome—the one whose Xist gene continues to be transcribed-becomes coated with Xist RNA and the other does not. The choice of the chromosome that becomes coated is apparently random. Although the coating mechanism is not yet understood, the consequence of coating is clear: most of the genes on the coated chromosome are repressed, and that chromosome becomes the inactive X chromosome. In the mammalian dosage compensation system, therefore, the X chromosome that remains active is, paradoxically, the one that represses its *Xist* gene.

Inactive X chromosomes are readily identified in mammalian cells. During interphase, they condense into a darkly staining mass associated with the nuclear membrane. This mass, the Barr body, decondenses during S phase to allow the inactive X chromosome to be replicated. However, because decondensation takes some time, the inactive X replicates later than the rest of the chromosomes.



■ **FIGURE 19.14** Three mechanisms of dosage compensation for X-linked genes: inactivation, hyperactivation, and hypoactivation.



■ FIGURE 19.15 Expression of the XIST gene in the inactive X chromosome of human females. For comparison, the expression of the HPRT gene on the active X chromosome is shown. This gene encodes hypoxanthine phosphoribosyl transferase, an enzyme that plays a role in the metabolism of purines.

Inactive X chromosomes must therefore have a very different chromatin structure than that of other chromosomes. This difference is partly determined by the kinds of histones associated with the DNA. One of the four core histones, H4, can be chemically modified by the addition of acetyl groups to any of several lysines in the polypeptide chain. Acetylated H4 is associated with all the chromosomes in the human genome. However, on the inactive X it seems to be restricted to three fairly narrow bands, each corresponding to a region that contains some active genes. Acetylated H4 is also depleted in areas of heterochromatin on the other chromosomes. These findings suggest that the depletion of acetylated H4 is a key feature of the inactive X chromosome.

HYPERACTIVATION OF X CHROMOSOMES IN DROSOPHILA

In *Drosophila*, dosage compensation requires the protein products of at least five different genes. Null mutations in these genes result in male-specific lethality because the single X chromosome in males is not hyperactivated. Mutant males usually die during the late larval or early pupal stages. These dosage compensation genes are therefore called male-specific lethal (*msl*) loci, and their products are called the MSL proteins. Antibodies prepared against these proteins have been used as probes to localize the proteins inside cells. The remarkable finding is that each of the MSL proteins binds specifically to the X chromosome in males' genome, and they do not bind to any of the chromosomes, including the X's, in a female's genome. The binding of the MSL proteins to the male's X chromosome is facilitated by two types of RNA molecules called *roX1* and *roX2* (for *R*NA *on* the X chromosome) that are transcribed from genes on the X chromosome.

The current model proposes that the MSL proteins form a complex that is joined by the *roX* RNAs. This complex then binds to 30 to 40 sites along the male's X chromosome, including the loci that contain the two *roX* genes. From each of these entry sites, the MSL/*roX* complex spreads bidirectionally until it reaches all the genes on the male's X chromosome that need to be hyperactivated. The process of hyperactivation may involve chromatin remodeling by the MSL/*roX* complex. One of the MSL proteins is a histone acetyl transferase, and a particular acetylated version of histone H4 is exclusively associated with hyperactivated X chromosomes.

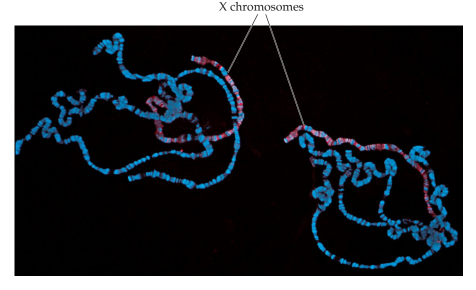


FIGURE 19.16 Binding of the protein product of one of the *Drosophila msl* genes to the single X chromosome in males.

HYPOACTIVATION OF X CHROMOSOMES IN CAENORHABDITIS

In *C. elegans*, dosage compensation involves the partial repression of X-linked genes in the somatic cells of hermaphrodites. The mechanism is not fully understood, but the products of several genes are involved. Like the MSL proteins in *Drosophila*, the proteins encoded by these genes bind specifically to the X chromosome. However, unlike the situation in *Drosophila*, they bind only when two X chromosomes are present. The proteins apparently do not bind to the single X chromosome in males, nor do they bind to any of the autosomes in either males or hermaphrodites. Dosage compensation in *C. elegans* therefore seems to involve a mechanism exactly opposite to the one in *Drosophila*. A protein complex binds to the X chromosomes and represses rather than enhances transcription.

- Inactivation of an X chromosome in XX female mammals is mediated by a noncoding RNA transcribed from the XIST gene on that chromosome.
- Hyperactivation of the single X chromosome in male Drosophila is mediated by an RNAprotein complex that binds to many sites on that chromosome and stimulates the transcription of its genes.
- Hypoactivation of the two X chromosomes in C. elegans hermaphrodites is mediated by
 proteins that bind to these chromosomes and reduce the transcription of their genes.

Basic Exercises

Illustrate Basic Genetic Analysis

 Arrange the following events in chronological order, beginning with the earliest: (a) splicing of an RNA molecule, (b) migration of an mRNA molecule into the cytoplasm, (c) transcription of a gene, (d) degradation of an mRNA molecule, (e) polypeptide synthesis.

Answer: c-a-b-e-d.

2. What factor induces the expression of the *hsp70* gene in *Drosophila*?

Answer: The *hsp70* gene is induced by heat stress.

- **3.** Indicate whether each of the following phenomena related to the regulation of gene expression occurs in the nucleus or the cytoplasm of a eukaryotic cell.
 - (a) Stimulation of gene expression by a transcription factor.
 - (b) Alternate splicing of the primary transcript of a gene.
 - (c) Polyadenylation of a gene's primary transcript.
 - (d) Translation of a messenger RNA.
 - (e) Inhibition of translation by a microRNA binding to a messenger RNA.
 - (f) Degradation of a messenger RNA induced by a short interfering RNA.
 - (g) Binding of a peptide hormone to its receptor.
 - (h) Binding of a steroid hormone to its receptor.
 - (i) Silencing of gene expression by heterochromatin.
 - (j) Whole chromosome inactivation.

Answer: Item (h) may take place in the cytoplasm or the nucleus, depending on the particular steroid hormone. Items (a), (b), (c), (i), and (j) take place in the nucleus. All other items take place in the cytoplasm.

- **4.** What are some differences between euchromatin and heterochromatin?
- **Answer:** Heterochromatin stains darkly throughout the cell cycle; euchromatin does not stain darkly during interphase. Heterochromatin is rich in repeated DNA sequences and in transposable elements; euchromatin may contain repeated sequences and transposons, but usually not to the extent that heterochromatin does. Heterochromatin has few protein-coding genes; euchromatin has many protein-coding genes.
- 5. Indicate whether the following are associated with gene activity or inactivity: (a) DNA methylation, (b) histone acetylation, (c) histone methylation, (d) heterochromatin, (e) locus control region, (f) GAL4 protein, (g) DNase I sensitivity.
- **Answer:** (a) inactivity, (b) activity, (c) inactivity, (d) inactivity, (e) activity, (f) activity, (g) activity.
- 6. How is the level of X-linked gene expression equalized in the two sexes of (a) humans, (b) flies, (c) worms?
- Answer: (a) In humans, one of the two X chromosomes in females is randomly inactivated. (b) In flies, the single X chromosome in males is hyperactivated. (c) In worms, the two X chromosomes in hermaphrodites are hypoactivated.

KEY POINTS

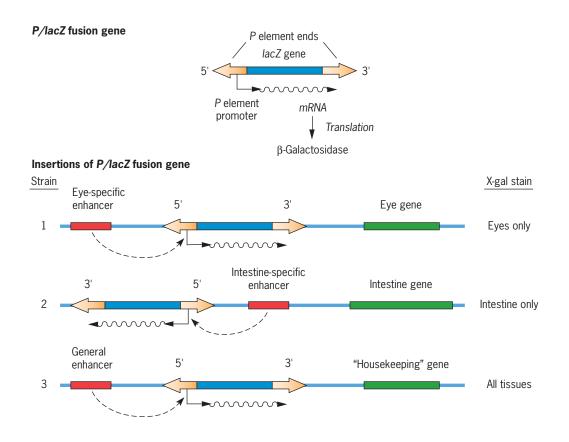
Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. The bacterial *lacZ* gene for β -galactosidase was inserted into a transposable P element from Drosophila (Chapter 17) so that it could be transcribed from the *P* element promoter. This fusion gene was then injected into the germ line of a Drosophila embryo along with an enzyme that catalyzes the transposition of *P* elements. During development, the modified P element became inserted into the chromosomes of some of the germ-line cells. Progeny from this injected animal were then individually mated to flies from a standard laboratory stock to establish strains that carried the *P/lacZ* fusion gene in their genomes. Three of these strains were analyzed for *lacZ* expression by staining dissected tissues from adult flies with X-gal, a chromogenic substrate that turns blue in the presence of β -galactosidase. In the first strain, only the eyes stained blue, in the second, only the intestines stained blue, and in the third, all the tissues stained blue. How do you explain these results?
- **Answer:** The three strains evidently carried different insertions of the P/lacZ fusion gene (see accompanying diagram). In each strain, the expression of the P/lacZ fusion gene must have come under the influence of a different regulatory sequence, or enhancer, capable of interacting with the P promoter and initiating transcription into the lacZ gene. In the first strain, the modified P element must have inserted near an eye-specific enhancer, which would drive

transcription only in eye tissue. In the second strain, it must have inserted near an enhancer that drives transcription in the intestinal cells, and in the third strain, it must have inserted near an enhancer that drives transcription in all, or nearly all, cells, regardless of tissue affiliation. Presumably each of these different enhancers lies near a gene that would normally be expressed under its control. For example, the eye-specific enhancer would be near a gene needed for some aspect of eye function or development. These results show that random insertions of the *P/lacZ* fusion gene can be used to identify different types of enhancers and, through them, the genes they control. These fusion gene insertions are therefore often called *enhancer traps*.

2. In their seminal paper on RNA interference, Andrew Fire and coworkers (1998 Nature 391: 806–811) describe the results of experiments in which RNA derived from the mex-3 gene was injected into C. elegans hermaphrodites. Embryos obtained from these injected hermaphrodites were analyzed by in situ hybridization using probes for mex-3 RNA. The probes were designed to bind to mex-3 messenger RNA, which normally accumulates in the gonads of hermaphrodites and in their embryos. Binding of the probe molecules to mRNA in the embryos is easily detected if the probe molecules have been labeled. When Fire and his colleagues performed these in situ



hybridization experiments, they found that embryos from worms that had been injected with double-stranded *mex-3* RNA were not labeled by the probe molecules, whereas embryos from worms that had been injected with singlestrandedRNAcomplementaryto*mex-3*mRNA—thatis,with antisense *mex-3* RNA—were labeled, though not quite as intensively as embryos from worms that had not been injected at all. What do these results indicate about the efficacy of double-stranded versus single-stranded antisense RNA to silence gene expression?

- **Answer:** The results of these *in situ* hybridization experiments indicate that double-stranded RNA is a strong silencer of *mex-3* gene expression in *C. elegans* embryos. By contrast, single-stranded antisense RNA barely has an effect on *mex-3* gene expression. The embryos from worms injected with double-stranded *mex-3* RNA did not carry any detectable *mex-3* messenger RNA. The absence of *mex-3* messenger RNA in these embryos is the result of RNA interference induced by the injected with single-stranded antisense *mex-3* RNA did carry some *mex-3* messenger RNA. Thus, single-stranded antisense *mex-3* RNA is not as effective as double-stranded *mex-3* RNA in the induction of RNA.
- **3.** The patchy phenotype of tortoiseshell cats (Chapter 5) results from random inactivation of X chromosomes in females that are heterozygous for different alleles of an X-linked gene for fur color; one allele leads to light-colored fur, the other to dark-colored fur. The patchy phenotype of gynandromorphs in *Drosophila* (Chapter 6)

results from nondisjunction of the X chromosomes during one of the early cleavage divisions. If an XX zygote is heterozygous for wild-type and mutant alleles of the X-linked *white* gene, nondisjunction can produce a lineage of XO cells that carry only the mutant allele, and if these cells form an eye, or part of an eye, that eye tissue will be white. By contrast, tissue derived from XX cells will be red because those cells carry the wild-type allele of the *white* gene. Is either of these patchy phenotypes an example of epigenetic regulation of gene expression? Explain your answer.

Answer: The patchy phenotype of tortoiseshell cats results from an epigenetic phenomenon-random inactivation of an X chromosome in each of the cells destined to form pigmentproducing cells in the adult. All the pigment-producing cells are genetically equivalent-that is, they have the same DNA content. The tortoiseshell phenotype is not due to a change in the underlying genotype during the animal's embryological development. Rather, it is due to a change in the state of one of the X chromosomes, the X that is inactivated, and this state is inherited clonally through cell division. Thus, the light and dark patches of fur in the cat differ epigenetically, not genetically. In contrast, the patchy phenotype of Drosophila gynandromorphs is due to a genetic change that occurs during development. One of the X chromosomes is lost. The red and white patches of tissue in a gynandromorph's eye are not genetically equivalent. The difference between them is therefore genetic rather than epigenetic.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **19.1** Operons are common in bacteria but not in eukaryotes. Suggest a reason why.
- **19.2** In bacteria, translation of an mRNA begins before the synthesis of that mRNA is completed. Why is this "coupling" of transcription and translation not possible in eukaryotes?
- **19.3** Muscular dystrophy in humans is caused by mutations in an X-linked gene that encodes a protein called dystrophin. What techniques could you use to determine if this gene is active in different types of cells, say skin cells, nerve cells, and muscle cells?
- **19.4** Why do steroid hormones interact with receptors inside the cell, whereas peptide hormones interact with receptors on the cell surface?
- **19.5** In the polytene chromosomes of *Drosophila* larvae (Chapter 6), some bands form large "puffs" when the larvae are subjected to high temperatures. How could you

show that these puffs contain genes that are vigorously transcribed in response to this heat-shock treatment?

- **19.6** How would you distinguish between an enhancer and a promoter?
- **19.7** Tropomyosins are proteins that mediate the interaction of actin and troponin, two proteins involved in muscle contractions. In higher animals, tropomyosins exist as a family of closely related proteins that share some amino acid sequences but differ in others. Explain how these proteins could be created from the transcript of a single gene.
- **19.8** A polypeptide consists of three separate segments of amino acids, A—B—C. Another polypeptide contains segments A and C, but not segment B. How might you determine if these two polypeptides are produced by translating alternately spliced versions of RNA from a single gene or by translating mRNA from two different genes?

- **19.9** What techniques could be used to show that a plant gene is transcribed when the plant is illuminated with light?
- **19.10** When introns were first discovered, they were thought to be genetic "junk"—that is, sequences without any useful function. In fact, they appeared to be worse than junk because they actually interrupted the coding sequences of genes. However, among eukaryotes, introns are pervasive and anything that is pervasive in biology usually has a function. What function might introns have? What benefit might they confer on an organism?
- **19.11** The GAL4 transcription factor in yeast regulates two adjacent genes, *GAL1* and *GAL10*, by binding to DNA sequences between them. These two genes are transcribed in opposite directions on the chromosome, one to the left of the GAL4 protein's binding site and the other to the right of this site. What property of enhancers does this situation illustrate?
- 19.12 **co** Using the techniques of genetic engineering, a researcher has constructed a fusion gene containing the heat-shock response elements from a *Drosophila hsp70* gene and the coding region of a jellyfish gene (*gfp*) for green fluorescent protein. This fusion gene has been inserted into the chromosomes of living *Drosophila* by the technique of transposon-mediated transformation (Chapter 17). Under what conditions will the green fluorescent protein be synthesized in these genetically transformed flies? Explain.
- **19.13** Suppose that the segment of the *hsp70* gene that was used to make the *hsp70/gfp* fusion in the preceding problem had mutations in each of its heat-shock response elements. Would the green fluorescent protein encoded by this fusion gene be synthesized in genetically transformed flies?
- **19.14** The polypeptide products of two different genes, *A* and *B*, each function as transcription factors. These polypeptides interact to form dimers: AA homodimers, BB homodimers, and AB heterodimers. If the A and B polypeptides are equally abundant in cells, and if dimer formation is random, what is the expected ratio of homodimers to heterodimers in these cells?
- **19.15** A particular transcription factor binds to enhancers in 40 different genes. Predict the phenotype of individuals homozygous for a frameshift mutation in the coding sequence of the gene that specifies this transcription factor.
- **19.16** The alternately spliced forms of the RNA from the *Drosophila doublesex* gene encode proteins that are needed to block the development of one or the other set of sexual characteristics. The protein that is made in female animals blocks the development of male characteristics, and the protein that is made in male animals blocks the development of female characteristics. Predict the phenotype of XX and XY animals homozygous for a null mutation in the *doublesex* gene.
- **19.17** The RNA from the *Drosophila Sex-lethal (Sxl)* gene is alternately spliced. In males, the sequence of the mRNA

derived from the primary transcript contains all eight exons of the *Sxl* gene. In females, the mRNA contains only seven of the exons because during splicing exon 3 is removed from the primary transcript along with its flanking introns. The coding region in the female's mRNA is therefore shorter than it is in the male's mRNA. However, the protein encoded by the female's mRNA is longer than the one encoded by the male's mRNA. How might you explain this paradox?

- **19.18** In *Drosophila*, expression of the *yellow* gene is needed for the formation of dark pigment in many different tissues; without this expression, a tissue appears yellow in color. In the wings, the expression of the *yellow* gene is controlled by an enhancer located upstream of the gene's transcription initiation site. In the tarsal claws, expression is controlled by an enhancer located within the gene's only intron. Suppose that by genetic engineering, the wing enhancer is placed within the intron and the claw enhancer is placed upstream of the transcription initiation site. Would a fly that carried this modified *yellow* gene in place of its natural *yellow* gene have darkly pigmented wings and claws? Explain.
- **19.19** A researcher suspects that a 550-bp-long intron contains an enhancer that drives expression of an *Arabidopsis* gene specifically in root-tip tissue. Outline an experiment to test this hypothesis.
- **19.20** What is the nature of each of the following classes of enzymes? What does each type of enzyme do to chromatin? (a) HATs, (b) HDACs, (c) HMTs.
- **19.21** In *Drosophila* larvae, the single X chromosome in males appears diffuse and bloated in the polytene cells of the salivary gland. Is this observation compatible with the idea that X-linked genes are hyperactivated in *Drosophila* males?
- **19.22** Suppose that the LCR of the β -globin gene cluster was deleted from one of the two chromosomes 11 in a man. What disease might this deletion cause?
- **19.23** Would double-stranded RNA derived from an intron be able to induce RNA interference?
- **19.24** RNA interference has been implicated in the regulation of transposable elements. In *Drosophila*, two of the key proteins involved in RNA interference are encoded by the genes *aubergine* and *piwi*. Flies that are homozygous for mutant alleles of these genes are lethal or sterile, but flies that are heterozygous for them are viable and fertile. Suppose that you have strains of *Drosophila* that are heterozygous for *aubergine* or *piwi* mutant alleles. Why might the genomic mutation rate in these mutant strains be greater than the genomic mutation rate in a wild-type strain?
- **19.25** Suppose female mice homozygous for the *a* allele of the *Igf2* gene are crossed to male mice homozygous for the *b* allele of this gene. Which of these two alleles will be expressed in the F_1 progeny?

- **19.26** Epigenetic states are transmitted clonally through cell division. What kinds of observations indicate that these states can be reversed or reset?
- **19.27** A researcher hypothesizes that in mice gene A is actively transcribed in liver cells, whereas gene B is actively transcribed in brain cells. Describe procedures that would allow the researcher to test this hypothesis.
- **19.28 (co)** Suppose that the hypothesis mentioned in the previous question is correct and that gene A is actively transcribed in liver cells whereas gene B is actively transcribed in brain cells. The researcher now extracts equivalent amounts of chromatin from liver and brain tissues and treats these extracts separately with DNase I for a limited period of time. If the DNA that remains after the treatment is then fractionated by gel electrophoresis, transferred to a membrane by Southern blotting, and hybridized with a radioactively labeled probe specific for gene A, which sample (liver or brain) will be expected to show the greater signal on the autoradiogram? Explain your answer.
- **19.29** Why do null mutations in the *msl* gene in *Drosophila* have no effect in females?
- **19.30** Suppose that a woman carries an X chromosome in which the *XIST* locus has been deleted. The woman's other X chromosome has an intact *XIST* locus. What pattern

of X-inactivation would be observed throughout the woman's body?

- 19.31 In *Drosophila*, the variegated phenotype of the *white mottled* allele is suppressed by a dominant autosomal mutation that knocks out the function of the gene for heterochromatin protein 1 (HP1), an important factor in heterochromatin formation. Flies with the *white mottled* allele and the suppressor mutation have an almost uniform red color in their eyes; without the suppressor mutation, the eyes are mosaics of red and white tissue. Can you suggest an explanation for the effect of the suppressor mutation?
- **19.32** The sheep Dolly (Chapter 2) was the first cloned mammal. Dolly was created by implanting a nucleus from a cell taken from the udder of a female sheep into an enucleated egg. This nucleus had two X chromosomes, and because it came from a differentiated cell, one of them must have been inactivated. If the udder cell was heterozygous for at least one X-linked gene whose expression you could assay, how could you determine if all of Dolly's cells had the same X chromosome inactivated? If, upon testing, Dolly's cells prove to be mosaic for X chromosome activity—that is, different X's are active in different clones of cells—what must have happened during her embryological development?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

The human β -globin genes are located in a cluster on the short arm of chromosome 11.

- 1. Search for the namesake gene of the cluster, the adult β -globin gene, in the human genome database. What is the official symbol of this gene? How many exons does it contain?
- 2. Use the Map Viewer function to locate the β -globin gene cluster on the ideogram of chromosome 11. In what cytological band does it reside? Is it closer to the telomere of the short arm or to the centromere?
- **3.** Use the Sequence Viewer to inspect the adult β-globin gene in detail. Is the gene transcribed toward the centromere or toward the telomere? How long is the transcript of the gene? How long is the mature mRNA? How many amino acids does the mRNA specify? What are the first three amino acids, and what codons specify them?
- **4.** Bring up the text sequence of the adult β -globin gene by clicking the ATGC button on the Sequence Viewer page. Locate the initiation codon for methionine in the first exon. Because the sequence in the window is that of the template strand of the DNA, this codon reads 5'-CAT-3' from left to right on the screen.
- **5.** GATA1 and MyoD are two transcription factors that recognize short sequences in mammalian genomes. The sequence recognized by GATA1 is 5'-TGATAG-3', and the sequence recognized by MyoD is 5'-CAAATG-3'. Copy the sequence of the transcribed portion of the adult β -globin gene into a text file and scan it for each of these recognition sequences. Where are they located? Which of these two transcription factors might be involved in regulating the expression of the adult β -globin gene?

The Genetic Control of Animal Development

CHAPTER OUTLINE

- A Genetic Perspective on Development
- Maternal Gene Activity in Development
- Zygotic Gene Activity in Development
- Genetic Analysis of Development in Vertebrates

Stem-Cell Therapy

Stem cells are in the news. Scientists are discussing their possible uses, and all sorts of peoplepoliticians, religious leaders, journalists, the victims of illnesses such as Parkinson's disease, diabetes, and arthritis, and even Hollywood celebrities-are joining the conversation. Though nondescript themselves, stem cells have the ability to produce offspring that can differentiate into special cell types, like muscle fiber, lymphocyte, neuron, or bone cell. They might, therefore, be used to regenerate wornout tissues, to replace lost organs or body parts, to correct injuries, or to alleviate biochemical deficits. These prospects point out the importance of understanding how different types of cells acquire their specialized functions, and how, in a multicelled organism, they form tissues and organs in an orderly manner over time. In other words, they point out the importance of understanding the process of development-from fertilized egg to embryo to adult. The possibility for stem-cell therapy also

Human fetus late during development.

raises important ethical questions. Must the stem cells be derived by destroying embryos? Should embryonic life be sacrificed to prolong and enhance adult life? Is it acceptable to produce embryos merely to harvest stem cells for therapeutic purposes? Around the world, people and their governments are debating these questions, while scientists continue to explore the properties of stem cells and how they might be used.

A Genetic Perspective on Development

The development of a multicelled animal from a fertilized egg demonstrates the power of controlled gene expression. Genes must be expressed carefully over time to bring about the specialization of cells, the orderly assembly of these cells into tissues and organs, and the formation of the animal's body. The process of animal develop*Drosophila* has been one of the premier model organisms for the genetic analysis of animal development.

ment therefore depends on the faithful execution of a genetic program encoded in the animal's DNA. It should come as no surprise, therefore, that genetics has contributed greatly to our understanding of this process.

Classical studies from anatomy and embryology provided detailed observations about the events of development-the division of the fertilized egg to form an embryo, the movement of cells within the embryo to form primitive tissues, and the subsequent differentiation of cells within these tissues to form different organs. For practical reasons, these classical studies focused on a few kinds of animals, especially sea urchins, frogs, and chickens. The eggs of such animals can be manipulated experimentally, and their embryos develop outside the mother's body. Embryologists could therefore see how an embryo developed in response to an experimental treatment. When geneticists began to study development, they focused on animals that were easy to breed, especially Drosophila and C. elegans. Their objective was to identify genes whose products are involved in important developmental events. The standard way for a geneticist to achieve such an objective is to collect mutations. Thus, for example, if a geneticist wishes to study the development of Drosophila wings, he or she would collect mutations that alter or prevent wing formation. These mutations would then be tested for allelism with one another and mapped on the chromosomes to define and position the relevant genetic loci. Once these loci have been identified, the geneticist would combine representative mutations from each locus in pairwise fashion to determine whether some of the mutations are epistatic over others. Such epistasis testing can provide valuable insights into how different genes contribute to a developmental process (see Chapter 4). Finally, to investigate the molecular basis of gene action and to elucidate the role that each gene's product plays in development, the geneticist would clone individual genes and study them with the full panoply of techniques now available-sequencing, RNA and protein blotting, RT-PCR, fluorescent labeling, the production of transgenics and so on (see Chapters 14 and 16).

Using this general strategy, geneticists have learned a great deal about the way that development proceeds in *Drosophila* and *C. elegans*. Much is now known about how cells become specialized, how tissues and organs form, and how the body plan is laid out. This knowledge has also provided an intellectual framework to guide the study of development in other animals, including vertebrates such as the mouse. The study of the mouse has, in turn, provided many insights into the process of development in humans. However, before exploring any of these topics, we need to discuss some of the basic features of development in one of the premier models for studying the genetic control of development, *Drosophila*.

Adult *Drosophila* develop from ellipsoidal eggs about 1 mm long and 0.5 mm wide at their maximum diameter (**Figure 20.1a**). Each egg is surrounded by a *chorion*, a tough shell-like structure that is made of materials synthesized by somatic cells in the ovary. The anterior end is distinguished by two filaments that help to bring oxygen into the egg. Sperm enter the egg through another anterior structure, the *micropyle*. The cell divisions that follow fertilization are rapid—so rapid that there is no time for membranes to form between daughter cells. Consequently, the early *Drosophila* embryo is actually a single cell with many identical nuclei; such a cell is called a *syncytium* (**Figure 20.1b**). After division cycle 9 within the syncytium, the 512 nuclei that have been created migrate to the cytoplasmic membrane on the periphery of the embryo, where they continue to divide four more times. In addition, a few of the nuclei migrate to the posterior pole of the embryo. At division cycle 13, all the nuclei in the syncytium become separated from each other by cell membranes, creating a single layer of cells on the embryo's surface. This single layer, called the *cellular*

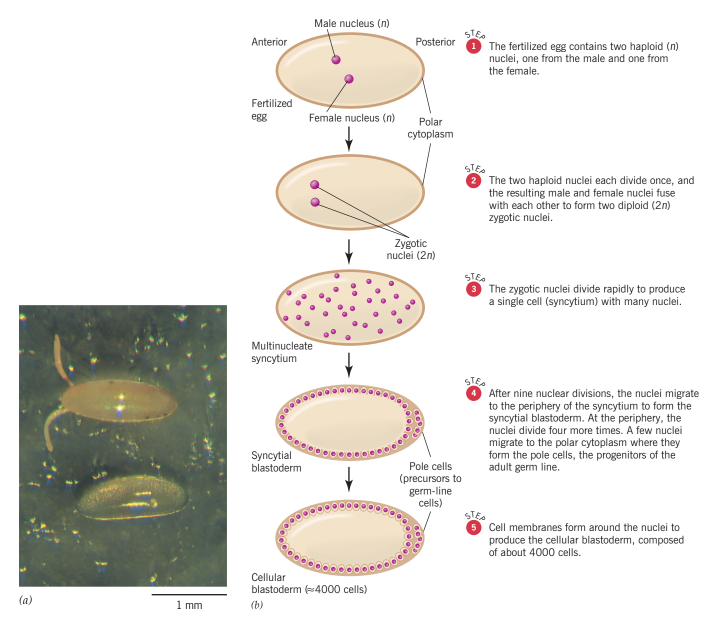


FIGURE 20.1 Basic features of *Drosophila* development. (a) Photograph of *Drosophila* eggs, with (top) and without (bottom) the surrounding chorion. (b) Early embryonic development in *Drosophila*.

blastoderm, will give rise to all the somatic tissues of the animal. Cellularization of the nuclei at the posterior pole creates the *pole cells*, which give rise to the adult germ line. Thus, at this very early stage of development, the somatic and germ-cell lineages of the future adult have already been separated.

It takes about a day for the *Drosophila* embryo to develop into a wormlike *larva*. This larva hatches by chewing its way through the egg shell and then begins feeding voraciously. It sheds its skin twice to accommodate increases in size and then, about five days after hatching, becomes immobile and hardens its skin, forming a *pupa*. During the next four days, many of the larval tissues are destroyed, and flat packets of cells that were sequestered during the larval stages expand and differentiate into adult structures such as antennae, eyes, wings, and legs. Because an adult insect is called an *imago*, these packets are referred to as *imaginal discs*. When this anatomical reorganization is completed, a radically different animal emerges from the pupal casing—one that can fly and reproduce!

- In Drosophila, the developmental sequence is egg, embryo, larva, pupa, and adult.
- The early Drosophila embryo is a syncytium—many nuclei in one cell.
- The structures in adult Drosophila develop from packets of cells called imaginal discs.

Maternal Gene Activity in Development

Important events occur in animal development even before an egg is fertilized. At this time, nutritive and determinative materials are transported into the egg from surrounding cells, laying up food stores and organizing

the egg for its subsequent development—the molecular equivalent of a mother's love. These materials are generated by the expression of genes in the female reproductive system, some being expressed in somatic reproductive tissues and others only in germline tissues. Collectively, these genes help to form eggs that can develop into embryos after fertilization. In some species, these maternal gene products lay out the basic body plan of the embryo, distinguishing head from tail and back from belly. These maternally supplied materials therefore establish a molecular coordinate system to guide an embryo's development. To illustrate how maternal gene activity influences development, we focus on events in *Drosophila*.

MATERNAL-EFFECT GENES

Mutations in genes that contribute to the formation of healthy eggs may have no effect on the viability or appearance of the female making those eggs. Instead, their effects may be seen only in the next generation. Such mutations are called **maternal-effect mutations** because the mutant phenotype in the offspring is caused by a mutant genotype in its mother.

Genes identified by such mutations are called **maternal-effect genes**. The *dorsal* (*dl*) gene in *Drosophila* is a good example (**Trigure 20.2**). Matings between flies homozygous for recessive mutations in this gene produce inviable progeny. This lethal effect is strictly maternal. A cross between homozygous mutant females and homozygous wild-type males produces inviable progeny, but the reciprocal cross (homozygous mutant males \times homozygous wild-type females) produces viable progeny. The lethal effect of the *dorsal* mutation is therefore manifested only if females are homozygous for it. The male genotype is irrelevant.

Molecular characterization of the *dorsal* gene has revealed the basis for this maternal effect. The *dorsal* gene encodes a transcription factor that is produced during oogenesis and stored in the egg. Early in development, this transcription factor plays an important role in the differentiation of the dorsal and ventral parts of the embryo. When it is missing, the ventral parts incorrectly differentiate as if they were on the dorsal side, creating an embryo with two dorsal surfaces. This lethal condition cannot be prevented by a wild-type *dorsal* allele inherited from the father because *dorsal* is not transcribed in the embryo. Expression of the *dorsal* gene is, in fact, limited to the female germ line. Mutations in *dorsal* are therefore strict maternal-effect lethals. To explore a case in which the maternal effect of a mutation can be mitigated by other factors, work through Solve It: A Maternal-Effect Mutation in the *cinnamon* Gene.

Mutant embryo due to maternal effect



Wild-type embryo



FIGURE 20.2 The maternal effect of a mutation in the *dorsal* (*dl*) gene of *Drosophila*. The mutant phenotype is an embryo that lacks ventral tissues; that is, it is dorsalized.



Materials transported into the egg during oogenesis

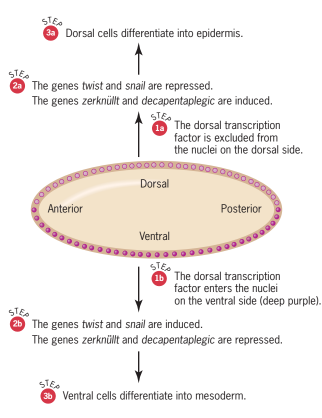
play a major role in embryonic development.

Solve It!

A Maternal-Effect Mutation in the *cinnamon* Gene

The cinnamon (cin) gene is located at the left end of the X chromosome in Drosophila. Animals that are homozygous or hemizygous for a mutation in this gene are abnormal only if their mother was homozygous for the mutation. In the best case, the abnormality in these mutant animals from mutant mothers is a reddish-brown eye color-that is, they have cinnamon-colored eyes; most often, however, they simply die during embryogenesis. A homozygous cin/cin female was crossed to a wild-type cin^+ male. Almost all the offspring were females with normal-colored eyes. The few males that appeared had cinnamon-colored eyes. Propose an explanation for these results?

To see the solution to this problem, visit the Student Companion site.



■ **FIGURE 20.3** Determination of the dorsal-ventral axis in *Drosophila* by the dorsal protein. This protein is a transcription factor that acts only in the nuclei on the ventral side of the embryo. The genes *twist, snail, zerknüllt,* and *decapentaplegic* are regulated by dorsal protein.

DETERMINATION OF THE DORSAL-VENTRAL AND ANTERIOR-POSTERIOR AXES

Animals with bilateral symmetry have two primary body axes, one distinguishing back from belly (dorsal from ventral) and the other distinguishing head from tail (anterior from posterior). Both of these axes are established very early in development, in some species even before fertilization. In *Drosophila*, the processes of axis formation have been dissected genetically by collecting mutations that affect early embryonic development.

In the 1970s and 1980s, massive searches for such mutations were carried out by Christiane Nüsslein-Volhard, Eric Weischaus, Trudi Schüpbach, Gerd Jurgens, and others. These researchers used chemical mutagens to induce mutations in each of the *Drosophila* chromosomes. Many mutations were identified, including maternal-effect lethals in genes such as *dorsal*. Molecular and genetic analyses of these mutations have provided a great deal of insight into the events of early *Drosophila* development.

Formation of the Dorsal-Ventral Axis

Differentiation of a *Drosophila* embryo along the dorsal-ventral axis hinges on the action of the transcription factor encoded by the *dorsal* gene (**Figure 20.3**). This protein is synthesized maternally and stored in the cytoplasm of the egg. At the time of blastoderm formation, the dorsal protein enters the nuclei on the ventral side of the embryo, inducing the transcription of two genes called *twist* and *snail* (whimsically named for their mutant phenotypes). In these same nuclei, it represses the genes *zerknüllt* (from the German for "crumpled") and *decapentaplegic* (from the Greek words for "15" and "stroke"). The selective induction and repression of these genes cause the ventral cells to differentiate into a primitive embryonic layer of tissue called

the mesoderm. On the opposite side of the embryo, where the dorsal protein is excluded from the nuclei, *twist* and *snail* are not induced and *zerknüllt* and *decapentaplegic* are not repressed. Consequently, these cells differentiate into a different primitive tissue, the embryonic epidermis. The entrance of the dorsal transcription factor into the ventral nuclei and its exclusion from the dorsal nuclei therefore initiate differentiation along the dorsal-ventral axis.

But what triggers the dorsal protein to move into the nuclei on only one side of the embryo? The answer is an interaction between two proteins on the ventral surface of the developing embryo (**Figure 20.4**). One protein, the product of the *Toll* gene (from the German word for "tuft"), is distributed uniformly over the embryo's surface; this protein is embedded in the plasma membrane that surrounds the embryo. The other protein, the product of the *spätzle* gene (from the German word for "little dumpling"), is found in the perivitelline space, a fluid-filled cavity between the plasma membrane and the external vitelline membrane. Through the action of a protease encoded by a gene called *easter* (because it was discovered on Easter Sunday), the spätzle protein is cleaved to produce a polypeptide that interacts with the Toll protein. However, because of a pattern established by the cells that had surrounded the egg inside the ovary, cleavage of the spätzle protein occurs only in the perivitelline space on the ventral side of the embryo. When the Toll protein interacts with the ventrally generated spätzle polypeptide, it initiates a cascade of events within the embryo that ultimately sends the dorsal protein into the embryonic nuclei. There the dorsal protein functions as a transcription factor to regulate the expression of the genes twist, snail, decapentaplegic, and zerknüllt. Thus, the membranebound Toll protein acts as a receptor for the determinative spätzle polypeptide, and the physical interaction between these two molecules acts as a signal to trigger a genetic program for the differentiation of the embryo along its dorsal-ventral axis.

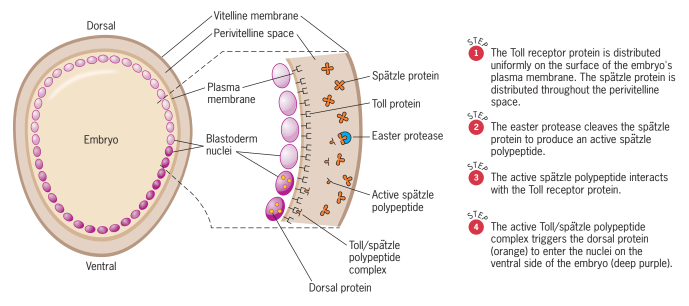


FIGURE 20.4 Differentiation of the dorsal-ventral axis in a *Drosophila* embryo. The cross section shows the interaction between the membrane-bound Toll receptor protein and a polypeptide from the spätzle protein that induces differentiation along the dorsal-ventral axis. Formation of the interacting spätzle polypeptide occurs in the space between the plasma membrane and the vitelline membrane on the ventral side of the embryo.

Formation of the Anterior-Posterior Axis

The anterior-posterior axis in *Drosophila* is created by the regional synthesis of transcription factors encoded by the *hunchback* and *caudal* genes (**Figure 20.5**). These two genes are transcribed in the nurse cells of the maternal germ line. These special cells support the growth and development of the oocyte. The maternal transcripts of the *hunchback* and *caudal* genes are then carried from the nurse cells into the oocyte where they become uniformly distributed in the cytoplasm. However, both types of transcripts are translated in different parts of the embryo. The hunchback RNA is translated only in the anterior part, and the caudal RNA is translated only in the posterior part. This differential translation produces concentration gradients of the proteins encoded by these two genes; hunchback protein is concentrated in the anterior part of the embryo, and caudal protein is concentrated in the posterior part. These two proteins then function to activate or repress transcription of the genes whose products are involved in the differentiation of the embryo along its anterior-posterior axis.

What limits the translation of hunchback RNA to the anterior part of the embryo and caudal RNA to the posterior part? It turns out that two maternally supplied RNAs are involved, one transcribed from the *bicoid* gene and the other from the *nanos* gene. Both of these RNAs are synthesized in the nurse cells of the maternal germ line and are then transported into the oocyte. The bicoid RNA becomes anchored at the anterior end of the developing oocyte, and the nanos RNA becomes anchored at the posterior end. After fertilization, each type of RNA is translated locally, and the resulting protein products diffuse through the embryo to form concentration gradients; bicoid protein is concentrated at the anterior end, and nanos protein is concentrated at the posterior end.

The bicoid protein has two functions. First, it acts as a transcription factor to stimulate the synthesis of RNAs from several genes, including *hunchback*. These RNAs are then translated into proteins that control the formation of the anterior structures of the embryo. Second, bicoid protein prevents the translation of caudal RNA by binding to sequences in the 3' untranslated region of that RNA. Thus, wherever bicoid protein is abundant (that is, in the anterior of the embryo), caudal RNA is not translated into protein. Conversely, wherever bicoid protein is scarce (that is, in the posterior of the embryo), caudal RNA is translated into protein. The translational regulation of caudal RNA by bicoid protein is therefore responsible for the gradient of caudal protein that forms in the embryo. Because caudal protein is a specific activator of

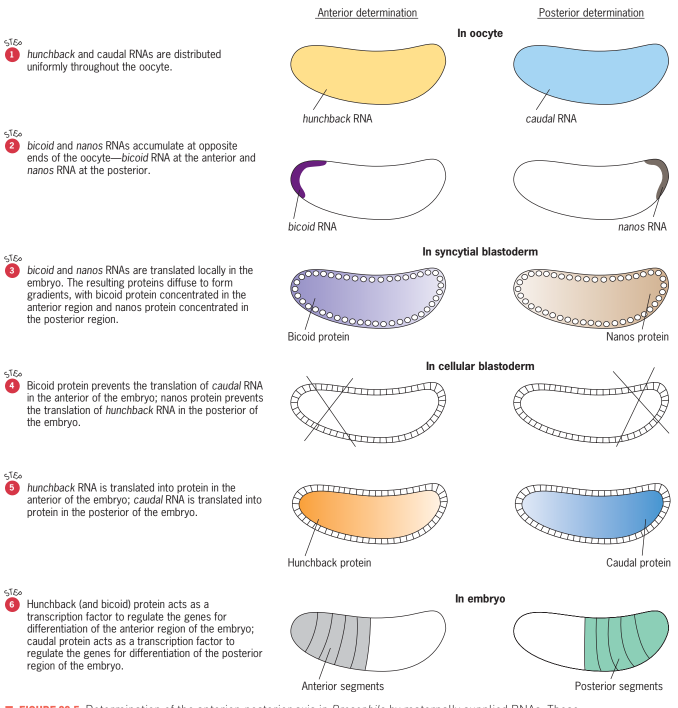


FIGURE 20.5 Determination of the anterior-posterior axis in *Drosophila* by maternally supplied RNAs. These RNAs come from the *hunchback, caudal, bicoid,* and *nanos* genes. For each oocyte or embryo, anterior is at the left and posterior is at the right.

genes that control posterior differentation, the part of the embryo that has the highest concentration of caudal protein develops posterior structures.

Unlike bicoid protein, nanos protein does not function as a transcription factor. However, like bicoid protein, it does function as a translational regulator. Nanos protein is concentrated in the posterior of the embryo, and there it binds to the 3' untranslated region of hunchback RNA and causes the degradation of that RNA. Consequently, hunchback protein is not synthesized in the posterior of the embryo. Instead, its synthesis is restricted to the anterior of the embryo where it acts as a transcription factor to regulate the expression of genes involved in anterior-posterior differentiation. Wherever hunchback protein is synthesized, the embryo develops anterior structures. The bicoid and nanos proteins are examples of **morphogens**—substances that control developmental events in a concentration-dependent manner. The concentration gradients of these two morphogens are the reverse of each other; where bicoid protein is abundant, nanos protein is scarce, and vice versa. Thus, in *Drosophila* the anteriorposterior axis is defined by high concentrations of these morphogens at opposite ends of the early embryo.

- The proteins and RNAs encoded by maternal-effect genes such as dorsal, hunchback, bicoid, and nanos are transported into Drosophila eggs during oogenesis.
- Maternal-effect gene products are involved in the determination of the dorsal-ventral and anterior-posterior axes in Drosophila embryos.
- Recessive mutations in maternal-effect genes are expressed only in embryos produced by females homozygous for these mutations.

Zygotic Gene Activity in Development

The earliest events in animal development are controlled by maternally synthesized factors. However, at some point, the genes in the embryo are selectively activated, and new materials are made. This process is referred to as *zygotic gene expression* because it occurs after the egg has been fertilized. The initial

wave of zygotic gene expression is a response to maternally synthesized factors. In *Drosophila*, for example, the maternally supplied dorsal transcription factor activates the zygotic genes *twist* and *snail*. As development proceeds, the activation of other zygotic genes leads to complex cascades of gene expression. We will now examine how these zygotic genes carry the process of development forward. Again, we focus on events in *Drosophila*.

BODY SEGMENTATION

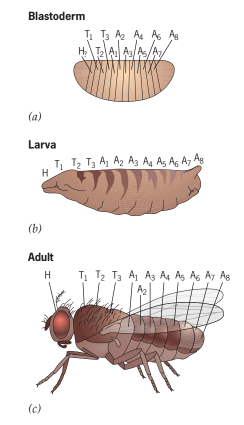
In many invertebrates the body consists of an array of adjoining units called *segments*. An adult *Drosophila*, for example, has a head, three distinct thoracic segments, and eight abdominal segments. Within the thorax and abdomen, each segment can be identified by coloration, bristle pattern, and the kinds of appendages attached to it. These segments can also be identified in the embryo and the larva (**Figure 20.6**). In vertebrates, a segmental pattern is not so evident in the adult, but it can be recognized in the embryo from the way that nerve fibers grow from the central nervous system, from the formation of branchial arches in the head, and from the organization of muscle masses along the anterior-posterior axis. Later in development, these features are modified, and the original segmental pattern becomes obscured. Nonetheless, in both vertebrates and many invertebrates, segmentation is a key aspect of the overall body plan.

Homeotic Genes

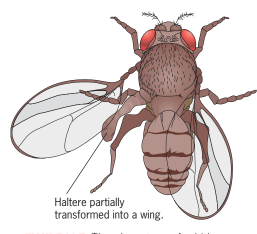
Interest in the genetic control of segmentation began with the discovery of mutations that transform one segment into another. The first such mutation was found in *Drosophila* in 1915 by Calvin Bridges. He named it *bithorax* (*bx*) because it affected two thoracic segments. In this mutant, the third thoracic segment was transformed, albeit weakly, into the second, creating a fly with a small pair of rudimentary wings in place of the small balancing structures called halteres (**■** Figure 20.7). Later, other segment-transforming mutations were found in *Drosophila*—for example, *Antennapedia*

The differentiation of cell types and the formation of organs depend on genes being activated in particular spatial and temporal patterns.

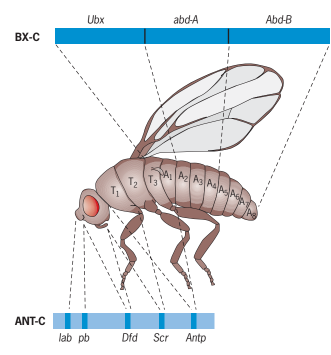
KEY POINTS



■ FIGURE 20.6 Segmentation in *Drosophila* at the (*a*) blastoderm, (*b*) larval, and (*c*) adult stages of development. Although segments are not visible in the blastoderm, its cells are already committed to form segments as shown; H, head segment; T, thoracic segment; A, abdominal segment.



■ **FIGURE 20.7** The phenotype of a *bithorax* mutation in *Drosophila*.



■ **FIGURE 20.8** The homeotic genes in the bithorax complex (BX-C) and Antennapedia complex (ANT-C) of *Drosophila*. The body regions in which each gene is expressed are indicated.

(*Antp*), a mutant that partially transforms the antennae on the head into legs, which normally grow from the thorax. These mutations have come to be called **homeotic mutations** because they cause one body part to look like another. The word "homeotic" comes from William Bateson, who coined the term *homeosis* to refer to cases in which "something has been changed into the likeness of something else." Like so many other words Bateson coined, this one has become a standard term in the modern genetics vocabulary.

The bithorax and Antennapedia phenotypes result from mutations in **homeotic** genes. Several such genes have now been identified in *Drosophila*, where they form two large clusters on one of the autosomes (**Figure 20.8**). The *bithorax complex*, usually denoted *BX-C*, consists of three genes, *Ultrabithorax* (*Ubx*), *abdominal-A* (*abd-A*), and *Abdominal-B* (*Abd-B*); the *Antennapedia complex*, denoted *ANT-C*, consists of five genes, *labial* (*lab*), *proboscipedia* (*pb*), *Deformed* (*Dfd*), *Sex combs reduced* (*Scr*), and *Antennapedia* (*Antp*). Molecular analysis of these genes has demonstrated that they all encode helix-turn-helix transcription factors with a conserved region of 60 amino acids. This region, called the **homeodomain**, is involved in DNA binding.

The BX-C was the first of the two homeotic gene complexes to be dissected genetically. Analysis of this complex began in the late 1940s with the work of Edward Lewis. By studying mutations in the BX-C, Lewis showed that the wildtype function of each part of the complex is restricted to a specific region in the developing animal. Molecular analyses later reinforced and refined this conclusion. Study of the ANT-C began in the 1970s, principally through the work of Thomas Kaufman, Matthew Scott, and their collaborators. Through a combination of genetic and molecular analyses, these investigators showed that the genes of the ANT-C are also expressed in a regionally specific fashion. However, the ANT-C genes are expressed more anteriorly than the BX-C genes. Curiously, the pattern of expression of the ANT-C and BX-C genes along the anterior-posterior axis corresponds exactly to the order of the genes along the chromosome (Figure 20.8); it is not yet clear why this is so. The developmental pathway that each cell takes seems to depend simply on the set of homeotic genes that are expressed within it. Because the homeotic genes play such a key role in selecting the segmental identities of individual cells, they are often called selector genes.

The proteins encoded by the homeotic genes are homeodomain transcription factors. These proteins bind to regulatory sequences in the DNA, including some within the bithorax and Antennapedia complexes themselves. For example, the UBX and ANTP proteins bind to a sequence within the promoter of the Ubx gene—a suggestion that the homeotic genes can regulate themselves and each other. Other gene targets of the homeodomain transcription factors have been identified, including some that encode other types of transcription factors. The homeotic genes therefore seem to control a regulatory cascade of target genes, which in turn act to determine the segmental identities of individual cells. However, the homeotic genes do not stand at the top of this regulatory cascade. Their activities are controlled by another group of genes expressed earlier in development.

Segmentation Genes

Most of the homeotic genes were identified by mutations that alter the phenotype of the adult fly. However, these same mutations also have phenotypic effects in the embryonic and larval stages. This finding suggested that other genes involved in segmentation might be discovered by screening for mutations that cause embryonic and larval defects. In the 1970s and 1980s, Christiane Nüsslein-Volhard and Eric Wieschaus carried out such screens (see A Milestone in Genetics: Mutations that Disrupt Segmentation in *Drosophila* on the Student Companion site). They found a whole new set of genes required for segmentation along

the anterior-posterior axis. Nüsslein-Volhard and Wieschaus classified these **segmentation genes** into three groups based on embryonic mutant phenotypes.

- 1. Gap Genes. These genes define segmental regions in the embryo. Mutations in the gap genes cause an entire set of contiguous body segments to be missing; that is, they create an anatomical gap along the anterior-posterior axis. Four gap genes have been well characterized: Krüppel (from the German for "cripple"), giant, bunchback, and knirps (from the German for "dwarf"). Each is expressed in characteristic regions in the early embryo under the control of the maternal-effect genes bicoid and nanos. The gap genes encode transcription factors.
- Pair-Rule Genes. These genes define a pattern of segments within the embryo. The pair-rule genes are regulated by the gap genes and are expressed in seven alternating bands, or stripes, along the anterior-posterior axis, in effect dividing the embryo into 14 distinct zones, or *parasegments* (■ Figure 20.9). Some of the mutations in pair-rule genes produce embryos with only half

0.1 mm

■ **FIGURE 20.9** The seven-stripe pattern of RNA expression of the pair-rule gene *fushi tarazu* (*ftz*) in a *Drosophila* blastoderm embryo. The RNA was detected by *in situ* hybridization with a *ftz*-specific probe. Anterior is at the left; dorsal is at the top. Other pair-rule genes show a different seven-stripe pattern.

as many parasegments as wild-type have. In each mutant, every other parasegment is missing, although the missing parasegments are not the same in different pair-rule mutants. Examples of pair-rule genes are *fushi tarazu* (from the Japanese for "something missing") and *even-skipped*. In *fushi tarazu* mutants, each of the odd-numbered parasegments is missing; in *even-skipped* mutants, each of the even-numbered parasegments is missing. The pair-rule genes also encode transcription factors.

3. *Segment-Polarity Genes.* These genes define the anterior and posterior compartments of individual segments along the anterior-posterior axis. Mutations in segment-polarity genes cause part of each segment to be replaced by a mirror-image copy of an adjoining half-segment. For example, mutations in the segment-polarity gene *gooseberry* cause the posterior half of each segment to be replaced by a mirror-image copy of the adjacent anterior half-segment. Many of the segment-polarity genes are expressed in 14 narrow bands along the anterior-posterior axis. Thus, they refine the segmental pattern established by the pair-rule genes. Two of the best-studied segment-polarity genes are *engrailed* and *wingless; engrailed* encodes a transcription factor, and *wingless* encodes a signaling molecule.

These three groups of genes form a regulatory hierarchy (**Figure 20.10**). The gap genes, which are regionally activated by the maternal-effect genes, regulate the expression of the pair-rule genes, which in turn regulate the expression of the segment-polarity genes. Concurrent with this process, the homeotic genes are activated under the control of the gap and pair-rule genes to give unique identities to the segments that form along the anterior-posterior axis. Interactions among the products of all these genes then refine and stabilize the segmental boundaries. In this way, the *Drosophila* embryo is progressively subdivided into smaller and smaller developmental units.

ORGAN FORMATION

When many different types of cells are organized for a specific purpose, they form an organ. The heart, stomach, kidney, liver, and eye are all examples of organs. One of the remarkable features of an organ is that it forms in a specific part of the body. The development of a heart in the head or an eye in the thorax of a fly, for example, would be extremely abnormal, and we would wonder what had gone wrong. Anatomically correct organ formation is obviously under tight genetic control.

Geneticists have obtained insights into the nature of this control from the study of another gene in *Drosophila*. This gene is called *eyeless* after the phenotype of flies that are mutant for it (**■** Figure 20.11). The wild-type *eyeless* gene encodes a

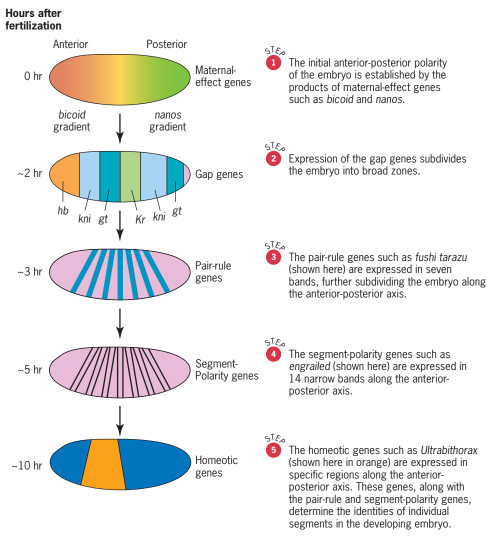


FIGURE 20.10 Cascade of gene expression to produce segmentation in Drosophila embryos.

homeodomain transcription factor whose action switches on a developmental pathway that involves several thousand genes. Initially, several subordinate regulatory genes are activated. Their products then trigger a cascade of events that create specific cell types within the developing eye.

The role of the *eyeless* gene has been demonstrated by expressing it in tissues that normally do not form eyes (**Figure 20.12**). Walter Gehring and colleagues did this by creating transgenic flies in which the *eyeless* gene was fused to a promoter that could be activated in specific tissues. Activation of this promoter caused transcription of the *eyeless* gene outside its normal domain of expression. This, in turn, caused eyes to form in unorthodox places such as wings, legs, and antennae. These extra (or ectopic) eyes were anatomically well developed and functional; in fact, their photoreceptors responded to light.

An even more remarkable finding is that a mammalian homologue of the *eyeless* gene, called *Pax6*, also produces these extra eyes when it is inserted into *Drosophila* chromosomes. Gehring and coworkers used the mouse homologue of *eyeless* to transform *Drosophila*, and they got the same result as they did with the *eyeless* gene itself. This showed that the mouse gene, which also encodes a homeodomain protein, is functionally equivalent to the *Drosophila* gene; that is, it regulates the pathway for eye development. However, when the mouse gene is put into *Drosophila*, it produces *Drosophila* eyes, not mouse eyes. *Drosophila* eyes develop because the genes that respond to the regulatory command of the inserted mouse gene are normal *Drosophila*



FIGURE 20.11 The phenotype of an *eyeless* mutant in *Drosophila*.

genes, which must, of course, specify the formation of a *Drosophila* eye. In mice, mutations in the homologue of the *eyeless* gene reduce the size of the eyes; for that reason, the mutant phenotype is called *Small eye*. A homologue of *eyeless* and *Small eye* has also been found in humans. Mutations in this gene cause a syndrome of eye defects called *aniridia* in which the iris is reduced or missing.

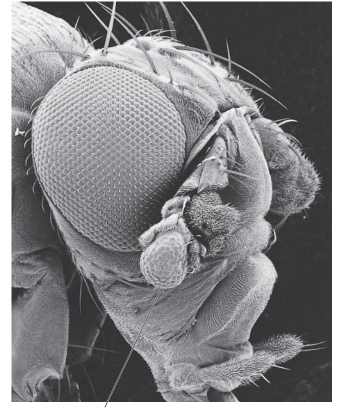
The discovery of homologous genes that control eye development in different organisms has profound evolutionary implications. It suggests that the function of these genes is very ancient, dating back to the common ancestor of flies and mammals. Perhaps the eyes in this ancestral organism were nothing more than a cluster of light-sensitive cells organized through the regulatory effects of a primitive *eyeless* gene. Over evolutionary time, this gene continued to regulate the increasingly more complicated process of eye development, so that today, eyes as different as those in insects and those in mammals are still formed under its control. Solve It: Cave Blindness challenges you to think about the genetic situation in organisms that have permanently lost the ability to form eyes.

SPECIFICATION OF CELL TYPES

Within organs, cells differentiate in specific ways. For example, some cells become neurons whereas others become neuronal support cells. The mechanisms that regulate this differentiation have been analyzed by studying very simple situations involving a few distinct cell types. One such situation occurs in the development of the *Drosophila* eye (**Figure 20.13**).

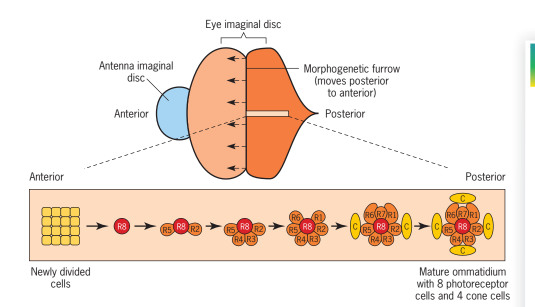
Each of the large compound eyes in *Drosophila* originates as a flat sheet of cells in one of the imaginal discs. Initially, all the cells in this epithelial sheet look the same, but late during the larval stage, a furrow

forms near the posterior margin of the disc. As this furrow moves in the anterior direction across the disc, it triggers a wave of cell divisions in its wake. The newly divided cells then differentiate into specific cell types to form the 800 individual



Extra eye

FIGURE 20.12 An extra eye produced by expressing the wild-type *Drosophila eyeless* gene in the antenna of a fly.



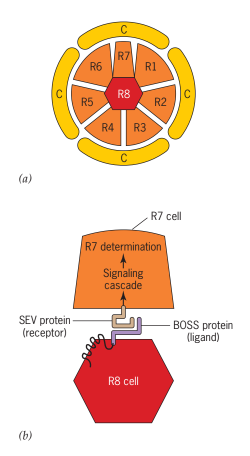
■ FIGURE 20.13 Development of the *Drosophila* eye. As the morphogenetic furrow moves toward the anterior of the eye-antenna imaginal disc, a wave of cell divisions follows in its wake. The newly divided cells then begin to differentiate into specific types. The insert shows the differentiation of the photoreceptor (R1–R8) and cone (C) cells that form each ommatidium (facet) of the compound eye.

Solve It!

Cave Blindness

The *Drosophila eyeless* gene and the mouse *Pax6* gene are master regulators of eye development. Sequence analysis has demonstrated that these two genes are homologous—that is, they are derived from a gene that was present in the common ancestor of flies and mammals. Other animals with eyes seem to have a derivative of this gene too. Some cavedwelling animals—for example, the blind cave fish, have lost the ability to form eyes. What hypothesis could you propose to explain why these animals are eyeless? How could you test this hypothesis?

To see the solution to this problem, visit the Student Companion site.



■ FIGURE 20.14 Determination of the R7 photoreceptor of an ommatidium (facet) in the *Drosophila* compound eye. (a) Arrangement of the eight photoreceptors (1–8) and four cone cells (C) in an ommatidium. (b) Signaling between the differentiated R8 cell and the presumptive R7 cell. The bride of sevenless (BOSS) protein on the R8 cell is the ligand for the sevenless (SEV) receptor protein on the surface of the R7 cell. Activation of this receptor initiates a signaling cascade within the R7 cell that induces it to differentiate. facets of the adult eye. Each facet consists of 20 cells. Eight are photoreceptor neurons designed to absorb light; four are cone cells that secrete a lens to focus light into the photoreceptors; six are sheath cells to provide insulation and support; and the two remaining cells form sensory hairs on the eye's surface. Thus, a highly patterned array of intricately differentiated facets develops from what had been a flat sheet of identical cells. What brings this transformation about?

Gerald Rubin and his collaborators have attempted to answer this question by collecting mutations that disrupt eye development. Their research has led to the idea that the specification of cell types within each facet depends on a series of cell–cell interactions. This is illustrated in the differentiation of the eight photore-ceptor cells, denoted R1, R2, . . . R8 (**■** Figure 20.14). In a fully formed facet, six of the photoreceptors (R1–R6) are arranged in a circle around the other two (R7, R8). One of the central cells, R8, is the first to differentiate in the developing facet. Its appearance is followed by the differentiation of the peripheral cells R2 and R5, then by R3 and R4, and R1 and R6; finally, the second central cell, R7, differentiates into a photoreceptor.

This last event has been studied in great detail. Rubin and his colleagues have shown that the differentiation of the R7 cell depends on reception of a signal from the already differentiated R8 cell. To receive this signal, the R7 cell must synthesize a specific receptor, a membrane-bound protein encoded by a gene called *sevenless* (sev). Mutations in this gene abolish the function of the receptor and prevent the R7 cell from differentiating as a neuron; instead, it differentiates as a cone cell. The signal for the R7 receptor is produced by a gene called *bride of sevenless (boss)* and is specifically expressed on the surface of the R8 cell. Contact between the differentiated R8 cell and the undifferentiated R7 cell allows the R8 signal, or ligand as it is technically called, to interact with the R7 receptor and activate it. This activation induces a cascade of changes within the R7 cell that ultimately prompt it to differentiate as a light-receiving neuron. This differentiation is presumably mediated by one or more transcription factors acting on genes within the R7 nucleus. Thus, the signal from the R8 cell is "transduced" into the R7 nucleus, where it alters the pattern of gene expression. The analysis of eye development in Drosophila therefore shows that *induction*, the process of determining the fate of an undifferentiated cell by a signal from a differentiated cell, can play an important role in the specification of cell types.

The protein encoded by the *sev* gene is a tyrosine kinase—that is, a protein that phosphorylates tyrosine residues in other proteins. Once the SEV protein has been activated by contact with the BOSS ligand, it phosphorylates other proteins inside the R7 cell. These intracellular proteins are downstream effectors of the BOSS signal. Ultimately, they activate transcription factors to stimulate the expression of the genes that are involved in the differentiation of the R7 cell as a photoreceptor. To explore the BOSS-SEV interaction further, work through Problem-Solving Skills: The Effects of Mutations during Eye Development.

KEY POINTS

- The zygotic genes are activated after fertilization in response to maternal gene products.
- In Drosophila, the products of the segmentation genes regulate the subdivision of the embryo into a series of segments along the anterior-posterior axis.
- The identity of each body segment is determined by the products of genes in the bithorax and Antennapedia homeotic gene complexes.
- The formation of an organ may depend on the product of a master regulatory gene, such as the eyeless gene in Drosophila.
- In Drosophila specific cell types differentiate after segmental identities have been established.
- Differentiation events may involve a signal produced by one cell and a receptor produced by another cell.

PROBLEM-SOLVING SKILLS



The Effects of Mutations during Eye Development

THE PROBLEM

In *Drosophila*, the interaction between the SEV and BOSS proteins signals R7 cells to differentiate as photoreceptors in the ommatidia of the compound eyes; when this interaction does not occur, the R7 cells differentiate as cone cells. Neither the SEV nor the BOSS proteins appear to be needed for any other developmental event in the fly. (a) Predict the phenotypes of flies that are homozygous for recessive, loss-of-function mutations in either the *sev* or the *boss* genes. (b) Predict the phenotype of a fly that is heterozygous for a dominant, gain-of-function mutation that constitutively activates the SEV protein. (c) Suppose that one copy of this dominant, gain-of-function was introduced into a fly that was homozygous for a recessive, loss-of-function mutation in the *boss* gene. What would the phenotype of that fly be?

FACTS AND CONCEPTS

- **1.** A loss-of-function mutation in a gene abolishes the function of that gene's protein product.
- **2.** A gain-of-function mutation in a gene endows that gene's product with a new function.
- **3.** A protein that is constitutively active carries out its function all the time.

ANALYSIS AND SOLUTION

This problem focuses on a developmental event in the Drosophila eye-differentiation of the R7 photoreceptor cell. A key step in the process that leads to this event is signaling between the BOSS ligand molecule, which is located in the membrane of the already differentiated R8 cell, and the SEV receptor, which is located in the membrane of the still undifferentiated R7 cell (see Figure 20.14). The failure of either protein to function will prevent the signal from "going through." (a) Recessive, loss-of-function mutations in either the sev and/or boss genes will therefore lead to flies that do not have R7 photoreceptors in the ommatidia of their eyes. (b) However, a dominant, gain-of-function mutation that constitutively activates the SEV protein would be expected to lead to R7 differentiation. (c) Furthermore, this differentiation would be expected to occur even if the fly is homozygous for a recessive, loss-of-function mutation in the boss gene, because with a constitutively activated SEV protein, BOSS function is irrelevant.

For further discussion visit the Student Companion site.

Genetic Analysis of Development in Vertebrates

Much of the knowledge about the genetic control of development comes from the study of model invertebrates. Geneticists would like to apply and extend this knowledge to vertebrates. The ultimate goal would be to learn about the genetic control of development in our own species. One strategy for achieving this goal is to use the information obtained from the study of invertebrate genes to identify developmentally significant genes in vertebrates. Another is to study model vertebrate species with techniques similar to those that are being used in invertebrates. Geneticists can study development in vertebrates by applying knowledge gained from the study of model invertebrates, by analyzing mutations in model vertebrates such as mice, and by examining the differentiation of stem cells.

VERTEBRATE HOMOLOGUES OF INVERTEBRATE GENES

Once a gene has been isolated and sequenced, researchers can screen DNA sequence databases for homologous genes in other organisms. If the gene's sequences have been reasonably well conserved over evolutionary time, this procedure works even for distantly related species. Thus, it has been possible to identify genes from various vertebrate species that are homologous to genes from *Drosophila* and *C. elegans*. The identification of a vertebrate gene then makes many kinds of experimental analyses possible, including assays for the gene's expression at both the RNA and protein levels.

One of the most dramatic applications of this approach has shown that vertebrates contain homologues of the homeotic genes of *Drosophila*. These so-called *Hox* genes were initially identified by probing Southern blots of mouse and human genomic DNA with segments of the *Drosophila* homeotic genes. Subsequently, the cross-hybridizing DNA fragments were cloned, mapped with restriction enzymes, and sequenced. The results of all these analyses have established that mice, humans, and

many other vertebrates so far examined have 38 *Hox* genes in their genomes. These genes are usually organized in four clusters, each about 120 kb long; in mice and humans each cluster is located on a different chromosome. It seems that the four *Hox* gene clusters were created by the quadruplication of a primordial cluster very early in the evolution of the vertebrates, probably 500 to 600 million years ago.

The genes within each *Hox* cluster are transcribed in the same direction, and their expression proceeds from one end of the cluster to the other end, both spatially (anterior to posterior in the embryo) and temporally (early to late in development). There is, therefore, a close parallel with the expression profiles of the ANT-C and BX-C genes of *Drosophila*. Comparative studies indicate that the *Hox* genes play important roles in establishing the identities of specific regions in many different types of vertebrate embryos.

THE MOUSE: RANDOM INSERTION MUTATIONS AND GENE-SPECIFIC KNOCKOUT MUTATIONS

The genetic control of development cannot be studied in vertebrates with the same thoroughness as it can in invertebrates such as *Drosophila*. Obviously, there are technical and logistical constraints. Vertebrates have comparatively long life cycles, their husbandry is expensive, and it is difficult to obtain and analyze mutant strains, especially those with a developmental significance. In spite of these shortcomings, geneticists have been able to make headway in the genetic analysis of development in some vertebrate species, especially the mouse.

Over 500 loci responsible for genetic diseases have been identified in the mouse, and some of them are involved in developmental processes. Most of these loci were discovered through ongoing projects to collect spontaneous mutations. Such work requires that very large numbers of mice be reared and examined for phenotypic differences, and that whatever differences are found be tested for genetic transmission. This is painstaking, costly work that can be supported only at a few facilities in the entire world. Once a mutation is detected, it can be mapped on the chromosomes, and then the mutant gene can be identified and analyzed at the molecular level. Techniques for inducing mutations by inserting known DNA sequences into genes have expedited this process. Insertion mutations are much easier to map and analyze than spontaneous mutations because they have been tagged by the inserted DNA. Furthermore, because the inserting agent-either a transposon or an inactivated retrovirus—is usually not too specific about where it lands in the genome, these techniques are fairly indiscriminant about which genes they mutate. Many of the genes that are relevant to a developmental process under study can therefore be "hit" by an insertion and subsequently identified.

Mouse geneticists have also invented procedures to mutate specific genes. In these procedures, which are discussed in Chapter 16, the integrity of a gene is disrupted by an insertion that is specifically targeted to that gene. Such a disruption, called a **knockout mutation**, can help a researcher determine what role the normal gene plays during development. For example, mice that are homozygous for a knockout mutation in the *Hoxc8* gene develop an extra pair of ribs posterior to the normal set of ribs; they also have clenched toes on their forepaws. The extra-rib phenotype in these mutant mice is reminiscent of the segmental transformations that are seen with homeotic mutations in *Drosophila*. Thus, the mouse's *Hoxc8* gene appears to be involved in establishing the identities of tissues along the anterior-posterior axis and also within the digits.

The genetic analysis of development in mice is providing clues about development in our own species. For example, mutations in at least two different mouse genes mimic the development of abnormal left-right asymmetries in humans. Normally, humans, mice, and other vertebrates exhibit structures that are asymmetric along the left-right body axis. The heart tube always loops to the right, and the liver, stomach, and other viscera are shifted either to the left or right away from the body's midline. In mutant individuals, these characteristic asymmetries are not seen, perhaps because of a defect in the mechanisms that establish the basic body plan. Studying these types of mutants in the mouse may therefore help to elucidate how the organs are positioned in humans.

STUDIES WITH MAMMALIAN STEM CELLS

The terminally differentiated cells in the human body—lymphocytes, neurons, muscle fibers, and so on—usually do not divide. When these types of cells are lost through death, they must be replenished or the tissue they belong to will atrophy. Replenishment occurs when unspecialized cells present in the tissue divide to produce cells that subsequently differentiate into the specialized cell type. These unspecialized precursors of specialized cells are called **stem cells**. For example, the marrow in a human femur contains undifferentiated cells that can replenish various types of blood cells. These *hematopoietic stem cells* keep the circulatory system supplied with lymphocytes, erythrocytes, and platelets. The tissues in some organs such as the heart appear to have very few stem cells; consequently, their ability to regenerate lost or damaged material is limited. Other tissues, such as the gut lining and the skin, have large populations of stem cells that vigorously replace differentiated cells as they are lost. Because these types of stem cells are found in developed organisms, they are called *adult stem cells*.

Stem cells are also found in developing organisms. In fact, during the earliest stages of development, all or most of the cells have the properties of stem cells. Cells taken from a mouse embryo, for example, can be cultured *in vitro* and subsequently transplanted into another mouse embryo, where they will divide and ultimately contribute to the formation of many kinds of tissues and organs. *Embryonic stem (ES) cells* therefore have tremendous developmental potential; that is, they are **pluripotent**—able to develop in many ways.

No matter if they are derived from embryonic or adult tissue, stem cells provide an opportunity to study the mechanisms involved in the differentiation of special cell types. Stem cells can be obtained from a variety of mammals, including mice, monkeys, and humans. They can be cultured *in vitro* and examined for differentiation while growing there or after being transplanted into a host organism. While in culture, stem cells can be treated in various ways to ascertain what triggers their development in a specific direction. Molecular techniques, including gene-chip technologies, allow researchers to determine which genes the cells are expressing as their developmental programs unfold.

Because embryonic stem cells have the greatest developmental potential, they are ideally suited for this kind of analysis. These cells are usually derived from the inner cell mass of embryos that had been created by *in vitro* fertilization. Cells isolated from this mass are plated on a layer of mitotically inactive "feeder cells," which provide growth factors to stimulate division. For mouse ES cells growing in culture, the doubling time is about 12 hours; for human ES cells, it is about 36 hours. After the isolated embryonic cells have grown for a while on the feeder cells, they are dissociated and replated to establish clonal stem-cell populations, which may then be frozen for long-term storage. A clonal cell population is one that has come from a single progenitor cell.

ES cells begin to differentiate when they are transferred from feeder cell cultures to suspension cultures supplied with an appropriate medium. Under these conditions they form **embryoid bodies**, which are multicellular aggregates consisting of differentiated and undifferentiated cells. For some species, the embryoid bodies resemble early embryos. The cells in these bodies may differentiate into the types of specialized cells that are derived from each of the three primary tissue layers—ectoderm, mesoderm, and endoderm. For example, they may form neurons, which are derived from ectoderm; smooth muscle cells or rhythmically contracting cardiac cells, which are derived from mesoderm; or pancreatic islet cells, which are derived from endoderm. By observing this process in different cell lines—for instance, in lines in which particular genes have been mutated—it may be possible to dissect the genetic network of interactions involved in the differentiation of various cell types. The issue of procuring and analyzing human ES cells is, of course, controversial. The human ES cell lines now in use were derived from embryos that were donated by people who had sought medical help to have children through *in vitro* fertilization. Typically, many more embryos are created through this process than are eventually used to produce children. A couple may then decide to donate its unused embryos for research purposes. The derivation of ES cells from such embryos necessarily requires that the embryos be destroyed. Some people view the destruction of early embryos as an acceptable practice; others consider it immoral. The controversy surrounding this practice has caused some governments to withhold or restrict financial support for research on human embryonic stem cells.

The debate on funding for human embryonic stem-cell research has been intensified by the prospect of using human ES cells to cure diseases that result from the loss of specific cell types, such as diabetes mellitus (in which the pancreatic islet cells have been lost) and Parkinson's disease (in which certain types of neurons in a particular region of the brain have been lost). ES cell therapy has also been proposed to treat disabilities such as those resulting from spinal cord damage. The idea is to transplant cells derived from ES cells into the diseased or injured tissue and allow these cells to regenerate the lost or damaged parts of the tissue. Experiments with mice and rats suggest that this strategy might work in humans. However, many technical problems have yet to be solved. For instance, it is not yet possible to obtain pure cultures of a particular differentiated cell type. When human ES cells develop in culture, they differentiate into many kinds of cells; isolating one kind—say, for example, cardiac cells—is a formidable technical challenge.

The proponents of human stem-cell therapy also have to solve other kinds of problems. Cells derived from an *in vitro* culture might divide uncontrollably and form tumors upon being transplanted into a host, or they might be wiped out by the host's immune system. To circumvent the latter problem, researchers have proposed transplanting cells that are genetically identical to the host's cells. Such genetically identical cells could be created by using one of the host's somatic cells to generate the ES cell population. A somatic cell from the host could be fused with an enucleated egg cell obtained from a female donor (not necessarily the host). If the genetically altered egg, which is diploid, divides to form an embryo, cells could be isolated from that embryo to establish an ES cell line, which could then provide genetically identical material for transplantation back into the host.

The production of ES cells by transferring the nucleus of a somatic cell into an enucleated egg is called **therapeutic cloning**. Stem cells might also be obtained by inducing somatic cells to revert to an undifferentiated state. Recent experiments in the United States and Japan indicate that this approach might be feasible. Differentiated skin cells were induced to become pluripotent cells by genetically transforming them with a mixture of four cloned genes. However, some of the genes that were used in these experiments are associated with tumor formation when they are expressed inappropriately. Thus, more research is needed before induced pluripotent cells can be used in stem-cell therapy.

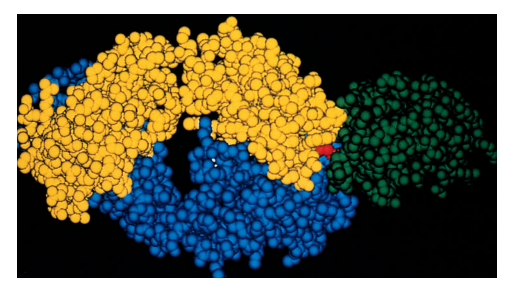
REPRODUCTIVE CLONING

Therapeutic cloning is different from **reproductive cloning**, which aims to produce a complete individual by transferring a somatic-cell nucleus from a donor into an enucleated egg and then allowing that egg to develop into a genetically identical copy of the donor. In 1997, researchers at the Roslin Institute in Scotland produced the first cloned mammal—a sheep named Dolly (see the opening essay in Chapter 2). Dolly was created by replacing the nucleus of an egg with the nucleus from a cell that had been taken from the udder of an adult female sheep. The transplanted nucleus evidently contained all the genetic information needed to direct Dolly's development even though it came from a differentiated cell. Since the creation of Dolly, scientists have produced many other animals by reproductive cloning—mice, cats, cows, and goats. Differentiated cells therefore seem to have the genetic potential to direct development. However, animals produced by reproductive cloning sometimes have developmental abnormalities and shortened lifespans. Frequently, they fail to thrive. This lack of vigor suggests that the somatic nuclei used in reproductive cloning are different from the zygotic nuclei produced by ordinary fertilization. Perhaps the somatic nuclei have accumulated mutations, or perhaps they have undergone changes associated with genetic imprinting or chromosome inactivation—methylation of some nucleotides, acetylation of histones, and so on. Such changes would have to be reversed for a somatic nucleus to function as a zygotic nucleus. Because of the problems encountered in the reproductive cloning of animals, the international scientific community does not consider reproductive cloning of humans to be safe. Consequently, there is widespread agreement that the reproductive cloning of humans should not be attempted.

GENETIC CHANGES IN THE DIFFERENTIATION OF VERTEBRATE IMMUNE CELLS

Although evidence from reproductive cloning suggests that differentiated cells may have the same DNA content as a fertilized egg, we know of some types of differentiated vertebrate cells that do not. These cells are components of the system that protects animals against infection by viruses, bacteria, fungi, and protists—the immune system.

In mammals, where most of the research has been focused, the immune system comprises several distinct types of cells, all derived from stem cells that reside in the bone marrow. These stem cells divide to produce more of their own kind, as well as precursors of specialized immune cells. Two important classes of specialized immune cells participate directly in the fight against invading pathogens. The plasma B cells produce and secrete proteins called immunoglobulins, also known as antibodies, and the killer T cells produce proteins that project from their surfaces and act as receptors for a variety of substances. Both the B-cell antibodies and the T-cell receptors are able to recognize other molecules—for example, the foreign materials introduced by a pathogen-through a lock-and-key mechanism. The foreign molecule, called an **antigen**, is the key that fits precisely into the lock formed by the B-cell antibody or the T-cell receptor (**Figure 20.15**). This specificity of fit is the basis of an animal's ability to defend itself against pathogens. However, because there are many different potential pathogens, an animal must be able to produce many different types of antibodies and T-cell receptors in order to ward off infection.



■ FIGURE 20.15 The three-dimensional structure of an antigen-antibody complex. Only one of the two antigen-binding sites of a typical antibody is shown. The antigen (green) is the enzyme lysozyme. The antigen-binding site of the antibody is formed by the aminoterminal portions of a light chain (yellow) and a heavy chain (blue). A glutamine residue that protrudes from lysozyme where the antibody binds is shown in red. The structure is based on X-ray diffraction data.

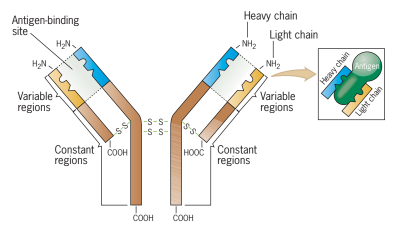


FIGURE 20.16 Structure of an antibody molecule. The inset shows the lock-and-key interaction between the antibody and the antigen that it recognizes.

Antibodies and T-cell receptors are proteins, and proteins are encoded by genes. Therefore, to produce the large array of antibodies and T-cell receptors needed to counter all possible pathogens, it would seem that an animal would have to possess an enormous number of genes-too many to fit even in a large genome such as our own. This predicament perplexed geneticists for years. In the last quarter of the twentieth century, however, researchers discovered how an animal could produce a large number of different antibodies and T-cell receptors by recombining small genetic elements into functional genes. The coding potential achieved by this combinatorial shuffling of gene segments is truly astounding. With a modest amount of DNA dedicated to immune system functions, an animal can produce hundreds of thousands, if not millions, of antibodies and T-cell receptors, each with a different ability to lock on to a foreign molecule from an invading organism.

To see how this recombination system works, we'll focus on the production of antibodies. Each antibody is a tetramer composed of four polypeptides, two identical *light chains* and two identical *heavy chains*, joined by disulfide bonds (**Figure 20.16**). The light chains are about 220 amino acids long, and the heavy chains are about 445 amino acids long. Every chain, light or heavy, has an amino-terminal *variable region*, within which the amino acid sequence varies among the different kinds of antibodies that an animal produces, and a carboxy-terminal *constant region*, within which the amino acid sequence is the same for all antibodies of a particular class.

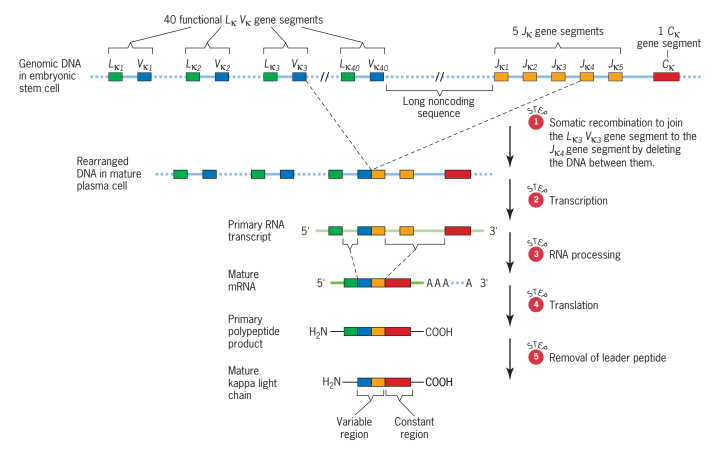


FIGURE 20.17 The genetic control of human antibody kappa light chains. Each kappa light chain is encoded by a gene assembled from different types of gene segments within the immunoglobulin kappa locus (*IGK*) on chromosome 2. This assembly occurs during the differentiation of the plasma B cells of the immune system.

The light and heavy chains of an antibody are encoded by different loci in the genome. In humans, there are two light chain loci, the kappa (κ) locus on chromosome 2 and the lambda (λ) locus on chromosome 22, and there is one heavy chain locus, located on chromosome 14. Each of these loci consists of a long array of gene segments. We'll focus on the kappa locus to see how these segments are organized and how they are recombined into coherent coding sequences to produce different polypeptides.

A kappa polypeptide is encoded by three types of gene segments:

- An L_kV_k gene segment, which encodes a *l*eader peptide and the aminoterminal 95 amino acids of the variable region of the kappa light chain; the leader peptide is removed from the kappa light chain by cleavage after it guides the nascent polypeptide through the membrane of the endoplasmic reticulum in an antibody-synthesizing plasma cell.
- **2.** A \mathcal{J}_{κ} gene segment, which encodes the last 13 amino acids of the variable region of the kappa light chain; the symbol \mathcal{J}_{κ} is used for this gene segment because the peptide it encodes *j*oins the amino-terminal peptide encoded by the $L_{\kappa}V_{\kappa}$ segment to a carboxy-terminal peptide encoded by the next type of gene segment.
- **3.** A C_{κ} gene segment, which encodes the constant region of the kappa light chain.

In humans, the kappa locus contains 76 $L_{\kappa}V_{\kappa}$ gene segments (although only 40 are functional), five \mathcal{J}_{κ} gene segments, and a single C_{κ} gene segment. The \mathcal{J}_{κ} gene segments are located between the $L_{\kappa}V_{\kappa}$ gene segments and the C_{κ} gene segment. In germ-line cells, the five \mathcal{J}_{κ} segments are separated from the $L_{\kappa}V_{\kappa}$ segments by a long noncoding sequence, and from the C_{κ} gene segment by another noncoding sequence approximately 2 kb long (**Figure 20.17**). During the development of a particular B cell, the kappa light chain gene that will be expressed is assembled from one $L_{\kappa}V_{\kappa}$ segment, one \mathcal{J}_{κ} segment, and the single C_{κ} segment by a process of somatic recombination. Any one of the 40 functional $L_{\kappa}V_{\kappa}$ gene segments can be joined with any one of the five \mathcal{J}_{κ} segments in this process; the DNA between the joined segments is simply deleted (**Figure 20.18**). The joining event is mediated by sites called recombination signal sequences (RSS), which are adjacent to each of the gene segments. These sites are composed of 7- or 9-base pair-long repeats separated by 12- or 23-base pair-long spacers. The repeats within the RSS immediately downstream of an $L_{\kappa}V_{\kappa}$ gene segment are complementary to the repeats within the RSS immediately upstream of a \mathcal{J}_{κ} gene segment. When these repeats pair, a protein complex can catalyze recombination between them, joining the $L_{\kappa}V_{\kappa}$ segment to the 7, segment. The recombination activating gene proteins 1 and 2 (RAG1 and RAG2) are important components of this complex; together, they control the specificity of the recombination event.

The $L_{\kappa}V_{\kappa}\mathcal{J}_{\kappa}$ fusion that is produced by this recombination event encodes the variable portion of the kappa light chain. The entire DNA sequence— $L_{\kappa}V_{\kappa}\mathcal{J}_{\kappa}$ -noncoding stretch- C_{κ} —in the rearranged kappa locus is then transcribed. The noncoding sequence between the fused $L_{\kappa}V_{\kappa}\mathcal{J}_{\kappa}$ segments and the C_{κ} segment is removed during RNA processing, just as are the introns of other genes, and the resulting mRNA is translated into a polypeptide. The amino-terminal leader peptide is cleaved from this polypeptide to create the finished kappa light chain. The total number of functional kappa light chains that can be produced by this mechanism is 40 (the number of functional $L_{\kappa}V_{\kappa}$ gene segments) \times 5 (the number of \mathcal{J}_{κ} gene segments) \times 1 (the number of C_{κ} gene segments) = 200. In a similar manner, recombination of gene segments can create 120 different lambda light chains and 6600 different heavy chains. The combinatorial assembly of all these chains then makes it possible for a human to produce 320 (200 + 120) \times 6600 = 2,112,000

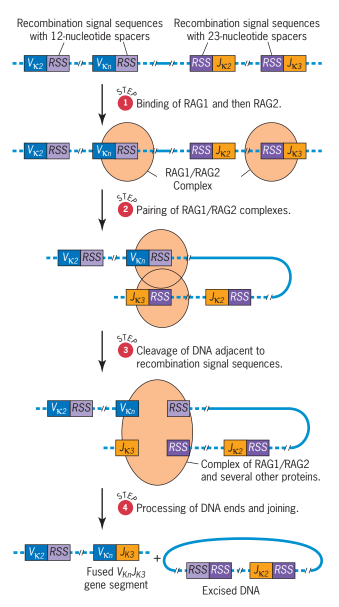


FIGURE 20.18 Simplified model of $V\kappa$ - $J\kappa$ joining. The joining process is mediated by the specific binding of RAG1 and RAG2 to the recombination signal sequences (RSS) adjacent to the $V\kappa$ and $J\kappa$ gene segments. The RSS adjacent to each $V\kappa$ segment contains 12-nucleotide spacers; those adjacent to $J\kappa$ segments contain 23-nucleotide spacers. The RAG1/RAG2 complex catalyzes recombination only when one RSS contains a 12-nucleotide spacer.

different antibodies. However, the actual number of different antibodies is even greater because of slight variations in the sites where the recombination events take place, and because of hypermutability in the sequences that encode the variable regions of the antibody chains. All these events occur independently in the precursors of the plasma B cells. Thus, as these cells differentiate, each one acquires the ability to produce a different antibody.

KEY POINTS •

- Many vertebrate genes—for example, the Hox genes—have been identified by homology with genes isolated from model organisms such as Drosophila and C. elegans.
 - Among vertebrates, the mouse provides opportunities to study mutations that affect development.
 - Mammalian stem cells, especially those derived from embryos, can be cultured in vitro to study the mechanisms that underlie differentiation.
 - Animals produced by reproductive cloning suggest that differentiated cells have the same genetic potential as the zygote.
 - Recombination between gene segments during immune cell differentiation creates the sequences that encode the light and heavy chains of antibodies.

Basic Exercises

Illustrate Basic Genetic Analysis

1. Arrange the following developmental stages in *Drosophila melanogaster* in chronological order from earliest to latest: pupa, blastoderm, zygote, unfertilized egg, larva, adult.

Answer: unfertilized egg, zygote, blastoderm, larva, pupa, adult.

2. *Drosophila* females homozygous for a newly discovered recessive, autosomal mutation lay eggs that do not hatch into larvae, regardless of the genotype of their mates. However, the females themselves show no obvious abnormality. What type of gene does this new mutation define?

Answer: The new mutation defines a maternal-effect gene.

- **3.** Predict the eye phenotype of a fly homozygous for a recessive loss-of-function mutation in the *sevenless* gene. Would a fly homozygous for a recessive loss-of-function mutation in the *bride of sevenless* gene have the same phenotype?
- **Answer:** A fly homozygous for the *sevenless* mutation would not develop the R7 photoreceptor in each of the ommatidia in its compound eyes. The *sevenless* gene encodes the membrane-bound receptor for the extracellular ligand that triggers the R7 cell to differentiate; the ligand is encoded by the *bride of sevenless* gene. A fly homozygous for the *bride of sevenless* mutation would have the same phenotype.
- **4.** Suppose that an antibody light chain gene is assembled from three different gene segments. How many different chains can be produced if the genome contains 5, 20, and 200 copies of the three gene segments?
- **Answer:** If each gene is assembled using one copy of each gene segment, $5 \times 20 \times 200 = 20,000$ different genes are possible.

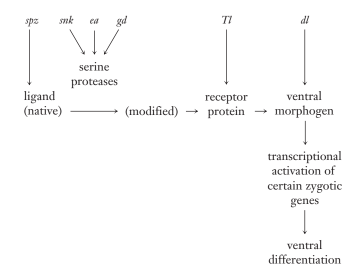
Testing Your Knowledge

Integrate Different Concepts and Techniques

1. The protein product of the *dorsal* (*dl*) gene in *Drosophila* has been called a ventral morphogen—that is, a substance that brings about the formation of ventral structures in the embryo by virtue of its high concentration in the nuclei on the ventral side of the blastoderm. However, the dorsal protein

can enter these ventral nuclei only if a receptor on the embryo's ventral surface has been activated. This receptor is encoded by the *Toll* (*Tl*) gene. The extracellular ligand for the Toll receptor is encoded by the *spätzle* (*spz*) gene. This ligand can exist in two states, "native" and "modified," and the modified state is needed for the activation of the Toll receptor. The products of three genes, *snake* (*snk*), *easter* (*ea*), and *gastrulation defective* (*gd*), are required to convert the native ligand into the modified ligand. All three of these gene products are serine proteases, proteins capable of cleaving other proteins at certain serines in the polypeptide chain. Using these facts, diagram the developmental pathway that ultimately causes the dorsal protein to induce the formation of ventral structures in the *Drosophila* embryo.

Answer: Here is one representation.



The protein product of the *spz* gene is modified by the serine proteases made by the *snk*, *ea*, and *gd* genes. In its

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **20.1** During oogenesis, what mechanisms enrich the cytoplasm of animal eggs with nutritive and determinative materials?
- **20.2** Predict the phenotype of a fruit fly that develops from an embryo in which the posterior pole cells had been destroyed by a laser beam.
- **20.3** Outline the main steps in the genetic analysis of development in a model organism such as *Drosophila*.
- 20.4 Why is the early *Drosophila* embryo a syncytium?
- **20.5** In *Drosophila*, what larval tissues produce the external organs of the adult?
- **20.6 (co)** Like *dorsal, bicoid* is a strict maternal-effect gene in *Drosophila;* that is, it has no zygotic expression. Recessive mutations in *bicoid* (*bcd*) cause embryonic death by preventing the formation of anterior structures. Predict the phenotypes of (a) *bcd/bcd* animals produced by mating heterozygous males and females; (b) *bcd/bcd* animals produced by mating *bcd/bcd* females with *bcd/+* males;

modified form, this ligand is able to activate the Toll receptor protein, but the activation is restricted to the ventral side of the embryo. When the Toll receptor has been activated (presumably, by binding the modified spätzle ligand), it transduces a signal into the cytoplasm of the embryo. This signal ultimately causes the dorsal protein to move into the nuclei on the ventral side of the embryo, where it acts as a transcription factor to regulate the expression of the zygotic genes involved in the differentiation of ventral fates.

- 2. Considering the pathway described above, what would be the phenotypes of recessive loss-of-function mutations in the *spz* and *Tl* genes?
- Answer: For reference, we should note that loss-of-function mutations in *dl* are maternal effect lethals; that is, embryos from *dl/dl* mothers die during development. When these dying embryos are examined, they are found to lack ventral structures. Geneticists say that they are "dorsalized." This peculiar phenotype is due to the failure of the dorsal transcription factor to induce appropriate development in the ventral nuclei of the embryo. In the absence of this induction, the ventral cells differentiate as if they were on the dorsal side of the embryo. Mutations in spz and Tl might be expected to have the same phenotypic effect because they would block steps in the pathway that ultimately causes the dorsal protein to induce ventral differentiation. Recessive mutations in spz and Tl are therefore maternal-effect lethals. Females homozygous for these mutations produce dorsalized embryos that die during development.

(c) *bcd/+* animals produced by mating *bcd/bcd* females with *bcd/+* males; (d) *bcd/bcd* animals produced by mating *bcd/+* females with *bcd/bcd* males; (e) *bcd/+* animals produced by mating *bcd/+* females with *bcd/bcd* males.

- **20.7** Why do women, but not men, who are homozygous for the mutant allele that causes phenylketonuria produce children that are physically and mentally retarded?
- **20.8** In *Drosophila*, recessive mutations in the dorsal-ventral axis gene *dorsal* (*dl*) cause a dorsalized phenotype in embryos produced by *dl/dl* mothers; that is, no ventral structures develop. Predict the phenotype of embryos produced by females homozygous for a recessive mutation in the anterior-posterior axis gene *nanos*.
- **20.9** A researcher is planning to collect mutations in maternaleffect genes that control the earliest events in *Drosophila* development. What phenotype should the researcher look for in this search for maternal-effect mutations?
- **20.10** A researcher is planning to collect mutations in the gap genes, which control the first steps in the segmentation

of *Drosophila* embryos. What phenotype should the researcher look for in this search for gap gene mutations?

- **20.11** How do the somatic cells that surround a developing *Drosophila* egg in the ovary influence the formation of the dorsal-ventral axis in the embryo that will be produced after the egg is fertilized?
- **20.12** What events lead to a high concentration of hunchback protein in the anterior of *Drosophila* embryos?
- 20.13 Diagram a pathway that shows the contributions of the *sevenless (sev)* and *bride of sevenless (boss)* genes to the differentiation of the R7 photoreceptor in the ommatidia of *Drosophila* eyes. Where would *eyeless (ey)* fit in this pathway?
- **20.14** The *sev^{B4}* allele is temperature-sensitive; at 22.7 °C, flies that are homozygous for it develop normal R7 photoreceptors, but at 24.3 °C, they fail to develop these photoreceptors. *sos²⁴* is a recessive, loss-of-function mutation in the *son of sevenless (sos)* gene. Flies with the genotype *sev^{B4}/sev^{B4}*; *sos^{24/+}* fail to develop R7 photoreceptors if they are raised at 22.7 °C. Therefore *sos²⁴* acts as a dominant enhancer of the *sev^{B4}* mutant phenotype at this temperature. Based on this observation, where is the protein product of the wild-type *sos* gene—called SOS—likely to act in the pathway for R7 differentiation?
- 20.15 When the mouse *Pax6* gene, which is homologous to the *Drosophila eyeless* gene, is expressed in *Drosophila*, it produces extra compound eyes with ommatidia, just like normal *Drosophila* eyes. If the *Drosophila eyeless* gene were introduced into mice and expressed there, what effect would you expect? Explain.

- **20.16** Would you expect to find homologues of *Drosophila's* BX-C and ANT-C genes in animals with radial symmetry such as sea urchins and starfish? How could you address this question experimentally?
- **20.17** How might you show that two mouse *Hox* genes are expressed in different tissues and at different times during development?
- **20.18** Distinguish between therapeutic and reproductive cloning.
- 20.19 What is the scientific significance of reproductive cloning?
- **20.20** The methylation of DNA, the acetylation of histones, and the packaging of DNA into chromatin by certain kinds of proteins are sometimes referred to as epigenetic modifications of the DNA. These modifications portend difficulties for reproductive cloning. Do they also portend difficulties for therapeutic cloning and for the use of stem cells to treat diseases or injuries that involve the loss of specific cell types?
- **20.21** Assume that an animal is capable of producing 100 million different antibodies and that each antibody contains a light chain 220 amino acids long and a heavy chain 450 amino acids long. How much genomic DNA would be needed to accommodate the coding sequences of these genes?
- **20.22** Each $L_{\kappa}V_{\kappa}$ gene segment in the kappa light chain locus on chromosome 2 consists of two coding exons, one for the leader peptide and one for the variable portion of the kappa light chain. Would you expect to find a stop codon at the end of the coding sequence in the second (V_{ν}) exon?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

- 1. Images showing the anatomy and developmental stages of *Drosophila* are archived on the Flybase web site. Follow the links from the NCBI web site to the Flybase web site and click on the ImageBrowse feature. Then click on the Embryo icon and browse through the images. When do the syncytial nuclei in the early embryo migrate to the cell membrane? When are these nuclei separated from one another by the formation of membranes between them?
- 2. The Flybase web site also has movies of *Drosophila* development. Click on the Movies icon and explore embryogenesis by looking at the film that shows the cell migration events called gastrulation from a lateral perspective—that is, from a side view. Then look at the film that shows gastrulation in an embryo that is homozygous for a mutation in the pair-rule gene *fushi tarazu* (*ftz*). Describe what is abnormal in the *ftz* embryo.

The Genetic Basis of Cancer



CHAPTER OUTLINE

- Cancer: A Genetic Disease
- Oncogenes
- Tumor Suppressor Genes
- Genetic Pathways to Cancer

was removed surgically, and after a lengthy convalescence, Allison returned to seventh grade, healthy and happy, and imbued with an interest in learning about the disease that had afflicted her. In high school, the courses Allison took reinforced this interest. She read a lot and met several students who enjoyed studying biology. Then another adrenal tumor appeared, but this time not in Allison. Rather, the tumor was found in her father. Louis Romano's tumor—the size of a golf ball—was successfully removed, and Louis recovered fully.

After this incident, the oncologist suspected that both Louis and Allison had developed adrenal tumors—a rare form of cancer called pheochromocytoma—because they carried a mutation in the VHL gene, located in the short arm of chromosome 3. Published research had shown that such mutations are sometimes associated with this type of cancer. The oncologist therefore sent DNA samples from Louis and Allison to a genetics laboratory. DNA tests showed that both Louis and Allison were heterozygous for a mutant VHL allele. At nucleotide 490 in the VHL gene, a G:C base pair had been changed into an A:T base pair, causing serine to be substituted for glycine at position 93 in the polypeptide encoded by the gene.

When Allison learned of this result, she resolved to study genetics. Her older sister, who showed no sign of pheochromocytoma, asked to be tested for the mutant allele and was found to have it. Her doctor then advised her to have regular screenings for any sign of pheochromocytoma. Louis Romano's two siblings—both asymptomatic—were also informed about the *VHL* mutation, but neither of them opted for testing. Allison subsequently majored in biology at a large university and worked for two semesters in a cancer genetics lab. Her project, on the identification of cancerrelated genes in mice, was presented as a poster at the university's annual undergraduate research symposium, where her father and sister could see how she had found purpose in their family's molecular connection.

A Molecular Family Connection

When Allison Romano started looking at colleges and universities, she wanted to find a school where she could study genetics in depth, maybe even do some hands-on research. Her plans were, in a sense, genetically motivated. At age 12 she was diagnosed with a tumor on one of her adrenal glands. This tumor



Colored X-ray image of a pheochromocytoma showing excessive blood vessel growth into the tumor area.

Cancer: A Genetic Disease

Mutations in genes that control cell growth and division are responsible for cancer.

Cancerous tumors kill several hundred thousand Americans every year. What causes tumors to form, and what causes some of them to spread? Why do some types of tumors tend to be found in families? Is the tendency to develop cancer inherited? Do environmental fac-

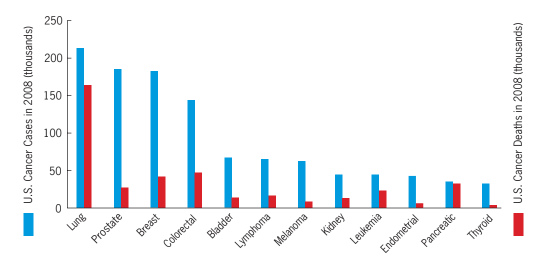
tors contribute to the development of cancer? These and other questions have stimulated an enormous amount of research on the basic biology of cancer. Although many details are still unclear, the fundamental finding is that cancers result from genetic malfunctions. In some instances, these malfunctions may be triggered or exacerbated by environmental factors such as diet, excessive exposure to sunlight, or chemical pollutants. Cancers arise when critical genes are mutated. These mutations can cause biochemical processes to go awry and lead to the unregulated proliferation of cells. Without regulation, cancer cells divide ceaselessly, piling up on top of each other to form tumors. When cells detach from a tumor and invade the surrounding tissues, the tumor is *malignant*. When the cells do not invade the surrounding tissues, the tumor is *benign*. Malignant tumors may spread to other locations in the body, forming secondary tumors. This process is called **metastasis**, from Greek words meaning to "change state." In both benign and malignant tumors, something has gone wrong with the systems that control cell division. Researchers have now firmly established that this loss of control is due to underlying genetic changes.

THE MANY FORMS OF CANCER

Cancer is not a single disease, but rather a group of diseases. Cancers can originate in many different tissues of the body. Some grow aggressively, others more slowly. Some types of cancer can be stopped by appropriate medical treatment; others cannot. **Figure 21.1** shows the frequencies of new cases of different types of cancer in the United States, as well as the number of fatalities attributed to each type. Lung cancer is the most prevalent type, in large measure due to the effects of cigarette smoking. Breast cancer and prostate cancer are also fairly common.

The most prevalent types of cancer are derived from cell populations that divide actively, for example, from epithelial cells in the intestines, lungs, or prostate gland. Rarer forms of cancer develop from cell populations that typically do not divide, for example, from differentiated muscle or nerve cells.

Although the death rate from cancer is still high, enormous progress has been made in detecting and treating different types of cancer. The techniques of molecular genetics have enabled scientists to characterize cancers in ways that were not previously possible, and they have allowed them to devise new strategies for cancer therapy. There is little doubt that the large investment in basic cancer research is paying off.



■ FIGURE 21.1 Estimated number of new cases and deaths from specific types of cancer in the United States in 2008.

Cancer cells can be obtained for experimental study by removing tissue from a tumor and dissociating it into its constituent cells. With appropriate nutrients, these dissociated tumor cells can be cultured *in vitro*, sometimes indefinitely. Cancer cells can also be derived from cultures of normal cells by treating the cells with agents that induce the cancerous state. Radiation, mutagenic chemicals, and certain types of viruses can irreversibly transform normal cells into cancerous cells. The agents that cause this type of transformation are called **carcinogens**.

The abiding characteristic of all cancer cells is that their growth is unregulated. When normal cells are cultured *in vitro*, they form a single cell layer—a monolayer on the surface of the culture medium. Cancer cells, by contrast, overgrow each other, piling up on the surface of the culture medium to form masses. This unregulated pileup occurs because cancer cells do not respond to the chemical signals that inhibit cell division and because they cannot form stable associations with their neighbors.

The external abnormalities that are apparent in a culture of cancer cells are correlated with profound intracellular abnormalities. Cancer cells often have a disorganized cytoskeleton, they may synthesize unusual proteins and display them on their surfaces, and they frequently have abnormal chromosome numbers—that is, they are an euploid.

CANCER AND THE CELL CYCLE

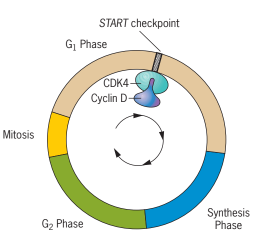
The cell cycle consists of periods of growth, DNA synthesis, and division. The length of this cycle and the duration of each of its components are controlled by external and internal chemical signals. The transition from each phase of the cycle requires the integration of specific chemical signals and precise responses to these signals. If the signals are incorrectly sensed or if the cell is not properly prepared to respond, the cell could become cancerous.

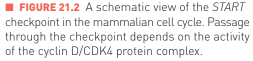
The current view of cell-cycle control is that transitions between different phases of the cycle (G_1 , S, G_2 , and M; see Chapter 2) are regulated at "checkpoints." A **checkpoint** is a mechanism that halts progression through the cycle until a critical process such as DNA synthesis is completed, or until damaged DNA is repaired. When a checkpoint is satisfied, the cell cycle can progress. Two types of proteins play important roles in this progression: the *cyclins* and the *cyclin-dependent kinases*, often abbreviated CDKs. Complexes formed between the cyclins and the CDKs cause the cell cycle to advance.

The CDKs are the catalytically active components of the cell-cycling mechanism. These proteins regulate the activities of other proteins by transferring phosphate groups to them. However, the phosphorylation activity of the CDKs depends on the presence of the cyclins. The cyclins enable the CDKs to carry out their function by forming cyclin/CDK complexes. When the cyclins are absent, these complexes cannot form, and the CDKs are inactive. Cell cycling therefore requires the alternate formation and degradation of cyclin/CDK complexes.

One of the most important cell-cycle checkpoints, called *START*, is in mid- G_1 (**Figure 21.2**). The cell receives both external and internal signals at this checkpoint to determine when it is appropriate to move into the S phase. This checkpoint is regulated by D-type cyclins in conjunction with CDK4. If a cell is driven past the *START* checkpoint by the cyclin D/CDK4 complex, it becomes committed to another round of DNA replication. Inhibitory proteins with the capability of sensing problems in the late G_1 phase, such as low levels of nutrients or DNA damage, can put a brake on the cyclin/CDK complex and prevent the cell from entering the S phase. In the absence of such problems, the cyclin D/CDK4 complex drives the cell through the end of the G_1 phase and into the S phase, thereby initiating the DNA replication that is a prelude to cell division.

In tumor cells, checkpoints in the cell cycle are typically deregulated. This deregulation is due to genetic defects in the machinery that alternately raises and lowers the abundance of the cyclin/CDK complexes. For example, the genes encoding the cyclins or the CDKs may be mutated, or the genes encoding the proteins that respond to specific cyclin/CDK complexes or that regulate the abundance of these complexes may be mutated. Many different types of genetic defects can deregulate the cell cycle, with the ultimate consequence that the cells may become cancerous.





Cells in which the *START* checkpoint is dysfunctional are especially prone to become cancerous. The *START* checkpoint controls entry into the S phase of the cell cycle. If DNA within a cell has been damaged, it is important that entry into the S phase be delayed to allow for the damaged DNA to be repaired. Otherwise, the damaged DNA will be replicated and transmitted to all the cell's descendants. Normal cells are programmed to pause at the *START* checkpoint to ensure that repair is completed before DNA replication commences. By contrast, cells in which the *START* checkpoint is dysfunctional move into S phase without repairing their damaged DNA. Over a series of cell cycles, mutations that result from the replication of unrepaired DNA may accumulate and cause further deregulation of the cell cycle. A clone of cells with a dysfunctional *START* checkpoint may therefore become aggressively cancerous.

CANCER AND PROGRAMMED CELL DEATH

Every cancer involves the accumulation of unwanted cells. In many animals, superfluous cells can be disposed of by mechanisms that are programmed into the cells themselves. Programmed cell death is a fundamental and widespread phenomenon among animals. Without it, the formation and function of organs would be impaired by cells that simply "get in the way."

Programmed cell death is also important in preventing the occurrence of cancers. If a cell with an abnormal ability to replicate is killed, it cannot multiply to form a potentially dangerous tumor. Thus, programmed cell death is a check against renegade cells that could otherwise proliferate uncontrollably in an organism.

Programmed cell death is called **apoptosis**, from Greek roots that mean "falling away." The events that trigger cell death are only partially understood; we will investigate some of them later in this chapter. However, the actual killing events are known in some detail. A family of proteolytic enzymes called *caspases* plays a crucial role in the cell death phenomenon. The caspases remove small parts of other proteins by cleaving peptide bonds. Through this enzymatic trimming, the target proteins are inactivated. The caspases attack many different kinds of proteins, including the lamins, which make up the inner lining of the nuclear envelope, and several components of the cytoskeleton. The collective impact of this proteolytic cleavage is that cells in which it occurs lose their integrity; their chromatin becomes fragmented, blebs of cytoplasm form at their surfaces, and they begin to shrink. Cells undergoing this kind of disintegration are usually engulfed by phagocytes, which are scavenger cells of the immune system, and are then destroyed. If the apoptotic mechanism has been impaired or inactivated, a cell that should otherwise be killed can survive and proliferate. Such a cell has the potential to form a clone that could become cancerous if it acquires the ability to divide uncontrollably.

A GENETIC BASIS FOR CANCER

The recent great advances in understanding cancer have come through application of molecular genetic techniques. However, before these techniques were available to researchers, there was strong evidence that the underlying causes of cancer are genetic. First, it was known that the cancerous state is clonally inherited. When cancer cells are grown in culture, their descendants are all cancerous. The cancerous condition is therefore transmitted from each cell to its daughters at the time of division a phenomenon indicating that cancer has a genetic (or epigenetic) basis. Second, it was known that certain types of viruses can induce the formation of tumors in experimental animals. The induction of cancer by viruses implies that the proteins encoded by viral genes are involved in the production of the cancerous state. Third, it was known that cancer can be induced by agents capable of causing mutations. Mutagenic chemicals and ionizing radiation had been shown to induce tumors in experimental animals. In addition, a wealth of epidemiological data had implicated these agents as the causes of cancer in humans. Fourth, it was known that certain types of cancer tend to run in families. In particular, susceptibility to retinoblastoma, a rare cancer of the eye, and susceptibility to some forms of colon cancer appeared to be inherited as simple dominant

conditions, albeit with incomplete penetrance and variable expressivity. Because susceptibility to these special types of cancer is inherited, it seemed plausible that all cancers might have their basis in genetic defects—either inherited mutations or somatic mutations acquired during a person's lifetime. Finally, it was known that certain types of white blood cell cancers (leukemias and lymphomas) are associated with particular chromosomal aberrations. Collectively, these diverse observations strongly suggested that cancer is caused by genetic malfunctions.

In the 1980s, when molecular genetic techniques were first used to study cancer cells, researchers discovered that the cancerous state is, indeed, traceable to specific genetic defects. Typically, however, not one but several such defects are required to convert a normal cell into a cancerous cell. Cancer researchers have identified two broad classes of genes that, when mutated, can contribute to the development of a cancerous state. In one of these classes, mutant genes actively promote cell division; in the other class, mutant genes fail to repress cell division. Genes in the first class are called **oncogenes**, from the Greek word for "tumor." Genes in the second class are called **tumor suppressor genes**. In the sections that follow, we discuss the discovery, characteristics, and significance of each of these classes of cancer-related genes.

- Cancer is a group of diseases in which the cellular cycle of growth and division is unregulated.
- Cancers may develop if the mechanism for programmed cell death (apoptosis) is impaired.
- Cancers are due to the occurrence of mutations in genes whose protein products are involved in the control of the cell cycle.

Oncogenes

Oncogenes comprise a diverse group of genes whose products play important roles in the regulation of biochemical activities within cells, including those activities related to cell division. These genes were first discovered in the genomes of RNA viruses that Many cancers involve the overexpression of certain genes or the abnormal activity of their mutant protein products.

are capable of inducing tumors in vertebrate hosts. Later, the cellular counterparts of these viral oncogenes were discovered in many different organisms, ranging from *Drosophila* to humans.

TUMOR-INDUCING RETROVIRUSES AND VIRAL ONCOGENES

Fundamental insights into the genetic basis of cancer have come from the study of tumor-inducing viruses. Many of these viruses have a genome composed of RNA instead of DNA. After entering a cell, the viral RNA is used as a template to synthesize complementary DNA, which is then inserted at one or more positions in the cell's chromosomes. The synthesis of DNA from RNA is catalyzed by the viral enzyme reverse transcriptase. This reversal of the normal flow of genetic information from DNA to RNA has prompted biologists to call these pathogens **retroviruses** (see Chapter 17).

The first tumor-inducing virus was discovered in 1910 by Peyton Rous; it caused a special kind of tumor, or sarcoma, in the connective tissue of chickens and has since been called the Rous sarcoma virus. Modern research has shown that the RNA genome of this retrovirus contains four genes: *gag*, which encodes the capsid protein of the virion; *pol*, which encodes the reverse transcriptase; *env*, which encodes a protein of the viral envelope; and *v-src*, which encodes a protein kinase that inserts into the plasma membranes of infected cells. The distinguishing feature of a kinase is that it can phosphorylate other proteins. Of these four genes, only the *v-src* gene is responsible for the virus's ability to form tumors. A virus in which the *v-src* gene has been deleted is infectious but unable to induce tumors. Genes such as *v-src* that cause cancer are called oncogenes.

KEY POINTS

TABLE 21.1

Retroviral Oncogenes

Oncogene	Virus	Host Species	Function of Gene Product	
abl	Abelson murine leukemia virus	Mouse	Tyrosine-specific protein kinase	
erbA	Avian erythroblastosis virus	Chicken	Analog of thyroid hormone receptor	
erbB	Avian erythroblastosis virus	Chicken	Truncated version of epidermal growth- factor (EGF) receptor	
fes	ST feline sarcoma virus	Cat	Tyrosine-specific protein kinase	
fgr	Gardner-Rasheed feline sarcoma virus	Cat	Tyrosine-specific protein kinase	
fms	McDonough feline sarcoma virus	Cat	Analog of colony stimulating growth-factor (CSF-1) receptor	
fos	FJB osteosarcoma virus	Mouse	Transcriptional activator protein	
fps	Fuginami sarcoma virus	Chicken	Tyrosine-specific protein kinase	
jun	Avian sarcoma virus 17	Chicken	Transcriptional activator protein	
mil (mht)	MH2 virus	Chicken	Serine/threonine protein kinase	
mos	Moloney sarcoma virus	Mouse	Serine/threonine protein kinase	
myb	Avian myeloblastosis virus	Chicken	Transcription factor	
тус	MC29 myelocytomatosis virus	Chicken	Transcription factor	
raf	3611 murine sarcoma virus	Mouse	Serine/threonine protein kinase	
H-ras	Harvey murine sarcoma virus	Rat	GTP-binding protein	
K-ras	Kirsten murine sarcoma virus	Rat	GTP-binding protein	
rel	Reticuloendotheliosis virus	Turkey	Transcription factor	
ros	URII avian sarcoma virus	Chicken	Tyrosine-specific protein kinase	
sis	Simian sarcoma virus	Monkey	Analog of platelet-derived growth factor (PDGF)	
src	Rous sarcoma virus	Chicken	Tyrosine-specific protein kinase	
yes	Y73 sarcoma virus	Chicken	Tyrosine-specific protein kinase	

Studies with other tumor-inducing retroviruses have uncovered at least 20 different viral oncogenes, usually denoted *v-onc* (Table 21.1). Each type of viral oncogene appears to encode a protein that could theoretically play a role in regulating the expression of cellular genes, including those involved in the processes of growth and division. Some of these proteins may act as signals to stimulate certain types of cellular activity; others may act as receptors to pick up these signals or as intracellular agents to convey them from the plasma membrane to the nucleus; yet another category of viral oncogene proteins may act as transcription factors to stimulate gene expression. To explore the functions of two of these proteins, use your research skills to answer the questions in Solve It: The *v-erbB* and *v-fms* Viral Oncogenes.

Solve It!

The *v-erbB* and *v-fms* Viral Oncogenes

The *v*-*erbB* gene encodes a truncated version of the receptor for epidermal growth factor (EGF), and the *v*-*fms* gene encodes an analog of the receptor for colony stimulating growth factor (CSF-1). Both of these receptors are transmembrane proteins with a growth-factor-binding domain on the outside of the cell and a protein kinase domain on the inside. How might these proteins transfer a signal from outside the cell to inside the cell?

To see a solution to this problem, visit the Student Companion site.

CELLULAR HOMOLOGUES OF VIRAL ONCOGENES: THE PROTO-ONCOGENES

The proteins encoded by viral oncogenes are similar to cellular proteins with important regulatory functions. Many of these cellular proteins were identified by isolating the cellular homologue of the viral oncogene. For example, the cellular homologue of the *v-src* gene was obtained by screening a genomic DNA library made from uninfected chicken cells. For this screening, the *v-src* gene was used as a hybridization probe to detect recombinant DNA clones that could base-pair with it. Analysis of these clones established that chicken cells contain a gene that is similar to *v-src*—indeed, that is related to it in an evolutionary sense. However, this gene is not associated with an integrated sarcoma virus, and it differs from the *v-src* gene in a very important respect: it contains introns. There are, in fact, 11 introns in the chicken homologue of *v-src*, compared to zero in the *v-src* gene itself. This startling discovery suggested that perhaps *v-src* had evolved from a normal cellular gene and that, concomitantly, it had lost its introns.

The cellular homologues of viral oncogenes are called **proto-oncogenes**, or sometimes, *normal cellular oncogenes*, denoted *c-onc*. The cellular homologue of *v-src* is therefore *c-src*. The coding sequences of these two genes are very similar, differing only in 18 nucleotides; *v-src* encodes a protein of 526 amino acids, and *c-src* encodes a protein of 533 amino acids. By using *v-onc* genes as probes, other *c-onc* genes have been isolated from many different organisms, including humans. As a rule, these cellular oncogenes show considerable conservation in structure. *Drosophila*, for example, carries very similar homologues of the vertebrate cellular oncogenes *c-abl*, *c-erbB*, *c-fps*, *c-raf*, *c-ras*, and *c-myb*. The similarity of oncogenes from different species strongly suggests that the proteins they encode are involved in important cellular functions.

Why do *c-onc*s have introns whereas *v-onc*s do not? The most plausible answer is that *v-onc*s were derived from *c-onc*s by the insertion of a fully processed *c-onc* mRNA into the genome of a retrovirus. A virion that packaged such a recombinant molecule would then be able to transduce the *c-onc* gene whenever it infected another cell. During infection, the recombinant RNA would be reverse-transcribed into DNA and then integrated into the cell's chromosomes. What could be of greater value to a virus than to have a new gene that stimulates increased growth of its host, while its integrated genome goes along for the ride?

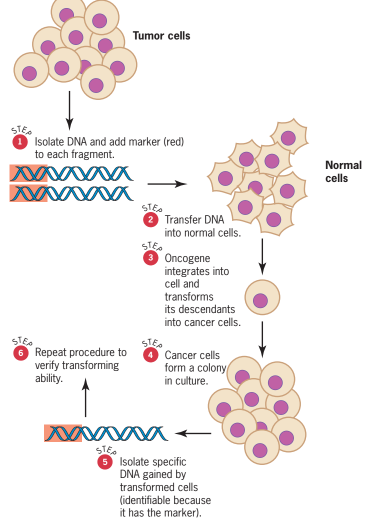
In many cases, the acquisition of an oncogene by a retrovirus has been accompanied by the loss of some viral genetic material. Because this lost material is needed for viral replication, these oncogenic viruses are able to reproduce only if a helper virus is present. In this respect, they resemble the defective transducing bacteriophages we discussed in Chapter 8.

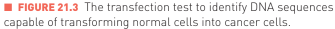
Why do *v-oncs* induce tumors, whereas normal *c-oncs* do not? In some cases it appears that the viral oncogene produces much more protein than its cellular counterpart, perhaps because it has been transcriptionally activated by enhancers embedded in the viral genome. In chicken tumor cells, for example, the *v-src* gene produces 100 times as much tyrosine kinase as the *c-src* gene. This vast oversupply of the kinase evidently upsets the delicate signaling mechanisms that control cell division, causing unregulated growth. Other *v-onc* genes may induce tumors by expressing their proteins at inappropriate times, or by expressing altered—that is, mutant—forms of these proteins.

MUTANT CELLULAR ONCOGENES AND CANCER

The products of the *c-onc*s play key roles in regulating cellular activities. Consequently, a mutation in one of these genes can upset the biochemical balance within a cell and put it on the track to becoming cancerous. Studies of many different types of human cancer have demonstrated that mutant cellular oncogenes are associated with the development of a cancerous state.

The first evidence linking cancer to a mutant *c-onc* came from the study of a human bladder cancer. The mutation responsible for this bladder cancer was isolated by Robert Weinberg and colleagues using a *transfection test* (**Figure 21.3**). DNA was extracted from the cancerous tissue and fragmented into small pieces; then each of these pieces was joined to a segment of bacterial DNA, which served as a molecular marker. The marked DNA fragments were then introduced, or transfected, into cells growing in culture to determine if any of them could transform the cells into a cancerous state. This state could be recognized by the tendency of the cancer cells to form small clumps, or foci, when grown on soft agar plates. The DNA from such cells was extracted and screened to see if it carried the molecular marker that was linked to the original transfecting fragments. If it did, this DNA was retested for its ability to induce the cancerous state. After several





tests, Weinberg's research team identified a DNA fragment from the original bladder cancer that reproducibly transformed cultured cells into cancer cells. This fragment carried an allele of the *c*-*H*-*ras* oncogene, a homologue of an oncogene in the Harvey strain of the rat sarcoma virus. DNA sequence analysis subsequently showed that a nucleotide in codon 12 of this allele had been mutated, with a substitution of a valine for the glycine normally found at this position in the c-H-ras protein.

Geneticists now have some understanding of how this mutation causes cells to become cancerous. Unlike viral oncogenes, the mutant c-H-ras gene does not synthesize abnormally large amounts of protein. Instead, the valine-for-glycine substitution at position 12 impairs the ability of the mutant c-H-ras protein to hydrolyze one of its substrates, guanosine triphosphate (GTP). Because of this impairment, the mutant protein is kept in an active signaling mode, transmitting information that ultimately stimulates the cells to divide in an uncontrolled way (**■** Figure 21.4).

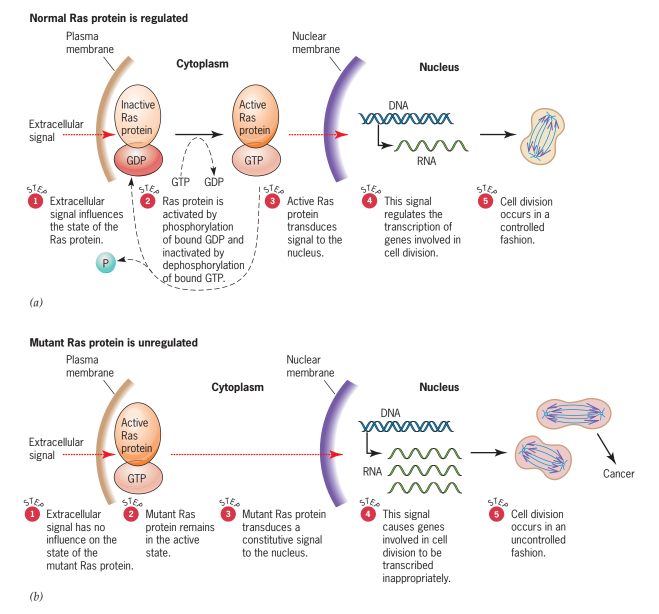


FIGURE 21.4 Ras protein signaling and cancer. (a) The normal protein product of the *ras* gene alternates between inactive and active states, depending on whether it is bound to GDP or GTP. Extracellular signals such as growth factors stimulate the conversion of inactive Ras to active Ras. Through active Ras, these signals are transmitted to other proteins and eventually to the nucleus, where they induce the expression of genes involved in cell division. Because this signaling is intermittent and regulated, cell division occurs in a controlled manner. (b) Mutant Ras proteins exist mainly in the active state. These proteins transmit their signals more or less constantly, leading to uncontrolled cell division, the hallmark of cancer.

Mutant versions of the *c-ras* oncogenes have now been found in a large number of different human tumors, including lung, colon, mammary, prostate, and bladder tumors, as well as neuroblastomas (nerve cell cancers), fibrosarcomas (cancers of the connective tissues), and teratocarcinomas (cancers that contain different embryonic cell types). In all cases, the mutations involve amino acid changes in one of three positions—12, 59, or 61. Each of these amino acid changes impairs the ability of the mutant Ras protein to switch out of its active signaling mode. These types of mutations therefore stimulate cells to grow and divide.

In these types of cancer, only one of the two copies of the *c-ras* gene has been mutated. The single mutant allele is dominant in its ability to bring about the cancerous state. Mutations in *c-ras* and other cellular oncogenes that lead to cancer in this way are therefore *dominant activators* of uncontrolled cell growth.

Dominant activating mutations in cellular oncogenes are seldom inherited through the germ line; rather, the vast majority of them occur spontaneously in the soma during the course of cell division. Because the number of cell divisions in a human life is very large—more than 10¹⁶—thousands of potentially oncogenic mutations are bound to occur, and if each one functioned as a dominant activator of uncontrolled cell growth, the development of a tumor would be inevitable. However, many people lead long lives without developing tumors. The explanation for this paradox is that each individual oncogene mutation is, by itself, seldom able to induce a cancerous state. However, when several different growth-regulating genes have been mutated, the cell cannot compensate for their separate effects, its growth becomes unregulated, and cancer ensues. In many tumors, at least one of these deleterious mutations is in a cellular oncogene. Thus, this group of genes plays an important role in the etiology of human cancer.

CHROMOSOME REARRANGEMENTS AND CANCER

Certain types of human cancer are associated with chromosome rearrangements. For example, chronic myelogenous leukemia (CML) is associated with an aberration of chromosome 22. This abnormal chromosome was originally discovered in the city of Philadelphia and thus is called the *Philadelphia chromosome*. Initially it was thought to have a simple deletion in its long arm; however, subsequent analysis using molecular techniques has shown that the Philadelphia chromosome is actually the result of a reciprocal translocation between chromosomes 9 and 22. (For a general discussion of translocations, see Chapter 6.) In the Philadelphia translocation, the tip of the long arm of chromosome 22 has been joined to the body of chromosome 22, and the distal portion of the long arm of chromosome 22 has been joined to the body of chromosome 9 is in the *c-abl*

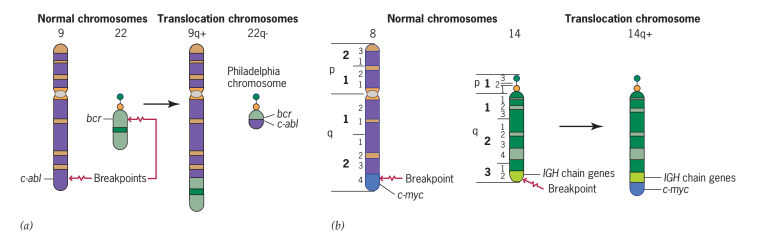


FIGURE 21.5 Translocations implicated in human cancers. (a) The reciprocal translocation involved in the Philadelphia chromosome that is associated with chronic myelogenous leukemia. (b) A reciprocal translocation involved in Burkitt's lymphoma. Only the translocation chromosome (14q+) that carries both the *c-myc* oncogene and the immunoglobulin heavy chain genes (*IGH*) is shown.

oncogene, which encodes a tyrosine kinase, and the breakpoint on chromosome 22 is in a gene called *bcr*. Through the translocation, the *bcr* and *c-abl* genes have been physically joined, creating a fusion gene whose polypeptide product has the amino terminus of the Bcr protein and the carboxy terminus of the c-Abl protein. Although it is not understood precisely why, this fusion polypeptide causes white blood cells to become cancerous. The mechanism may involve the tyrosine kinase activity of the c-Abl protein, which is tightly controlled in normal cells but is deregulated in cells that produce the fusion polypeptide. In effect, the tyrosine kinase function of the c-Abl protein has been constitutively activated by the *bcr/c-abl* gene fusion. This fusion is therefore a dominant activator of the c-Abl tyrosine kinase. Deregulation of the c-Abl tyrosine kinase leads to abnormal phosphorylation of other proteins, including some that are involved in controlling the cell cycle. In their phosphorylated state, these proteins cause cells to grow and divide uncontrollably.

Burkitt's lymphoma is another example of a white blood cell cancer associated with reciprocal translocations. These translocations invariably involve chromosome 8 and one of the three chromosomes (2, 14, and 22) that carry genes encoding the polypeptides that form immunoglobulins (also known as antibodies; see Chapter 20). Translocations involving chromosomes 8 and 14 are the most common (**Translocations**, the *c-myc* oncogene on chromosome 8 is juxtaposed to the genes for the immunoglobulin heavy chains (*IGH*) on chromosome 14. This rearrangement results in the overexpression of the *c-myc* oncogene in cells that produce immunoglobulin heavy chains—that is, in the B cells of the immune system. The *c-myc* gene encodes a transcription factor that activates genes involved in promoting cell division. Consequently, the overexpression of *c-myc* that occurs in cells that carry the *IGH/ c-myc* fusion created by the t8;14 translocation causes those cells to become cancerous.

KEY POINTS

• Some viruses carry genes (oncogenes) that can induce the formation of tumors in animals.

- Viral oncogenes are homologous to cellular genes (proto-oncogenes), which can induce tumors when they are overexpressed or when they are mutated to produce abnormally active protein products.
- Mutations in proto-oncogenes actively promote cell proliferation.
- Some cancers are associated with chromosome rearrangements that enhance the expression of proto-oncogenes or that alter the nature of their protein products.

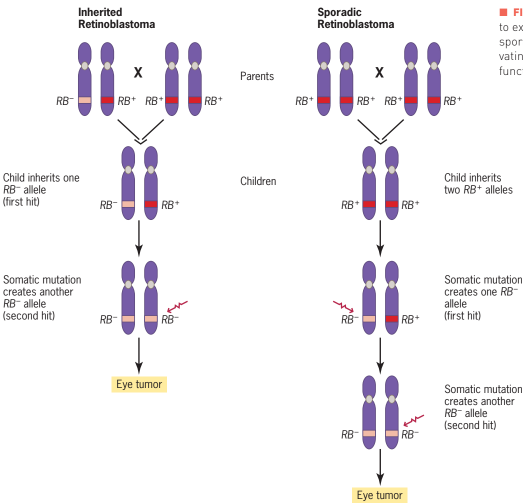
Tumor Suppressor Genes

Many cancers involve the inactivation of genes whose products play important roles in regulating the cell cycle. The normal alleles of genes such as *c-ras* and *c-myc* produce proteins that regulate the cell cycle. When these genes are overexpressed, or when they produce proteins that function as dominant activators, the cell is predisposed to become cancerous. However, the full development of a cancerous state usually requires additional mutations,

and typically these mutations affect genes that are normally involved in the restraint of cell growth. These mutations therefore define a second class of cancer-related genes— the anti-oncogenes, or, as they are more often called, the tumor suppressor genes.

INHERITED CANCERS AND KNUDSON'S TWO-HIT HYPOTHESIS

Many of the tumor suppressor genes were initially discovered through the analysis of rare cancers in which a predisposition to develop the cancer follows a dominant pattern of inheritance. This predisposition is due to heterozygosity for an inherited loss-of-function mutation in the tumor suppressor gene. A cancer develops only if a second



■ **FIGURE 21.6** Knudson's two-hit hypothesis to explain the occurrence of inherited and sporadic cases of retinoblastoma. Two inactivating mutations are required to eliminate the function of the *RB* gene.

mutation occurs in the somatic cells and if this mutation knocks out the function of the wild-type allele of the tumor suppressor gene. Thus, development of the cancer requires two loss-of-function mutations—that is, two inactivating "hits," one in each of the two copies of the tumor suppressor gene.

In 1971 Alfred Knudson proposed this explanation for the occurrence of *retinoblastoma*, a rare childhood cancer of the eye. In most human populations, the incidence of retinoblastoma is about 5 in 100,000 children. Pedigree analysis indicates that approximately 40 percent of the cases involve an inherited mutation that predisposes the individual to develop the cancer. The other 60 percent of the cases cannot be traced to a specific inherited mutation. These noninherited cases are said to be *sporadic*. On the basis of statistical analyses, Knudson proposed that both the inherited and sporadic cases of retinoblastoma occur because the two copies of a particular gene have been inactivated (**Figure 21.6**). In the inherited cases, one of the inactivating mutations has been transmitted through the germ line, and the other occurs during the development of the somatic tissues of the eye. In the sporadic cases, both of the inactivating mutational "hits" are required to knock out a gene that normally functions to suppress tumor formation in the eye.

Subsequent research findings have verified the correctness of Knudson's two-hit hypothesis. First, several cases of retinoblastoma were found to be associated with a small deletion in the long arm of chromosome 13. The gene that normally prevents retinoblastoma—symbolized *RB*—must therefore be located in the region defined by this deletion. More refined cytogenetic mapping subsequently placed the *RB* gene in locus 13q14.2. Second, positional cloning techniques were used to isolate a

TABLE 21.2

Inherited Cancer Syndromes

Syndrome	Primary Tumor	Gene	Chromosomal Location	Proposed Protein Function
Familial retinoblastoma	Retinoblastoma	RB	13q14.3	Cell cycle and transcriptiona regulation
Li-Fraumeni syndrome	Sarcomas, breast cancer	TP53	17p13.1	Transcription factor
Familial adenomatous polyposis (FAP)	Colorectal cancer	APC	5q21	Regulation of β -catenin
Hereditary nonpolyposis	Colorectal cancer	MSH2	2p16	DNA mismatch repair
colorectal cancer (HNPCC)		MLH1	3p21	
		PMS1	2q32	
		PMS2	7p22	
Neurofibromatosis type 1	Neurofibromas	NF1	17q11.2	Regulation of Ras-mediated signaling
Neurofibromatosis type 2	Acoustic neuromas, meningiomas	NF2	22q12.2	Linkage of membrane proteins to cytoskeleton
Wilms' tumor	Wilms' tumor	WT1	11p13	Transcriptional repressor
Familial breast cancer 1	Breast cancer	BRCA1	17q21	DNA repair
Familial breast cancer 2	Breast cancer	BRCA2	13q12	DNA repair
von Hippel-Lindau disease	Renal cancer	VHL	3p25	Regulation of transcriptional elongation
Familial melanoma	Melanoma	p16	9p21	Inhibitor of CDKs
Ataxia telangiectasia	Lymphoma	ATM	11q22	DNA repair
Bloom's syndrome	Solid tumors	BLM	15q26.1	DNA helicase

candidate *RB* gene. Once isolated, the gene's structure, sequence, and expression patterns were determined. Third, the structure of the candidate gene was examined in cells taken from tumorous eye tissue. As predicted by Knudson's two-hit hypothesis, both copies of this gene were inactivated in retinoblastoma cells. Thus, the candidate gene appeared to be the authentic *RB* gene. Finally, cell culture experiments demonstrated that a cDNA from the wild-type allele of the candidate gene could revert the cancerous properties of cultured tumor cells. These cancer reversion experiments proved beyond a doubt that the candidate gene was the authentic *RB* tumor suppressor gene. The protein product of this gene—denoted pRB—was subsequently found to be a ubiquitously expressed protein that interacts with a family of transcription factors involved in regulating the cell cycle.

Knudson's two-hit hypothesis has since been applied to other inherited cancers, including Wilms' tumor, Li-Fraumeni syndrome, neurofibromatosis, von Hippel-Lindau disease, and certain types of colon and breast cancer (Table 21.2). In each case, a different tumor suppressor gene is involved. For example, in Wilms' tumor, a cancer of the urogenital system, the relevant tumor suppressor gene is the *WT1* gene located in the short arm of chromosome 11; in neurofibromatosis, a disease characterized by benign tumors and skin lesions, it is the *NF1* gene located in the long arm of chromosome 17; and in familial adenomatous polyposis, a condition characterized by the occurrence of numerous tumors in the colon, it is the *APC* gene located in the long arm of chromosome 5. Like retinoblastoma, these three diseases are rare, and only a fraction of the observed cases involve an inherited mutation in the relevant tumor suppressor gene. The other cases are caused either by two independent somatic mutations in that gene or by mutations in other, as-yet-unidentified tumor suppressor genes. To explore the genetic dimensions of the two-hit hypothesis, work through Problem-Solving Skills: Estimating Mutation Rates in Retinoblastoma.

PROBLEM-SOLVING SKILLS



Estimating Mutation Rates in Retinoblastoma

THE PROBLEM

Alfred Knudson based his two-hit hypothesis of cancer on a statistical analysis of retinoblastoma. Patients with retinoblastoma (RB) may have tumors in one eye (unilateral RB) or in both eyes (bilateral RB), and within each eye, there may be more than one tumor. Among patients that had inherited an *RB* gene mutation from a parent, Knudson found that the average total number of tumors that formed was 3. Furthermore, he estimated that the total number of retinoblasts—the cells that form the embryonic retina—was about 2 million in each eye. If each tumor in this group of patients is due to the occurrence of another *RB* gene mutation within the first two years of life—the second hit in Knudson's hypothesis—what is the somatic mutation rate for the *RB* gene per year?

FACTS AND CONCEPTS

- 1. Retinoblastoma occurs when both *RB* genes have been inactivated by mutations.
- 2. One of these inactivating mutations may be inherited from a parent.

3. Sporadic cases of retinoblastoma occur when both of the inactivating mutations arise during eye development.

4. When two events are independent, we multiply their probabilities to obtain the probability that they will both occur.

ANALYSIS AND SOLUTION

To estimate the somatic mutation rate, we need to count the number of mutational events in comparison to the total number of chances for such events. The average number of tumors (3) is an estimate of the average number of mutational events. The number of chances for such events is a function of the total number of genes that can mutate to produce a tumor: $1 RB^+$ gene per cell in a patient that has already inherited one RB^- mutation from a parent $\times 2 \times 10^6$ cells per eye $\times 2$ eyes per patient = 4×10^6 chances for a mutational event. Thus, the mutation rate is $3/(4 \times 10^6) = 7.5 \times 10^{-7}$ mutations, or, on an annualized basis, 7.5×10^{-7} mutations/2 years = 3.7×10^{-7} mutations/year.

For further discussion visit the Student Companion site.

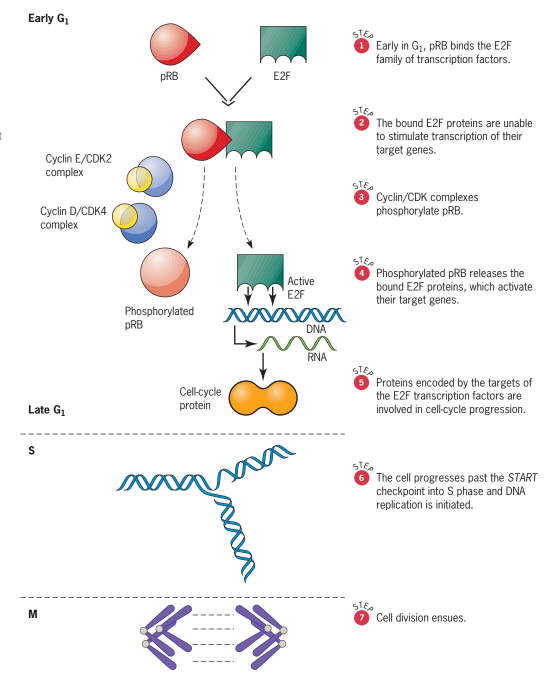
CELLULAR ROLES OF TUMOR SUPPRESSOR PROTEINS

Only about 1 percent of all cancers are hereditary. However, more than 20 different inherited cancer syndromes have been identified, and in nearly all of them the underlying defect is in a tumor suppressor gene rather than in an oncogene. The proteins encoded by these tumor suppressor genes function in a diverse array of cellular processes, including division, differentiation, programmed cell death, and DNA repair. In the following sections, we discuss some of the tumor suppressor proteins that have been studied intensively.

pRB

Recent research has revealed that the RB tumor suppressor protein plays a key role in regulation of the cell cycle. Although the *RB* gene was discovered through its association with retinoblastoma, mutations in this gene are also associated with other types of cancer, including small-cell lung carcinomas, osteosarcomas, and bladder, cervical, and prostate carcinomas. Furthermore, mice that are homozygous for an *RB* knockout mutation die during embryonic development. Thus, the RB gene product is essential for life.

The RB gene product, symbolized pRB, is a 105-kilodalton nuclear protein that is involved in cell-cycle regulation. Two genes homologous to *RB* have been found in mammalian genomes, and their protein products, p107 and p130 (each named for its mass in kilodaltons), may also play key roles in cell-cycle regulation. No human tumors are known to have inactivating mutations in either of these two genes, and mice homozygous for a knockout mutation in either of them do not show abnormal phenotypes. However, mice that are homozygous for knockout mutations in both of these genes die shortly after birth. Thus, together the p107 and p130 members of the RB family of proteins are involved in important cellular processes. ■ FIGURE 21.7 Role of pRB in progression of the cell cycle. Through its negative interaction with E2F transcription factors, pRB stalls the cell cycle in the G₁ phase. Phosphorylation of pRB by the cyclin/CDK complexes frees E2F proteins to activate their target genes, which encode proteins that are instrumental in moving the cell past the *START* checkpoint into the S phase.



Molecular and biochemical analyses have elucidated the role of pRB in cell-cycle regulation (**Figure 21.7**). Early in the G_1 phase of the cell cycle, pRB binds to the E2F proteins, a family of transcription factors that control the expression of several genes whose products move the cell through its cycle. When E2F transcription factors are bound to pRB, they cannot bind to specific enhancer sequences in their target genes. Consequently, the cell-cycle factors encoded by these genes are not produced, and the machinery for DNA synthesis and cell division remains quiescent. Later in G_1 , pRB is phosphorylated through the action of cyclin-dependent kinases. In this changed state, pRB releases the E2F transcription factors that have bound to it. These released transcription factors are then free to activate their target genes, which encode proteins that induce the cell to progress through S phase and into mitosis. After mitosis, pRB is dephosphorylated, and each of the daughter cells enters the quiescent phase of a new cell cycle.

This orderly and rhythmic progression through the cell cycle is disrupted in cancer cells. In many types of cancer—not just retinoblastoma—both copies of the *RB* gene have been inactivated, either by deletions or by mutations that impair or abolish the ability of the RB protein to bind E2F transcription factors. The inability of pRB to bind to these transcription factors leaves them free to activate their target genes, thereby setting in motion the machinery for DNA synthesis and cell division. In effect, one of the natural brakes on the process of cell division has been released. In the absence of this brake, cells have a tendency to move through their cycle quickly. If other cell-cycle brakes fail, the cells divide ceaselessly to form tumors.

p53

The 53-kilodalton tumor suppressor protein p53 was discovered through its role in the induction of cancers by certain DNA viruses. This protein is encoded by a tumor suppressor gene called *TP53*. Inherited mutations in *TP53* are associated with the Li-Fraumeni syndrome, a rare dominant condition in which any of several different types of cancer may develop. Somatic mutations that inactivate both copies of the *TP53* gene are also associated with a variety of cancers. In fact, such mutations are found in a majority of all human tumors. Loss of p53 function is therefore a key step in carcinogenesis.

The p53 protein is a 393-amino-acid-long transcription factor that consists of three distinct domains: an N-terminal transcription-activation domain (TAD), a central DNA-binding core domain (DBD), and a C-terminal homo-oligomerization domain (OD) (**True 21.8a**). Most of the mutations that inactivate p53 are located in the DBD. These mutations evidently impair or abolish the ability of p53 to bind to specific DNA sequences that are embedded in its target genes, thereby preventing the transcriptional activation of these genes. Thus, mutations in the DBD are typically recessive loss-of-function mutations. Other types of mutations are found in the OD portion of the polypeptide. Molecules of p53 with these types of mutations dimerize with wild-type p53 polypeptides and prevent the wild-type polypeptides from functioning as transcriptional activators. Thus, mutations in the OD have a *dominant negative* effect on p53 function.

The p53 protein plays a key role in cellular responses to stress (**Figure 21.8b**). In normal cells the level of p53 is low, but when the cells are treated with a DNAdamaging agent such as radiation, the level of p53 increases dramatically. This response to DNA damage is mediated by a pathway that decreases the degradation of p53. In response to DNA damage, p53 is phosphorylated, converting it into a stable and active form. Once activated, p53 either stimulates the transcription of genes whose products arrest the cell cycle, thereby allowing the damaged DNA to be repaired, or it activates another set of genes whose products ultimately cause the damaged cell to die.

One prominent factor in the response that arrests the cell cycle is p21, a protein encoded by a gene that is activated by the p53 transcription factor. The p21 protein is an inhibitor of cyclin/CDK protein complexes. When p21 is synthesized in response to cell stress, the cyclin/CDK complexes are inactivated and the cell cycle is arrested. During this timeout, the cell's damaged DNA can be repaired. Thus, p53 is responsible for activating a brake on the cell cycle, and this brake allows the cell to maintain its genetic integrity. Cells that lack functional p53 have difficulty applying this brake. If these cells progress through the cell cycle and proceed into subsequent divisions, additional mutations that cause them to be unregulated may accumulate. Mutational inactivation of p53 is therefore often a key step in the pathway to cancer. Solve It: Downstream of p53 challenges you to consider what might happen if p21 were inactivated by mutations.

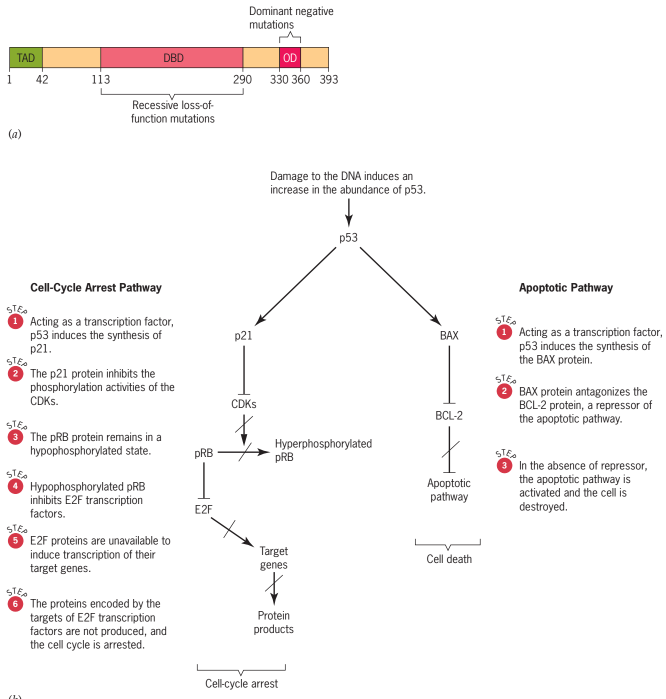
The p53 protein can also mediate another response to cell stress. Instead of orchestrating efforts to repair damage within a cell, p53 may trigger a suicidal response in which the damaged cell is programmed for destruction. The way in which p53 programs cell death is not well understood. One mechanism seems to involve the protein product of the *BAX* gene. The BAX protein is an antagonist of another protein called BCL-2, which normally suppresses the apoptotic, or cell-death, pathway. When the *BAX* gene is activated by p53, its protein product releases the BCL-2 protein from its

Solve It!

Downstream of p53

The p53 protein controls two pathways that respond to damage in a cell's DNA. One pathway arrests the cell cycle to permit repair of the damaged DNA. This pathway is triggered when p53 activates the gene for p21, a protein that inhibits the phosphorylation activities of the cyclin-dependent kinases (CDKs). Would this pathway operate in a cell that has loss-of-function mutations in both of its p21 genes? Explain your answer. Would you classify the p21 gene as a tumor suppressor gene?

To see a solution to this problem, visit the Student Companion site.



(b)

■ FIGURE 21.8 (a) Principal domains within p53. TAD = transcription-activation domain; DBD = DNA-binding domain; OD = oligomerization domain. The numbers refer to amino acid positions in the polypeptide. (b) Role of p53 in the cellular response to DNA damage. Two response pathways have been identified. Within each pathway, a pointed arrow indicates a positive influence or a directional change (e.g., a protein is synthesized or phosphorylated, a protein catalyzes a reaction, or a gene is expressed), and a blunted arrow indicates a negative influence (e.g., repression of protein synthesis or protein activity, or repression of a pathway). A slash through an arrow indicates that the influence—positive or negative—is blocked. suppressing mode. This release then opens the apoptotic pathway, and the cell proceeds to its own destruction.

Curiously, the p53 protein does not seem to play a significant role in the programmed cell death that occurs during embryogenesis. Mice that are homozygous for knockout mutations in *TP53* develop normally, although they have a tendency to develop tumors as they age. Thus, despite its pivotal role in regulating cellular responses to stress, p53 does not seem to influence the course of embryonic development.

pAPC

The 310-kilodalton pAPC protein was discovered through the study of *adenomatous polyposis coli*, an inherited condition that often leads to colorectal cancer. This large protein, 2843 amino acids long (**Figure 21.9a**), plays a key role in regulating the renewal of cells in the lining, or epithelium, of the large intestine. Although

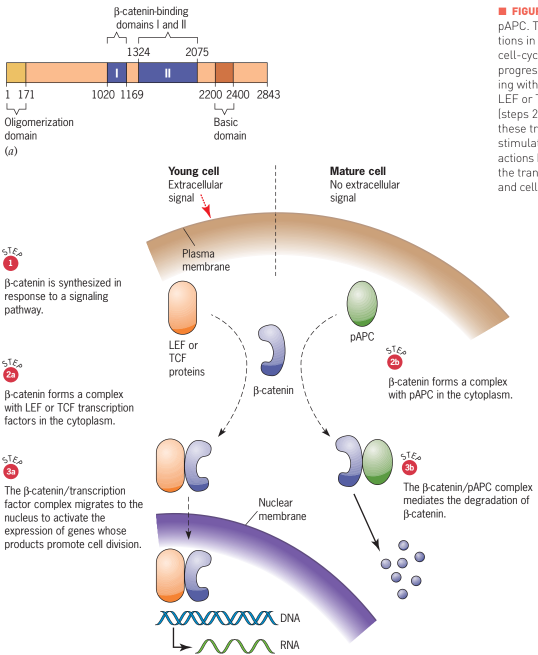


FIGURE 21.9 (a) Principal domains within pAPC. The numbers refer to amino acid positions in the polypeptide. (b) Role of pAPC in cell-cycle control. The pAPC protein influences progression through the cell cycle by interacting with β -catenin, a protein that can activate LEF or TCF transcription factors. In young cells (steps 2a, 3a), an extracellular signal activates these transcription factors and cell division is stimulated. In mature cells (steps 2b, 3b), interactions between pAPC and β -catenin prevent the transcription factors from being activated and cell division is inhibited.

the mechanisms that regulate this process are not fully understood, current information suggests that pAPC controls the proliferation and differentiation of cells in the epithelium of the intestine. When pAPC function is lost, the cells that generate the fingerlike projections on the intestinal epithelium remain in an undifferentiated state. As these cells continue to divide, they produce more of their own kind, and the resulting increase in cell number causes many small, benign tumors to form in the intestinal epithelium. These tumors are called *polyps* or *adenomas*, and the predisposition to form them is inherited as a rare autosomal dominant condition called *familial adenomatous polyposis (FAP)*. In Western countries, its population frequency is about 1 in 7000.

Patients with FAP develop multiple adenomas during their teens and early twenties. Although the adenomas are initially benign, there is a high probability that at least one of them will become a malignant tumor. Thus, at a relatively early age—in the United States, the median is 42—carriers of an FAP mutation develop full-fledged colorectal cancer.

Multiple adenomas develop in the intestines of people who are heterozygous for an FAP mutation because the wild-type *APC* allele they carry mutates multiple times during the natural regeneration of the intestinal epithelium. When such mutations occur, the cells lose their ability to synthesize functional pAPC protein. The absence of this protein releases an important brake on cell proliferation, and cell division proceeds unchecked. Thus, the formation of numerous benign tumors in the intestines of FAP heterozygotes results from the independent occurrence of second mutational "hits" in the cells of the intestinal epithelium. Individuals who do not carry an FAP mutation seldom form multiple adenomas. However, they may produce one or a few adenomas if by chance both of their *APC* genes are inactivated by somatic mutations.

The pAPC protein appears to regulate cell division through its ability to bind β -catenin, a protein that is present inside cells. β -catenin naturally binds to other proteins as well, including certain transcription factors that stimulate the expression of genes whose protein products promote cell division. The interactions with these transcription factors are favored when signals impinging on the cell surface cue the cell to divide (**Figure 21.9***b*). Signal-induced cell proliferation is a necessary process in the intestinal epithelium because this tissue loses an enormous number of cells every day—in humans, about 10¹¹—and the lost cells must be replaced by fresh cells generated by division. Normally, the newly created cells lose their ability to divide as they move away from the generative part of the epithelium and assume their roles in the mature part of the epithelium. This shift from a dividing to a nondividing state occurs because the mature epithelial cells do not receive the extracellular signals that stimulate cells to divide. In the absence of these signals, pAPC forms a complex with the β -catenin in the cells' cytoplasm, and the complexed β -catenin is targeted for degradation. Because pAPC keeps β -catenin levels low in the mature cells of the intestinal epithelium, there is little chance for β -catenin to combine with and activate the transcription factors that stimulate cell division. Cells with mutations in pAPC lose their ability to control β -catenin levels. Without this control, they retain their vigor for division and fail to differentiate properly into mature epithelial cells. The result is that a tumor begins to form in the intestinal lining. Thus, normal pAPC molecules play an important role in suppressing tumor formation in the intestine.

phMSH2

The phMSH2 protein is the human homologue of a DNA repair protein called MutS found in bacteria and yeast. Its involvement in human cancer was elucidated through the study of *bereditary nonpolyposis colorectal cancer (HNPCC)*, a dominant autosomal condition with a population frequency of about 1 in 500. Unlike FAP, HNPCC is characterized by the occurrence of a small number of adenomas, one of which eventually progresses to a cancerous condition. In the United States, the median age at

which the cancer occurs is 42, the same age at which malignant cancer occurs in FAP patients.

The *bMSH2* gene was implicated in the inheritance of HNPCC after researchers found that cells in HNPCC tumors suffer from a general genetic instability. In these cells, di- and trinucleotide microsatellite repeat sequences (see Chapter 13) throughout the genome exhibit frequent changes in length. This instability is reminiscent of the types of DNA sequence changes observed in bacteria with mutations in the genes that control DNA mismatch repair (see Chapter 13). The human homologue of one of these bacterial genes maps to the short arm of chromosome 2, a chromosome that had previously been implicated in HNPCC by linkage analysis. Sequence analysis of this gene—denoted *bMSH2*—indicated that it was inactivated in tumors removed from some HNPCC patients. Thus, loss of *bMSH2* function was causally connected to the genome-wide instability observed in HNPCC tumors. Subsequent analysis has demonstrated that germ-line mutations in *bMSH2*, or in three other human homologues of bacterial mismatch repair genes, account for the inherited cases of HNPCC.

pBRCA1 AND pBRCA2

Mutant versions of the tumor suppressor genes *BRCA1* and *BRCA2* genes have been implicated in hereditary breast and ovarian cancer. *BRCA1* was mapped to chromosome 17 in 1990 and isolated in 1994 (see A Milestone in Genetics: The Identification of the *BRCA1* Gene on the Student Companion site), and *BRCA2* was mapped to chromosome 13 in 1994 and isolated in 1995. Both genes encode large proteins; pBRCA1 is a 220-kilodalton polypeptide, and pBRCA2 is a 384-kilodalton polypeptide. Cellular and biochemical studies have shown that each of these proteins is located within the nuclei of normal cells and that each contains a transcriptional activation domain. The pBRCA1 and pBRCA2 proteins also contain a domain that allows them to interact physically with other proteins, in particular with pRAD51, a eukaryotic homologue of the bacterial DNA repair protein known as RecA. Thus, pBRCA1 and pBRCA2 likely participate in one of the many systems that repair damaged DNA in human cells.

Both pBRCA1 and pBRCA2 carry out important functions within cells. Mice that are homozygous for a knockout mutation in either gene die early during embryogenesis. In the etiology of human cancers, mutant pBRCA1 and pBRCA2 proteins appear to compromise a cell's ability to detect or repair damaged DNA.

Mutations in the *BRCA1* and *BRCA2* genes account for about 7 percent of all cases of breast cancer and about 10 percent of all cases of ovarian cancer in the United States. For each gene, the predisposition to develop these cancers is inherited as a dominant allele with high penetrance. Carriers have a 10- to 25-fold greater risk than noncarriers of developing breast or ovarian cancer, and in some families, the risk of developing colon or prostate cancer is also increased. Because many different inactivating mutations in *BRCA1* and *BRCA2* are found in the human population, genetic counseling for families that are segregating these mutations can be difficult (see the Focus on Cancer and Genetic Counseling).

- Tumor suppressor genes were discovered through their association with rare, inherited cancers
 such as retinoblastoma.
- Mutational inactivation of various tumor suppressor genes is characteristic of most forms of cancer.
- Two mutational hits are required to eliminate both functional copies of a tumor suppressor gene within a cell.
- The proteins encoded by tumor suppressor genes play key roles in regulating the cell cycle.

KEY POINTS

FOCUS ON

CANCER AND GENETIC COUNSELING¹

The identification of inherited mutations in tumor suppressor genes has opened a new era in genetic counseling. The carriers of such mutations are often at high risk to develop potentially life-threatening tumors, sometimes at a relatively early age. If molecular tests reveal that an individual carries a mutant tumor suppressor gene, medical treatment can be given to reduce the chance that he or she will develop a lethal cancer. For example, a child who carries a mutation in the *APC* gene could be checked periodically by endoscopy and suspicious lesions in the intestine could be removed, or a woman who carries a mutation in either of the *BRCA* genes could undergo a prophylactic mastectomy (removal of the breasts) or oophorectomy (removal of the ovaries).

A negative result from a test for a mutant tumor suppressor gene would, of course, be a cause for celebration—at least to the extent that the test can be trusted. For a large gene with many different mutant alleles segregating in the population, it is difficult to design a cost-effective test to detect mutations located anywhere in the gene. Typically, these tests are based on the polymerase chain reaction, and most of them are designed to detect specific mutant alleles. An individual who is at risk to carry a mutant tumor suppressor gene can be tested for the known mutations—at least the most frequent ones. However, a negative result is not definitive because that individual could carry a "private" mutation—that is, one that has not previously been identified in the population.

The existence of private alleles makes counseling for inherited cancers difficult. For example, over 300 different mutations have been identified in the *BRCA1* gene, and about 50 percent of them are private. If an individual with a family history of breast cancer comes to a genetic counselor for evaluation, which mutations should the counselor look for? Sometimes data from other family members or information collected from the individual's ethnic group can provide clues. If other individuals in the family have been found to carry a particular mutant allele, then the counselor should test for that allele. If certain mutant alleles are characteristic of the individual's ethnic group, then the counselor should test for them. In Ashkenazi Jewish populations, for example, some *BRCA1* and *BRCA2* mutant



alleles have frequencies as high as 2.5 percent. By comparison, the combined frequency of all mutant alleles in non-Jewish Caucasian populations is only 0.1 percent. Thus, an Ashkenazi Jew at risk for inherited breast or ovarian cancer should be tested for the mutant alleles that are likely to be segregating in Ashkenazi Jewish families.

Genetic testing for mutant tumor suppressor genes raises a host of psychological issues. In cases where therapeutic medical treatment is not available, an individual might choose not to be tested because the psychological burden of living with the knowledge that one carries a potentially lethal mutant gene could be overwhelming. Knowledge that one is a carrier might be expected to influence career plans and decisions about marriage and child-bearing. The prospect of an early death might dissuade an individual from seeking permanent commitments—to a spouse, to children, or to a vocation-and the chance of transmitting a mutant allele to children might deter the individual from reproducing. Knowledge that one is a carrier can also influence other people-family members, friends, and coworkers. A young daughter whose mother has tested positively for a BRCA1 mutation must herself begin to grapple with the prospect of being a carrier, and a husband whose wife carries a BRCA1 mutation must share in the decision of whether or not she should undergo a prophylactic oophorectomy and preclude the couple from ever having children of their own.

Testing for mutant tumor suppressor genes raises many ethical issues. To whom should the test results be revealed? the patient? the patient's family? parents? children? employer? landlord? insurance agent? What measures should society take to safeguard the privacy of genetic test results? What policies should governments adopt to protect individuals from discrimination on the basis of their genotypes? How should insurance and employment policies be modified? Should the reproductive rights of individuals who carry harmful mutations be limited? As with any technological advance, the ability to detect mutations in tumor suppressor genes leaves us with many questions about how we should proceed. Currently, the answers to these questions are far from clear.

¹Ponder, Bruce. 1997. Genetic testing for cancer risk. Science 278: 1050–1054.

Genetic Pathways to Cancer

Cancers develop through an accumulation of somatic mutations in proto-oncogenes and tumor suppressor genes.

In most cancer cases, the formation of a malignant tumor is not attributable to the uncontrolled activation of a single protooncogene or to the inactivation of a single tumor suppressor gene. Rather, tumor formation, growth, and metastasis usually depend on the accumulation of mutations in several different genes. Thus, the genetic pathways to cancer are diverse and complex.

We can see this diversity and complexity in the formation and development of different types of tumors. For example, benign tumors of the large intestine develop in individuals with inactivating mutations in the APC gene. However, the progression of these tumors to potentially lethal cancers requires mutations in several other genes. This mutational pathway is summarized in **Figure 21.10a**. Inactivating mutations

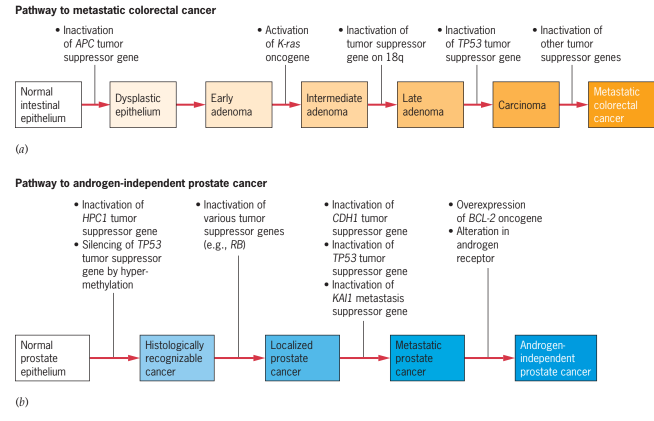


FIGURE 21.10 Genetic pathways to cancer.

in the APC gene initiate the process of tumor formation by causing the development of abnormal tissues within the intestinal epithelium. These abnormal tissues contain dysplastic cells—cells with unusual shapes and enlarged nuclei—that may grow into early-stage adenomas. If the *K*-*ras* proto-oncogene is activated in one of these adenomas, that adenoma may grow and develop more fully. Inactivating mutations in any of several tumor suppressor genes located in the long arm of chromosome 18 may then induce the adenoma to progress further, and inactivating mutations in the *TP53* tumor suppressor gene on chromosome 17 may transform it into a vigorously growing carcinoma. Additional tumor suppressor gene mutations may allow carcinoma cells to break away and invade other tissues. Thus, no less than seven independent mutations (two inactivating hits in the *APC* gene, one activating mutation in the *K*-*ras* gene, two inactivating hits in a tumor suppressor gene on chromosome 18, and two inactivating hits in the *TP53* gene) are required for the development of an intestinal carcinoma, and still more mutations are probably required for the metastasis of that carcinoma to other parts of the body.

The genetic pathways to prostate cancer has also been elucidated (**Tigure 21.10b**). Mutations in *HPC1*, a gene for hereditary prostate cancer located in the long arm of chromosome 1, have been implicated in the origin of prostate tumors. Mutations in other tumor suppressor genes located in chromosomes 13, 16, 17, and 18 can transform prostate tumors into metastatic cancers, and overexpression of the *BCL-2* protooncogene gene can make these cancers immune to androgen deprivation therapy, a standard technique for the treatment of prostate cancer. The steroid hormone androgen is required for the proliferation of cells in the prostate epithelium. In the absence of androgen, these cells are programmed to die. However, prostate tumor cells may acquire the ability to survive in the absence of androgen, probably because an excess of the *BCL-2* gene product represses the programmed cell death pathway. Prostate cancers that have progressed to the stage of androgen independence are almost always fatal.

Douglas Hanahan and Robert Weinberg have proposed six hallmarks of the pathways leading to malignant cancer:

- **1.** *Cancer cells acquire self-sufficiency in the signalling processes that stimulate division and growth.* This self-sufficiency may arise from changes in the extracellular factors that cue cells to divide, or from changes in any part of the system that transduces these cues or translates their instructions into action inside the cell. In the most extreme case, self-sufficiency occurs when cells respond to growth factors that they themselves produce, thereby creating a positive feedback loop that stimulates ceaseless cell division.
- **2.** *Cancer cells are abnormally insensitive to signals that inhibit growth.* Cell division is stimulated by a variety of biochemical signals; however, other signals inhibit cell division. In normal cells, these countervailing factors balance each other with the result that growth occurs in a regulated manner. In cancer cells, growth is unregulated because the stimulatory signals have the upper hand. During the progression to malignancy, cancer cells lose their ability to respond appropriately to signals that inhibit growth. For example, cells in intestinal adenomas often no longer respond to TGF β , a protein that instructs pRB to block progression through the cell cycle. When this block fails, the cells advance from G₁ into S, replicate their DNA, and divide. These cells are then on their way to forming a malignant tumor.
- **3.** *Cancer cells can evade programmed cell death.* As we have seen, p53 plays a key role in protecting an organism from the accumulation of damaged cells that could endanger its life. Through mechanisms that are still incompletely understood, p53 sends damaged cells into an autodestruct pathway that clears them from the organism. When p53 malfunctions, this autodestruct pathway is blocked, and the damaged cells survive and multiply. Such cells are likely to produce descendants that are even more abnormal than they are. Consequently, lineages derived from damaged cells are prone to advance to a cancerous state. The ability to evade programmed cell death is therefore a key characteristic in the progression to malignant cancer.
- **4.** *Cancer cells acquire limitless replicative potential.* Normal cells are able to divide around 60 to 70 times. This limitation arises from the minute, but inexorable, loss of DNA from the ends of chromosomes every time the DNA is replicated (Chapter 10). The cumulative effect of this loss enforces a finite reproductive ability on every cell lineage. Cells that go past the reproductive limit become genetically unstable and die. Cancer cells manage to transcend this limit by replenishing their lost DNA. They do so by increasing the activity of the enzyme telomerase, which adds DNA sequences to the ends of chromosomes. When cells have acquired limitless replicative potential by overcoming the loss of DNA at the ends of chromosomes, they are said to be *immortalized*.
- **5.** *Cancer cells develop ways to nourish themselves.* Any tissue in a complex, multicellular organism needs a vascular system to bring nutrients to it. In humans and other vertebrate animals, the circulatory system provides this function. The cells in premalignant tumors fail to grow aggressively because they are not directly fed by the circulatory system. However, when blood vessels are induced to grow among these cells—through a process called *angiogenesis*—the tumor is nourished and can then expand. Thus, a key step in the progression to malignant cancer is the induction of blood vessel growth by the cells of the tumor. Many factors that induce or inhibit angiogenesis are known. In normal tissues, these factors are kept in balance so that blood vessels grow appropriately in the body; in cancerous tissues, the balance is tipped in favor of the inducing factors, which act to stimulate blood vessel development. Once capillaries have grown into a tumor, a reliable means of nourishment is at hand. The tumor can then feed itself and grow to a size where it becomes a danger to the organism.
- **6.** *Cancer cells acquire the ability to invade other tissues and colonize them.* More than 90 percent of all cancer deaths are caused by metastasis of the cancer to other parts of the body. When tumors metastasize, the cancer cells detach from the primary tumor and travel through the bloodstream to another location, where they estab-

lish a new, lasting, and, in the end, lethal, relationship with the surrounding cells. Profound changes must take place on the surfaces of the cancer cells for this process to occur. When it does, secondary tumors may develop in tissues far removed from the primary tumor. Cancers that have spread in this fashion are extremely difficult to control and eradicate. Metastasis is therefore the most serious occurrence in the progression of a cancer.

Numerous studies have established that somatic mutation is the basis for the development and progression of all types of cancer. As a cancer progresses on the pathway to malignancy, its cells become increasingly unregulated. Mutations accumulate, and whole chromosomes or chromosome segments may be lost. This genetic instability increases the likelihood that the cancer will develop each of the hallmarks discussed above.

Because of the importance of somatic mutations in the etiology of cancer, factors that increase the mutation rate are bound to increase the incidence of cancer. Today many countries maintain research programs to identify mutagenic and carcinogenic agents (see Chapter 13 for a discussion of the Ames test to identify chemical mutagens). When such agents are identified, public health authorities devise policies to minimize human exposure to them. However, no environment is carcinogen-free, and human behaviors that contribute to the risk of cancer such as smoking, excessive exposure to sunlight, and consumption of fatty foods that contain little fiber are difficult to change. Understanding of the processes that cause cancer has advanced significantly. In the future, we can expect this understanding to lead to more effective strategies for cancer prevention and treatment.

- Different types of cancer are associated with mutations in different genes.
- Cancer cells may stimulate their own growth and division.
- *Cancer cells do not respond to factors that inhibit cell growth.*
- Cancer cells can evade the natural mechanisms that kill abnormal cells.
- Immortalized cancer cells can divide endlessly.
- Tumors can expand when they induce the in-growth of blood vessels to nourish their cells.
- Metastatic cancer cells can invade other tissues and colonize them.

Basic Exercises

Illustrate Basic Genetic Analysis

1. Which cell-cycle checkpoint prevents a cell from replicating damaged DNA?

Answer: The *START* checkpoint in mid-G₁ of the cell cycle.

2. (a) In which class of genes do dominant gain-of-function mutations cause cancer? (b) In which class of genes do recessive loss-of-function mutations cause cancer?

Answer: (a) Oncogenes. (b) Tumor suppressor genes.

- 3. Why do some chromosomal rearrangements lead to cancer?
- **Answer:** The breakpoints of these rearrangements often juxtapose a cellular oncogene to a promoter that stimulates the vigorous expression of the oncogene. Overexpression of the gene product can lead to excessive cell division and growth.

- 4. Intestinal cancer occurs in individuals with inactivating mutations in the *APC* gene. Explain how it might also occur in individuals with mutations in the β -catenin gene.
- Answer: A mutation that specifically prevented β-catenin from binding to pAPC might lead to cancer. β-catenin that cannot bind to pAPC would be available to bind to the transcription factors that stimulate the expression of genes whose products promote cell division and growth.
- 5. Which tumor suppressor gene is most frequently mutated in human cancers?

Answer: *TP53*, the gene that encodes p53.

KEY POINTS

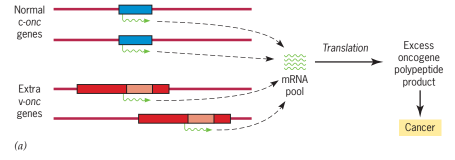
Testing Your Knowledge

Integrate Different Concepts and Techniques

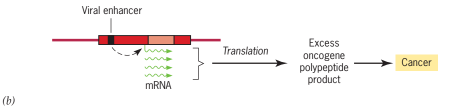
- 1. An oncogene within the genome of a retrovirus has a high probability of causing cancer, but an oncogene in its normal chromosomal position does not. If these two oncogenes encode exactly the same polypeptide, how can we explain their different properties?
- **Answer:** There are at least three possibilities. One (*a*) is that the virus simply adds extra copies of the oncogene to the cell and that collectively these produce too much of the polypeptide. An excess of polypeptide might cause uncontrolled cell division; that is, cancer. Another possibility (*b*) is that the viral oncogene is expressed inappropriately under the control of enhancers in the

viral DNA. These enhancers might trigger the oncogene to be expressed at the wrong time or to be overexpressed constitutively. In either case, the polypeptide would be inappropriately produced and might thereby upset the normal controls on cell division. A third possibility (c) is that integration of the virus into the chromosomes of the infected cell might put the viral oncogene in the vicinity of an enhancer in the chromosomal DNA and that this enhancer might elicit inappropriate expression. All three explanations stress the idea that the expression of an oncogene must be correctly regulated. Misexpression or overexpression could lead to uncontrolled cell division.

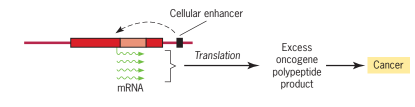
The virus adds extra copies of the oncogene to the cell







The viral oncogene is expressed inappropriately under the control of a cellular enhancer



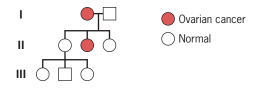
(c)

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **21.1** Many cancers seem to involve environmental factors. Why, then, is cancer called a genetic disease?
- **21.2** Both embryonic cells and cancer cells divide quickly. How can these two types of cells be distinguished from each other?
- **21.3** Most cancer cells are aneuploid. Suggest how aneuploidy might contribute to deregulation of the cell cycle.
- **21.4** Would you ever expect to find a tumor-inducing retrovirus that carried a processed cellular tumor suppressor gene in its genome?

- **21.5** How do we know that normal cellular oncogenes are not simply integrated retroviral oncogenes that have acquired the proper regulation?
- **21.6** How might the absence of introns in a retroviral oncogene explain that gene's overexpression in the tissues of an infected animal?
- **21.7** When cellular oncogenes are isolated from different animals and compared, the amino acid sequences of the polypeptides they encode are found to be very similar. What does this suggest about the functions of these polypeptides?
- **21.8** The majority of the *c-ras* oncogenes obtained from cancerous tissues have mutations in codon 12, 59, or 61 in the coding sequence. Suggest an explanation.
- **21.9** When a mutant *c-H-ras* oncogene with a valine for glycine substitution in codon 12 is transfected into cultured NIH 3T3 cells, it transforms those cells into cancer cells. When the same mutant oncogene is transfected into cultured embryonic cells, it does not transform them. Why?
- **21.10** A mutation in the *ras* cellular oncogene can cause cancer when it is in heterozygous condition, but a mutation in the *RB* tumor suppressor gene can cause cancer only when it is in homozygous condition. What does this difference between dominant and recessive mutations imply about the roles that the *ras* and *RB* gene products play in normal cellular activities?
- **21.11** Explain why individuals who develop nonhereditary retinoblastoma usually have tumors in only one eye, whereas individuals with hereditary retinoblastoma usually develop tumors in both eyes.
- **21.12** Approximately 5 percent of the individuals who inherit an inactivated RB gene do not develop retinoblastoma. Use this statistic to estimate the number of cell divisions that form the retinal tissues of the eye. Assume that the rate at which somatic mutations inactivate the RB gene is 1 mutation per 10⁶ cell divisions.
- **21.13** Inherited cancers like retinoblastoma show a dominant pattern of inheritance. However, the underlying genetic defect is a recessive loss-of-function mutation—often the result of a deletion. How can the dominant pattern of inheritance be reconciled with the recessive nature of the mutation?
- **21.14** The following pedigree shows the inheritance of familial ovarian cancer caused by a mutation in the *BRCA1* gene. Should II-1 be tested for the presence of the predisposing mutation? Discuss the advantages and disadvantages of testing.



- **21.15** In what sense is pRB a negative regulator of E2F transcription factors?
- 21.16 A particular E2F transcription factor recognizes the sequence TTTCGCGC in the promoter of its target gene. A temperature-sensitive mutation in the gene encoding this E2F transcription factor alters the ability of its protein product to activate transcription; at 25°C the mutant protein activates transcription normally, but at 35°C, it fails to activate transcription at all. However, the ability of the protein to recognize its target DNA sequence is not impaired at either temperature. Would cells heterozygous for this temperature-sensitive mutation be expected to divide normally at 25°C? at 35°C? Would your answers change if the E2F protein functions as a homodimer?
- 21.17 During the cell cycle, the p16 protein is an inhibitor of cyclin/CDK activity. Predict the phenotype of cells homozygous for a loss-of-function mutation in the gene that encodes p16. Would this gene be classified as a proto-oncogene or as a tumor suppressor gene?
- **21.18** The *BCL-2* gene encodes a protein that represses the pathway for programmed cell death. Predict the phenotype of cells heterozygous for a dominant activating mutation in this gene. Would the *BCL-2* gene be classified as a proto-oncogene or as a tumor suppressor gene?
- **21.19** The protein product of the *BAX* gene negatively regulates the protein product of the *BCL-2* gene—that is, BAX protein interferes with the function of the BCL-2 protein. Predict the phenotype of cells homozygous for a loss-of-function mutation in the *BAX* gene. Would this gene be classified as a proto-oncogene or as a tumor suppressor gene?
- **21.20** Cancer cells frequently are homozygous for loss-of-function mutations in the *TP53* gene, and many of these mutations map in the portion of *TP53* that encodes the DNA-binding domain of p53. Explain how these mutations contribute to the cancerous phenotype of the cells.
- **21.21** Suppose that a cell is heterozygous for a mutation that caused p53 to bind tightly and constitutively to the DNA of its target genes. How would this mutation affect the cell cycle? Would such a cell be expected to be more or less sensitive to the effects of ionizing radiation?
- **21.22** Mice homozygous for a knockout mutation of the *TP53* gene are viable. Would they be expected to be more or less sensitive to the killing effects of ionizing radiation?
- **21.23** Would cancer-causing mutations of the *APC* gene be expected to increase or decrease the ability of pAPC to bind β -catenin?
- 21.24
 Mice that are heterozygous for a knockout mutation in the *RB* gene develop pituitary and thyroid tumors. Mice that are homozygous for this mutation die during embryonic development. Mice that are homozygous for a knockout mutation in the gene encoding the p130 homologue of RB and heterozygous for a knockout mutation

in the gene encoding the p107 homologue of RB do not have a tendency to develop tumors. However, homozygotes for knockout mutations in both of these genes die during embryonic development. What do these findings suggest about the roles of the *RB*, *p139*, and *p107* genes in embryos and adults?

- **21.25** It has been demonstrated that individuals with diets poor in fiber and rich in fatty foods have an increased risk to develop colorectal cancer. Fiber-poor, fat-rich diets may irritate the epithelial lining of the large intestine. How could such irritation contribute to the increased risk for colorectal cancer?
- **21.26** Messenger RNA from the *KAI1* gene is strongly expressed in normal prostate tissues but weakly expressed in cell lines derived from metastatic prostate cancers. What does this finding suggest about the role of the KAI1 gene product in the etiology of prostate cancer?
- **21.27** The p21 protein is strongly expressed in cells that have been irradiated. Researchers have thought that this strong expression is elicited by transcriptional activation of the *p21* gene by the p53 protein acting as a transcription factor. Does this hypothesis fit with the observation that p21 expression is induced by radiation treatment in mice homozygous for a knockout mutation in the *TP53* gene? Explain.

Genomics on the Web at http://www.ncbi.nlm.nih.gov

The von Hippel-Lindau syndrome is characterized by the occurrence of cancer in the kidney. Often the *VHL* tumor suppressor gene has been mutated in this type of cancer.

- 1. Search the NCBI databases for information on the *VHL* gene. Where is it located in the genome? How long is its polypeptide product? Are different isoforms of the VHL protein created by alternate splicing?
- 2. The VHL protein physically interacts with other proteins inside cells. One interactant is the von Hippel-Lindau binding protein, VBP1. Search the databases for the gene encoding this protein. Where is this gene located? How long is the VBP1 polypeptide? How is this polypeptide thought to function inside cells?
- **3.** The VHL protein plays a role in biochemical pathways inside cells. Find the Pathways section on the *VHL* page and click on KEGG pathway: renal cell carcinoma to see where the VHL protein functions. What is its role in renal cells? What proteins does it interact with?
- 4. Homologues of the *VHL* gene exist in the genomes of the rat and mouse. Use the Map Viewer function under the Homology section on the *VHL* page to locate these homologues. What chromosomes are they on? Is the region around these homologues similar in all three organisms—rat, mouse, and human? What does the structure of this chromosomal region in these three organisms suggest about the evolutionary process?

Inheritance of Complex Traits



CHAPTER OUTLINE

- Complex Traits
- Statistics of Quantitative Genetics
- Analysis of Quantitative Traits
- Correlations between Relatives
- Quantitative Genetics of Human Behavioral Traits



Color-enhanced angiogram of the heart showing narrowing in one of the coronary arteries (center left). If left untreated, this situation can lead to a heart attack.

Cardiovascular Disease: A Combination of Genetic and Environmental Factors

Near the end of December, Paul Reston, a 47-year-old biology teacher in a suburban high school outside Pittsburgh, Pennsylvania, was spending his Saturday morning grading examinations. He was somewhat tired that day and felt a bit of stomach distress. He also had a slight pain in his left arm and shoulder. These symptoms had persisted for a few days. At first, Mr. Reston thought he had a mild case of the flu, but the arm and shoulder pain suggested another possibility: that he was having a heart attack. This possibility seemed more real when he remembered that his father had died from a sudden heart attack many years earlier at the relatively young age of 45. After a telephone conversation with a nurse in his health care clinic, Mr. Reston had his son drive him to a nearby hospital, where he spent two hours in the emergency room. The attending physician gave Mr. Reston a battery of tests to evaluate his condition. His heartbeat was regular, his blood pressure was normal, and an electrocardiogram revealed no abnormalities. Biochemical tests for telltale signs of heart damage were also negative. In addition, except for a family history of heart disease, Mr. Reston did not present other major risk factors. He was not overweight, he did not smoke, and he exercised regularly. The physician released Mr. Reston but advised him to return to the hospital the following week for a cardiac stress test. The following Monday, he was tested for heart function while running on a treadmill. The test results were good. Based on his performance, the supervising cardiologist concluded that Mr. Reston had less than a 1 percent chance of suffering a fatal heart attack.

In spite of his family history of heart disease, Mr. Reston's risk to develop this disease was low. The cardiologist explained that heart disease is a complex trait influenced by many factors: diet, physical activity, and smoking, for example, as well as a fairly large number of genes. Because Mr. Reston's father had succumbed to a heart attack, Mr. Reston may have inherited genes that put him at risk. However, the cardiologist emphasized that heart disease is not inherited as a simple Mendelian trait; rather, it involves the interplay of many different genetic and environmental factors.

Complex Traits

Breeding experiments and comparisons between relatives reveal that complex phenotypes may be influenced by a combination of genetic and environmental factors.

Many traits, such as disease susceptibility, body size, and various aspects of behavior, do not show simple patterns of inheritance. Nonetheless, we know that genes influence these types of traits. One indication is that genetically related individuals resemble one another. We see these resemblances between siblings, between parents and offspring, and sometimes between more distant relatives. The extreme case is monozygotic twins—twins that have

developed from a single fertilized egg. Such twins are often strikingly similar, in behavior as well as in appearance. Another indication for a genetic influence is that these types of traits respond to selective breeding. In agriculture, crops and livestock have been shaped by propagating individuals with desirable features—greater protein content, reduced body fat, greater productivity, resistance to disease, and so forth. This ability to change phenotypes through selective breeding indicates that the traits have a genetic basis. Usually, however, this genetic basis is complex. Several to many genes are involved, and their individual effects are difficult to discern through conventional genetic analysis. Consequently, other techniques are needed to study the inheritance of complex traits.

QUANTIFYING COMPLEX TRAITS

Many complex traits vary continuously in a population. One phenotype seems to blend imperceptibly into the next. Examples are body size, height, weight, enzyme activity, blood pressure, and reproductive ability. The phenotypic variation in these types of traits can be quantified by measuring the trait in a sample of individuals from the population. We might, for example, capture mice in a barn and weigh each of them, or we might collect corncobs from a field and count the number of kernels on each. With such a quantitative approach, the phenotype of every individual in the sample is reduced to a number. These numbers can be analyzed with a variety of statistical techniques, enabling us to study the trait and, ultimately, to investigate its genetic basis. Traits that are amenable to this kind of treatment are called **quantitative traits.** Their essential characteristic is that they can be measured.

GENETIC AND ENVIRONMENTAL FACTORS INFLUENCE QUANTITATIVE TRAITS

The Danish biologist Wilhelm Johannsen was one of the first people to show that variation in a quantitative trait is due to a combination of genetic and environmental factors. Johannsen studied the weight of seeds from the broad bean, *Phaseolus vulgaris*. Among the plants available to him, seed weight varied from 150 mg to 900 mg. Johannsen established lines from individual seeds across this range and maintained each line by self-fertilization for several generations. The seeds from each of these "pure" lines tended to resemble the seed from which they were founded. This ability to establish lines of beans with characteristically different seed weights indicated that some variation in this trait is due to genetic differences. However, Johannsen observed that seed weight also varied within each of the pure lines. This residual variation was not likely to be due to genetic differences because each line had been systematically inbred to make it homozygous for its genes. Rather, it must have been due to variation in uncontrolled factors in the environment. Johannsen's work, published in 1903 and 1909, therefore led to the realization that phenotypic variation in a quantitative trait has two components—one genetic, the other environmental.

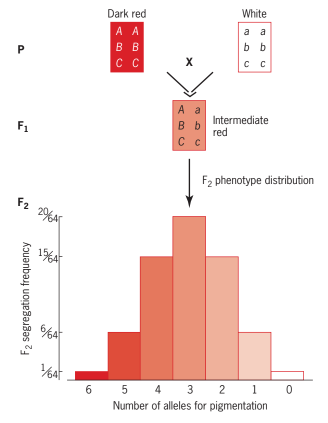
MULTIPLE GENES INFLUENCE QUANTITATIVE TRAITS

Another Scandinavian, Herman Nilsson-Ehle, provided evidence that the genetic component of this variation could involve the contributions of several different genes.

Nilsson-Ehle studied color variation in wheat grains. When he crossed a white-grained variety with a dark red-grained variety, he obtained an F_1 with an intermediate red phenotype (**Figure 22.1**). Self-fertilization of the F_1 produced an F_2 with seven distinct classes, ranging from white to dark red. The number of F2 classes and the phenotypic ratio that Nilsson-Ehle observed suggested that three independently assorting genes were involved in the determination of grain color. Nilsson-Ehle hypothesized that each gene had two alleles, one causing red grain color and the other white grain color, and that the alleles for red grain color contributed to pigment intensity in an additive fashion. Based on this hypothesis, the genotype of the white-grained parent could be represented as *aa bb cc*, and the genotype of the red-grained parent could be represented as AA BB CC. The F₁ genotype would be Aa Bb Cc, and the F_2 would contain an array of genotypes that would differ in the number of pigment-contributing alleles present. Each phenotypic class in the F_2 would carry a different number of these pigmentcontributing alleles. The white class, for example, would carry none, the intermediate red class would carry three, and the dark red class would carry six. Nilsson-Ehle's work, published in 1909, showed that a complex inheritance pattern could be explained by the segregation and assortment of multiple genes.

The American geneticist Edward M. East extended Nilsson-Ehle's studies to a trait that did not show simple Mendelian ratios in the F_2 . East studied the length of the corolla in tobacco flowers (**I** Figure 22.2a). In one pure line, the corolla length averaged 41 mm; in another, it averaged 93 mm. Within each pure line, East observed some phenotypic variation—presumably the result of environmental influences (**I** Figure 22.2b). By

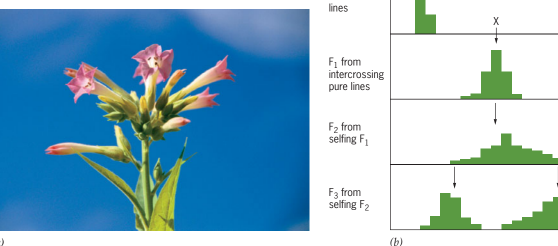
crossing the two lines, East obtained an F_1 that had intermediate corolla length and approximately the same amount of variation that he had seen within each of the parental strains. When East intercrossed the F_1 plants, he obtained an F_2 with about the same corolla length, on average, that he saw in the F_1 ; however, the F_2 plants were much more variable than the F_1 . This variability was due to two sources: (1) the segregation and independent assortment of different pairs of alleles controlling corolla length, and (2) environmental factors. East inbred some of the F_2 plants to produce an F_3 and observed less variation within the different F_3 lines than in the F_2 . The reduced



■ **FIGURE 22.1** Inheritance of grain color in wheat. Three independently assorting genes (*A*, *B*, and *C*) are assumed to control grain color. Each gene has two alleles. The alleles that contribute additively to pigmentation are represented by uppercase letters.

Sources of variation

amount of variation within the F_3 lines was presumably due to the segregation of fewer allelic differences. Thus, the complex inheritance pattern that East observed with corolla length could be explained by a combination of genetic segregation and environmental influences.



Pure lines F_1 from intercrossing pure lines F_2 from selfing F_1 F_3 from selfing F_2 F_2 from selfing F_2 F_2 from selfing F_2

Corolla length (mm)

(a)

FIGURE 22.2 Corolla length as a quantitative trait. (*a*) Tobacco flowers showing the long corolla. (*b*) Inheritance of corolla length in tobacco. At least five genes appear to be involved.

How many genes were involved in determining corolla length in East's strains of tobacco? We can make a crude guess by comparing the F, plants with each of the inbred parental strains. Let's suppose that the strain with the shorter corollas was homozygous for one set of alleles and that the strain with the longer corollas was homozygous for another set of alleles. Furthermore, let's suppose that the long-corolla alleles act additively, that all length-controlling genes assort independently, and that each gene makes an equal contribution to the phenotype. If corolla length were determined by one gene, with alleles a (for short corolla) and A (for long corolla), we would expect 1/4 of the F, plants to have short corollas (like the short parental strain) and 1/4 to have long corollas (like the long parental strain). If two genes determined corolla length, we would expect 1/16 of the F, plants to resemble the short-corolla parent and 1/16 to resemble the long-corolla parent. If three genes were involved, the frequency of each parental type in the F_2 would be 1/64, and if four genes were involved, it would be 1/256. With five genes, the parental frequencies in the F, would each be 1/1024. East studied 444 F, plants and failed to find even one with either of the parental phenotypes. This failure would seem to rule out the hypothesis of four or fewer genes controlling corolla length. Thus, we can conclude that at least five genes are responsible for the difference in corolla length between East's two inbred strains.

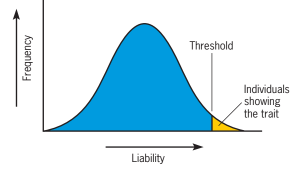
THRESHOLD TRAITS

Continuously varying traits such as bean size, grain color, and corolla length are controlled by multiple factors, both genetic and environmental. Geneticists have found that some traits that do not vary continuously in the population also appear to be influenced by multiple factors. For example, many people develop heart disease in their fifth or sixth decade of life. Heart disease is not a quantitative trait in the usual sense; individuals either have it or they don't. However, many factors predispose an individual to develop heart disease: body weight, amount of exercise, diet, blood cholesterol level, whether or not the individual smokes, and the presence of heart disease in close relatives such as parents or siblings. These underlying risk factors contribute to a variable called the *liability*. Geneticists theorize that when the liability exceeds a certain level, or threshold, the trait appears. This type of trait is therefore called a **threshold trait** (**Teigure 22.3**).

In humans, the evidence that threshold traits are influenced by genetic factors comes from comparisons between relatives, especially twins. Occasionally a fertilized human egg splits and forms two genetically identical zygotes. The individuals who develop from these zygotes are referred to as one-egg, or **monozygotic (MZ)**, twins; they share 100 percent of their genes. More frequently, two independently fertilized eggs

develop at the same time in the mother's womb. These two-egg, or **dizygotic** (DZ), twins are as closely related as ordinary siblings; thus, they share 50 percent of their genes. Because of their genetic identity, we would expect MZ twins to be phenotypically more similar than DZ twins.

Similarity with respect to a threshold trait is assessed by determining the **concordance rate**—the fraction of twin pairs in which both twins show the trait among pairs in which at least one of them does. For cleft lip, a congenital condition due to an error in embryological development, the concordance rate has been estimated to be about 40 percent for MZ twins and about 4 percent for DZ twins. The much greater concordance rate for MZ twins strongly suggests that genetic factors influence an individual's likelihood of being born with cleft lip. Mental illnesses such as schizophrenia and bipolar disorder can also be regarded as threshold traits. For schizophrenia, the concordance rate ranges from 30 to 60 percent for MZ twins and from 6 to 18 percent for DZ twins; for bipolar disorder, the concordance rate is 70–80 percent for MZ twins and about 20 percent for DZ twins. Thus, twin studies suggest that both of these mental illnesses are influenced by genetic factors.



■ FIGURE 22.3 A model for expression of a threshold trait. When the underlying variable, the liability, reaches a threshold value, the trait is expressed. This variable is assumed to be continuously distributed in the population.

- Resemblances between relatives and responses to selective breeding indicate that complex traits
 have a genetic basis.
- Some complex traits can be quantified to permit genetic analysis.
- Many genetic and environmental factors influence the variation observed in quantitative traits.
- Phenotypic segregations may provide a way to estimate the number of genes that influence a quantitative trait.
- Traits that are manifested when an underlying continuous variable (the liability) reaches a threshold value may be influenced by genetic factors.
- In humans, evidence that a threshold trait has a genetic basis comes from studies with twins.
- The concordance rate is the fraction of twin pairs in which both twins show a trait among pairs in which at least one of them does.

KEY POINTS

Statistics of Quantitative Genetics

The hallmark of quantitative traits is that they vary continuously in a population of individuals. This type of variation poses a formidable problem for the geneticist. Segregation ratios are difficult, if not impossible,

to discern because the number of phenotypes is large and one phenotype blends imperceptibly into the next. For quantitatively varying traits, routine genetic analyses of the sort that we have done with eye color in *Drosophila* and with human disorders such as albinism are out of the question. For these types of traits we must resort to a different kind of analysis, one that is based on statistical descriptions of the phenotype in a population. In the sections that follow, we introduce the basic statistical concepts that are needed for this type of analysis.

FREQUENCY DISTRIBUTIONS

The first step in the study of any quantitative trait is to collect measurements of the trait from individuals in a population. Usually, only a small fraction of all the individuals in the population can be measured. We call this group the **sample**. The data from the sample can be presented graphically as a **frequency distribution**. In the graph the horizontal or *x*-axis measures values of the trait. This axis is divided into regular intervals that allow each individual in the population to be categorized for the trait. Thus, each observation in the sample can be placed into one of the intervals on the *x*-axis. The vertical or *y*-axis measures the frequency of the observations within each interval.

Figure 22.4 shows frequency distributions that were obtained in a genetic study of wheat. The investigators measured the time that wheat takes to mature. Four different populations of wheat were grown in test plots in the same season, and 40 plants from each population were monitored until the heads of grain matured. The time to maturity for each plant was recorded in days. Two of the populations (A and B) were inbred strains, and one was an F_1 produced by crossing these two strains. The fourth population was an F_2 produced by intercrossing the F_1 plants.

The two parental strains A and B were highly inbred varieties that were completely or almost completely homozygous. As the frequency distributions indicate, strain A matured quickly and strain B matured slowly. The lack of phenotypic overlap between the samples from these two strains demonstrates their genetic distinctiveness. Apparently, strains A and B were homozygous for different alleles of genes controlling maturation time. Within each strain, however, there was still some phenotypic variation, presumably the result of microenvironmental differences within the test plots.

The distributions of the F_1 and F_2 samples indicate that these populations had intermediate maturation times. Their intermediate position on the *x*-axis suggests

The frequency distributions of quantitative traits can be characterized by summary statistics.

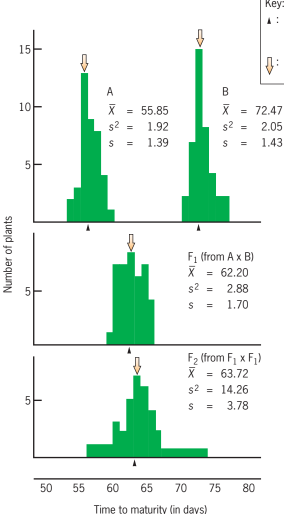


FIGURE 22.4 Frequency distributions and descriptive statistics of time to maturity in four populations of wheat. A and B are inbred strains that were crossed to produce F_1 hybrids. The F_1 plants were intercrossed to produce an F_2 . Seed from all four populations was planted in the same season to determine the time to maturity. In each case, data were obtained from 40 plants. The mean (\overline{X}) , mode, variance (s^2) , and standard deviation (s) are given.



that the alleles controlling maturation time contribute additively to the trait. Notice that the distribution of the F_2 sample is considerably broader than that of the F_1 . The additional variability seen in the F_2 population reflects the genetic segregation that occurred when the F_1 plants reproduced. We now explore ways in which quantitative geneticists summarize the data in a frequency distribution.

THE MEAN AND THE MODAL CLASS

The essential characteristics of a frequency distribution can be summarized by simple statistics calculated from the data. One of these summary statistics is called the **mean** or average. It gives us the "center" of the distribution—the "typical" value. We calculate the sample mean (\overline{X}) by summing all the data in the sample and dividing by the total number of observations (*n*). In mathematical notation, the mean is:

$$X = (\Sigma X_k)/n$$

The Greek letter Σ in this formula is a mathematical shorthand for the sum of all the individual measurements in the sample; thus, $\Sigma X_i = (X_1 + X_2 + X_3 + ... X_n)$, where X_k represents the *k*th of the *n* individual observations. In Figure 22.4 the positions of the sample means are indicated by triangles beneath the distributions; the numerical values of these means are given on the right. The means of the F₁ and F₂ samples are 62.20 and 63.72 days, respectively; both are a little less than the average of the means of the two inbred parental strains (64.16 days).

The **modal class** in a sample is the class that contains the most observations. Like the mean, it also captures the "center" of the distribution. In Figure 22.4 the modal classes are indicated by short arrows. We see that in each of the distributions the mean is within or very close to the modal class. This coincidence reflects the symmetry of the distributions; in each case, roughly equal numbers of observations are above and below the mean and the modal class. Not all distributions have this feature. Some are skewed, with

most of the observations clustered at one end and only a few at the other end forming a long tail. Statisticians have developed an extensive theory about a particular type of symmetrical distribution called a *normal distribution* (**■** Figure 22.5). In this bell-shaped distribution, the mean and the modal class are located exactly in the center. Often distributions of sample data approximate the shape of a normal distribution. Thus, we can apply the extensive theory about normal distributions to analyze such data.

THE VARIANCE AND THE STANDARD DEVIATION

The data in a frequency distribution could be dispersed, or they could be clustered. To measure the spread of data in a frequency distribution, we use a statistic called the **variance**. Data that are widely dispersed produce a large value for the variance, whereas data that are tightly clustered produce a small value. The sample variance, denoted s^2 , is calculated from the formula

$$s^2 = \sum (X_{\mu} - \overline{X})^2 / (n-1)$$

In this formula, $(X_k - \overline{X})^2$ is the squared difference between the *k*th observation and the sample mean (often called the *squared deviation from the mean*), and the Greek letter Σ indicates that all such squared deviations are summed. The sum of the squared deviations is averaged by dividing by n - 1. (For technical reasons, the divisor is one less than the sample size.) The exponent 2 in the symbol s^2 is a reminder that we have used squared differences in calculating the sample variance. We should note two features of the variance. First, it measures the dispersion of the data around the *mean*. When we calculate the variance, we take the mean to be the central value of the distribution and find the difference between it and each of the observations in the sample. Second, the variance is always positive. When we calculate the variance, we square the difference between each observation and the mean, and then sum the squared differences. Because each of the squared differences is positive, the variance, calculated by summing these squared differences, is also positive.

Although the variance has desirable mathematical properties, it is difficult to interpret because the units of measurement are squared (for example, $s^2 = 2.88 \text{ days}^2$). Consequently, another statistic, called the **standard deviation**, is often used to describe the variability of a sample. The standard deviation (*s*) is the square root of the sample variance

$$s = \sqrt{s^2}$$

This statistic is easier to interpret than the variance because it is expressed in the same units as the original measurements.

The variances and standard deviations of the four wheat populations are given in Figure 22.4. The F_2 population has the greatest variance and standard deviation, no doubt because it is segregating for genes that control maturation time. In the F_2 plants, both genetic and environmental differences produce the observed variability. In the other populations, most if not all of the observed variation is due to environmental factors alone. Each of the two parental strains is highly inbred and is therefore expected to be homozygous for most of its genes. The F_1 plants are heterozygous for the alleles that are different in the two parental strains, but they all have the same genotype. Thus, in neither the parental strains nor the F_1 do we expect to find much genetic variation among plants. In a later section we will see how to estimate that part of the variance in a quantitative trait that is due to genetic differences among individuals in a population.

As mentioned above, the distribution of a quantitative trait often looks like a normal distribution. The shape and the position of a normal distribution are completely specified by its mean and standard deviation. Thus, if we know only the mean and standard deviation of a quantitative trait, and assume that the trait is normally distributed, we can construct the approximate shape of the trait's distribution. In this distribution, 66 percent of the measurements will lie within one standard deviation of the mean, 95 percent will lie within two standard deviations of the mean, and 99 percent will lie within three standard deviations of the mean (Figure 22.5).

- The mean $(\overline{X} = (\Sigma X_{k})/n)$ and modal class point to the center of a frequency distribution.
- The variance $(s^2 = \Sigma (X_k \overline{X})^2/(n 1)$ and standard deviation $s = \sqrt{s^2}$ are statistics that indicate the extent to which data are scattered around the mean in a frequency distribution.

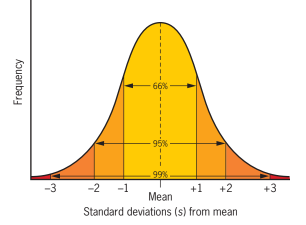


FIGURE 22.5 A normal frequency distribution showing the percentage of measurements within 1, 2, and 3 standard deviations of the mean.

KEY POINTS

Analysis of Quantitative Traits

In this section we will see how statistics are used in the genetic analysis of quantitative traits. The thrust of the analysis is to partition the observed variation in the trait into genetic and environmental components, Quantitative geneticists focus their analyses on phenotypic variability as measured by the variance.

and then to use the genetic component to make predictions about the phenotypes of the offspring of particular crosses.

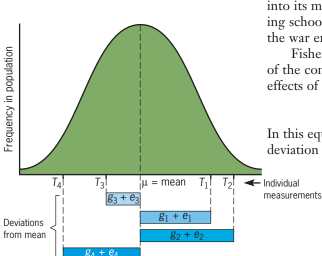


FIGURE 22.6 Quantitative phenotypes and the deviations of individual measurements from the population mean. Each individual's deviation is hypothesized to consist of a deviation due to its genotype (g) and a deviation due to its environment (e).

THE MULTIPLE FACTOR HYPOTHESIS

The key idea in quantitative genetics is that traits are controlled by many different factors in the environment and in the genotype. This **Multiple Factor Hypothesis** emerged in the second decade of the twentieth century through the experimental investigations of E. M. East, W. Johannsen, H. Nilsson-Ehle, and others. However, it was a theoretician, R. A. Fisher, who crystallized the Multiple Factor Hypothesis into its modern form. Fisher did this work during World War I while he was teaching school in Great Britain. His theoretical analysis was published in 1918, the year the war ended.

Fisher hypothesized that a particular value of a quantitative trait, T, is the result of the combined influence of genetic and environmental factors. He represented the effects of these factors as deviations from the overall population mean:

$$T = \mu + g + e$$

In this equation, the Greek letter μ represents the population mean, *g* represents the deviation from the mean that is due to genetic factors, and *e* represents the deviation

from the mean that is due to environmental factors. In Fisher's scheme, the position of a particular value of the trait, T, in the population depends on the genetic and environmental factors that have affected it (**Figure 22.6**). Some factors produce large values of T, and some produce small values of T. For each individual, these factors are different. Furthermore, Fisher emphasized that a multitude of factors are involved. He hypothesized that many genes contribute to a quantitative trait, and he assumed that many aspects of the environment also make contributions. Today we say that a trait that is controlled by many genes is **polygenic**.

PARTITIONING THE PHENOTYPIC VARIANCE

With these simple ideas, Fisher was able to develop a procedure to analyze the variability of a quantitative trait in terms of the contributing genetic and environmental factors. To measure the variability of the trait, he focused on the statistic we have called the variance. Specifically, he discovered how to split the overall variance of the trait into two component variances, one measuring the effects of genetic differences among individuals and the other measuring the effects of environmental differences. Thus, in Fisher's analysis, the variance of a quantitative trait, symbolized V_{τ} , is equal to the sum of a genetic variance, symbolized V_{e} , and an environmental variance, symbolized V_{e} :

$$V_T = V_g + V_e$$

In this variance equation, the variance of the quantitative trait, V_T , is often referred to as the *total phenotypic variance*.

A discussion of Fisher's method of splitting the total phenotypic variance into its genetic and environmental components is beyond the scope of this book. However, this method has since been used in many different contexts and has given rise to a general statistical technique called *analysis of variance*.

To see the basic idea, let's partition the variance of maturation time in the F_2 population of wheat shown in Figure 22.4. The total phenotypic variance of this population (V_T) is 14.26 days². In terms of Fisher's variance equation, this total can be represented as the sum of a genetic variance (V_g) and an environmental variance (V_e) , bloth of which must be estimated using other data. To estimate the environmental variance, we can use the data from the parental and F_1 populations. The parental populations are genetically uniform because they are both inbred. The F_1 population is also genetically uniform because it was created by crossing the two inbred populations; every F_1 plant is expected to be identically heterozygous for the genes that differ in the inbred parental populations. Because of this genetic uniformity, the variability that we see in each of these three populations must reflect differences due to environmental effects.

To obtain a representative value for V_{ϕ} we can average the variances of these groups:

$$V_e = (V_A + V_B + V_{FI})/3$$

= (1.92 days² + 2.05 days² + 2.88 days²)/3
= 2.28 days²

With this estimate of the environmental variance, we can now estimate V_g by subtraction from the total variance V_T :

$$V_g = V_T - V_e$$

= 14.26 days² - 2.28 days²
= 11.98 days²

Thus, the total phenotypic variance for maturation time in the F_2 wheat population has been split into two components:

$$V_T = V_g + V_e$$

14.26 days² = 11.98 days² + 2.28 days²

From this partition, we see that most of the variance in maturation time in the F_2 wheat population is due to genetic differences among the individuals. This genetic variability arose from the segregation and assortment of genes when the F_1 plants reproduced. These plants were heterozygous for the genes that differed in the parental populations. When they reproduced, segregation and assortment produced an array of genotypes—three distinct genotypes for each heterozygous gene. The variation that we see in the F_2 is due primarily to phenotypic differences among these genotypes. To reinforce your understanding of how the total phenotypic variance is partitioned into genetic and environmental components, work through Solve It: Estimating Genetic and Environmental Variance Components.

BROAD-SENSE HERITABILITY

Often it is informative to calculate the proportion of the total phenotypic variance that is due to genetic differences among individuals in a population. This proportion is called the **broad-sense heritability**, symbolized H^2 . In terms of Fisher's variance components,

$$\begin{split} H^2 &= V_g/V_T \\ &= V_g/(V_g + V_e) \end{split}$$

The symbol for the broad-sense heritability, H^2 , is written with the exponent 2 to remind us that this statistic is calculated from variances, which are squared quantities.

Because of the way it is calculated, the broad-sense heritability must lie between 0 and 1. If it is close to 0, little of the observed variability in the population is attributable to genetic differences among individuals. If it is close to 1, most of the observed variability is attributable to genetic differences. The broad-sense heritability therefore summarizes the relative contributions of genetic and environmental factors to the observed variability in a population. However, it is important to note that this statistic is population-specific. For a given trait, different populations may have different values of the broad-sense heritability. Thus, the broad-sense heritability of one population cannot automatically be assumed to represent the broad-sense heritability of another population.

In the F_2 wheat population, $H^2 = 11.98/14.26 = 0.84$. This result tells us that in this population 84 percent of the observed variability in wheat maturation time is due to genetic differences among individuals. However, it does not tell us what these differences are. The genetic variance upon which the broad-sense heritability depends includes all

Solve It!

Estimating Genetic and Environmental Variance Components

E. M. East studied variation in the length of flowers in two inbred strains of tobacco plants and in the F_1 and F_2 populations derived from crosses between these strains:

Population	Mean length (mm)	Variance (mm²)
Inbred 1	41	6
Inbred 2	93	7
F₁ hybrids	63	8
F_2 from $F_1 \times F_1$	68	43

From these data, estimate the genetic and environmental components of variance in the F_2 population.

• To see the solution to this problem, visit the Student Companion site.

the factors that cause genotypes to have different phenotypes: the effects of individual alleles, the dominance relationships between alleles, and the epistatic interactions among different genes. In Chapter 4 we saw how these factors influence phenotypes. In the next two sections, we will see that by breaking out these components of genetic variability and by focusing on the component that involves the effects of individual alleles, we can predict the phenotypes of offspring from the phenotypes of their parents.

NARROW-SENSE HERITABILITY

The ability to make predictions in quantitative genetics depends on the amount of genetic variation that is due to the effects of individual alleles. Genetic variation that is due to the effects of dominance and epistasis has little predictive power.

To see how dominance limits the ability to make predictions, consider the ABO blood types in humans (Table 4.1 in Chapter 4). This trait is determined strictly by the genotype; environmental variation has essentially no effect on the phenotype. However, because of dominance, two individuals with the same phenotype can have different genotypes. For example, a person with type A blood could be either $I^A I^A$ or $I^A i$. If two people with type A blood produce a child, we cannot predict precisely what phenotype the child will have. It could be either type A or type O, depending on the genotypes of the parents; however, we know that it will not have type B or type AB blood. Thus, although we can make some kind of prediction about the child's phenotype, dominance prevents us from making a precise prediction.

Our ability to make predictions about an offspring's phenotype is improved in situations where the genotypes are not confused by dominance. Consider, for example, the inheritance of flower color in the snapdragon, Antirrhinum majus. Flowers in this plant are white, red, or pink, depending on the genotype (Figure 4.1 in Chapter 4). As with the ABO blood types, variation in flower color has essentially no environmental component; all the variance is the result of genetic differences. However, for the flower color trait, the genotype of an individual is not obscured by the complete dominance of one allele over the other. A plant with two w alleles has white flowers, a plant with one w allele and one W allele has pink flowers, and a plant with two W alleles has red flowers. In this system, the phenotype depends simply on the number of W alleles present; each Wallele intensifies the color by a fixed amount. Thus, we can say that the color-determining alleles contribute to the phenotype in a strictly *additive* fashion. This kind of allele action improves our ability to make predictions in crosses between different plants. A mating between two red plants produces only red offspring; a mating between two white plants produces only white offspring; and a mating between red and white plants produces only pink offspring. The only uncertainty is in a cross involving heterozygotes, and in this case the uncertainty is due to Mendelian segregation, not to dominance.

Quantitative geneticists distinguish between genetic variance that is due to alleles that act additively (such as those in the flower color example just discussed) and genetic variance that is due to dominance. These different variance components are symbolized as:

> V_a = additive genetic variance V_d = dominance variance

In addition, geneticists define a third variance component that measures variation due to epistatic interactions between alleles of different genes:

$$V_i$$
 = epistatic variance

Epistatic interactions, like dominance, are of little help in predicting phenotypes. Altogether, these three variance components constitute the total genetic variance:

$$V_g = V_a + V_d + V$$

If we recall that $V_T = V_g + V_e$, we can express the total phenotypic variance as the sum of four components:

$$V_T = V_a + V_d + V_i + V$$

Of these four variance components, only the additive genetic variance, V_a , is useful in predicting the phenotypes of offspring from the phenotypes of their parents. This variance, as a fraction of the total phenotypic variance, is called the **narrow-sense** heritability, symbolized b^2 . Thus,

$$b^{2} = V_{a}/V_{7}$$

Like the broad-sense heritability, b^2 lies between 0 and 1. The closer it is to one, the greater is the proportion of the total phenotypic variance that is additive genetic variance, and the greater is our ability to predict an offspring's phenotype. Table 22.1 gives some estimates for the narrow-sense heritability for several traits. Human stature is highly heritable, but litter size in pigs is not. Thus, if we knew the parental phenotypes, we would be better able to predict the height of a human's offspring than the litter size of a pig's offspring.

PREDICTING PHENOTYPES

To gain insight into the meaning of the narrow-sense heritability, let's consider the situation diagrammed in Figure 22.7. Michael (M) and Frances (F) have taken a standardized intelligence test, and their Intelligence Quotients (IQs) have been determined. Michael's score is 110 and Frances's score is 120. The mean IQ score in the population is 100. Michael and Frances had an infant son Oswald (O), who was given up for adoption when he was born, and the adoptive parents wish to predict Oswald's IQ. If IQ had no genetic component, our best estimate for Oswald's IQ would be 100, the mean of the population. We would have no way of predicting what kind of home environment Oswald will receive and therefore cannot predict what kind of nongenetic factors will influence his mental development. Nor could we use the IQs of Michael and Frances to predict anything about Oswald's IQ, since, by assumption, the genes they gave to him would have nothing to do with mental development. However, several studies have indicated that variation in IQ scores does have a genetic component. In fact, the narrow-sense heritability of IQ has been estimated to be about 0.4—that is, about 40 percent of the observed variation in IQ scores is due to the additive effects of alleles. Can we use this statistic along with the parental IQs to predict Oswald's IQ score?

Let's symbolize the IQs of Oswald, Michael, and Frances as T_0 , T_M , and T_F , respectively, and let's symbolize the population mean as μ . The best prediction for Oswald's IQ is

$$T_o = \mu + b^2 [(T_M + T_F)/2 - \mu]$$

The expression with parentheses, $(T_M + T_F)/2$, is usually called the *midparent value*. It is the average of the phenotypes of the two parents. If we denote the midparent value with the symbol T_P , the prediction equation for Oswald's phenotype simplifies to

$$T_O = \mu + b^2 [T_P - \mu]$$

The expression in brackets, $[T_p - \mu]$, is the difference between the midparent value and the mean of the population. The product of this difference and the narrow-sense heritability is the predicted deviation of the offspring's phenotype from the mean of the population. In effect, the narrow-sense heritability translates the difference between the midparent value and the mean of the population into a "heritable" difference that we can expect to see in the offspring. By adding this heritable difference to the mean, we can predict the offspring's phenotype.

Let's now substitute the known quantities for each of the terms in the prediction equation: $\mu = 100$, $T_P = (110 + 120)/2 = 115$, and $b^2 = 0.4$. Thus, the predicted value of T_O is

$$T_o = 100 + (0.4)[115 - 100]$$

= 106

This result tells us that Oswald's IQ is expected to be between the midparent value (115) and the mean of the population (100). In fact, it is at a point 40 percent

TABLE 22.1

Estimates of Narrow-Sense Heritability (*h*²) for Quantitative Traits

Trait	h^2
Stature in human beings	0.65
Milk yield in dairy cattle	0.35
Litter size in pigs	0.05
Egg production in poultry	0.10
Tail length in mice	0.40
Body size in <i>Drosophila</i>	0.40

Source: D. S. Falconer. 1981. *Introduction to Quantitative Genetics*, 2nd ed., p. 51. Longman, London.

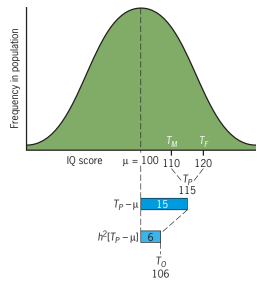


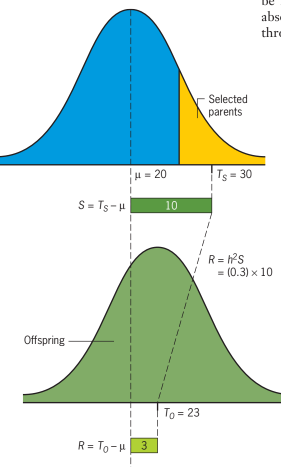
FIGURE 22.7 Predicting an offspring's phenotype based on the phenotypes of its parents and the narrow-sense heritability of the trait. Only a portion of the deviation of the midparent (T_p) value from the population mean is heritable. The magnitude of this portion is determined by the narrow-sense heritability.

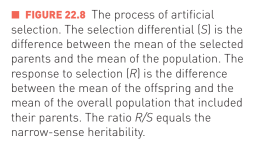
Solve It!

Using the Narrow-Sense Heritability

Linda and William have IQ scores of 120 and 90, respectively. In the general population, the mean IQ is 100 and the narrowsense heritability is estimated to be 0.4. If Linda and William have a child and the child is reared in an average environment, what is the child's predicted IQ?

• To see the solution to this problem, visit the Student Companion site.





of the distance between the population mean and the midparent value. This 40 percent corresponds to the narrow-sense heritability (0.4). If the narrow-sense heritability of IQ were greater than 0.4, the predicted value of Oswald's IQ would be closer to the midparent value. For a perfectly heritable trait, $b^2 = 1$ and the predicted value of the offspring's phenotype would equal the average of the two parents' phenotypes. Thus, the narrow-sense heritability is a critical statistic. It tells us how closely the offspring will resemble the average of their parents. We should emphasize, however, that the IQ score we have calculated for Oswald is a predicted value—not one we know for certain. If we were to look at thousands of couples, each having a midparent IQ value of 115, the IQs of their children would be expected to form a frequency distribution. The mean IQ of this distribution would be 106; however, most children would have higher or lower IQs—some even higher than the IQs of either parent and some even lower than the population mean of 100. The variability in this distribution comes from Mendelian segregation of the alleles that influence IQ and from factors in the environment. If, for example, Oswald were raised in a home with little or no intellectual stimulation, with a poor diet, and with other unfavorable conditions, his IQ might turn out to be considerably lower than 106. Conversely, in a nurturing home environment, Oswald's IQ might turn out to be much greater than 106. We have predicted Oswald's IQ to be 106; however, we should keep in mind that this number is a prediction, not an absolutely determined value. To test your understanding of these concepts, work through Solve It: Using the Narrow-Sense Heritability.

ARTIFICIAL SELECTION

In addition to predicting an offspring's phenotype, the narrow-sense heritability has another use: to predict the outcome of a program of selective breeding in a population. The ideas are summarized in Figure 22.8, which shows the frequency distributions of a quantitative trait among parents and their offspring. In the parental generation, the mean value of the trait is 20 units. To form the next generation, we select the individuals in the upper tail of the distribution to be parents; let's suppose that the mean of these selected individuals is 30 units. Can we predict the mean value of the trait in the offspring of these selected parents? The answer is yes, providing we know the narrow-sense heritability of the trait. The prediction equation is

$$T_o = \mu + h^2 [T_s - \mu]$$

where T_o is the mean of the offspring, μ is the mean of the overall population, T_s is the mean of the selected parents, and b^2 is the narrow-sense heritability. Notice that this equation is the same as the prediction equation for the phenotype of a single offspring, except that T_s has been substituted for T_p . In effect, we have adapted the single-offspring prediction equation to a situation in which many parents (albeit *selected* parents) produce a whole group of offspring, which then forms the population in the next generation. Thus, the new equation allows us to predict how the mean of the population will change by selecting the individuals that will be parents. We call this process **artificial selection**. It is a practice common in plant and animal breeding, and to a large extent, it is responsible for the highly productive strains of crop and livestock species that are used in agriculture today.

We can see more clearly how selection changes the mean of a quantitative trait in a population by rearranging the terms in the selection equation. After subtracting μ from both sides of the equation and introducing brackets around the term on the left, we have

$$[T_o - \mu] = b^2 [T_s - \mu]$$

The bracketed term on the right, $[T_s - \mu]$, is called the *selection differential*; it is the difference between the mean of the selected parents and the mean of the population

from which they were selected. The selection differential measures the intensity of artificial selection. The bracketed term on the left, $[T_o - \mu]$, is called the *response to selection*; it is the difference between the mean of the offspring and the mean of the entire population in the previous generation. Thus, the response to selection measures how much the mean of the trait has changed in one generation. We can put this in even simpler terms if we denote the response to selection by R and the selection differential by S; then

 $R = b^2 S$

Thus, the response to selection is the product of the selection differential and the narrow-sense heritability. Let's now return to our example; $\mu = 20$, $T_s = 30$, and let's suppose that $b^2 = 0.3$. With these values, S = 10 and $R = (0.3) \times 10 = 3$; thus, $T_o = 20 + 3 = 23$. If the selection process were repeated generation after generation, we would expect the mean of the population to increase incrementally. The feature Focus on Artificial Selection shows how this is accomplished in practice.

FOCUS ON

ARTIFICIAL SELECTION

rtificial selection is a standard practice to improve crop plants and livestock. However, improvement is usually slow because the generation time of agriculturally significant species is typically measured in years rather than weeks or months. To study the efficacy of artificial selection, Franklin Enfield and his colleagues carried out extensive experiments with a laboratory animal, the flour beetle, Tribolium castaneum. In these experiments, Enfield selected for increased body size. He measured the weight of the animals at the pupal stage and selected the heaviest pupae to be the parents of the next generation. This process was continued for 125 generations. At the start of the experiment, the weight of the individual pupae ranged from 1800 to 3000 μ g, the mean was 2400 μ g, and the variance was 40,000 μ g². After 125 generations of selection, the mean pupa weight had increased to 5800 μ g, more than twice the mean of the starting population. Moreover, none of the individuals in the selected population was as small as the largest individuals in the original starting population (**Figure 1**). This complete lack of overlap in the frequency distributions indicates that the genetic makeup of the population had been radically altered.

To achieve this stunning result, Enfield used a selection differential of 200 μ g in each generation. Initially, the narrow-sense

heritability for pupa weight was estimated to be about 0.3; thus, the predicted response to selection was $0.3 \times 200 \ \mu g = 60 \ \mu g$ per generation. For the first 40 generations, this was approximately what Enfield observed. However, the cumulative response during this time was 2000 μ g, a little less than the 2400 μ g that was expected ($60 \ \mu g$ /generation $\times 40$ generations). This discrepancy was due to factors that reduced the selection efficiency, including such things as infertility among the selected individuals. Thus, although the narrow-sense heritability is a reasonably good predictor of the response to selection over a few generations, in the long term it tends to overestimate this response.

The later generations of Enfield's project dramatically demonstrate this point. Between generations 40 and 125, the cumulative response was 1400 μ g, which, though impressive, is much less than the expected response of 5100 μ g (60 μ g/generation \times 85 generations). A detailed analysis demonstrated that during these generations, the efficiency of selection was severely reduced by a negative correlation between size and reproductive ability—after a certain point, the larger the beetle, the less reproductively successful it is. This reduced the effective selection differential and made it difficult to select for further increases in size.

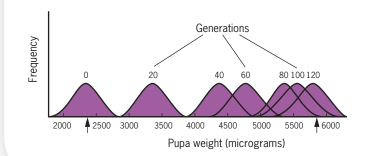


FIGURE 1 Frequency distributions of pupa weight in *Tribolium* populations selected for increased size. The shape of the distributions is only approximate. The means at generations 0 and 120 are indicated by arrows.

Now let's suppose that we select for a change in another trait whose narrow-sense heritability is unknown. For this trait, the mean of the population is 100 and the mean of the selected parents is 120. Among the offspring of these parents, we find that the mean is 104. What is the narrow-sense heritability? From the equation for the response to selection, we see that $R/S = b^2$, and in this example, R = 104 - 100 = 4, and S = 120 - 100 = 20. Thus, $R/S = 4/20 = 0.2 = b^2$, the narrow-sense heritability. From this example, we see that the response to an artificial selection experiment can be used to estimate the narrow-sense heritability.

QUANTITATIVE TRAIT LOCI

Statistical analysis has been a mainstay of quantitative genetics since Fisher's 1918 paper. With this type of analysis, quantitative geneticists have studied many different traits in many different organisms, and recently they have developed techniques to identify individual genes that influence complex traits. A gene's position in a chromosome is called a *locus* (plural, loci), and the locus for a gene that influences a quantitative trait is called a **quantitative trait locus**—abbreviated QT locus, or more simply QTL.

Modern molecular techniques have made it possible to search genomes for QT loci. These loci have been identified and mapped on specific chromosomes in model laboratory organisms such as the fruit fly and the mouse, in agriculturally signifi-

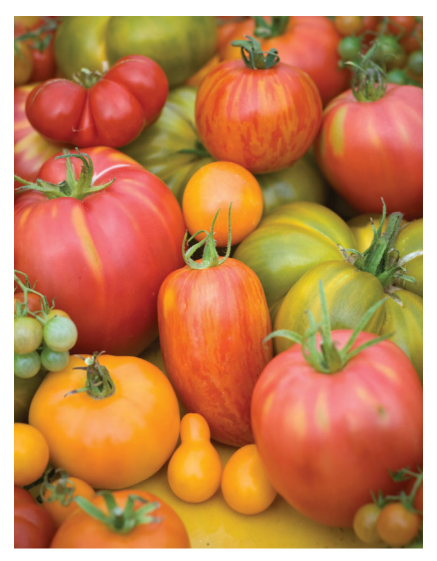


FIGURE 22.9 Variation in fruit size, shape, and color in tomatoes.

cant plants such as corn and rice, in livestock such as pigs and cows, and in our own species. The traits that have been studied include bristle number in the fruit fly, obesity in the mouse, crop yield in rice and corn, milk production in dairy cattle, fatness and growth rate in pigs, and susceptibility to illnesses such as diabetes, cancer, cardiovascular disease, and schizophrenia in human beings.

To illustrate the methods used to identify QT loci in organisms where breeding experiments are possible, let's consider a study on fruit weight in tomatoes conducted by Steven Tanksley and colleagues. Cultivated tomatoes belong to the species Lycopersicon esculentum. There are many different varieties, and in each variety the fruits have a characteristic size, shape, and color (**Figure 22.9**). All these varieties were derived by artificial selection from wild tomatoes, which are native to South America. L. pimpinellifolium, which has small, berry-like fruits, is thought to be the genetic ancestor of cultivated tomatoes. A fruit from L. pimpinellifolium weighs about 1 gram, whereas a fruit from the cultivated variety Giant Heirloom may weigh as much as 1000 grams-a dramatic indication of the power of artificial selection.

Tanksley and colleagues began their efforts to identify the loci responsible for variation in tomato fruit weight by constructing detailed molecular maps for each of the tomato's 12 chromosomes. They exploited the fact that *L. pimpinellifolium* and *L. esculentum* differ in the sites where restriction enzymes cleave genomic DNA. For example, *Eco*RI may cleave at a particular site in the DNA of *L. pimpinellifolium*, but not cleave at this site in the DNA of *L. esculentum* because the *Eco*RI recognition sequence there (GAATTC) had mutated. Differences of this sort create *restriction fragmentlength polymorphisms* (RFLPs) that can be analyzed by

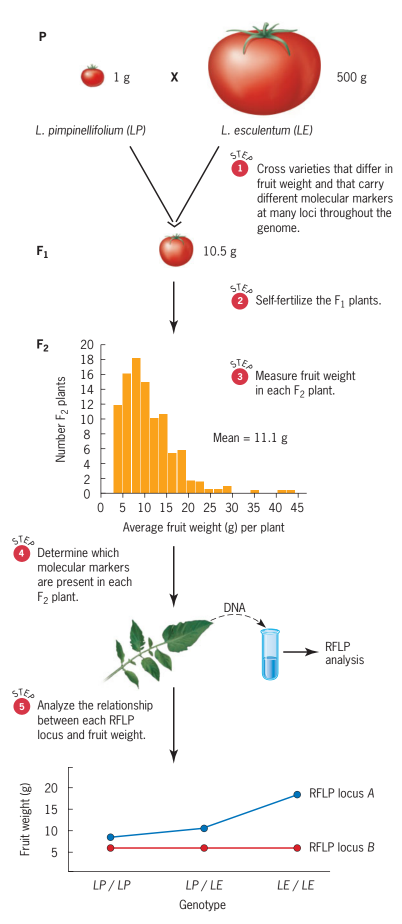
Analysis of Quantitative Traits 621

■ FIGURE 22.10 Methods to identify QT loci for fruit weight in tomatoes. Two different species of tomatoes were crossed to produce an F₁ plant, which was self-fertilized to produce many F₂ plants, each of which was characterized for the quantitative trait fruit weight and a battery of loci whose alleles are defined by restriction fragmentlength polymorphisms (RFLPs). The resulting data were analyzed to determine if fruit weight was related to the genotypes at any of the RFLP loci. The LP allele is derived from L. pimpinellifolium, and the LE allele is derived from L. esculentum. For one RFLP locus (A), the LE allele increases fruit weight when it is homozygous. For the other RFLP locus (B), the LE allele has no effect on fruit weight. A QTL for fruit weight therefore appears to be located near RFLP locus A. Data from Lippman, Z. and S. Tanksley. 2001. Dissecting the genetic pathway to extreme fruit size in tomato using a cross between the small-fruited wild species Lycopersicon pimpinellifolium and L. esculentum var. Giant Heirloom. Genetics 158: 413-422.

Southern blotting (see Figure 15.3 and the associated discussion in Chapter 15). Tanksley and colleagues catalogued a large number of RFLPs in the tomato genome and then positioned them on the genetic maps of the chromosomes by observing the frequency of recombination in hybrids created by crossing the two different species. In effect, they treated the RFLPs as molecular genetic markers and performed recombination experiments similar to the ones using phenotypic markers that we discussed in Chapter 7. Altogether, 88 RFLP loci were positioned on the maps of the tomato chromosomes. Then Tanksley and Zachary Lippman carried out an experiment to determine which of these loci were associated with differences in fruit weight. The experimental procedures are outlined in **Figure 22.10**.

L. pimpinellifolium plants were crossed to the Giant Heirloom variety of L. esculentum, and a single F_1 plant was self-fertilized to produce F₂ progeny. At each stage in the experiment, the fruits produced by each plant were weighed. The parental strains differed dramatically in fruit weight: 1 gram for L. pimpinellifolium and 500 grams for L. esculentum. The fruit of the F₁ plant averaged 10.5 grams, and the fruit of the 188 F₂ plants that were generated averaged 11.1 grams. However, among the F₂ plants, fruit weight varied considerably, with some plants bearing fruit that averaged more than 20 grams. This variation is due to the segregation of genes affecting fruit weight. To locate these genes-or QT locion the genetic map, Tanksley and Lippman determined the RFLP genotypes of the F₂ plants. DNA was extracted from individual plants, digested with restriction enzymes, and analyzed by Southern blotting to determine what RFLP markers were present. For a particular RFLP locus, an F₂ plant could be homozygous for the marker from L. pimpinellifolium, it could be homozygous for the marker from L. esculentum, or it could be heterozygous-that is, carry a marker from each species. We can designate these genotypes as LP/LP, LE/LE, and LP/LE, respectively. Each F₂ plant was genotyped for the LP and LE markers at each of the 88 RFLP loci-a heroic undertaking.

Then Tanksley and Lippman studied the relationship between the genotypes at each *RFLP* locus and fruit weight. For example, at the TG167 RFLP locus on chromosome 2, they



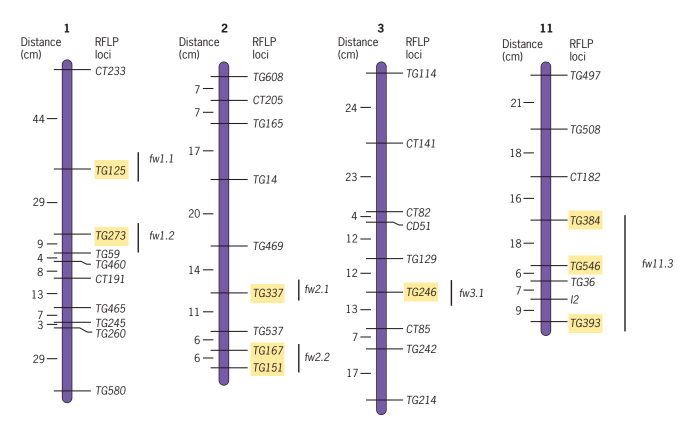


FIGURE 22.11 RFLP and QT loci for fruit weight on four chromosomes in the tomato genome. The highlighted RFLP loci are associated with effects on fruit weight. The QT loci, which are designated with the letters *fw*, are situated nearby. Data from Lippman, Z. and S. Tanksley. 2001. Dissecting the genetic pathway to extreme fruit size in tomato using a cross between the small-fruited wild species *Lycopersicon pimpinellifolium* and *L. esculentum* var. Giant Heirloom. *Genetics* 158: 413–422.

found that plants that were homozygous for the LP marker had fruits that weighed 8.4 grams, that plants that were heterozygous for the LP and LE markers had fruits that weighed 10.0 grams, and that plants that were homozygous for the LE marker had fruits that weighed 17.5 grams. Thus, at this RFLP locus it seems that the LE marker is associated with increased fruit weight, which suggests that in L. esculentum there is an allele for increased fruit weight somewhere near the TG167 locus. However, we cannot conclude that the allele for increased fruit weight is actually at the TG167 locus—only that it is nearby. Thus, this analysis points to the existence of a QTL affecting fruit weight near TG167 on chromosome 2. Tanksley and Lippman designated this QTL as fw2.2.

After examining the relationship between fruit weight and the genotypes at all the other RFLP loci, Tanksley and Lippman concluded that there are five additional fruit weight loci, including one more on chromosome 2, two on chromosome 1, and one each on chromosomes 3 and 11 (**Figure 22.11**). More detailed mapping studies ultimately allowed Tanksley and colleagues to pinpoint the *fw2.2* QTL and show that it is a single gene, *ORFX*. This gene is expressed early in floral development and is structurally similar to the human *c-ras* oncogene. Thus, its product might be involved in signal transduction within cells (see Chapters 20 and 21). To delve deeper into the analysis of QT loci in the tomato, work through Problem-Solving Skills: Detecting Dominance at a QTL.

Tanksley's research shows that identifying and mapping QT loci can be an elaborate and time-consuming enterprise. Fortunately, newer technologies such as gene chips that detect single-nucleotide polymorphisms have speeded up the work. These

PROBLEM-SOLVING SKILLS

Detecting Dominance at a QTL

THE PROBLEM

- **a.** Figure 22.10 shows how Zachary Lippman and Steven Tanksley identified QT loci for fruit weight in tomatoes. The parents in the initial cross differed dramatically in the average weights of their fruits—1 gram versus 500 grams. The F_1 fruits averaged 10.5 grams, and the F_2 fruits averaged 11.1 grams. Why do these data indicate that dominance plays a role in determining fruit weight in tomatoes?
- **b.** Lippman and Tanksley identified six QT loci affecting fruit weight. One locus, *fw11.3*, was located near the RFLP locus *TG36* on chromosome 11. Another locus, *fw2.2*, was located near the RFLP locus *TG167* on chromosome 2. When F_2 plants were genotyped for these two loci, Lippman and Tanksley found the following relationship between the genotypes and average fruit weight (all values in grams)¹:

		Genotype of F ₂ Plants		
QTL	RFLP Locus	LP/LP	LP/LE	LE/LE
fw11.3	TG36	6.2	12.2	20.0
fw2.2	TG167	8.4	10.0	17.5

Which QTL shows dominance for the trait fruit weight? Which of the alleles, *LE* or *LP*, is dominant?

¹Data from Lippman, Z. and S. Tanksley. 2001. Dissecting the genetic pathway to extreme fruit size in tomato using a cross between the small-fruited wild species *Lycopersicon pimpinellifolium* and *L. esculentum* var. Giant Heirloom. *Genetics* 158: 413–422.

FACTS AND CONCEPTS

- **1.** When alleles act additively, the phenotype of the heterozygote is midway between the phenotypes of the two homozygotes.
- **2.** To a quantitative geneticist, dominance exists when the alleles are not acting in a strictly additive fashion. Dominance is, therefore, a deviation from strict additivity.
- **3.** For a single locus acting on a trait, dominance is indicated when the phenotype of the heterozygote is not midway between the phenotypes of the two homozygotes.
- **4.** For many loci acting on a trait, dominance is indicated when the phenotype of the F₁ is not midway between the phenotypes of the two parents.

ANALYSIS AND SOLUTION

- a. The mean fruit weights in the P, F₁, and F₂ generations indicate that dominance plays a role in determining this quantitative trait. The F₁ and F₂ averages are much closer to the fruit weight of *L. pimpinellifolium* than *L. esculentum*. This skew is clear evidence for dominance.
- **b.** The *fw2.2* QTL shows dominance whereas the *fw11.3* QTL does not. For *fw2.2*, the heterozygote's phenotype is close to the phenotype of the *LP/LP* homozygote, not midway between the two homozygotes. This observation indicates that the *LP* allele of the *fw2.2* QTL is partially dominant over the *LE* allele. By contrast, the phenotype of the heterozygote at the *fw11.3* QTL is nearly midway between that of the two homozygotes. Thus, the alleles of this locus appear to act more or less additively to determine fruit weight.

For further discussion visit the Student Companion site.

technologies have also been used to find associations between molecular markers and various human diseases, including some that can be considered polygenic threshold traits. Sometimes the associations between the markers and the diseases are found in pedigrees, but more often, they are discovered in samples from the general population.

We began this chapter with a story about cardiovascular disease, which is a major cause of death among people in postindustrial societies. It has long been known that susceptibility to this disease is influenced by genetic factors. For example, relatives who share half their genes with people who have had coronary heart disease are seven times more likely to develop this disease themselves than are equivalent relatives of unaffected people. Furthermore, the risk of a monozygotic twin dying of coronary heart disease when its co-twin died of this disease before age 65 is three to seven times greater than the risk for dizygotic twins. These and other statistical data indicate that susceptibility to cardiovascular disease is under genetic control. Current research is focusing on efforts to identify specific genes that contribute to variation in the factors that put people at risk to develop this disease. These factors include plasma cholesterol level, obesity, blood pressure, high- and low-density lipoprotein levels, and triglyceride level. **Table 22.2** lists some of the QT loci that have been identified in these efforts.

TABLE 22.2

Quantitative Trait Loci That Contribute to Variation in Risk Factors for Cardiovascular Disease

Locus	Gene Product	Chromosome	Risk Factor
AGT	Angiotensin	1	Blood pressure
APOA-1	Apolipoprotein A1	11	HDL ^ª cholesterol
APOA-2	Apolipoprotein A2	1	HDL cholesterol
APOA-4	Apolipoprotein A4	11	HDL cholesterol, triglycerides
APOB	Apolipoprotein B	2	LDL ^b cholesterol
APOC-3	Apolipoprotein C3	11	Triglycerides
APOE	Apolipoprotein E	19	LDL cholesterol, triglycerides
CETP	Cholesterol ester transfer protein	16	HDL cholesterol
DCP	Dipeptidyl carboxypeptidase	17	HDL cholesterol, blood pressure
FGA/B	Fibrinogen A and B	4	Fibrinogen
HRG	Histidine-rich glycoprotein	3	Histidine-rich glycoprotein
LDLR	Low-density lipoprotein receptor	19	LDL cholesterol
LPA	Lipoprotein (a)	6	HDL cholesterol, triglycerides
LPL	Lipoprotein lipase	8	Triglycerides
PLAT	Plasminogen activator tissue-type	8	Tissue plasminogen activator level
PLANH1	Plasminogen activator inhibitor-1	7	PAI-1 level

Source: G. P. Vogler et al. 1997. Genetics and behavioral medicine: risk factors for cardiovascular disease. Behavioral Medicine 22:141–149.

- ^a High-density lipoprotein.
- ^b Low-density lipoprotein.

KEY POINTS

• The total phenotypic variance can be partitioned into genetic and environmental components: $V_T = V_g + V_e$.

- The phenotypic variance in a population that is genetically uniform estimates V_e.
- The broad-sense heritability is the proportion of the total phenotypic variance that is genetic variance: $H^2 = V_g / V_T$.
- The genetic variance can be subdivided into additive genetic, dominance, and epistatic variances: $V_g = V_a + V_d + V_i$.
- The narrow-sense heritability is the proportion of the total phenotypic variance that is due to the additive effects of alleles: $h^2 = V_a/V_T$.
- The narrow-sense heritability is used to predict the phenotypes of offspring (T_0) given the average phenotype of the parents (T_p) and the mean phenotype in the population (μ) from which the parents came: $T_0 = \mu + h^2(T_p \mu)$.
- The response to artificial selection can be predicted from the narrow-sense heritability and the selection differential: $R = h^2 S$.
- By using molecular markers, geneticists are able to identify and map quantitative trait loci.

Correlations between Relatives

Quantitative analyses of the resemblance between relatives can provide estimates of broad- and narrow-sense heritabilities. Much of classical genetic analysis involves comparisons between relatives—parents and offspring, siblings, half siblings, and so forth. The usual procedure is to follow a particular trait through a series of crosses or to trace it through a collection of pedigrees. By analyzing the data, it is possible to discern whether or not the trait has a genetic basis. If it does, further work may allow the researcher to identify the gene

or genes involved, to locate these genes on chromosomes and, ultimately, to analyze them at the molecular level. For complex traits that involve many genes and that are also influenced by a host of environmental factors, this type of analysis is extremely difficult. Nevertheless, comparisons between relatives can provide useful information about the underlying genetic variation in the trait.

CORRELATING QUANTITATIVE PHENOTYPES BETWEEN RELATIVES

Relatives often have similar phenotypes for a quantitative trait. As an example, let's consider data on the heights of monozygotic twins. **Figure 22.12a** shows such data, with each twin pair represented as a point in a graph. The height of one member of each pair is plotted on the horizontal or *x*-axis, and the height of its co-twin is plotted on the vertical or *y*-axis. From the graph it is clear that monozygotic twins are remarkably similar with respect to height. When one twin is short, the other tends to be short too; when one twin is tall, the other also tends to be tall. We refer to this

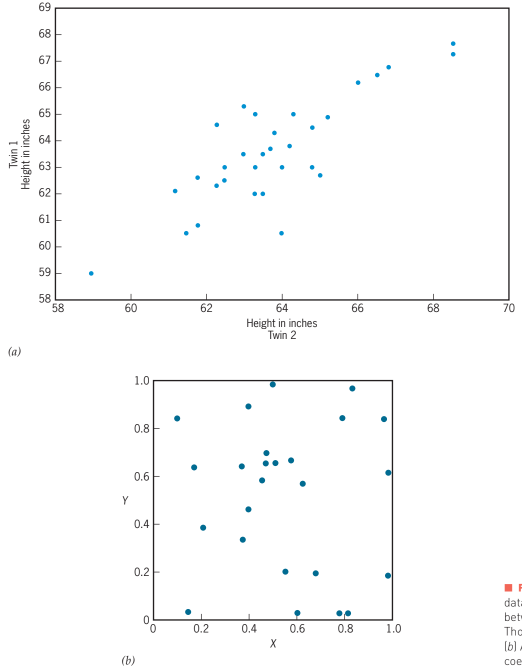


FIGURE 22.12 Correlations between paired data points. (a) Positive correlation for height between monozygotic twins (data courtesy of Thomas Bouchard, University of Minnesota).
 (b) A set of paired data in which the correlation coefficient is close to zero.

pattern of resemblance as a positive correlation, and we summarize it quantitatively by calculating a statistic called the correlation coefficient, usually symbolized by the letter r. Let's denote the height of the twin plotted on the x-axis by the letter X and that of its co-twin plotted on the y-axis by the letter Y; then the correlation coefficient for all the twin pairs in the graph is calculated from the expression

$$r = \sum [(X_k - \overline{X})(Y_k - \overline{Y})]/[(n-1)s_X s_Y]$$

In this formula, X and Y are the sample means of the twins plotted on the x- and y-axes, s_X and s_Y are the respective sample standard deviations, and n is the number of twin pairs. The Greek letter Σ indicates a summation on the index k over all the twin pairs. This formula provides researchers with a way of assigning a numerical score to a set of paired measurements such as the heights of twins in the graph. The value of the correlation coefficient can range from -1 to +1, with -1 indicating a perfect negative correlation between the X's and the Y's (high values on one axis consistently paired with low values on the other axis) and +1 indicating a perfect positive correlation. When the correlation is illustrated in Figure 22.12b, where there is no consistent relationship between the values plotted on the x- and y-axes. For the twin data in Figure 22.12a, the correlation coefficient is +0.84, which is very close to +1. Thus, monozygotic twins show a strong positive correlation with respect to height.

Correlation coefficients can be calculated for all sorts of quantitative phenotypes height, Weight, IQ score, and so forth. Furthermore, these coefficients can be calculated using data from different types of relatives—for example, from pairs of twins, pairs of full siblings, pairs of half siblings, and pairs of first cousins. We can also calculate correlation coefficients using data from unrelated individuals—for example, from pairs of college roommates. If some of the variation in a quantitative trait is due to genetic differences among individuals, we would expect the value of the correlation coefficient to increase with the closeness of the genetic relationship. Thus, monozygotic twins, who share 100 percent of their genes, should be more strongly correlated than first cousins, who share 12.5 percent of their genes.

INTERPRETING CORRELATIONS BETWEEN RELATIVES

We have already seen that variation in a quantitative trait can be partitioned into genetic and environmental components. The broad-sense heritability (H^2) is the proportion of the phenotypic variance that is due to genetic variation in a population, and the narrow-sense heritability (b^2) is the proportion of the phenotypic variance that is due to additive genetic variation in a population. If dominance and epistasis influence a trait, we expect the broad-sense heritability to be greater than the narrow-sense heritability. If these factors do not influence a trait, then the broad-sense heritability and the narrow-sense heritability are equivalent.

Correlation coefficients calculated by the formula given in the previous section can be interpreted in terms of broad- and narrow-sense heritabilties. Geneticists have analyzed the relationships among these quantities, beginning with the pioneering work of R. A. Fisher. This analysis assumes that T, the value of a trait in an individual, is equal to the mean of the population (μ) plus genetic (g) and environmental (e) deviations from the mean:

$$T = \mu + g + e$$

= $\mu + a + d + i + e$

The terms a, d, and i in this expression are, respectively, the additive, dominance, and epistatic components of the genetic deviation from the mean. It is also necessary to assume that the genetic factors influencing the phenotype are independent of the environmental factors and that the genetic and environmental factors do not interact in a nonadditive way. Under these assumptions, the correlation coefficient for a pair of relatives equals the proportion of the total variance in the trait that is due to the genetic and environmental factors *shared* by the relatives. Table 22.3

TABLE 22.3			
Theoretical Values of Correlation Coefficients for MZ and DZ Twins and Unrelated Individuals Reared Together or Apart			
Relationship	Theoretical Value of Correlation Coefficient (r)		
MZA	H ²		
MZT	$H^2 + C^2$		
DZA	$(1/2)h^2 + D^2$		
DZT	$(1/2)h^2 + D^2 + C^2$		
URA	0		
URT	C^2		

presents theoretical interpretations of correlation coefficients for different types of human twins.

Monozygotic twins reared apart (MZA) have identical genotypes. Thus, these twins share all the genetic factors that contribute to the term g in the expression for the value of a quantitative trait, including the additive effects of alleles, the effects of dominance, and the effects of epistasis. However, because MZA have had separate upbringings, they do not share the environmental effects represented by the term ein the expression. Consequently, a correlation between MZA depends only on their identical genotypes. In the theory of quantitative genetics, this correlation equals the proportion of the total phenotypic variance that is due to genetic differences among the twin pairs—that is, it equals the broad-sense heritability, H^2 .

Monozygotic twins reared together (MZT) have a common environment as well as identical genotypes. A correlation between them therefore equals the proportion of the total variance that is due to shared gentoypes (H^2), plus the proportion that is due to shared environmental factors. This latter component, which is denoted by the term C^2 in Table 22.3, is called the **environmentality**.

Dizygotic (DZ) twins are as closely related as ordinary siblings. To interpret a correlation coefficient between DZ twins, we must therefore discount its genetic component by a factor of 1/2, which is the fraction of genes that DZ twins (or siblings) share by virtue of common ancestry. Furthermore, although DZ twins experience the same additive effects of the genes they share, they experience only some of the same dominance and epistatic effects. This diminished similarity due to dominance and epistasis reflects the low probability that DZ twins will inherit specific combinations of alleles from their parents. The correlation coefficient for DZ twins is therefore greater than or equal to $(1/2)h^2$, but less than or equal to $(1/2)H^2$. If dominance and epistasis are negligible, then the correlation coefficient equals $(1/2)b^2$. If there is some dominance and epistasis, then it equals $(1/2)h^2$ plus a fraction of the difference between $(1/2)H^2$ and $(1/2)k^2$. In Table 22.3, this fraction is denoted by the term D^2 . For dizygotic twins reared together (DZT) the correlation coefficient will also include the effect of a shared environment (C^2). This effect will not contribute to the correlation between dizygotic twins reared apart (DZA) because these types of twins do not share a common environment.

Unrelated individuals reared apart (URA) or together (URT, for example, unrelated children adopted into the same family) do not share genes by virtue of common ancestry. Consequently, a correlation between these types of individuals does not involve a genetic component. However, it does involve the effect of a shared environment (C^2) if the individuals were reared together.

These and other theoretical results allow geneticists to use correlations between relatives to estimate the broad- and narrow-sense heritabilities for quantitative traits. The correlation between monozygotic twins reared apart provides an estimate of the broad-sense heritability, and the correlation between dizygotic twins reared apart provides a maximal estimate of the narrow-sense heritability. Correlations between other types of relatives—full siblings, half-siblings, and first cousins—also provide maximal estimates of the narrow-sense heritability. It should be emphasized, however, that all these estimates depend on several simplifying assumptions, which may or may not be met in the population under study. Thus, their interpretation is subject to considerable uncertainty.

- **KEY POINTS** The correlation coefficient summarizes the degree of association between paired measurements, X_k and Y_k : $r = \sum [(X_k - X)(Y_k - Y)]/[(n - 1)s_X s_V]$.
 - A correlation coefficient can be used to estimate the proportion of the total variance in a quantitative trait that is due to genetic and environmental factors shared by relatives.
 - The correlation between monozygotic twins reared apart provides an estimate of the broad-sense heritability.
 - The correlation between dizygotic twins reared apart provides a maximum estimate of the narrow-sense heritability.

Quantitative Genetics of Human Behavioral Traits

Quantitative genetics theory has been used to assess the heritability of intelligence and personality traits in humans.

Animals exhibit a wide range of behaviors associated with feeding, courtship, reproduction, and a host of other activities. The genetic determinants of these behaviors are only now beginning to be identified through experimental work. Studies with mutant strains of worms, fruit flies, and mice have revealed several genes that influence behavior. Research on

human beings has also indicated that behavior is affected by genetic factors. For example, people with Huntington's disease gradually lose motor control and mental function; as the disease progresses, they may become depressed, even psychotic. Huntington's disease is due to a dominant mutation that is manifested in adults, usually after age 30. At present, there is no cure. Phenylketonuria is another human genetic condition with a behavioral phenotype. People with this disease accumulate toxic metabolites in their nervous tissues, including the brain. Without treatment—which involves restricting the amount of phenylalanine consumed in the food-individuals with this disorder fail to develop normal mental abilities. Still another example of how the genotype can influence behavior is Down syndrome, a condition that arises from the presence of an extra chromosome 21. People with this condition have below-normal mental abilities, and if they survive to middle age, they invariably develop Alzheimer's disease, a form of dementia that also occurs in chromosomally normal individuals, although at a much lower rate and usually much later in life. People with Alzheimer's disease gradually, but inexorably, lose their memories and intellectual functions; they become progressively more forgetful and disoriented, and need to be monitored constantly to prevent them from hurting themselves or others. Researchers now believe that Alzheimer's disease may be caused by extra copies or mutant alleles of a gene located on chromosome 21. Mutant alleles of other genes may also lead to Alzheimer's disease.

Conditions such as Huntington's disease, phenylketonuria, and Down syndrome indicate that genetic factors can influence human behavior. However, these conditions do not offer much insight into the nature of the behavioral differences that we see in the general population. Does genetic variation account for some of these differences, and if it does, what proportion of the overall variability is due to genetic factors? These provocative questions fall within the purview of quantitative genetics. In the following sections, we apply quantitative genetics theory to the study of two complex human behavioral traits, intelligence and personality.

INTELLIGENCE

The term *intelligence* refers to an assortment of mental abilities, including verbal and mathematical skills, memory and recall, reasoning and problem solving,

Newman et al. 1937 0.71 Juel-Nielsen 1980 0.69 Shields 1962 0.75 Bouchard et al. 1990 0.83 0.75 Pedersen et al. 1992 0.80 0.78 0.22 0.3 Newman et al. 1998 0.4 0.4	TABLE 22.4				
Newman et al. 1937 0.71 Juel-Nielsen 1980 0.69 Shields 1962 0.75 Bouchard et al. 1990 0.83 0.75 Pedersen et al. 1992 0.80 0.78 0.22 0.3 Newman et al. 1998 0.4					
Juel-Nielsen 1980 0.69 Shields 1962 0.75 Bouchard et al. 1990 0.83 0.75 Pedersen et al. 1992 0.80 0.78 0.22 0.3 Newman et al. 1998 0.4	Study	MZT	MZA	DZT	DZA
^a Data and references from Bouchard, T. J. 1998. Genetic and environmental influences on adul	Juel-Nielsen 1980 Shields 1962 Bouchard et al. 1990 Pedersen et al. 1992 Newman et al. 1998 Average	0.80	0.69 0.75 0.75 0.78 0.75	0.22	0.32 0.47 0.38 s on adult

discrimination of different objects, and spatial perception. For more than a century, psychologists have tried to characterize and quantify these abilities by administering intelligence tests. The tests—and many different ones have been used—attempt to measure general reasoning ability. The score that an individual makes on one of these tests is converted into an *intelligence quotient*, or *IQ*, which is scaled so that the mean of the population is 100 and the standard deviation is 15. Although there is considerable debate about what an IQ score actually measures—is it a true reflection of a person's intelligence?—these scores have been used to assess whether variation in mental abilities has a genetic component. Some of the most revealing data have come from studies of monozygotic and dizygotic twins.

For IQ test scores, the correlation coefficients of MZ twins, reared together or apart, are very high—in the range of 0.7–0.8 (Table 22.4). By comparison, the correlation coefficients of DZ twins tend to be lower—presumably because they share only half their genes, and the correlation coefficients for unrelated individuals reared together are essentially zero. Such analyses strongly suggest that whatever an IQ test measures, it has a large genetic component. This conclusion is supported by other correlated with the IQs of their biological parents than with those of their adoptive parents. Thus, in the determination of IQ, the biological (that is, genetic) link between parents and children seems to be more influential than the environmental one.

What fraction of the variation among IQ scores is attributable to genetic differences among people? The most direct estimate comes from the correlation coefficient for MZ twins reared apart. Observed values of this correlation coefficient are around 0.7; thus, as much as 70 percent of the variation in IQ scores is attributable to genetic variability in the population. This estimate of the broad-sense heritability implies that, with respect to intelligence (as measured by IQ), people differ one from another more because of genetic factors than because of environmental factors.

PERSONALITY

Personality traits, like intelligence, can be assessed by testing. Psychologists use many different tests, some to measure personality characteristics and others to measure vocational and social interests. The results of these tests tend to be less reliable than those of IQ tests. Nevertheless, they quantify aspects of human personality in ways that allow them to be analyzed for genetic influences.

Perhaps the most thorough genetic analysis of personality in the general population has come from the Minnesota Study of Twins Reared Apart, a long-term research project carried out at the University of Minnesota. (See A Milestone in Genetics: The

TABLE 22.5

Mean Correlation Coefficients for MZ Twins Reared Together or Apart Who Were Evaluated for Personality Traits, Psychological Interests, and Social Attitudes as Part of the Minnesota Study of Twins Reared Apart^a

Test Instrument	MZT	MZA
Personality traits		
Multidimensional Personality Questionnaire	0.49	0.50
California Psychological Inventory	0.49	0.48
Psychological interests		
Strong Campbell Interest Inventory	0.48	0.39
Jackson Vocational Interest Survey	NA	0.43
Minnesota Occupational Interest Scales	0.49	0.40
Social attitudes		
Religiosity Scales	0.51	0.49
Nonreligious Social Attitude Items	0.28	0.34
MPQ Traditionalism Scale	0.50	0.53
^a Abstracted with permission from Bouchard et al. 1990. American Association for the Advancement of Science.	Science 250: 223–228. C	opyright 1990

Minnesota Study of Twins Reared Apart on the Student Companion site.) The results from this project suggest that genetic differences explain a significant fraction of the overall variation in human personality, perhaps as much as 50 percent (Table 22.5). The correlation coefficient for the personality and psychological interest test scores of MZ twins reared apart ranges from 0.39 to 0.50. Thus, the broad-sense heritability for these traits is reasonably high. Additional insight into the genetic control of personality has come from studying conditions such as manic depression, schizophrenia, and alcoholism. The occurrence of these traits in the members of MZ and DZ twin pairs has been estimated, and the general finding is that MZ twins are more similar than DZ twins. Thus, for example, among male MZ twin pairs with one member identified as alcoholic, the co-twin is alcoholic 41 percent of the time. By contrast, a male DZ co-twin is alcoholic only 22 percent of the time. The greater concordance for alcoholism between MZ twins suggests that this trait is influenced by genetic factors.

KEY POINTS

- Studying monozygotic and dizygotic twins, reared together or apart, has been useful in assessing the extent to which genes influence behavior in the general human population.
- The broad-sense heritability for intelligence, as measured by IQ tests, is estimated to be 70 percent.
- The broad-sense heritability for personality traits is estimated to be between 34 and 50 percent.

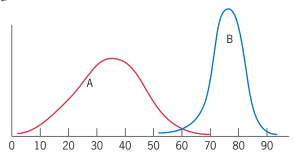
Basic Exercises

Illustrate Basic Genetic Analysis

1. In a plant species, stalk height is determined by four independently assorting genes, *A*, *B*, *C*, and *D*, each segregating two alleles; with each gene one allele, denoted by the superscript zero, adds nothing to the basic stalk height of 10 centimeters, whereas the other allele, denoted by the

superscript one, adds 1 centimeter to the basic stalk height. If all the alleles of these genes act additively to determine stalk height, (a) what is the phenotype of a plant with the genotype $A^0A^1 B^0B^1 C^0C^1 D^0D^1$, and (b) if this plant is selfed, what fraction of its offspring will be 10 cm tall?

- Answer: (a) The phenotype of the quadruple heterozygote should be the basic height (10 cm) plus the contributions of each of the one-superscript alleles (4 cm)—that is, 14 cm. (b) Among the progeny of the selfed plant, only those that are homozygous for all the zero-superscript alleles will manifest the basic phenotype of 10 cm. These quadruple zero-homozygotes will have a frequency of $(1/4)^4 = 1/256$.
- 2. For schizophrenia, the concordance for monozygotic twins is 60 percent and for dizygotic twins it is 10 percent. Do these facts argue that schizophrenia is a threshold trait with a genetic basis?
- **Answer:** The greater concordance for monozygotic twins, which are genetically identical, does argue that schizophrenia is a threshold trait with a genetic basis. The lower concordance for dizygotic twins presumably reflects the fact that they share only 50 percent of their genes.
- **3.** Which of the two frequency distributions shown below has (a) the greater mean, (b) the greater variance, (c) the greater standard deviation?



Answer: Distribution B has the greater mean. Distribution A has the greater variance and standard deviation.

- 4. Two phenotypically different highly inbred strains, P_1 and P_2 , were crossed to produce an F_1 population, which was intercrossed to produce an F_2 population. In which strain or population is the genetic variance for a quantitative trait expected to be greater than zero?
- **Answer:** The genetic variance is expected to be greater than zero in the F_2 population because it is segregating for the genetic differences introduced by the initial cross between P_1 and P_2 . The inbred strains themselves as well as the F_1 population created by crossing them are expected to have little, if any, genetic variability. Thus, in each of these populations the genetic variance should be essentially zero.
- 5. Distinguish between the broad- and narrow-sense heritabilities.
- **Answer:** The broad-sense heritability includes all the genetic variance as a fraction of the total phenotypic variance. The narrow-sense heritability includes only the additive genetic variance as a fraction of the total phenotypic variance.
- 6. Suppose that the correlation coefficient for height between human DZ twins reared apart is 0.30. What does this correlation suggest about the value of the narrow-sense heritability for height in this population?
- **Answer:** Theoretically, the correlation coefficient for DZ twins reared apart estimates $(1/2)b^2 + D^2$, where D^2 reflects correlations due to dominance and epistasis. If we assume that neither dominance nor epistasis causes variation in this trait, then the correlation coefficient estimates $(1/2)b^2$. Thus, if we double the correlation coefficient, we obtain a maximum estimate of the narrow-sense heritability; $b^2 < 2 \times 0.30 = 0.60$.

Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. A group of researchers studied variation in the number of abdominal bristles in female *Drosophila*. Two inbred strains that differed in bristle number were crossed to produce F_1 hybrids. The variance in bristle number among the F_1 flies was 3.33. These F_1 flies were intercrossed with one another to produce an F_2 population, in which the variance in bristle number was 5.44. Estimate the broad-sense heritability for bristle number in the F_2 population.
- **Answer:** Because the F_1 flies were produced by crossing two inbred strains, they are genetically uniform. The variance observed among these flies therefore estimates the environmental variance, V_e . The variance observed among the F_2 flies, V_T , is the sum of the genetic variance, V_g , and the environmental variance, V_e . Thus, we can estimate V_g by

subtracting the variance observed in the F₁ flies from that observed in the F₂ flies: $V_g = V_T - V_e = 5.44 - 3.33 = 2.11$. The broad-sense heritability, which is defined as V_g/V_T , is therefore 2.11/5.44 = 0.37.

- 2. The mean value of a trait is 100 units, and the narrowsense heritability is 0.3. A male and a female measuring 130 and 90 units, respectively, mate and produce a large number of offspring, which are reared in randomized environments. What is the expected value of the trait among these offspring?
- **Answer:** The midparent value (the average of the two parents) is (130 + 90)/2 = 110. This value deviates from the population mean (100) by 10 units. If the narrow-sense

heritability for the trait is 0.3, 30 percent of this deviation should be heritable. Consequently, the predicted value of the trait for the offspring of these two parents is $100 + (0.3 \times 10) = 103$.

- 3. In a study of MZ and DZ twins, reared together and apart, a group of Swedish researchers obtained the following correlation coefficients for IQ test scores: MZT, 0.80; MZA, 0.78; DZT, 0.22; DZA, 0.32. What do these correlations suggest about the extent to which variation in IQ scores is attributable to genetic variation? Are the results internally consistent?
- Answer: The correlation for MZ twins reared apart, 0.78, implies that 78 percent of the population's variability in IQ is due to genetic variation—that is, the broad-sense heritability is 0.78. The slightly higher correlation for MZ twins reared together reinforces this conclusion and suggests that the effect of a common environment on the correlation for IQ is negligible. Thus, common

environmental influences seem to account for a very small percentage of the overall variation in IQ within the population. The correlations for the DZ twins are generally in agreement with this view, but there is one inconsistency: the correlation for DZ twins reared together is less than that for DZ twins reared apart. We might have expected the correlation for DZ twins reared together to be as great as or greater than the correlation for DZ twins reared apart. This inconsistency is probably due to sampling error. If we accept the correlation for DZ twins reared apart at face value, then doubling it should provide a maximal estimate of the narrow-sense heritability; $2 \times 0.32 = 0.64$. The fact that this estimate is less than the broad-sense heritability estimated from the correlation between MZ twins reared apart (0.78) suggests (albeit not too strongly given all the statistical uncertainties associated with these data) that some of the genetic variation in IQ is due to nonadditive genetic factors such as dominance and epistasis.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

- **22.1** If heart disease is considered to be a threshold trait, what genetic and environmental factors might contribute to the underlying liability for a person to develop this disease?
- **22.2 (a)** A wheat variety with red kernels (genotype A'A'B'B') was crossed with a variety with white kernels (genotype AA BB). The F₁ were intercrossed to produce an F₂. If each primed allele increases the amount of pigment in the kernel by an equal amount, what phenotypes will be expected in the F₂? Assuming that the A and B loci assort independently, what will the phenotypic frequencies be?
- **22.3** For alcoholism, the concordance rate for monozygotic twins is 55 percent, whereas for dizygotic twins, it is 28 percent. Do these data suggest that alcoholism has a genetic basis?
- 22.4 💿 The height of the seed head in wheat at maturity is determined by several genes. In one variety, the head is just 9 inches above the ground; in another, it is 33 inches above the ground. Plants from the 9-inch variety were crossed to plants from the 33-inch variety. Among the F₁, the seed head was 21 inches above the ground. After self-fertilization, the F₁ plants produced an F₂ population in which 9-inch and 33-inch plants each appeared with a frequency of 1/256. (a) How many genes are involved in the determination of seed head height in these strains of wheat? (b) How much does each allele of these genes contribute to seed head height? (c) If a 21-inch F₁ plant were crossed to a 9-inch plant, how often would you expect 18-inch wheat to occur in the progeny?
- **22.5** Assume that size in rabbits is determined by genes with equal and additive effects. From a total of 2012 F_2 progeny

from crosses between true-breeding large and small varieties, eight rabbits were as small as the small variety and eight were as large as the large variety. How many sizedetermining genes were segregating in these crosses?

- **22.6** A sample of 20 plants from a population was measured in inches as follows: 18, 21, 20, 23, 20, 21, 20, 22, 19, 20, 17, 21, 20, 22, 20, 21, 20, 22, 19, and 23. Calculate (a) the mean, (b) the variance, and (c) the standard deviation.
- **22.7** Quantitative geneticists use the variance as a measure of scatter in a sample of data; they calculate this statistic by averaging the squared deviations between each measurement and the sample mean. Why don't they simply measure the scatter by computing the average of the deviations without bothering to square them?
- **22.8** Two inbred strains of corn were crossed to produce an F_1 , which was then intercrossed to produce an F_2 . Data on ear length from a sample of F_1 and F_2 individuals gave phenotypic variances of 15.2 cm² and 27.6 cm², respectively. Why was the phenotypic variance greater for the F_2 than for the F_1 ?
- **22.9** A study of quantitative variation for abdominal bristle number in female *Drosophila* yielded estimates of $V_T = 6.08$, $V_g = 3.17$, and $V_e = 2.91$. What was the broad-sense heritability?
- **22.10** A researcher has been studying kernel number on ears of corn. In one highly inbred strain, the variance for kernel number is 426. Within this strain, what is the broad-sense heritability for kernel number?

- **22.11** Measurements on ear length were obtained from three populations of corn—two inbred varieties and a randomly pollinated population derived from a cross between the two inbred strains. The phenotypic variances were 9.2 cm² and 9.6 cm² for the two inbred varieties and 26.4 cm² for the randomly pollinated population. Estimate the broad-sense heritability of ear length for these populations.
- **22.12** Figure 22.4 summarizes data on maturation time in populations of wheat. Do these data provide any insight as to whether or not this trait is influenced by dominance? Explain.
- **22.13** A person claims that the narrow-sense heritability for body mass in human beings is 0.7, while the broad-sense heritability is only 0.3. Why must there be an error?
- **22.14** The mean value of a trait is 100 units, and the narrowsense heritability is 0.4. A male and a female measuring 124 and 126 units, respectively, mate and produce a large number of offspring, which are reared in an average environment. What is the expected value of the trait among these offspring?
- **22.15** The narrow-sense heritability for abdominal bristle number in a population of *Drosophila* is 0.3. The mean bristle number is 12. A male with 10 bristles is mated to a female with 20 bristles, and a large number of progeny are scored for bristle number. What is the expected mean number of bristles among these progeny?
- **22.16** A breeder is trying to decrease the maturation time in a population of sunflowers. In this population, the mean time to flowering is 100 days. Plants with a mean flowering time of only 90 days were used to produce the next generation. If the narrow-sense heritability for flowering time is 0.2, what will the average time to flowering be in the next generation?
- **22.17** A fish breeder wishes to increase the rate of growth in a stock by selecting for increased length at six weeks after hatching. The mean length of six-week-old fingerlings

is currently 10 cm. Adult fish that had a mean length of 15 cm at six weeks of age were used to produce a new generation of fingerlings. Among these, the mean length was 12.5 cm. Estimate the narrow-sense heritability of fingerling length at six weeks of age and advise the breeder about the feasibility of the plan to increase growth rate.

- **22.18 (a)** Leo's IQ is 86 and Julie's IQ is 110. The mean IQ in the population is 100. Assume that the narrow-sense heritability for IQ is 0.4. What is the expected IQ of Leo and Julie's first child?
- **22.19** One way to estimate a maximum value for the narrowsense heritability is to calculate the correlation between half-siblings that have been reared apart and divide it by the fraction of genes that half-siblings share by virtue of common ancestry. A study of human half-siblings found that the correlation coefficient for height was 0.14. From this result, what is the maximum value of the narrowsense heritability for height in this population?
- **22.20** 0 A selection differential of 40 µg per generation was used in an experiment to select for increased pupa weight in *Tribolium*. The narrow-sense heritability for pupa weight was estimated to be 0.3. If the mean pupa weight was initially 2000 µg and selection was practiced for 10 generations, what was the mean pupa weight expected to become?
- **22.21** On the basis of the observed correlations for personality traits shown in Table 22.5, what can you say about the value of the environmentality (C^2 in Table 22.3)?
- 22.22 Correlations between relatives provide estimates of the broad and narrow-sense heritabilties on the assumption that the genetic and environmental factors influencing quantitative traits are independent of each other and that they do not interact in some peculiar way. In Chapter 19, we considered epigenetic modifications of chromatin that regulate genes and noted the possibility that some of these modifications might be induced by environmental factors. How could epigenetic influences on complex traits be incorporated into the basic theory of quantitative genetics?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

- 1. QTL mapping has been carried out for many organisms, including crop plants such as rice and maize. Follow the links to the *Zea mays* page, and then under Related Resources, go to the web site of "Gramene," a resource for comparative grass genomics. Explore the rice and maize QTL data by entering specific traits in the query box. For 100-grain weight in rice, how many QT loci have been mapped? How many of rice's 12 chromosomes contain at least one of these QT loci? For kernel row number in maize, how many QT loci have been mapped? On how many of maize's 10 chromosomes do these QT loci lie?
- 2. With many people now living into their seventh and eighth decades of life, Alzheimer's disease has become more frequent. Geneticists have found variants at several loci that seem to predispose people to develop this condition. These loci include *APOE*, *APP*, *PSEN1*, and *PSEN2*. Use the search function on the *Homo sapiens* web page to locate each of these loci in the human genome. On what chromosomes do they reside? Click on each locus to bring up a summary about the gene. How are the gene products thought to function in the etiology of Alzheimer's disease?



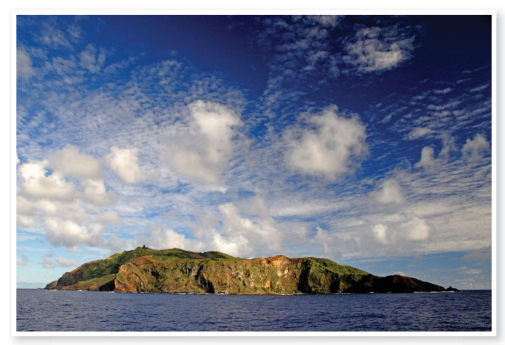
Population Genetics

CHAPTER OUTLINE

- The Theory of Allele Frequencies
- Natural Selection
- Random Genetic Drift
- Populations in Genetic Equilibrium

A Remote Colony

In September 1787, Lieutenant William Bligh and a crew of 45 men set sail from England aboard the ship H.M.S. Bounty. Their destination was the Pacific island of Tahiti, where they were to collect breadfruit tree saplings for transplantation to the Caribbean island of Jamaica. Because their passage around Cape Horn was blocked by ferociously bad weather, they sailed to Tahiti by crossing the south Atlantic, rounding the Cape of Good Hope, and then traversing the southern Indian Ocean and the western Pacific. Their voyage was long and difficult. When they finally reached Tahiti, they relaxed there and enjoyed the hospitality of the local people. After collecting the breadfruit saplings, Bligh and his crew departed Tahiti on April 6, 1789, bound for the Caribbean. Barely three weeks into the voyage, the crew mutinied. Led by Bligh's friend and chief subordinate Fletcher Christian, the mutineers put Bligh and his supporters into the ship's launch and set



Pitcairn Island in the south Pacific.

them adrift in the lonely waters of the south Pacific. Eventually Bligh and his men reached civilization. The mutineers initially returned to Tahiti, where some decided to stay, but nine of them, including Fletcher Christian, resolved to find another place to live. Along with a group of Polynesians—six men, twelve women, and a baby—they set sail in the *Bounty*, and on January 15, 1790, landed on Pitcairn Island, an uninhabited speck of land 1350 miles from Tahiti. Pitcairn Island had been discovered decades earlier, but because cartographers had put it in the wrong place on their charts, it held promise as a refuge for the mutineers. On January 23, 1790, Fletcher Christian and his followers burned the *Bounty* and set about establishing their new home.

Life on Pitcairn Island was not easy. The men fought over land and women, and the women murdered some of the men. In 1808, the island was visited by an American whaling ship, which found that only one of the original mutineers was still alive. British ships subsequently stopped at the island, and in 1838, Pitcairn Island was formally incorporated into the British Empire. By 1855 the population of the colony had increased to nearly 200, which was more than it could sustain, and in 1856 all the people were moved to Norfolk Island, a former British penal colony 3500 miles away. Two years later, 17 of the former inhabitants returned to Pitcairn Island to reestablish the colony, which has survived for over 150 years and today is home to about 50 people, all descendants of the original settlers.

The Theory of Allele Frequencies

The population on Pitcairn Island is the result of mixing two different groups of people, Britons and Polynesians. The offspring of the original settlers received genes from each of these groups, and when they reproduced, some of these genes were transmitted to their offspring and ultimately to the current members When the members of a population mate randomly, it is easy to predict the frequencies of the genotypes from the frequencies of their constituent alleles.

of the population. Which of the founding genes were passed down through time? How did factors such as the health, vigor, and reproductive ability of the people, and the ways in which they chose mates, influence the pathways of genetic descent? Did any of the genes mutate as they were transmitted through time? How did migration to and from the island affect its genetic composition? Has the island's genetic diversity increased, decreased, or remained the same? What is the significance of the population's size? Has the genetic composition of the population changed over time—that is, has it evolved?

These and other questions about the genetic makeup and history of the people on Pitcairn Island fall within the purview of *population genetics*, a discipline that studies genes in groups of individuals. Population genetics examines allelic variation among individuals, the transmission of allelic variants from parents to offspring generation after generation, and the temporal changes that occur in the genetic makeup of a population because of systematic and random evolutionary forces.

The theory of population genetics is a theory of allele frequencies. Each gene in the genome exists in different allelic states, and, if we focus on a particular gene, a diploid individual is either a homozygote or a heterozygote. Within a population of individuals, we can calculate the frequencies of the different types of homozygotes and heterozygotes of a gene, and from these frequencies we can estimate the frequency of each of the gene's alleles. These calculations are the foundation for population genetics theory.

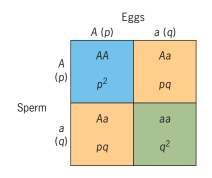
ESTIMATING ALLELE FREQUENCIES

Because an entire population is usually too large to study, we resort to analyzing a representative sample of individuals from it. **Table 23.1** presents data from a sample of people who were tested for the M-N blood types. These blood types are determined by two alleles of a gene on chromosome 4: L^M , which produces the M blood type, and L^N , which produces the N blood type (see Chapter 4). People who are $L^M L^N$ hetero-zygotes have the MN blood type.

To estimate the frequencies of the L^M and L^N alleles, we simply calculate the incidence of each allele among all the alleles sampled:

- 1. Because each individual in the sample carries two alleles of the blood-type locus, the total number of alleles in the sample is two times the sample size: $2 \times 6129 = 12,258$.
- **2.** The frequency of the L^M allele is two times the number of $L^M L^M$ homozygotes plus the number of $L^M L^N$ heterozygotes, all divided by the total number of alleles sampled: $[(2 \times 1787) + 3039]/12,258 = 0.5395.$
- **3.** The frequency of the L^N allele is two times the number of $L^N L^N$ homozygotes plus the number of $L^M L^N$ heterozygotes, all divided by the total number of alleles sampled: $[(2 \times 1303) + 3039]/12,258 = 0.4605.$

TABLE 23.1			
Frequency of the M-N Blood Types in a Sample of 6129 Individuals			
Blood Type	Genotype	Number of Individuals	
М	$L^{M}L^{M}$	1787	
MN	L ^M L ^N	3039	
Ν	L ^N L ^N	1303	



■ **FIGURE 23.1** Punnett square showing the Hardy–Weinberg principle.

Thus, letting p represent the frequency of the L^M allele and letting q represent the frequency of the L^N allele, we estimate that in the population from which the sample was taken, p = 0.5395 and q = 0.4605. Furthermore, because L^M and L^N represent 100 percent of the alleles of this particular gene, p + q = 1.

RELATING GENOTYPE FREQUENCIES TO ALLELE FREQUENCIES: THE HARDY-WEINBERG PRINCIPLE

Do the estimated allele frequencies have any predictive power? Can we use them to predict the frequencies of genotypes? In the first decade of the twentieth century, these questions were posed independently by G. H. Hardy, a British mathematician, and by Wilhelm Weinberg, a German physician. In 1908 Hardy and Weinberg each published papers describing a mathematical relationship between allele frequencies and genotype frequencies. This relationship, now called the Hardy-Weinberg principle, allows us to predict a population's genotype frequencies from its allele frequencies.

Let's suppose that in a population a particular gene is segregating two alleles, A and a, and that the frequency of A is p and that of a is q. If we assume that the members of the population mate randomly, then the diploid genotypes of the next generation will be formed by the random union of haploid eggs and haploid sperm (**Figure 23.1**). The probability that an egg (or sperm) carries A is p, and the probability that it carries a is q. Thus, the probability of producing an AA homozygote in the population is simply $p \times p = p^2$, and the probability of producing an aa homozygote is $q \times q = q^2$. For the Aa heterozygotes, there are two possibilities: An A sperm can unite with an a egg, or an a sperm can unite with an A egg. Each of these events occurs with probability $p \times q$, and because they are equally likely, the total probability of forming an Aa zygote is 2pq. Thus, on the assumption of random mating, the predicted frequencies of the three genotypes in the population are:

Genotype	Frequency
AA	p^2
Aa	2pq
aa	q^2

These predicted frequencies can be obtained by expanding the binomial expression $(p + q)^2 = p^2 + 2pq + q^2$. Population geneticists refer to them as the Hardy–Weinberg genotype frequencies.

The key assumption underlying the Hardy–Weinberg principle is that the members of the population mate at random with respect to the gene under study. This assumption means that the adults of the population essentially form a pool of gametes that, at fertilization, combine randomly to produce the zygotes of the next generation. If these zygotes have equal chances of surviving to the adult stage, then the genotype frequencies created at the time of fertilization will be preserved, and when the next generation reproduces, these frequencies will once again appear in the offspring. Thus, with random mating and no differential survival or reproduction among the members of the population, the Hardy–Weinberg genotype frequencies—and, of course, the underlying allele frequencies—persist generation after generation. This condition is referred to as the *Hardy–Weinberg equilibrium*. Later in this chapter we will consider forces that upset this equilibrium by altering allele frequencies; these forces—mutation, migration, natural selection, and random genetic drift—play key roles in the evolutionary process.

APPLICATIONS OF THE HARDY–WEINBERG PRINCIPLE

The intellectual roots of the Hardy–Weinberg principle are discussed in A Milestone in Genetics on the Student Companion site. Here, let's return to the M-N blood-type example to see how the Hardy–Weinberg principle applies to a real population. From

the sample data given in Table 23.1, the frequency of the L^M allele was estimated to be p = 0.5395, and the frequency of the L^N allele was estimated to be q = 0.4605. With the Hardy–Weinberg principle, we can now use these frequencies to predict the genotype frequencies of the M-N blood-type gene:

Genotype	Hardy–Weinberg Frequency
$L^{M}L^{M}$	$p^2 = (0.5395)^2 = 0.2911$
$L^{M}L^{N}$	2pq = 2(0.5395)(0.4605) = 0.4968
$L^{N}L^{N}$	$q^2 = (0.4605)^2 = 0.2121$

Do these predictions fit with the original data from which the two allele frequencies were estimated? To answer this question, we must compare the observed genotype numbers with numbers predicted by the Hardy–Weinberg principle. We obtain these predicted numbers by multiplying the Hardy–Weinberg frequencies by the size of the sample taken from the population. Thus,

Genotype	Predicted Number
$L^{M}L^{M}$	$0.2911 \times 6129 = 1784.2$
$L^{M}L^{N}$	$0.4968 \times 6129 = 3044.8$
$L^{N}L^{N}$	$0.2121 \times 6129 = 1300.0$

The results are extraordinarily close to the original sample data presented in Table 23.1. We can check for agreement between the observed and predicted numbers by calculating a chi-square statistic (see Chapter 3):

$$\chi^{2} = \frac{(1787 - 1784.2)^{2}}{1784.2} + \frac{(3039 - 3044.8)^{2}}{3044.8} + \frac{(1303 - 1300.0)^{2}}{1300.0} = 0.0223$$

This chi-square statistic has 3 - 2 = 1 degree of freedom because (1) the sum of the three predicted numbers is fixed by the sample size, and because (2) the allele frequency p was estimated directly from the sample data. (The frequency q can be estimated indirectly as 1 - p and therefore does not reduce the degrees of freedom any further.) The critical value for a chi-square statistic with one degree of freedom is 3.841 (see Table 3.2), which is much greater than the observed value. Consequently, we conclude that the predicted genotype frequencies are in agreement with the observed frequencies in the sample, and furthermore, we infer that in the population from which the sample was obtained, the M-N genotypes are in Hardy–Weinberg proportions—a finding that is not too surprising given that marriage is usually not based on blood type.

The preceding analysis indicates how we can use the Hardy–Weinberg principle to predict genotype frequencies from allele frequencies. Can we turn the Hardy–Weinberg principle around and use it to predict allele frequencies from genotype frequencies? For example, in the United States, the incidence of the recessive metabolic disorder phenylketonuria (PKU) is about 0.0001. Does this statistic allow us to calculate the frequency of the mutant allele that causes PKU?

We cannot proceed as before by counting the different types of alleles, mutant and normal, that are present in the population because heterozygotes and normal homozygotes are phenotypically indistinguishable. Instead, we must proceed by applying the Hardy–Weinberg principle in reverse to estimate the mutant allele frequency. The incidence of PKU, 0.0001, represents the frequency of mutant homozygotes in the population. Under the assumption of random mating, these individuals should occur with a frequency equal to the square of the mutant allele frequency. Denoting this allele frequency by q, we have

$$q^2 = 0.0001 q = \sqrt{0.0001} = 0.01$$

Thus, 1 percent of the alleles in the population are estimated to be mutant. Using the Hardy–Weinberg principle in the usual way, we can then predict the frequency of people in the population who are heterozygous carriers of the mutant allele:

Carrier frequency = 2pq = 2(0.99)(0.01) = 0.0198

Thus, approximately 2 percent of the population are predicted to be carriers.

The Hardy–Weinberg principle also applies to X-linked genes and to genes with multiple alleles. For an X-linked gene such as the one that controls color vision in humans, the allele frequencies are estimated from the frequencies of the genotypes in males, and the frequencies of the genotypes in females are obtained by applying the Hardy–Weinberg principle to these estimated allele frequencies. (We assume, of course, that the allele frequencies are the same in the two sexes.) In northern European populations, for example, about 88 percent of men have normal color vision and about 12 percent are color blind. Thus, in these populations, the frequency of the allele for normal color vision (*C*) is p = 0.88 and the frequency of the allele for color blindness (*c*) is q = 0.12. Under the assumptions of random mating and equal allele frequencies in the two sexes, we have:

Sex	Genotype	Frequency	Phenotype
Males	С	p = 0.88	Normal vision
	С	q = 0.12	Color blind
Females	CC	$p^2 = 0.77$	Normal vision
	Сс	2pq = 0.21	Normal vision
	СС	$q^2 = 0.02$	Color blind

For genes with multiple alleles, the Hardy–Weinberg genotype proportions are obtained by expanding a multinomial expression. For example, the A–B–O blood types are determined by three alleles I^A , I^B , and *i*. If the frequencies of these are *p*, *q*, and *r*, respectively, then the frequencies of the six different genotypes in the A–B–O blood-typing system are obtained by expanding the trinomial $(p + q + r)^2 = p^2 + q^2 + r^2 + 2pq + 2qr + 2pr$:

Blood Type	Genotype	Frequency
A	$I^{A}I^{A}$	<i>p</i> ²
	$I^{A}i$	2pr
В	$I^{\scriptscriptstyle B}I^{\scriptscriptstyle B}$	q^2
	$I^{\scriptscriptstyle B}i$	2qr
AB	$I^{A}I^{B}$	2 <i>pq</i>
0	ii	r^2

EXCEPTIONS TO THE HARDY-WEINBERG PRINCIPLE

There are many reasons why the Hardy–Weinberg principle might not apply to a particular population. Mating might not be random, the members of the population carrying different alleles might not have equal chances of surviving and reproducing, the population might be subdivided into partially isolated units, or it might be an amalgam of different populations that have come together recently by migration. We now briefly consider each of these exceptions to the Hardy–Weinberg principle.

1. *Nonrandom mating*. Random mating is the key assumption underlying the Hardy– Weinberg principle. If mating is not random, the simple relationship between allele frequencies and genotype frequencies breaks down. For example, individuals might mate with each other because they are genetically related. This type of nonrandom mating—called *consanguineous mating* (see Chapter 4)—reduces the frequency of heterozygotes and increases the frequency of homozygotes compared to the Hardy–Weinberg genotype frequencies. We can quantify this effect by using the inbreeding coefficient, F (see Chapter 4). Let's suppose that a gene has two alleles, A and a, with respective frequencies p and q, and that the population in which the gene is segregating has reached a level of inbreeding measured by F. (Recall from Chapter 4 that the range of F is between 0 and 1, with 0 corresponding to no inbreeding and 1 corresponding to complete inbreeding.) The genotype frequencies in this population are given by the following formulas:

Genotype	Frequency with Consanguineous Mating
AA	$p^2 + pqF$
Aa	2pq - 2pqF
aa	$q^2 + pqF$

From these formulas, it is clear that the frequencies of the two homozygotes have increased compared to the Hardy–Weinberg frequencies and that the frequency of the heterozygotes has decreased compared to the Hardy–Weinberg frequency. Notice that for each homozygote, the increase in frequency is exactly half the decrease in the frequency of the heterozygotes. Furthermore, each change in genotype frequency is directly proportional to the inbreeding coefficient. For a population that is completely inbred, F = 1, and the genotype frequencies become:

Genotype	Frequency with $F = 1$
AA	p
Aa	0
Aa	q

To see how the genotype frequencies change with different values of F, work through Solve It: The Effects of Inbreeding on Hardy–Weinberg Frequencies.

2. Unequal survival. If zygotes produced by random mating have different survival rates, we will not expect the genotype frequencies of the individuals that develop from these zygotes to conform to the Hardy–Weinberg predictions. For example, consider a randomly mating population of *Drosophila* that is segregating two alleles, A_1 and A_2 , of an autosomal gene. A sample of 200 adults from this population yielded the following data:

Genotype	Observed Number	Expected Number
A_1A_1	26	46.1
$A_{1}A_{2}$	140	99.8
$A_{2}A_{2}$	34	54.1

The expected numbers were obtained by estimating the frequencies of the two alleles among the flies in the sample; the frequency of the A_1 allele is $(2 \times 26 + 140)/(2 \times 200) = 0.48$, and the frequency of the A_2 allele is 1 - 0.48 = 0.52. Then the Hardy–Weinberg formulas were applied to these estimated frequencies. Obviously, the expected numbers are not in agreement with the observed numbers, which show an excess of heterozygotes and a dearth of both types of homozygotes. Here the disagreement is so obvious that a chi-square calculation to test the goodness of fit between the observed and expected numbers is unnecessary. The explanation for the disagreement probably lies with differential survival of the three genotypes during development from the zygote to the adult stage. The A_1A_2 heterozygotes survive better than either of the two homozygotes. Unequal survival rates can therefore lead to genotype frequencies that deviate from the Hardy–Weinberg predictions.

Solve It!

The Effects of Inbreeding on Hardy–Weinberg Frequencies

An autosomal gene is segregating two alleles, R and r, with respective frequencies 0.3 and 0.7. If mating is random, what are the expected frequencies of the genotypes? Now suppose that every individual in the population mates with a sibling. What will the genotype frequencies be among the offspring? Suppose instead that every individual mates with a first cousin. What will the genotype frequencies be among their offspring? Finally, suppose that after many generations of random mating, every individual in the population reproduces by selffertilization. What will the genotype frequencies be among the offspring of this kind of inbreeding?

To see the solution to this problem, visit the Student Companion site.

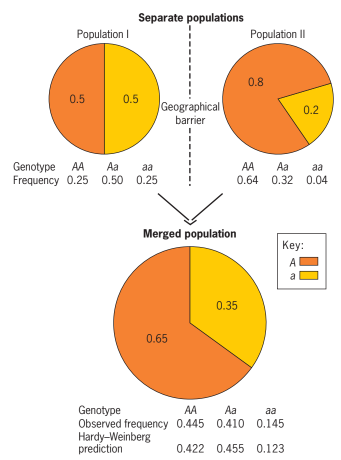
- **3.** *Population subdivision.* When a population is a single interbreeding unit, we say that it is **panmictic. Panmixis** (the noun) implies that any member of the population is able to mate with any other member—that is, there are no geographical or ecological barriers to mating in the population. In nature, however, populations are often subdivided. We can think of fish living in a group of lakes that are intermittently connected by streams, or of birds living on a chain of islands in an archipelago. Such populations are structured by geographical and ecological features that might be correlated with genetic differences. For example, the fish in one lake might have a high frequency of allele *A*, while those in another lake might have a low frequency of this allele. Although the genotype frequencies might conform to Hardy–Weinberg predictions within each lake, across the entire range of the fish population, they will not. Geographical subdivision makes the population genetically inhomogeneous, and such inhomogeneity violates a tacit assumption of the Hardy–Weinberg principle: that allele frequencies are uniform throughout the population.
- **4.** *Migration.* When individuals move from one territory to another, they carry their genes with them. The introduction of genes by recent migrants can alter allele and genotype frequencies within a population and disrupt the state of Hardy–Weinberg equilibrium. As an example, let's consider the situation in **Figure 23.2**. Two populations of equal size are separated by a geographical barrier. In population I the frequencies of *A* and *a* are both 0.5, whereas in population II the frequency of *A* is 0.8 and that of *a* is 0.2. With random mating within each population, the Hardy–Weinberg principle predicts that the two populations will have different genotype frequencies (see Figure 23.2).

Let's suppose that the geographical barrier between the populations breaks down and that the two populations merge completely. In the merged population, the allele frequencies will be the simple averages of the frequencies of the separate

populations; the frequency of A will be (0.5 + 0.8)/2 = 0.65, and the frequency of a will be (0.5 + 0.2)/2 = 0.35. Moreover, the genotype frequencies in the merged population will be the simple averages of the genotype frequencies in the separate populations: the frequency of AA will be (0.25 + 0.64)/2 = 0.445, that of Aa will be (0.50 + 0.32)/2 =0.410, and that of *aa* will be (0.25 + 0.04)/2 = 0.145. Notice, however, that these observed genotype frequencies are not equal to the frequencies predicted by the Hardy-Weinberg principle: $(0.65)^2 = 0.422$ for AA, 2(0.65)(0.35) = 0.455 for Aa, and $(0.35)^2 = 0.123$ for aa. The reason for this discrepancy is that the observed genotype frequencies were not created by random mating within the entire merged population. Rather, they were created by amalgamating genotype frequencies from separate randomly mating populations. Thus, the merger of two randomly mating populations does not produce a population with Hardy-Weinberg genotype frequencies. However, if the merged population mates randomly for just one generation, Hardy-Weinberg genotype frequencies will be established, and the allele frequencies of the merged population will allow prediction of these genotype frequencies. This example demonstrates that merging randomly mating populations temporarily upsets Hardy-Weinberg equilibrium. The migration of individuals from one population to another also causes a temporary upset in Hardy-Weinberg equilibrium. However, if a population that has received migrants mates randomly for just one generation, Hardy–Weinberg equilibrium will be restored.

USING ALLELE FREQUENCIES IN GENETIC COUNSELING

Genetic counselors sometimes use allele frequency data in conjunction with pedigree analysis to calculate the risk that an individual will develop a genetic disease. A simple case is shown in **Figure 23.3**. The man and





woman in generation I have had three children, the last of whom suffered from Tay-Sachs disease, which is caused by an autosomal recessive mutation (t_s) with a frequency of about 0.017 in certain populations. Assuming that the frequency of the mutant allele is 0.017 in II-1's ethnic group, her chance of being a carrier (TS ts) is obtained by using the Hardy-Weinberg principle: 2(0.017)(0.983) =0.033, which is approximately 1/30. The chance that her husband (II-2) is a carrier is determined by analyzing the pedigree. Because II-4 died of Tay-Sachs disease, we know that both I-1 and I-2 were heterozygous for the mutant allele. Either of them could have transmitted this allele to II-2. However, both of them did not transmit it to him because II-2 does not have the disease. Thus, the chance that II-2 is a carrier of the mutant allele is 2/3. To calculate the risk that II-1 and II-2 will have a child with Tay-Sachs disease, we combine the probabilities that each parent is a carrier (1/30 for II-1 and 2/3 for II-2) with the probability that if they are carriers, they will both transmit the mutant allele to their offspring $((1/2) \times (1/2) = 1/4)$. Thus, the risk for the child to have Tay-Sachs disease is $(1/30) \times (2/3) \times (1/4) = 1/180 = 0.006$, which is 20 times the risk for a random child in a population where the mutant allele frequency is 0.017.

- Allele frequencies can be estimated by enumerating the genotypes in a sample from a population.
- Under the assumption of random mating, the Hardy–Weinberg principle allows genotype frequencies for autosomal and X-linked genes to be predicted from allele frequencies.
- The Hardy–Weinberg principle does not apply to populations with consanguineous mating, unequal survival among genotypes, geographic subdivision, or migration.
- The Hardy–Weinberg principle is useful in genetic counseling.

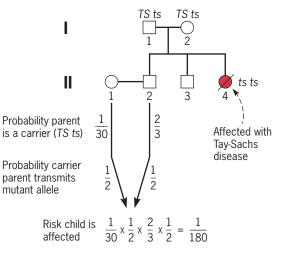


FIGURE 23.3 Pedigree analysis using population data to calculate the risk for Tay-Sachs disease in a child.

KEY POINTS

Natural Selection

Charles Darwin described the key force that drives evolutionary change in populations. He argued that organisms produce more offspring than the environment can support and that a struggle for survival ensues. In the face of this competition, the organ-

Allele frequencies change systematically in populations because of differential survival and reproduction among genotypes.

isms that survive and reproduce transmit to their offspring traits that favor survival and reproduction. After many generations of such competition, traits associated with strong competitive ability become prevalent in the population, and traits associated with weak competitive ability disappear. Selection for survival and reproduction in the face of competition is therefore the mechanism that changes the physical and behavioral characteristics of a species. Darwin called this process **natural selection**.

THE CONCEPT OF FITNESS

To put the mechanism of natural selection into a genetic context, we must recognize that the ability to survive and reproduce is a phenotype—arguably the most important phenotype of all—and that it is determined, at least partly, by genes. Geneticists refer to this ability to survive and reproduce as **fitness**, a quantitative variable they usually symbolize by the letter w. Each member of a population has its own fitness value: 0 if it dies or fails to reproduce, 1 if it survives and produces 1 offspring, 2 if it survives and produces 2 offspring, and so forth. The average of all these values is the average fitness of the population, usually symbolized \overline{w} .

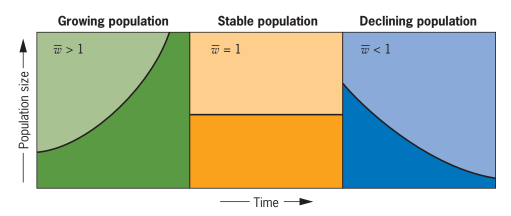


FIGURE 23.4 Significance of average fitness (\overline{w}) for population size as a function of time. Population size grows, is stable, or declines depending on the value of the average fitness.

For a population with a stable size, the average fitness is 1; each individual in such a population produces, on average, one offspring. Of course, some individuals will produce more than one offspring, and some will not produce any offspring at all. However, when the population size is not changing, the average number of offspring (that is, the average fitness) is 1. In a declining population, the average number of offspring is less than 1, and in a growing population it is greater than 1 (**Figure 23.4**).

NATURAL SELECTION AT THE LEVEL OF THE GENE

To see how fitness differences among individuals lead to change in the characteristics of a population, let's assume that fitness is determined by a single gene segregating two alleles, A and a, in a particular species of insect. Furthermore, let's assume that allele A causes the insects to be dark in color, that allele a causes them to be light in color, and that A is completely dominant to a. In a forest habitat, where plant growth is luxuriant, the dark form of the insect survives better than the light form. Consequently, the fitnesses of genotypes AA and Aa are greater than the fitness of genotype aa. By contrast, in open fields, where plant growth is scarce, the light form of the insect survives better than the dark form of the insect survives better than the dark form of the insect survives better than the light form of the insect survives better than the dark form of the insect survives better than the light form of the insect survives better than the dark form of the insect survives better than the light form of the insect survives better than the dark form of the insect survives better than the light form of the insect survives better than the dark form, and the fitness relationships are reversed.

We can express these relationships mathematically by applying the concept of **relative fitness**. In each of the two environments, we arbitrarily define the fitness of the competitively superior genotype(s) to be equal to 1 and express the fitness of the inferior genotype(s) as a deviation from 1. This fitness deviation, usually symbolized by the letter s, is called the **selection coefficient**; it measures the intensity of natural selection acting on the genotypes in the population. We can summarize the fitness relationships among the three insect genotypes in each of the two habitats in the following table:

Genotype:	AA	Aa	aa
Phenotype:	dark	dark	light
Relative fitness in forest habitat:	1	1	$1 - s_1$
Relative fitness in field habitat:	$1 - s_2$	$1 - s_2$	1

These relative fitnesses tell us nothing about the absolute reproductive abilities of the different genotypes in the two habitats. However, they do tell us how well each genotype competes with the other genotypes within a particular environment. Thus, for example, we know that *aa* is a weaker competitor than either *AA* or *Aa* in the forest habitat. How much weaker depends, of course, on the actual value of the selection coefficient, s_1 . If $s_1 = 1$, then *aa* is effectively a lethal genotype (its relative fitness is 0), and we would expect natural selection to reduce the frequency of the *a* allele in

the population. If s_1 were much smaller, say only 0.01, natural selection would still reduce the frequency of the *a* allele, but it would do so very slowly.

To see the effect of natural selection on allele frequencies, let's focus on an insect population in the forest habitat. We will assume that initially the frequency of A is p = 0.5, that the frequency of a is q = 0.5, and that $s_1 = 0.1$. Furthermore, let's assume that the population mates randomly and that the genotypes are present in Hardy– Weinberg frequencies at fertilization each generation. (Differential survival among the genotypes will change these frequencies as the insects mature.) Under these assumptions, the initial genetic composition of the population is:

Genotype:	AA	Aa	aa
Relative fitness:	1	1	1 - 0.1 = 0.9
Frequency (at fertilization):	$p^2 = 0.25$	2pq = 0.50	$q^2 = 0.25$

In forming the next generation, each genotype will contribute gametes in proportion to its frequency and relative fitness. Thus, the relative contributions of the three genotypes will be:

Genotype:	AA	Aa	aa
Relative contribution	$(0.25) \times 1$	$(0.50) \times 1$	$(0.25) \times (0.9)$
to next generation:	= 0.25	= 0.50	= 0.225

If we divide each of these relative contributions by their sum (0.25 + 0.50 + 0.225 = 0.975), we obtain the proportional contributions of each of the genotypes to the next generation:

Genotype:	AA	Aa	aa
Proportional contribution to next generation:	0.256	0.513	0.231

From these numbers we can calculate the frequency of the *a* allele after one generation of selection simply by noting that all the genes transmitted by the *aa* homozygotes are *a* and that half the genes transmitted by the *Aa* heterozygotes are *a*. In the next generation, the frequency of *a*, symbolized q', will be

$$q' = 0.231 + (1/2)(0.513) = 0.487$$

which is slightly less than the starting frequency of 0.5. Thus, in the forest habitat, natural selection, acting through the lower fitness of the *aa* homozygotes, has decreased the frequency of *a* from 0.5 to 0.487. In every subsequent generation, the frequency of *a* will be reduced slightly because of selection against the *aa* homozygotes, and eventually, this allele will be eliminated from the population altogether. **■** Figure 23.5*a* shows how natural selection will drive the *a* allele to extinction. To see what happens when the force of selection is stronger, work through Solve It: Selection against a Harmful Recessive Allele.

In the field habitat, *aa* homozygotes are selectively superior to the other two genotypes. Thus, starting with q = 0.5, Hardy–Weinberg genotype frequencies, and the selection coefficient $s_1 = 0.1$, we have:

Genotype:	AA	Aa	aa
Relative fitness:	1 - 0.1 = 0.9	1 - 0.1 = 0.9	1
Frequency:	0.25	0.50	0.25

After one generation of selection in the field habitat, the frequency of *a* will be q' = 0.513, which is slightly greater than the starting frequency. Every generation afterward, the frequency of *a* will rise, and eventually it will equal 1, at which point we can say that the allele has been fixed in the population. **■** Figure 23.5*b* shows the selection-driven path toward fixation of *a*.

Solve It!

Selection against a Harmful Recessive Allele

Suppose that the frequencies of the alleles *A* and *a* are each 0.5 in a randomly mating population. Predict the frequencies of the three genotypes in this population. Suppose that the *AA* and *Aa* genotypes are equally fit, but that the *aa* homozygotes survive only one-fourth as well as either the *AA* homozygotes or the *Aa* heterozygotes. What are the relative fitnesses of these genotypes? What is the value of the selection coefficient acting against *aa* homozygotes? Among the zygotes of the next generation, predict the frequency of the *a* allele.

To see the solution to this problem, visit the Student Companion site.

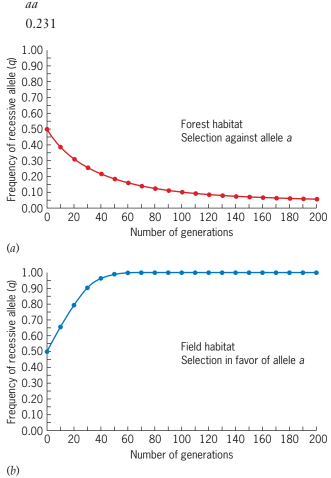


FIGURE 23.5 (a) Selection against the recessive allele a in the forest habitat. (b) Selection in favor of the recessive allele a in the field habitat.

These two scenarios illustrate selection for or against a recessive allele. In the forest habitat, the recessive allele a is deleterious in homozygous condition and selection acts against it. In the field habitat, a is selectively favored over the dominant allele A, which is deleterious in both homozygous and heterozygous condition.

Notice that selection *for* a recessive allele—and therefore against a harmful dominant allele—is more effective than selection *against* a recessive allele. The curve in Figure 23.5*b* shows the time course of selection in favor of a recessive allele. This curve rises steeply to the top of the graph, at which point the recessive allele is fixed in the population. The process shown in this graph efficiently changes the frequency of the recessive allele, and rather quickly gets it to a final value of 1, because every dominant allele in the population is exposed to the purifying action of selection. By virtue of their dominance, these alleles cannot "hide out" in heterozygous condition.

The curve in Figure 23.5*a* shows the time course of selection against a recessive allele. This curve changes more gradually than the curve in Figure 23.5*b* and asymptotically approaches a limit at the bottom of the graph, which represents the loss of the recessive allele. Selection is less effective in this case because it can only act against the recessive allele when it is homozygous. Once the recessive allele has been reduced in frequency, recessive homozygotes will be rare; most of the surviving recessive alleles will therefore be found in heterozygotes, where they are immune from the purifying effect of selection. By comparing the two graphs in Figure 23.5, we see that a harmful recessive allele can linger in a population much longer than a harmful dominant allele.

Studies of the moth *Biston betularia*, an inhabitant of wooded areas in Great Britain, have shown that selection of the type we have been discussing does operate to change allele frequencies in nature. This species, commonly known as the peppered moth, exists in two color forms, light and dark (**Figure 23.6**); the light form is homozygous for a recessive allele c, and the dark form carries a dominant allele C. From 1850 onward, the frequency of the dark form increased in certain areas of England, particularly in the industrialized Midlands section of the country. Around the heavily industrialized cities of Manchester and Birmingham, for example, the frequency of the dark form increased from 1 to 90 percent. This dramatic increase has been attributed to selection against the light form in the soot-polluted landscapes of industrialized areas. In recent times, the level of pollution has abated considerably and the light form of the moth has made a comeback, although not quite to its preindustrial frequencies. Whatever processes have been at work against the light form of the moth appear to have been reversed by environmental restoration in this region of England.



(a)

(b)

FIGURE 23.6 (a) The dark form of the peppered moth on tree bark covered with lichens. (b) The light form of the peppered moth on tree bark covered with soot from industrial pollution.

- Natural selection occurs when genotypes differ in the ability to survive and reproduce—that is, when they differ in fitness.
- The intensity of natural selection is quantified by the selection coefficient.
- At the level of the gene, natural selection changes the frequencies of alleles in populations.

Random Genetic Drift

In his book *The Origin of Species*, Darwin emphasized the role of natural selection as a systematic force in evolution. However, he also recognized that evolution is affected by random processes. New mutants appear

unpredictably in populations. Thus, mutation, the ultimate source of all genetic variability, is a random process that profoundly affects evolution; without mutation, evolution could not occur. Darwin also recognized that inheritance (which he did not understand) is unpredictable. Traits are inherited, but offspring are not exact replicas of their parents; there is always some unpredictability in the transmission of a trait from one generation to the next. In the twentieth century, after Mendel's principles were rediscovered, the evolutionary implications of this unpredictability were investigated by Sewall Wright and R. A. Fisher. From their theoretical analyses, it is clear that the randomness associated with the Mendelian mechanism profoundly affects the evolutionary process. In the following sections, we explore how the uncertainties of genetic transmission can lead to random changes in allele frequencies— a phenomenon called **random genetic drift**.

RANDOM CHANGES IN ALLELE FREQUENCIES

To investigate how the uncertainties associated with the Mendelian mechanism can lead to random changes in allele frequencies, let's consider a mating between two heterozygotes, $Cc \times Cc$, that produces two offspring, which is the number expected if each individual in the population replaces itself (**■** Figure 23.7). We can enumerate the possible genotypes of the two offspring and compute the probability associated with each of the possible combinations by using the methods discussed in Chapter 3. For example, the probability that the first offspring is CC is 1/4, and the probability

that the second offspring is *CC* is also 1/4; thus, the probability that both offspring are *CC* is $(1/4) \times (1/4) = 1/16$. The probability that one of the offspring is *CC* and the other is *Cc* is $(1/4) \times (1/2) \times 2$ (because there are two possible birth orders: *CC* then *Cc*, or *Cc* then *CC*); thus, the probability of observing the genotypic combination *CC* and *Cc* in the two offspring is 1/4. The entire probability distribution for the various genotypic combinations of offspring is given in Figure 23.7. This figure also gives the frequency of the *c* allele associated with each combination.

Among the parents, the frequency of *c* is 0.5. This frequency is the most probable frequency for *c* among the two offspring. In fact, the probability that the frequency of *c* will not change between parents and offspring is 6/16. However, there is an appreciable chance that the frequency of *c* will increase or decrease among the offspring simply because of the uncertainties associated with the Mendelian mechanism. The chance that the frequency of *c* will increase is 5/16, and the chance that it will decrease is also 5/16. Thus, the chance that the frequency of *c* will change in one direction or the other, 5/16 + 5/16 = 10/16, is actually greater than the chance that it will remain the same.

This situation illustrates the phenomenon of random genetic drift. For every pair of parents in the population that is segregating different alleles of a gene, there is a chance that the Mendelian mechanism will

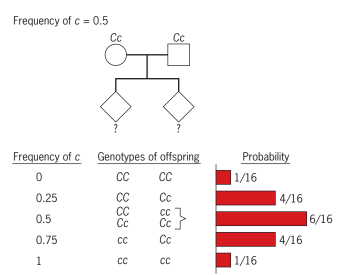


FIGURE 23.7 Probabilities associated with possible frequencies of the allele *c* among the two children of hetero-zygous parents.

KEY POINTS

Allele frequencies change unpredictably in populations because of uncertainties during reproduction. lead to changes in the frequencies of those alleles. When these random changes are summed over all pairs of parents, there may be aggregate changes in the allele frequencies. Thus, the genetic composition of the population can change even without the force of natural selection.

THE EFFECTS OF POPULATION SIZE

A population's susceptibility to random genetic drift depends on its size. In large populations, the effect of genetic drift is minimal, whereas in small ones, it may be the primary evolutionary force. Geneticists gauge the effect of population size by monitoring the frequency of heterozygotes over time. Let's focus, once again, on alleles C and c, with respective frequencies p and q, and let's assume that neither allele has any effects on fitness; that is, C and c are selectively neutral. Furthermore let's assume that the population mates randomly and that in any given generation, the genotypes are present in Hardy–Weinberg proportions.

In a very large population—essentially infinite in size—the frequencies of C and c will be constant and the frequency of the heterozygotes that carry these two alleles will be 2pq. In a small population of finite size N, the allele frequencies will change randomly as a result of genetic drift. Because of these changes, the frequency of heterozygotes, often called the **heterozygosity**, will also change. To express the magnitude of this change over one generation, let's define the current frequency of heterozygotes as H and the frequency of heterozygotes in the next generation as H'. Then the mathematical relationship between H' and H is

$$H' = \left(1 - \frac{1}{2N}\right)H$$

This equation tells us that in one generation, random genetic drift causes the heterozygosity to decline by a factor of $\frac{1}{2N}$. In a total of *t* generations, we would expect the heterozygosity to decline to a level given by the equation

$$H_t = \left(1 - \frac{1}{2N}\right)^t H$$

This equation enables us to see the cumulative effect of random genetic drift over many generations. In each generation, the heterozygosity is expected to decline by a factor of $\frac{1}{2N}$; over many generations, the heterozygosity will eventually be reduced to 0, at which point all genetic variability in the population will be lost. At this point the population will possess only one allele of the gene, and either p = 1 and q = 0, or p = 0 and q = 1. Thus, through random changes in allele frequencies, drift steadily erodes the genetic variability of a population, ultimately leading to the fixation and loss of alleles. It is important to recognize that this process depends critically on the population size (**■** Figure 23.8). Small populations are the most sensitive to the vari-

ability-reducing effects of drift. Large populations are less sensitive. To see how drift might have reduced genetic variability in the population of Pitcairn Island described at the beginning of this chapter, work through Problem-Solving Skills: Applying Genetic Drift to Pitcairn Island.

If selectively neutral alleles of the sort we have been discussing are ultimately destined for fixation or loss, can we determine the probabilities that are associated with these two ultimate outcomes? Let's suppose that at the current time, the frequency of *C* is *p* and that of *c* is *q*. Then, as long as the alleles are selectively neutral and the population mates randomly, the probability that a particular allele will ultimately be fixed in the population is its current frequency—*p* for allele *C* and *q* for allele *c*—and the probability that the allele will ultimately be lost from the population is 1 minus its current frequency, that is, 1 - p for allele *C* and 1 - q for allele *c*. Thus, when random genetic drift is the driving force in evolution, we can assign specific probabilities to the possible evolutionary outcomes, and, remarkably, these probabilities are independent of population size.

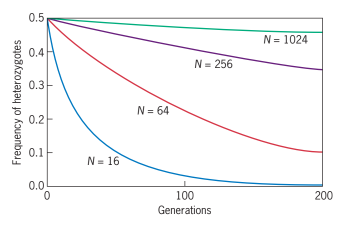


FIGURE 23.8 Decline in the frequency of heterozygotes due to random genetic drift in populations of different size *N*. The populations begin with p = q = 0.5.

PROBLEM-SOLVING SKILLS

Applying Genetic Drift to Pitcairn Island

THE PROBLEM

When Fletcher Christian and his fellow mutineers on the H.M.S. Bounty settled on Pitcairn Island, they didn't realize that they were beginning a genetic experiment. The founding group of men and women brought a finite sample of genes to the island-a sample from two larger populations, Britain and Polynesia. From its beginning in 1790, the Pitcairn Island colony has essentially been a closed system. Some people have left the island, but very few have migrated to it. Most of the alleles that are present on the island today are copies of alleles that were brought there by the colony's founders. Of course, not every allele that was present at the founding is present today. Some alleles were lost through the death or infertility of their carriers. Others have been lost through genetic drift. Let's suppose that the average population size of Pitcairn Island has been 20 and that when the colony was founded, H (the heterozygosity) was 0.20. Let's also suppose that 10 generations have elapsed since the founding of the colony. What is the expected value of *H* today?

FACTS AND CONCEPTS

1. The heterozygosity is a measure of genetic variability in a population.

- **2.** In a population of size N, genetic drift is expected to reduce the heterozygosity by a factor of 1/2N each generation.
- **3.** The loss in variability is cumulative; after t generations, the heterozygosity is given by $H_r = (1 1/2N)^r H_r$.

ANALYSIS AND SOLUTION

To predict the value of H today we can use the equation

 $H_t = [1 - 1/2N]^t H$ with t = 10, N = 20, and H = 0.20: $H_{10} = [1 - 1/2N]^{10} H$ $= [1 - 1/40]^{10} (0.20)$ = [0.78] (0.20)= 0.15

Genetic drift is therefore expected to have reduced the genetic variability on Pitcairn Island, as measured by the heterozygosity, by about 25 percent.

For further discussion visit the Student Companion site.

- Genetic drift, the random change of allele frequencies in populations, is due to uncertainties in Mendelian segregation.
- In diploid organisms, the rate at which genetic variability is lost by random genetic drift is 1/2N, where N is the population size.
- Small populations are more susceptible to drift than large ones.
- Drift ultimately leads to the fixation of one allele at a locus and the loss of all other alleles; the probability that an allele will ultimately be fixed is equal to its current frequency in the population.

Populations in Genetic Equilibrium

In a randomly mating population without selection or drift to change allele frequencies, and without migration or mutation to introduce new alleles, the Hardy– Weinberg genotype frequencies persist indefinitely. Such an idealized population is in a state of genetic The evolutionary forces of mutation, selection, and drift may oppose each other to create a dynamic equilibrium in which allele frequencies no longer change.

equilibrium. In reality, the situation is much more complicated; selection and drift, migration and mutation are almost always at work changing the population's genetic composition. However, these evolutionary forces may act in contrary ways to create a *dynamic equilibrium* in which there is no net change in allele frequencies. This type of equilibrium differs fundamentally from the equilibrium of the ideal Hardy–Weinberg population. In a dynamic equilibrium, the population simultaneously tends to change in opposite directions, but these opposing tendencies cancel each other and bring the

KEY POINTS



population to a point of balance. In the ideal Hardy–Weinberg equilibrium, the population does not change because there are no evolutionary forces at work. We now explore how opposing evolutionary forces can create a dynamic equilibrium within a population.

BALANCING SELECTION

One type of dynamic equilibrium arises when selection favors the heterozygotes at the expense of each type of homozygote in the population. In this situation, called *balancing selection* or *beterozygote advantage*, we can assign the relative fitness of the heterozygotes to be 1 and the relative fitnesses of the two types of homozygotes to be less than 1:

Genotype:	AA	Aa	aa
Relative fitness:	1 - s	1	1 - t

In this formulation, the terms 1 - s and 1 - t contain selection coefficients that are assumed to lie between 0 and 1. Thus, each of the homozygotes has a lower fitness than the heterozygotes. The superiority of the heterozygotes is sometimes referred to as *overdominance*.

In cases of heterozygote advantage, selection tends to eliminate both the A and a alleles through its effects on the homozygotes, but it also preserves these alleles through its effects on the heterozygotes. At some point these opposing tendencies balance each other, and a dynamic equilibrium is established. To determine the frequencies of the two alleles at the point of equilibrium, we must derive an equation that describes the process of selection, and then solve this equation for the allele frequencies when the opposing selective forces are in balance—that is, when the allele frequencies are no longer changing (Table 23.2). At the balance point, the frequency of A is p = t/(s + t), and the frequency of a is q = s/(s + t).

As an example, let's suppose that the AA homozygotes are lethal (s = 1) and that the aa homozygotes are 50 percent as fit as the heterozygotes (t = 0.5). Under these assumptions, the population will establish a dynamic equilibrium when p = 0.5/(0.5 + 1) = 1/3 and q = 1/(0.5 + 1) = 2/3. Both alleles will be maintained at appreciable frequencies by selection in favor of the heterozygotes—a condition known as a **balanced polymorphism**.

In humans, sickle-cell disease is associated with a balanced polymorphism. Individuals with this disease are homozygous for a mutant allele of the β -globin gene, denoted HBB^S , and they suffer from a severe form of anemia in which the hemoglobin molecules crystallize in the blood. This crystallization causes the red blood cells to assume a characteristic sickle shape. Because sickle-cell disease is usually fatal without medical treatment, the fitness of HBB^SHBB^S homozygotes has historically been 0. However, in some parts of the world, particularly in tropical Africa, the frequency of the HBB^S allele is as high as 0.2. With such harmful effects, why does the HBB^S allele remain in the population at all?

TABLE 23.2

Calculating Equilibrium Allele Frequencies with Balancing Selection

Genotypes: Relative fitnesses: Frequencies:	AA 1 - s p ²	Аа 1 2pq	aa 1 — t q²
Average relative fitness: $\overline{w} = p^2 \times (1 - s) + 2pq \times 1 + q^2 \times (1 - t)$ Frequency of A in the next generation after selection: $p' = [p^2(1 - s) + (1/2)2pq]/\overline{w} = p(1 - sp)/\overline{w}$			
Change in frequency of A	due to selecti	ion:	
$\Delta p = p' - p = pq(tq - sp)/\overline{w}$			
At equilibrium, $\Delta p = 0$; p	= t/(s + t) an	d q = s/(s + t)	

The answer is that there is moderate selection against homozygotes that carry the wild-type allele HBB^4 . These homozygotes are less fit than the HBB^SHBB^4 heterozygotes because they are more susceptible to infection by the parasites that cause malaria (**Figure 23.9**), a fitness-reducing disease that is widespread in regions where the frequency of the HBB^S allele is high. We can schematize this situation by assigning relative fitnesses to each of the genotypes of the β -globin gene:

Genotype:	HBB ^S HBB ^S	$HBB^{S}HBB^{A}$	$HBB^{A}HBB^{A}$
Relative fitness:	1 - s	1	1 - t

If we assume that the equilibrium frequency of HBB^{s} is p = 0.1—a typical value in West Africa—and if we note that s = 1 because the $HBB^{s}HBB^{s}$ homozygotes die, we can estimate the intensity of selection against the $HBB^{4}HBB^{4}$ homozygotes because of their greater susceptibility to malaria:

$$p = t/(s + t)$$

$$0.1 = t/(1 + t)$$

$$t = (0.1)/(0.9) = 0.11$$

This result tells us that the $HBB^{A}HBB^{A}$ homozygotes are about 11 percent less fit than the $HBB^{S}HBB^{A}$ heterozygotes. Thus, the selective inferiority of the $HBB^{S}HBB^{S}$ and $HBB^{A}HBB^{A}$ homozygotes compared to the heterozygotes creates a balanced polymorphism in which both alleles of the β -globin gene are maintained in the population.

Various other mutant *HBB* alleles are found at appreciable frequencies in tropical and subtropical regions of the world in which malaria is—or was—endemic. It is plausible that these alleles have also been maintained in human populations by balancing selection.

MUTATION-SELECTION BALANCE

Another type of dynamic equilibrium is created when selection eliminates deleterious alleles that are produced by recurrent mutation. For example, let's consider the case of a deleterious recessive allele *a* that is produced by mutation of the wild-type allele *A* at rate *u*. A typical value for *u* is 3×10^{-6} mutations per generation. Even though this rate is very low, over time, the mutant allele will accumulate in the population, and, because it is recessive, it can be carried in heterozygous condition without having any harmful effects. At some point, however, the mutant allele will become frequent enough for *aa* homozygotes to appear in the population, and these will be subject to the force of selection in proportion to their frequency and the value of the selection coefficient *s*. Selection against these homozygotes will counteract the force of mutation, which introduces the mutant allele into the population.

If we assume that the population mates randomly, and if we denote the frequency of A as p and that of a as q, then we can summarize the situation as follows:

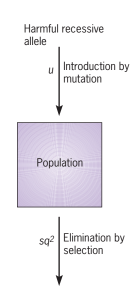
Mutation produces <i>a</i>		Select elimir	ion nates <i>a</i>	
$A \rightarrow a$	Genotype:	AA	Aa	aa
rate = u	Relative fitness:	1	1	1 - s
	Frequency:	p^2	2pq	q^2

Mutation introduces mutant alleles into the population at rate u, and selection eliminates them at rate sq^2 (**\blacksquare** Figure 23.10). When these two processes are in balance, a dynamic equilibrium will be established. We can calculate the frequency of the mutant allele at the equilibrium created by mutation–selection balance by equating the rate of mutation to the rate of elimination by selection:



aracita Placmodiu

FIGURE 23.9 The malaria parasite *Plasmodium falciparum* (yellow) emerging from red blood cells that it had infected.



■ **FIGURE 23.10** Mutation-selection balance for a deleterious recessive allele with frequency *q*. Genetic equilibrium is reached when the introduction of the allele into the population by mutation at rate *u* is balanced by the elimination of the allele by selection with intensity *s* against the recessive homozygotes.

$$u = sq^2$$

Thus, after solving for q, we obtain

$$q = \sqrt{u/s}$$

For a mutant allele that is lethal in homozygous condition, s = 1, and the equilibrium frequency of the mutant allele is simply the square root of the mutation rate. If we use the value for u that was given above, then for a recessive lethal allele the equilibrium frequency is q = 0.0017. If the mutant allele is not completely lethal in homozygous condition, then the equilibrium frequency will be higher than 0.0017 by a factor that depends on $1/\sqrt{s}$. or example, if s is 0.1, then at equilibrium the frequency of this slightly deleterious allele will be q = 0.0055, or 3.2 times greater than the equilibrium frequency of a recessive lethal allele.

Studies with natural populations of *Drosophila* have indicated that lethal alleles are less frequent than the preceding calculations predict. The discrepancy between the observed and predicted frequencies has been attributed to partial dominance of the mutant alleles—that is, these alleles are not completely recessive. Natural selection appears to act against deleterious alleles in heterozygous condition as well as in homozygous condition. Thus, the equilibrium frequencies of these alleles are lower than we would otherwise predict. Selection that acts against mutant alleles in homozygous or heterozygous condition is sometimes called *purifying selection*.

MUTATION-DRIFT BALANCE

We have already seen that random genetic drift eliminates variability from a population. Without any counteracting force, this process would eventually make all populations completely homozygous. However, mutation replenishes the variability that is lost by drift. At some point, the opposing forces of mutation and genetic drift come into balance and a dynamic equilibrium is established.

Previously we saw that genetic variability can be quantified by calculating the frequency of heterozygotes in a population—a statistic called the heterozygosity, which is symbolized by the letter H. The frequency of homozygotes in a population—often called the *homozygosity*—is equal to 1 - H. Over time, genetic drift decreases H and increases 1 - H, and mutation does just the opposite (**■** Figure 23.11). Let's assume that each new mutation is selectively neutral. In a randomly mating population of size N, the rate at which drift decreases H is $(\frac{1}{2N})H$ (see the earlier section, The Effects of Population Size). The rate at which mutation increases H is proportional to the frequency of the homozygotes in the population (1 - H) and the probability that one of the two alleles in a particular homozygote. This probability is simply the mutation rate u for each of the two alleles in the homozygote; thus, the total probability of mutation converting a particular homozygote into a heterozygote is 2u. The rate at which mutation increases H in a population is therefore equal to 2u(1 - H).

When the opposing forces of mutation and drift come into balance, the population will achieve an equilibrium level of variability denoted by \hat{H} . We can calculate this equilibrium value of H by equating the rate at which mutation increases H to the rate at which drift decreases it:

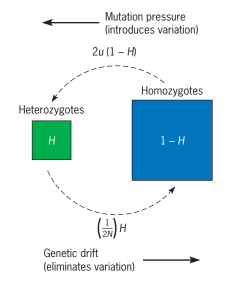
$$2u(1-H) = \left(\frac{1}{2N}\right)H$$

By solving for H, we obtain the equilibrium heterozygosity at the point of mutationdrift balance:

$$\hat{H} = 4Nu/(4Nu+1)$$

Thus, the equilibrium level of variability (as measured by the heterozygosity) is a function of the population size and the mutation rate.

If we assume that the mutation rate is $u = 1 \times 10^{-6}$, we can plot \hat{H} for different values of N (**I** Figure 23.12). For N < 10,000, the equilibrium frequency



■ **FIGURE 23.11** Mutation-drift balance for variability as measured by the frequency of heterozygotes *H* in a population of size *N*. An equilibrium frequency of heterozygotes is reached when the introduction of variability by mutation at rate *u* is balanced by the elimination of variability by genetic drift at rate $\frac{1}{2N}$.

of heterozygotes in the population is quite low; thus, drift dominates over mutation in small populations. For N equal to 1/u, the reciprocal of the mutation rate, the equilibrium frequency of heterozygotes is 0.8, and for even greater values of N, the frequency of heterozygotes increases asymptotically toward 1. Thus, in large populations, mutation dominates over drift; every mutational event creates a new allele, and each new allele contributes to the heterozygosity because the large size of the population protects the allele from being lost by random genetic drift.

Values of \hat{H} in natural populations vary among species. In the African cheetah, for example, \hat{H} is 1 percent or less among a sample of loci, suggesting that over evolutionary time, population size in this species has been small. In humans, \hat{H} is estimated to be about 12 percent, suggesting that over evolutionary time population size has averaged about 30,000 to 40,000 individuals. Estimates of population size that are derived from heterozygosity data are typically much smaller than estimates obtained from census data. The reason for this discrepancy is that the estimates

based on heterozygosity data are *genetically effective* population sizes—sizes that take into account restrictions on mating and reproduction, as well as temporal fluctuations in the number of mating individuals. The genetically effective size of a population is almost always less than the census size of a population.

- Selection involving heterozygote superiority (balancing selection) creates a dynamic equilibrium in which different alleles are retained in a population despite their being harmful in homozygotes.
- In humans sickle-cell disease is associated with balancing selection at the locus for β -globin.
- Selection against a deleterious recessive allele that is replenished in the population by mutation leads to a dynamic equilibrium in which the frequency of the recessive allele is a simple function of the mutation rate and the selection coefficient: $q = \sqrt{us}$.
- A population's acquisition of selectively neutral alleles through mutation is balanced by the loss of these alleles through genetic drift. At equilibrium, the frequency of heterozygotes involving these alleles is a function of the population's size and the mutation rate: H = 4Nu/(4Nu + 1).

1.0 At rospective for the second sec

■ FIGURE 23.12 Equilibrium frequency of heterozygotes (heterozygosity) under mutation-drift balance as a function of genetically effective population size. The mutation rate is assumed to be 10⁻⁶.

KEY POINTS

Basic Exercises

Illustrate Basic Genetic Analysis

1. Calculate the allele frequencies from the following population data:

Answer: The basic calculation	s are summarized in the following
table:	

Genotype	Number
AA	68
Aa	42
aa	24
Total	134

table: H–W Obs.–Exp. Genotype Obs No Frequency Exp. No No

Genotype	Obs. No.	Frequency	Exp. No.	No.
AA	68	$p^2 = 0.441$	59.1	8.9
Aa	42	2pq = 0.446	59.8	-17.8
aa	24	$q^2 = 0.113$	15.1	8.9

- Answer: The frequency of the *A* allele, *p*, is $[(2 \times 68) + 42]/(2 \times 134) = 0.664$. The frequency of the *a* allele, *q*, is $[(2 \times 24) + 42]/(2 \times 134) = 0.336$.
- 2. Predict the Hardy–Weinberg genotype frequencies using the allele frequencies calculated in Exercise 1. Are these frequencies in agreement with the observed frequencies?

To test for agreement between the observed and expected numbers, we calculate a χ^2 test statistic with 1 degree of freedom: $\chi^2 = \Sigma (\text{Obs.} - \text{Exp.})^2/\text{Exp} = 12.0$, which exceeds the critical value for this test statistic. Thus, we reject the hypothesis that the genotype frequencies calculated from the Hardy–Weinberg principle agree with the observed

frequencies. Evidently, the population is not in Hardy-Weinberg equilibrium.

- 3. In a population that has been mating randomly for many generations, two phenotypes are segregating; one is due to a dominant allele G, the other to a recessive allele g. The frequencies of the dominant and recessive phenotypes are 0.7975 and 0.2025, respectively. Estimate the frequencies of the dominant and recessive alleles.
- **Answer:** The frequency of the dominant phenotype represents the sum of two Hardy–Weinberg genotype frequencies: $p^2(GG) + 2pq(Gg)$. The frequency of the recessive phenotype represents just one Hardy–Weinberg genotype frequency, $q^2(gg)$. To estimate the frequency of the recessive allele, we take the square root of the observed frequency of the recessive phenotype: $q = \sqrt{0.2025} = 0.45$. The frequency of the dominant allele is obtained by subtraction: p = 1 - q = 0.55.
- **4.** A gene with two alleles is segregating in a population. The fitness of the recessive homozygotes is 90 percent that of the heterozygotes and the dominant homozygotes. What is the value of the selection coefficient that measures the intensity of natural selection against the recessive allele?
- **Answer:** Using *s* to represent the selection coefficient, the fitness scheme is

Genotype	Relative Fitnes	
AA	1	
Aa	1	
aa	1 - s	

Because the recessive homozygotes are 90 percent as fit as either of the other genotypes, the expression 1 - s = 0.9; thus, s = 0.1.

- 5. Suppose that the alleles of the T gene are selectively neutral. In a population of 50 individuals, currently 34 are heterozygotes. Predict the frequency of heterozygotes in this population 10 generations in the future. Assume that the population size is constant and that mating is completely random (including the possibility of self-fertilization).
- **Answer:** For a selectively neutral gene, evolution occurs by random genetic drift. The governing equation is $H_t = (1 \frac{1}{2N})^t H$, where H_t is the frequency of heterozygotes *t* generations in the future, *N* is the population size, and *H* is the frequency of heterozygotes now. From the data given in the problem, N = 50, H = 34/50 = 0.68, and t = 10. Thus, $H_t = (0.99)^{10} \times (0.68) = 0.615$.
- 6. Purifying selection eliminates deleterious alleles from a population, but recurrent mutation replenishes them. Suppose that recessive lethal alleles of the *B* gene are created at the rate of 2×10^{-6} per generation. What is the expected frequency of lethal alleles in a population in mutation–selection equilibrium?
- **Answer:** The frequency of lethal alleles is given by the equation $q = \sqrt{u/s}$, where *u* is the mutation rate (from dominant normal allele to recessive lethal allele) and *s* is the intensity of selection against the deleterious allele (in this case, *s* = 1). Thus, the expected frequency of lethal alleles in the population is $q = \sqrt{2 \times 10^{-6}} = 0.0014$.

Testing Your Knowledge

Integrate Different Concepts and Techniques

1. The A–B–O blood types of 1000 people from an isolated village were determined to obtain the following data:

Blood Type	Number of People
A	42
В	672
AB	36
0	250

Estimate the frequencies of the I^A , I^B , and i alleles of the A–B–O blood group gene from these data.

Answer: Let's symbolize the frequencies of the *I*⁴, *I*⁸, and *i* alleles of the *I* gene as *p*, *q*, and *r*, respectively, and let's assume that the genotypes of this gene are in Hardy–Weinberg proportions. We begin by estimating *r*, the frequency of the *i*

allele. To obtain this estimate, we note that the frequency of the O blood type, which is 250/1000 = 0.25 in the data, should correspond to the Hardy–Weinberg frequency of the *ii* genotype, r^2 . Thus, if we use the Hardy–Weinberg principle in reverse, we can estimate the frequency of the *i* allele as $r = \sqrt{0.250} = 0.500$.

To estimate *p*, the frequency of the I^A allele, we note that $(p + r)^2 = p^2 + 2pr + r^2$ corresponds to the combined frequencies of the A $(p^2 + 2pr)$ and O (r^2) blood types. From the data, these combined frequencies are estimated to be (42 + 250)/1000 = 0.292. If we set $(p + r)^2 = 0.292$ and take the square root, we obtain p + r = 0.540; then, by subtracting *r*, we can estimate the frequency of the I^A allele as p = 0.540 - 0.500 = 0.040. To estimate *q*, the frequency of the I^B allele, we note that p + q + r = 1. Thus, q = 1 - p - r = 1 - 0.040 - 0.500 = 0.460.

- 2. A man and a woman who both have normal color vision have had three children, including a male who is color blind. The incidence of color-blind males in the population from which this couple came is 0.30, which is unusually high for X-linked color blindness. If the color-blind male marries a female with normal color vision, what is the chance that their first child will be color blind?
- **Answer:** Clearly, the risk that the couple will have a color-blind child depends on the female's genotype. If the female is heterozygous for the allele for color blindness, she has a probability of 1/2 of transmitting this allele to her first child. The male will transmit either an X chromosome, which carries the mutant allele, or a Y chromosome; in either case, the female's contribution to the zygote will be determinative. To obtain the probability that the female is heterozygous for the mutant allele, we note that the incidence of color blindness among males in the population is 0.30; this number provides an estimate of the frequency of the mutant allele, q, in the population. Furthermore, because q = 0.30, the frequency of the wild-type allele, p, is 1 - q = 0.70. If the genotypes in the population are in Hardy-Weinberg proportions, then the frequency of heterozygous females is $2pq = 2 \times (0.7) \times (0.3) = 0.42$. However, among females who have normal color vision, the frequency of heterozygotes is greater because homozygous mutant females have been excluded from the total. To adjust for this effect, we calculate the ratio of heterozyotes to wild-type homozygotes plus heterozygotes and specifically exclude the mutant homozygotes-that is, we compute $2pq/(p^2 + 2pq) = 2pq/[p(p + 2q)] = 2q/(p + q + q) =$ 2q/(1 + q). Substituting q = 0.3 into the last expression, we estimate the frequency of heterozygotes among females

with normal color vision (wild-type homozygotes plus heterozygotes) to be $2 \times (0.3)/(1 + 0.3) = 0.46$. This number is the chance that the female in question is a heterozygous carrier of the mutant allele. The probability that her first child will be color blind is the chance that she is a carrier (0.46) times the chance that she will transmit the mutant allele to her child (1/2); thus, the risk for the child to be color blind is (0.46) \times (1/2) = 0.23.

- 3. The HBB^{s} allele responsible for sickle-cell disease is maintained in many human populations because in heterozygous condition it confers some resistance to infection by malaria parasites; however, in homozygous condition, this allele is essentially lethal. Thus, as malaria is eradicated we might expect the HBB^{s} allele to disappear from human populations. If the normal allele HBB^{4} mutates to HBB^{s} at a rate of 10^{-8} per generation, what ultimate frequency would you predict for the HBB^{s} allele in a malaria-free world?
- **Answer:** In a malaria-free world, the advantage of maintaining the *HBB^s* allele in a balanced polymorphism would disappear. *HBB^sHBB⁴* heterozygotes would have the same fitness as *HBB⁴HBB⁴* homozygotes, and *HBB^sHBB^s* homozygotes would continue to have very low fitness—essentially zero compared to the other two genotypes. Under these circumstances, the frequency of the *HBB^s* allele (q) would be determined by a balance between selection against it in homozygous condition (selection coefficient s = 1) and introduction into the population by mutation at rate u = 10^{-8} per generation. The equilibrium frequency of the *HBB^s* allele would be $q = \sqrt{u/s} = 0.0001$, a thousandfold less than its current frequency in malaria-infested regions of the world.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

23.1 The following data for the M-N blood types were obtained from native villages in Central and North America:

Group	Sample Size	Μ	MN	Ν
Central American	86	53	29	4
North American	278	78	61	139

Calculate the frequencies of the L^M and L^N alleles for the two groups.

- **23.2** The frequency of an allele in a large randomly mating population is 0.2. What is the frequency of heterozygous carriers?
- **23.3** The incidence of recessive albinism is 0.0004 in a human population. If mating for this trait is random in the population, what is the frequency of the recessive allele?

- **23.4** In a sample from an African population, the frequencies of the *L^M* and *L^N* alleles were 0.78 and 0.22, respectively. If the population mates randomly with respect to the M-N blood types, what are the expected frequencies of the M, MN, and N phenotypes?
- **23.5** Human beings carrying the dominant allele T can taste the substance phenylthiocarbamide (PTC). In a population in which the frequency of this allele is 0.4, what is the probability that a particular taster is homozygous?
- **23.6** oo A gene has three alleles, A_1, A_2 , and A_3 , with frequencies 0.6, 0.3, and 0.1, respectively. If mating is random, predict the combined frequency of all the heterozygotes in the population.
- **23.7** Hemophilia is caused by an X-linked recessive allele. In a particular population, the frequency of males with

hemophilia is 1/4000. What is the expected frequency of females with hemophilia?

- **23.8** In *Drosophila* the ruby eye phenotype is caused by a recessive, X-linked mutant allele. The wild-type eye color is red. A laboratory population of *Drosophila* is started with 25 percent ruby-eyed females, 25 percent homozygous red-eyed females, 5 percent ruby-eyed males, and 45 percent red-eyed males. (a) If this population mates randomly for one generation, what is the expected frequency of ruby-eyed males and females? (b) What is the frequency of the recessive allele in each of the sexes?
- **23.9** A trait determined by an X-linked dominant allele shows 100 percent penetrance and is expressed in 36 percent of the females in a population. Assuming that the population is in Hardy–Weinberg equilibrium, what proportion of the males in this population express the trait?
- **23.10** A phenotypically normal couple has had one normal child and a child with cystic fibrosis, an autosomal recessive disease. The incidence of cystic fibrosis in the population from which this couple came is 1/500. If their normal child eventually marries a phenotypically normal person from the same population, what is the risk that the newlyweds will produce a child with cystic fibrosis?
- **23.11** What frequencies of alleles *A* and *a* in a randomly mating population maximize the frequency of heterozygotes?
- **23.12** In an isolated population, the frequencies of the I^A , I^B , and *i* alleles of the A–B–O blood type gene are, respectively, 0.15, 0.25, and 0.60. If the genotypes of the A–B–O blood type gene are in Hardy–Weinberg proportions, what fraction of the people who have type A blood in this population are expected to be homozygous for the I^A allele?
- **23.13** In a survey of moths collected from a natural population, a researcher found 51 dark specimens and 49 light specimens. The dark moths carry a dominant allele, and the light moths are homozygous for a recessive allele. If the population is in Hardy–Weinberg equilibrium, what is the estimated frequency of the recessive allele in the population? How many of the dark moths in the sample are likely to be homozygous for the dominant allele?
- **23.14** A population of Hawaiian *Drosophila* is segregating two alleles, P^1 and P^2 , of the phosphoglucose isomerase (*PGI*) gene. In a sample of 100 flies from this population, 30 were P^1P^1 homozygotes, 60 were P^1P^2 heterozygotes, and 10 were P^2P^2 homozygotes. (a) What are the frequencies of the P^1 and P^2 alleles in this sample? (b) Perform a chi-square test to determine if the genotypes in the sample are in Hardy–Weinberg proportions. (c) Assuming that the sample is representative of the population, how many generations of random mating would be required to establish Hardy–Weinberg proportions in the population?
- **23.15** In a large population that reproduces by random mating, the frequencies of the genotypes *GG*, *Gg*, and *gg* are 0.04, 0.32, and 0.64, respectively. Assume that a change in the

climate induces the population to reproduce exclusively by self-fertilization. Predict the frequencies of the genotypes in this population after many generations of selffertilization.

- **23.16** \bigcirc The frequencies of the alleles *A* and *a* are 0.6 and 0.4, respectively, in a particular plant population. After many generations of random mating, the population goes through one cycle of self-fertilization. What is the expected frequency of heterozygotes in the progeny of the self-fertilized plants?
- **23.17** Each of two isolated populations is in Hardy–Weinberg equilibrium with the following genotype frequencies:

Genotype:	AA	Aa	aa
Frequency in Population 1:	0.04	0.32	0.64
Frequency in Population 2:	0.64	0.32	0.04

- (a) If the populations are equal in size and they merge to form a single large population, predict the allele and genotype frequencies in the large population immediately after merger.
- (b) If the merged population reproduces by random mating, predict the genotype frequencies in the next generation.
- (c) If the merged population continues to reproduce by random mating, will these genotype frequencies remain constant?
- **23.18** A population consists of 25 percent tall individuals (genotype TT), 25 percent short individuals (genotype tt), and 50 percent individuals of intermediate height (genotype Tt). Predict the ultimate phenotypic and genotypic composition of the population if, generation after generation, mating is strictly assortative (that is, tall individuals mate with tall individuals, short individuals mate with short individuals, and intermediate individuals mate with intermediate individuals).
- **23.19** In controlled experiments with different genotypes of an insect, a researcher has measured the probability of survival from fertilized eggs to mature, breeding adults. The survival probabilities of the three genotypes tested are: 0.92 (for GG), 0.90 (for Gg), and 0.56 (for gg). If all breeding adults are equally fertile, what are the relative fitnesses of the three genotypes? What are the selection coefficients for the two least fit genotypes?
- 23.20 In a large randomly mating population, 0.84 of the individuals express the phenotype of the dominant allele A and 0.16 express the phenotype of the recessive allele a.(a) What is the frequency of the dominant allele? (b) If the aa homozygotes are 5 percent less fit than the other two genotypes, what will the frequency of A be in the next generation?
- **23.21** Because individuals with cystic fibrosis die before they can reproduce, the coefficient of selection against them is s = 1. Assume that heterozygous carriers of the recessive mutant allele responsible for this disease are as fit as

wild-type homozygotes and that the population frequency of the mutant allele is 0.02. (a) Predict the incidence of cystic fibrosis in the population after one generation of selection. (b) Explain why the incidence of cystic fibrosis hardly changes even with s = 1.

23.22 For each set of relative fitnesses for the genotypes AA, Aa, and aa, explain how selection is operating. Assume that 0 < t < s < 1.

	AA	Aa	aa
Case 1	1	1	1 - s
Case 2	1 - s	1 - s	1
Case 3	1	1 - t	1 - s
Case 4	1 - s	1	1 - t

- **23.23** The frequency of newborn infants homozygous for a recessive lethal allele is about 1 in 25,000. What is the expected frequency of carriers of this allele in the population?
- **23.24** A population of size 50 reproduces in such a way that the population size remains constant. If mating is random, how rapidly will genetic variability, as measured by the frequency of heterozygotes, be lost from this population?
- **23.25** A population is segregating three alleles, A_1 , A_2 , and A_3 , with frequencies 0.2, 0.5, and 0.3, respectively. If these alleles are selectively neutral, what is the probability that A_2 will ultimately be fixed by genetic drift? What is the probability that A_3 will ultimately be lost by genetic drift?

- **23.26** A small island population of mice consists of roughly equal numbers of males and females. The Y chromosome in one-fourth of the males is twice as long as the Y chromosome in the other males because of an expansion of heterochromatin. If mice with the large Y chromosome have the same fitness as mice with the small Y chromosome, what is the probability that the large Y chromosome will ultimately be fixed in the mouse population?
- 23.27 In some regions of West Africa, the frequency of the *HBB^s* allele is 0.2. If this frequency is the result of a dynamic equilibrium due to the superior fitness of *HBB^sHBB⁴* heterozygotes, and if *HBB^sHBB^s* homo-zygotes are essentially lethal, what is the intensity of selection against the *HBB⁴HBB⁴* homozygotes?
- 23.28 Mice with the genotype *Hb* are twice as fit as either of the homozygotes *HH* and *bb*. With random mating, what is the expected frequency of the *b* allele when the mouse population reaches a dynamic equilibrium because of balancing selection?
- **23.29** A completely recessive allele g is lethal in homozygous condition. If the dominant allele G mutates to g at a rate of 10^{-6} per generation, what is the expected frequency of the lethal allele when the population reaches mutation-selection equilibrium?
- **23.30** Individuals with the genotype bb are 20 percent less fit than individuals with the genotypes BB or Bb. If B mutates to b at a rate of 10^{-6} per generation, what is the expected frequency of the allele b when the population reaches mutation–selection equilibrium?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

The mutant allele that causes sickle-cell disease is prevalent in areas where people have a high probability of contracting malaria, which is caused by a parasite transmitted by mosquitoes. Click on the links for Malaria and Mosquito on the Genomic biology page to find information on the malaria parasite *Plasmodium falciparum* and on the mosquito vector *Anopheles gambiae*.

1. How large is the *Plasmodium* genome? How many chromosomes does it comprise? How large is the *Anopheles* genome? How many chromosomes does it comprise? Have the genomes of these organisms been sequenced completely?

2. On the *Plasmodium* web page, click on the overview link to bring up a page with summary information on this parasite. Under related resources, click on WHO/Malaria info to bring up a page with links to information about various aspects of malaria. How widespread is the disease? How is it being treated today? How is the *Plasmodium* parasite transmitted from one person to another?



Evolutionary Genetics

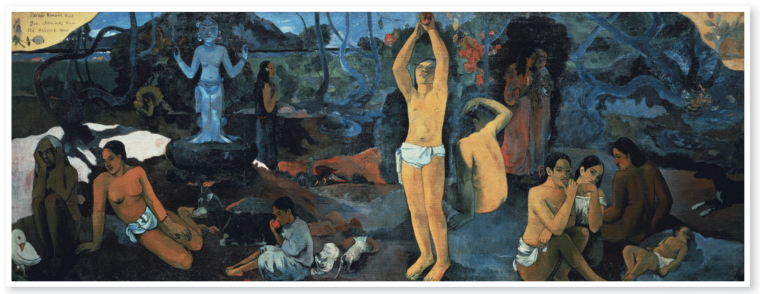
CHAPTER OUTLINE

- The Emergence of Evolutionary Theory
- Genetic Variation in Natural Populations
- Molecular Evolution
- Speciation
- Human Evolution

D'ou venons nous? Que sommes nous? Ou allons nous?

In 1897 in Tahiti, the French artist Paul Gauguin created an enormous painting with a provocative title: "Where do we come from? What are we? Where are we going?" The painting, now on display in the Boston Museum of Fine Arts, shows a group of Polynesian people, both young and old, reclining, sitting, walking, and eating in a strangely colored landscape. The figures are forlorn and abstracted, and a few of them seem to stare interrogatively at the viewer, posing, as it were, those three haunting questions that Gauguin inscribed in the painting's margin. This melancholy canvas, created near the end of Gauguin's life, seems to depict the artist's personal search for answers to some of life's deep questions. However, it is more than the statement of an individual who sought inspiration, freedom, and fulfillment in the South Seas. Gauguin's painting reflects a universal quest for what it means to be human. During the nineteenth century,

people began to see this issue in a new light, especially with the emergence of evolutionary theory. Charles Darwin's *The Origin of Species*, first published in 1859, advanced the ideas that species are not fixed and that populations of organisms change over time. In a later book, *The Descent of Man*, Darwin proposed that the human species was also subject to evolutionary forces. Darwin's ideas have troubled many people.



"Where do we come from? What are we? Where are we going?" An 1897 painting by the French artist Paul Gauguin.

The Emergence of Evolutionary Theory

The publication of *The Origin of Species* in 1859 provoked a storm of controversy—not because the idea that species evolve was new, but rather because Darwin made the case for it so well. Darwin's book was cogently

The theory of evolution, initially enunciated by Charles Darwin, is based on genetic principles.

written and rich in evidence. He argued that species change gradually over long periods of time. Some species split into two or more separate species; other species become extinct. Darwin's ideas were unsettling to many people who held to the notion that each species was divinely created and that except for trivial variations among individuals, species do not change—that is, they are immutable. Darwin's book contested this view. Although he did not say much about the origin of the first organisms on Earth, he argued that during millions of years they had changed and diversified to produce the plethora of species now alive. Furthermore, Darwin argued that this change and diversification, what he called "divergence of character," was the result of purely natural processes.

DARWIN'S THEORY OF EVOLUTION

Darwin proposed that a species changes as a result of generations of competition among individuals. Within a species individuals vary with respect to heritable characteristics that influence the ability to survive and reproduce. Individuals that possess these characteristics will, on average, have more offspring than individuals that do not possess them. Because of this unequal contribution to the next generation, the characteristics that enhance survival and reproduction will tend to become more frequent within the species. Over many generations, this process, which Darwin called *natural selection*, changes the characteristics of the species—that is, the species evolves. In his book, Darwin summarized his thoughts about evolution by natural selection:

Again, it may be asked, how is it that varieties, which I have called incipient species, become ultimately converted into good and distinct species which in most cases obviously differ from each other far more than do the varieties of the same species? How do those groups of species, which constitute what are called distinct genera, and which differ from each other more than do the species of the same genus, arise? All these results . . . follow from the struggle for life. Owing to this struggle, variations, however slight and from whatever cause proceeding, if they be in any degree profitable to the individuals of a species, in their infinitely complex relations to other organic beings and to their physical conditions of life, will tend to the preservation of such individuals, and will generally be inherited by the offspring. The offspring, also, will thus have a better chance of surviving, for, of the many individuals of any species which are periodically born, but a small number can survive. I have called this principle, by which each slight variation, if useful, is preserved, by the term Natural Selection.

Darwin hypothesized that selection was the driving force of evolution in nature because he was powerfully aware of how artificial selection has changed the characteristics of domesticated species. He recognized the impact that artificial selection has had in creating different breeds of cattle, dogs, and fowl (**Figure 24.1**); he also knew of its role in shaping horticultural and agricultural varieties of plants.

Darwin was also a first-rate naturalist. As a young man, he served for five years on the British survey ship H.M.S. *Beagle*.



Cocker spaniel



Golden Laced Wyandotte



English bulldog



Light Brahma's Bantam

FIGURE 24.1 Variation among breeds of dogs and chickens.



Warbler finch (Geospiza olivace)

Common cactus finch (G. scandens)

Medium ground finch (G. fortis)

FIGURE 24.2 Finches on the Galapagos Islands.

The *Beagle* departed from England in 1831, traveled to South America, and returned to England in 1836. The lengthy sojourn along the coast of South America afforded Darwin many opportunities to observe plants, animals, and geological formations. For example, on the Galapagos Islands off the coast of Ecuador he observed several species of birds that were different from each other in appearance and behavior, but that he subsequently recognized were related to each other and to birds on the South American mainland (**Tigure 24.2**). From these and other observations, Darwin was led to the view that species are not fixed entities. Rather, he inferred that they change over time, and that some—exemplified by the fossils he saw during his travels—even became extinct.

Darwin spent more than 20 years analyzing and interpreting the data that he collected on the voyage of the *Beagle*. In addition, at his country estate in Kent, England, he performed experiments with a variety of plants and domesticated animals. The observations that he made in this experimental work, along with his extensive reading and analysis of the data that he collected on the *Beagle's* voyage, gave Darwin the insights that eventually led to the publication of *The Origin of Species*.

EVOLUTIONARY GENETICS

Darwin's theory of evolution had one major gap. It offered no explanation for the origin of variation among individuals, and it could not explain how particular variants are inherited. Eventually, Darwin did propose a theory of inheritance based on the transmission of acquired characteristics. However, his theory was flawed. Biologists who were attracted to Darwin's ideas on evolution struggled, as he did, to explain how the variants that natural selection favors are transmitted from parents to offspring. In 1900 the rediscovery of Mendel's principles provided the long-sought-after explanation: traits are determined by genes, which segregate different alleles, and genes are transmitted to the offspring in gametes produced by their parents. The analysis of genetic transmission in experimental crosses and pedigrees quickly gave rise to a new type of analysis that involved whole populations. The discipline of evolutionary genetics was born, and by 1930, especially through the contributions of Sewall Wright, R. A. Fisher, and J. B. S. Haldane, it had become the foundation for Darwinian theory.

KEY POINTS

• Charles Darwin formulated a theory in which species evolve through natural selection.

• After the rediscovery of Mendel's work, Darwin's ideas became grounded on Mendelian principles of inheritance.

Genetic Variation in Natural Populations

Darwin's *The Origin of Species* begins with a discussion of variation. Without variation, populations cannot evolve. Soon after Mendel's principles were rediscovered, biologists began to document genetic variation in natural populations. Initially, these efforts focused on conspicuous features of the phenotype pigmentation, size, and so forth. Later, they emphasized charMany different experimental approaches provide information about genetic variation in populations of organisms.

acteristics that are more directly related to chromosomes and genes. In the following sections, we discuss variation at the phenotypic, chromosomal, and molecular levels.

VARIATION IN PHENOTYPES

Naturalists have described phenotypic variation within many species. For example, land snails have different colored bands on their shells, squirrels and other small mammals have different coat colors, and butterflies and moths have different patterns in their wings (**Figure 24.3**). In the plant kingdom, phenotypic variation may be manifested by different kinds of flowers. All these sorts of phenotypic differences are called *polymorphisms*, from



Brown-banded snail (Liguus fasciatus)



Yellow-banded snail



Gray squirrel (Sciurus carolinensis)



Albino squirrel



Yellow tiger swallowtail (Papilio glaucus)



Black tiger swallowtail

FIGURE 24.3 Naturally occurring phenotypic variation in land snails, squirrels, and butterflies.

. т.		_	ົ່	4
	$\Delta \mathbf{R}$	-	14	

Allele	Korea	South Africa	England
Fya	0.995	0.060	0.421
Fy ^b	0.005	0.940	0.579
<i>Source:</i> Data from Cavalli-Sforza, L. L., and A. W. F. Edwards. 1967. Phylogenetic analysis: models and estimation procedures. <i>Evolution</i> 21: 550–570.			

Greek roots meaning "many forms." To elucidate the underlying genetic basis of a polymorphism, it is necessary to bring the organisms into the laboratory and cross them with one another. Unfortunately, for many organisms this approach is not feasible. Thus, geneticists have tended to focus their investigations of naturally occurring phenotypic variation on organisms that can be reared and bred in the laboratory.

Some of the classic studies were carried out in Russia, where researchers collected *Drosophila* from natural populations, inbred them, and examined the progeny for characteristics associated with mutant genes—for example, white eyes (instead of red eyes) and yellow bodies (instead of gray bodies). This work documented the presence of mutant alleles in natural populations.

Humans are also polymorphic. Pedigree analysis and population sampling have enabled researchers to identify many human polymorphisms. The classic data come from the study of blood types, which are determined by antigens on the surfaces of cells. The alleles that encode these antigens are often polymorphic. For example, the Duffy blood-typing system identifies two antigens, each encoded by a different allele of a gene on chromosome 1. The two Duffy alleles, denoted Fy^a and Fy^b have different frequencies among different populations (Table 24.1). In England, both Fy^a and Fy^b are common, but in Korea, only Fy^a is common, and in southern Africa, only Fy^b is common. Thus, the status of the Duffy polymorphism varies among human ethnic groups.

VARIATION IN CHROMOSOME STRUCTURE

Phenotypic variation can be a reflection of underlying genetic variability. Is there a way to detect variability by looking at the genetic material itself? The polytene chromosomes from the salivary glands of *Drosophila* larvae afford researchers an unparalleled opportunity to look for variation in chromosome structure. Flies captured in the wild can be brought into the laboratory and bred to produce larvae, which can then be examined for alterations in the banding patterns of their polytene chromosomes. For more than 25 years, Theodosius Dobzhansky and his collaborators performed this type of analysis on several species of *Drosophila* native to North and South America. The most thorough studies involved three closely related species, *D. pseudoobscura*, *D. persimilis*, and *D. miranda*, which are found in western North America.

Dobzhansky and his collaborators identified many different arrangements of the banding patterns in the polytene chromosomes of these species. Each arrangement consists of one or more inversions of the most common banding pattern. For example, in the third chromosome of *D. pseudoobscura*, they found 17 different arrangements in natural populations. The Standard banding pattern, denoted ST, was most frequent in populations along the coast of California and in northern Mexico; in these areas 48 to 58 percent of all third chromosomes in the sample of captured flies showed the ST banding pattern. Different arrangements predominated in other areas. For example, the arrangement known as Arrowhead (AR) was found in 88 percent of chromosomes sampled from Arizona, Utah, and Nevada, and the arrangement known as Pike's Peak (PP) was found in 71 percent of chromosomes sampled from Texas. Repeated sampling of selected populations established that the frequencies of the arrangements changed seasonally. For example, at Piñon Flats, California, the frequency of the ST arrangement declined from greater than 50 percent in March to around 30 percent in June.

This shift in frequency was observed in each of several years in which samples were collected. In addition, Dobzhansky and his coworkers observed long-term changes in the frequencies of arrangements in some populations. At Lone Pine, California, for instance, the ST arrangement increased from a frequency of 21 percent in 1938 to 65 percent in 1963. These researchers also performed laboratory experiments to measure the competitive abilities of flies carrying different chromosome arrangements. Their experiments suggested that balancing selection plays an important role in maintaining these chromosomal polymorphisms in nature.

VARIATION IN PROTEIN STRUCTURE

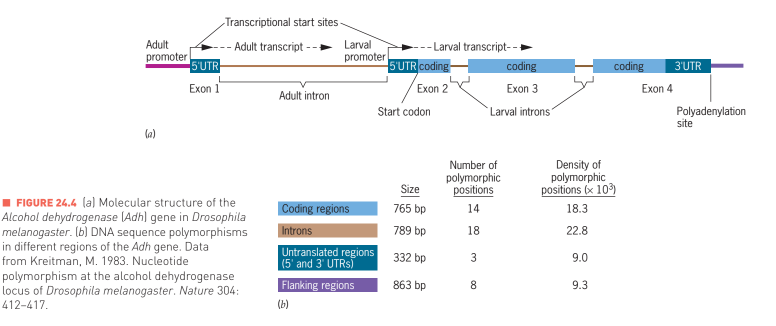
In 1966 R. C. Lewontin, J. L. Hubby, and H. Harris initiated a new era in the study of genetic variation in natural populations when they applied the technique of gel electrophoresis to detect amino acid differences in proteins. Lewontin and Hubby studied protein variation in *Drosophila*, and Harris studied it in humans. Their technique proved to be so successful that it was quickly applied to study genetic variation in all sorts of organisms, including creatures as diverse as starfish, wild oats, and spittle bugs. With this technique, a researcher can distinguish between different forms of a particular protein because each form moves at a specific rate through the electrophoretic gel. These forms reveal that the gene for that protein has different alleles, some "fast" and others "slow." Thus we can identify which alleles are present in an individual, and by analyzing many individuals, we can ascertain their frequencies in a population.

Protein gel electrophoresis provided the first extensive evidence of genetic variation at the molecular level. In many species one-fourth to one-third of all genes that encode soluble proteins exhibit electrophoretic polymorphisms, and for a given polymorphic gene, about 12 to 15 percent of individuals within a population are heterozygous for that gene. These two statistics—the proportion of genes that are polymorphic and the proportion of individuals that are heterozygous—are simple and convenient measures of the amount of genetic variability within a population.

VARIATION IN NUCLEOTIDE SEQUENCES

DNA sequencing provides the ultimate data on genetic variation. Any sequence coding, noncoding, genic, nongenic—can be analyzed. The first efforts to study genetic variation by DNA sequencing used material that had been cloned from the genomes of different individuals. The clones were then sequenced, and the sequences were compared to identify differences along their lengths.

As an example of this type of analysis, consider the results of a study of sequence variability in the gene for alcohol dehydrogenase, Adh, in Drosophila melanogaster performed by Martin Kreitman. Eleven cloned Adh genes from different populations were sequenced to obtain the data for the study. The *Adh* gene consists of four exons and three introns. Transcription of the Adh gene can be initiated from either of two promoters—one that functions in the adult and another that functions in the larva. The adult promoter is located upstream of the larval promoter. Thus, adult transcripts of the Adh gene contain all four exons and all three introns, whereas larval transcripts contain only the last three exons and the last two introns. The coding sequences of the Adh gene begin in the second exon; therefore, all the coding sequences are present in the larval transcript as well as in the adult transcript. Kreitman catalogued the differences among the *Adh* genes that he sequenced (**Figure 24.4**). Altogether, 43 nucleotide positions were polymorphic. The majority of the polymorphisms were in noncoding regions of the Adh gene-in introns, or in the 3' and 5' untranslated regions-and in the DNA flanking the gene. Some polymorphisms were also found within the gene's coding sequences; however, only one of these polymorphisms caused an amino acid difference in the Adh polypeptide. This difference, a lysine versus a threonine at position 192, alters the mobility of the Adh protein during gel electrophoresis; the polypeptide with lysine moves faster than the one with threonine. All the other nucleotide differences in the coding sequence of the Adh gene have no effect on the amino acid sequence of the polypeptide. Geneticists refer to them as *silent* 412-417.



polymorphisms; they arise from the degeneracy of the genetic code-that is, more than one codon being able to specify the incorporation of a particular amino acid into a polypeptide.

Today, obtaining DNA sequence data to study naturally occurring genetic variation is not nearly as difficult as it used to be. Particular regions of the genome can be amplified by PCR, and the resulting DNA products can be sequenced by machine. Sophisticated computer programs can then be used to analyze the sequence data and identify variation among individuals. This technique permits researchers to assess the level of variation in functionally different regions of DNA-for instance, in exons compared to introns.

Gene chip technologies (see Chapter 15) provide another means of documenting variation at the DNA level. These technologies allow researchers to screen genomic DNA for single-nucleotide polymorphisms (SNPs), which are found every 1-2 kb. Many different genomic DNA samples can be analyzed in parallel, and a great many SNPs can be detected on a single chip.

- **KEY POINTS** • Genetic variation in natural populations can be detected at the phenotypic, chromosomal, and molecular levels.
 - Classic studies established the existence of genetic polymorphisms for conspicuous phenotypic traits and for blood types.
 - Polymorphisms in chromosome structure have been documented in various species of Drosophila by analyzing banding patterns in the polytene chromosomes.
 - Polymorphisms in polypeptide structure have been detected by using the technique of protein gel electrophoresis.
 - Polymorphisms in DNA structure have been detected by sequencing cloned or PCR-amplified DNA and by using diagnostic gene chips.

Molecular Evolution

DNA and protein sequences provide information on the phylogenetic relationships among different organisms, and on their evolutionary history.

The ability to clone, amplify, manipulate, and sequence DNA molecules from any type of organism has had an enormous impact on the study of evolution. In The Origin of Species, Darwin repeatedly referred to evolution as a process of "descent with modification." His focus was on the traits of organisms, which are passed on more or less faithfully to their offspring every generation, but which also undergo modifications as the organisms adapt to changing environmental conditions. Today, knowing that heredity depends on the sequence of nucleotides in DNA, we understand the molecular basis of Darwin's concept. DNA molecules are passed from parents to offspring generation after generation. However, this process of genetic transmission is not perfect. Mutations occur, and when they do, modified DNA molecules are transmitted to the offspring. Over long periods of time, mutations accumulate and the DNA sequence is changed; segments of DNA molecules may also be duplicated or rearranged. This process of molecular evolution must underlie the evolution of organisms that Darwin wrote about.

MOLECULES AS "DOCUMENTS OF EVOLUTIONARY HISTORY"

One body of evidence that led Darwin to propose that species evolve came from the study of rocks in the ground. Fossils—the mineralized remains of animals and plants long since dead—were avidly collected in Darwin's day. These unusual rocks were curios for display in Victorian drawing rooms, but they were also evidence of organisms that had once lived on Earth. From the detailed study of fossils naturalists could reconstruct, at least crudely, what ancient organisms looked like and how they might have behaved. Comparisons between living organisms and the fossilized remains of extinct organisms stimulated speculation about the origin of species. Thus, with the perspective gained from studying fossils, naturalists began to think about life in historical terms.

DNA molecules, like fossils, contain information about life's history. The DNA molecules in creatures today are derived from their ancestors—parents, grandparents, and so on—going back in time to the very first organisms. Each DNA molecule is the end result of a long historical process involving mutation, recombination, selection, and genetic drift. In metaphorical terms, the sequence of nucleotides in a DNA molecule is the current version of an ancient text that, in the course of being copied generation after generation, has been altered (mutated), cut and pasted (recombined), preserved for its value (selected), and randomly disseminated (subjected to drift). Emile Zuckerkandl, one of the pioneers in the study of molecular evolution, put it this way: DNA molecules are "documents of evolutionary history."

So, too, are protein molecules. Polypeptides are encoded by genes, which are segments of DNA molecules. As the genes evolve, so do the proteins they encode. Geneticists can therefore investigate evolution at the molecular level either by studying nucleotide sequences in DNA or amino acid sequences in proteins.

The analysis of DNA and protein sequences has several advantages over more traditional methods of studying evolution based on comparative anatomy, physiology, and embryology. First, DNA and protein sequences follow simple rules of heredity. By contrast, anatomical, physiological, and embryological traits are subject to all the vicissitudes of complex heredity (see Chapter 22). Second, molecular sequence data are easy to obtain, and they are also amenable to quantitative analyses framed in the context of evolutionary genetics theory. The interpretation of these analyses is usually much more straightforward than the interpretation of analyses based on morphological data. Third, molecular sequence data allow researchers to investigate evolutionary relationships among organisms that are phenotypically very dissimilar. For instance, DNA and protein sequences from bacteria, yeast, protozoa, and humans can be compared to study the evolutionary relationships among them.

One problem with the molecular approach to evolution is that researchers usually cannot obtain DNA or protein sequence data from extinct organisms. In a few exceptional cases, such data have been obtained from fossils. However, in none of these cases was the fossilized specimen more than a few tens of thousands of years old. Thus, truly ancient organisms are beyond the reach of any molecular investigation. Another problem is that it is not always clear how molecular sequence data bear on questions about evolution at the phenotypic level.

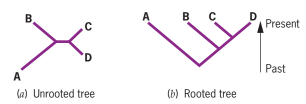


FIGURE 24.5 Difference between unrooted (*a*) and rooted (*b*) phylogenetic trees.

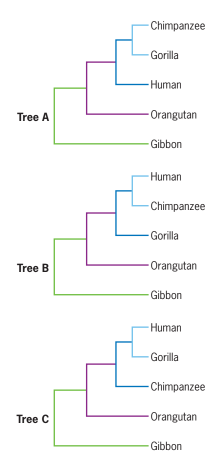


FIGURE 24.6 Phylogenetic trees of hominoid primates constructed from the analysis of an 896-base-pair-long sequence of mitochondrial DNA.

MOLECULAR PHYLOGENIES

The evolutionary relationships among organisms are summarized in diagrams called **phylogenetic trees**, or more simply, **phylogenies**. These trees may only show the relationships among the organisms, or they may superimpose the relationships on a time line to indicate how each of the organisms evolved. A phylogeny that only shows the relationships is an *unrooted tree*, whereas one that shows their derivation is a *rooted tree* (**Figure 24.5**). In both rooted and unrooted trees, the lineages bifurcate to produce branches. The branches at the tips of the tree—called terminal branches—lead to the organisms that are under study. Each bifurcation in a tree represents a common ancestor of the organisms farther out in the tree.

In molecular analyses of evolutionary relationships, the organisms are represented by DNA or protein sequences. Some analyses are based on a single gene or gene product. Other analyses combine data obtained by sequencing different genes or gene products. Sometimes the analyses utilize nongenic DNA sequences to ascertain the relationships among organisms.

The descendants of an ancestral DNA or protein sequence are said to be *homologous*, even if they have diverged significantly from the ancestor and are different from each other. Two sequences that come to resemble each other even though they are derived from entirely different ancestral sequences are said to be *analogous*. The construction of phylogenetic trees should always be based on the analysis of homologous sequences.

Many methods are now available to construct phylogenetic trees from DNA or protein sequence data. These methods usually have four features in common: (1) aligning the sequences to allow comparisons among them; (2) ascertaining the amount of similarity (or difference) between any two sequences; (3) grouping the sequences on the basis of similarity; and (4) placing the sequences at the tips of a tree.

Figure 24.6 shows trees constructed by comparing mitochondrial DNA sequences from a human, a chimpanzee, a gorilla, an orangutan, and a gibbon. The mitochondrial DNA (mtDNA) of each of these hominoid primates is a circular molecular containing about 16,600 base pairs. An 896-base-pair-long segment was cloned from each type of mitochondrial DNA and sequenced. The resulting sequences were then compared to determine the extent of their similarities (and differences). In all, sequence differences were found at 283 of the 896 nucleotide positions that were analyzed. The trees in Figure 24.6 were constructed by minimizing the number of mutational events needed to explain how the different mtDNA sequences were derived from a common ancestor. This criterion for tree construction is called the principle of parsimony. The number of mutational events required for tree A was 145, for tree B it was 147, and for tree C it was 148. All other possible trees required at least several more mutational events. Thus, the principle of parsimony yields three phylogenetic trees that plausibly explain the evolutionary relationships among the hominoid primates. From these trees we can see that gibbons and orangutans are clearly less related to humans than are chimpanzees and gorillas; however, we cannot readily discern how humans, chimpanzees, and gorillas are related to each other. Trees A, B, and C present the three possibilities. A more sophisticated analysis of the data employing statistical techniques that estimate the lengths of each of the branches favors tree B. This tree is also supported by the analysis of other types of DNA sequences. Thus, humans are more closely related to chimpanzees than to gorillas, they are less closely related to orangutans, and they are least closely related to gibbons. To apply the tree-building procedure to mtDNA sequences from different individuals within a particular human ethnic group, work through Problem-Solving Skills: Using Mitochondrial DNA to Establish a Phylogeny.

RATES OF MOLECULAR EVOLUTION

Molecular phylogenetic trees tell us about the evolutionary relationships among DNA or protein sequences. If we can link the branch points of a tree to specific times in the evolutionary history of the sequences, then we can determine the rate at which the sequences have been evolving. As an example of this kind of analysis, consider α -globin, which is one of two kinds of polypeptides found in the blood protein hemoglobin. The

PROBLEM-SOLVING SKILLS



Using Mitochondrial DNA to Establish a Phylogeny

THE PROBLEM

Derbeneva et al. (2002 *Am. J. Hum. Genet.* 71: 415–421) sequenced the mitochondrial DNA (mtDNA) obtained from a sample of 30 Aleut people from the Commander Islands, the westernmost islands of the Aleutian chain. Nine distinct types of mtDNA, each denoted by a Roman numeral, were identified. Each entry in the following table indicates the position of a nucleotide that differs from the nucleotide found in Type VIII, which was used as a standard. These differences are consistent across the various types—that is, the different nucleotide at position 9667 in Type I is the same as the different nucleotide at position 9667 in Type II. From these data, construct a diagram that shows the phylogenetic relationships among the different types of Aleut mtDNA.

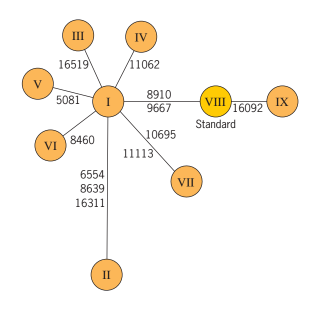
Туре	Number in Sample	Nucleotide Positions Different from Type VIII
	13	8910, 9667
	4	6554, 8639, 8910, 9667, 16311
	3	8910, 9667, 16519
IV	3	8910, 9667, 11062
V	1	8460, 8910, 9667
VI	1	5081, 8910, 9667
VII	1	8910, 9667, 10695, 11113
VIII	3	Standard
IX	1	16092

FACTS AND CONCEPTS

- 1. Human mtDNA is a circular molecule consisting of around 16,570 nucleotide pairs (see Chapter 15).
- 2. When two mtDNA sequences are compared, each single basepair difference represents a mutation.
- **3.** Phylogenetic trees are constructed by grouping the most similar DNA sequences near one another and by minimizing the number of mutations needed to explain the differences among all the DNA sequences.

ANALYSIS AND SOLUTION

The standard type of Aleut mtDNA differs from type IX at one nucleotide position (16092). It differs from all the other types at two or more nucleotide positions—positions 8910 and 9667 in Type I, and these two positions plus at least one other position in all the other types. The standard type is therefore one mutational step removed from Type IX, and it is two mutational steps removed from Type IX, and it is two mutational steps are one or more mutational step removed from Type I. We can summarize the relationships among the types in a phylogenetic diagram, which, however, does not tell us which mtDNA is ancestral to the others—that is, it is an unrooted tree:



For further discussion visit the Student Companion site.

 α -globin polypeptide consists of 141 amino acids. We can compare the sequence of α -globin from one organism with the sequence of α -globin from another organism and count the number of amino acids that differ between the two sequences. Such differences are tabulated in **Table 24.2**. The α -globins from humans and mice have the fewest differences (16); those from carp and sharks have the greatest number of differences (85).

The fossil record provides information about key events in the evolutionary history of the six types of organisms included in Table 24.2. For instance, the evolutionary lines that gave rise to humans and mice diverged about 80 million years ago (mya), near the end of the Mesozoic Era, and the lines that gave rise to carp and sharks diverged at least 440 mya near the end of the Ordovician Period in the Paleozoic Era. These and other branch points in the evolutionary history of the six different organisms are depicted in **Figure 24.7**.

The tree in Figure 24.7 was constructed by using evidence from the fossil record. However, its structure is consistent with the molecular data presented in Table 24.2. Humans and mice show the fewest amino acid differences in α -globin, and they are closest

TABLE 24.	2				
Number of Dissimilar Amino Acids in the $lpha$ -Globins of Representative Vertebrates					
	Mouse	Chicken	Newt	Carp	Shark
Human Mouse Chicken Newt Carp	6	35 39	62 63 63	68 68 72 74	79 79 83 84 85

together—that is, separated by the shortest evolutionary time—in Figure 24.7. The chicken's α -globin is the next closest to the α -globins of the two mammals, followed by the newt's, the carp's, and the shark's. The extent to which the amino acid sequences of these six organisms differ can be used to estimate the rate at which α -globin has been evolving.

To obtain this rate, we first need to determine the average number of amino acid changes that have occurred since any two of the lineages split from a common ancestor. We can start with the two most closely related organisms, humans and mice, which differ in 16 of the 141 amino acid sites in α -globin. The proportion of different sites in the α -globins of these two species is therefore 16/141 = 0.11, which we can also interpret as the average number of differences per amino acid site. Now consider two very distantly related organisms, humans and carp. The α -globins of these two organisms differ in 68 of the 141 amino acid sites; thus, the proportion of different sites is 68/141 = 0.48—that is, almost half the sites have changed during the evolution of the lineages that produced these two species. With such a high frequency of changed sites, we might expect that some of the sites have changed multiple times. The observed proportion of different sites, 0.48, must therefore underestimate the average number of changes that have occurred during the long time since the human and carp lineages split. Fortunately, we can adjust the observed proportion upward to account for multiple amino acid substitutions at particular sites. This adjustment

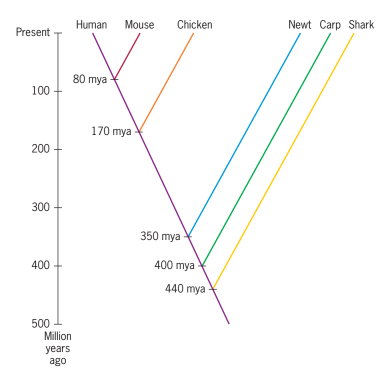


FIGURE 24.7 Phylogeny of representative vertebrates constructed from the fossil record.

involves a statistical procedure called the *Poisson correction*, which is explained in Appendix E: Evolutionary Rates. **Table 24.3** gives the Poisson-corrected differences for each pair of organisms. Each value estimates the average number of changes that have occurred per amino acid site in α -globin during the time since the evolving lineages split from a common ancestor. Notice that for the human and carp lineages, the average number of changes per amino acid site is 0.66, which is almost 1.4 times the observed proportion of amino acid differences between the human and carp α -globins.

With the average number of changes per amino acid site for each pair of organisms, we can now calculate the rate at which α -globin has evolved. This rate is the average number of changes per amino acid site divided by the total time that the two lineages have been evolving. For example, the lineages that produced humans and mice split from a common ancestor 80 mya. The total time that these lineages have been evolving is therefore 2 imes80 million years = 160 my. If we divide the average number of amino acid changes per site by this length of time, we obtain an estimate of the evolutionary rate of α -globin in the human–mouse lineages. Using the Poisson-corrected average number of amino acid changes per site from Table 24.3, we find that the average number of amino acid changes per site during the total evolutionary time is 0.12 amino acid changes per site/160 my = 0.74×10^{-9} amino acid changes per site/year. From this rate and all the other rates presented in Table 24.3, we see that α -globin has been evolving at a little less than one amino acid change per site every billion years.

TABLE 24.3

Poisson-Corrected Average Number of Amino Acid Differences per Site in the α -Globins of Representative Vertebrates and Associated Evolutionary Rates^a

	Mouse	Chicken	Newt	Carp	Shark
Human	0.12	0.28	0.58	0.66	0.82
	0.74	0.84	0.83	0.82	0.93
Mouse		0.33	0.59	0.66	0.82
		0.95	0.85	0.82	0.93
Chicken			0.59	0.72	0.89
			0.85	0.89	1.01
Newt				0.74	0.91
				0.93	1.03
Carp					0.92
					1.05

^aThe top number is the average number of amino acid differences between the α -globins of the two organisms. The bottom number is the annualized rate of amino acid substitution per site during the evolution of the α -globins in the lineages that produced these organisms (×10⁹ years).

THE MOLECULAR CLOCK

The values calculated for each pair of organisms in Table 24.3 imply that α -globin has evolved at more or less the same rate in all the evolutionary lineages analyzed. This apparent constancy of rate has been observed for other proteins as well. To evolutionary biologists, it suggests that amino acid substitutions occur in clocklike fashion over time. Thus, they sometimes metaphorically speak of the evolutionary process as one that follows a *molecular clock*. Extensive analyses have indicated that the rate of molecular evolution actually varies somewhat among different lineages. We see a hint of this variation in the data in Table 24.3, where the calculated rate of evolution in the mammalian lineages is slightly less than the rates in the other lineages. Therefore, a universal molecular clock probably does not keep the same time in all evolving lines. However, within some lines, local clocks may be operating—that is, within them the rate of molecular evolutionary change is approximately constant.

Calculations based on the assumption of a molecular clock can be very helpful in estimating when, in historical time, lineages diverged from a common ancestor. This approach has been used to date events in the evolution of our own species, for which fossil evidence is scarce. For instance, the lines that gave rise to humans and chimpanzees are estimated to have diverged between 5 and 6 mya. To see an application of this kind of analysis, work through Solve It: Calculating Divergence Times.

VARIATION IN THE EVOLUTION OF PROTEIN SEQUENCES

The α -globin polypeptide seems to be evolving at a rate of slightly less than one amino acid substitution per site every billion years. Do other proteins evolve at this rate too? Extensive analyses have shown that some do, but others evolve either faster or slower. The observed rates of amino acid sequence evolution range over three orders of magnitude. At the extremes, fibrinopeptide, which is derived from a protein involved in blood clotting, evolves at a rate of greater than 8 amino acid substitutions per site every billion years, whereas the histones, which interact intimately with DNA, evolve at a rate of only 0.01 amino acid substitutions per site every billion years. We can also see variation in evolutionary rates within some polypeptides. For example, amino acids on the surface of α -globin change at a rate of about 1.3 substitutions per site every billion years, whereas amino acids in the interior of the molecule change at a rate of only 0.17 substitutions per site every billion years.

Preproinsulin, the precursor of the peptide hormone insulin, provides another example of intramolecular variation in evolutionary rate. This polypeptide consists of

Solve It!

Calculating Divergence Times

The Poisson-corrected average number of amino acid differences per site in the α -globins of humans and mice is 0.12; for humans and kangaroos it is 0.20. The lineages leading to humans and mice diverged from a common ancestor about 80 million years ago. How long ago did the human lineage diverge from the kangaroo lineage?

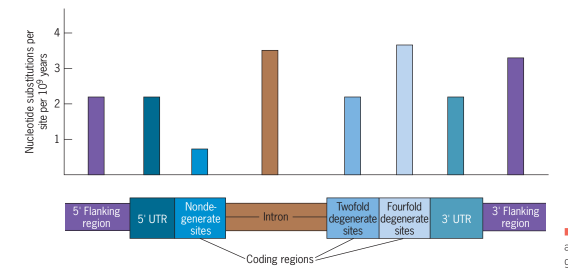
To see the solution to this problem, visit the Student Companion site. four segments. The first segment is a signal peptide, the second and fourth segments form the active insulin molecule, and the third segment is a peptide bridge that initially links the two active segments. When active insulin is formed, this bridge segment is deleted and the two active segments are joined together covalently by disulfide bonds. The signal and bridge segments evolve at a rate of slightly more than one amino acid substitution per site every billion years; however, the two active segments evolve at a rate of only 0.2 substitutions per site every billion years. Thus, within the preproinsulin polypeptide, the evolutionary rate varies significantly.

What might explain the observed variation in evolutionary rates? Geneticists hypothesize that in more rapidly evolving proteins, the exact amino acid sequence is not as important as it is in more slowly evolving proteins. They speculate that in some proteins, amino acid changes can occur with relative impunity, whereas in others, they are rigorously selected against. According to this view, the rate of evolution depends on the degree to which the amino acid sequence of a protein is constrained by selection to preserve that protein's function. Slowly evolving proteins are more constrained than rapidly evolving proteins. Variation in evolutionary rates is therefore explained by the amount of *functional constraint* on the amino acid sequence. This idea also applies to parts of proteins. For example, the specific amino acids at or near the active sites of enzymes might be expected to be more rigorously constrained by selection than amino acids that simply take up space, such as those in the bridge segment of preproinsulin, which is discarded during the formation of the active insulin molecule. Thus, functionally more important proteins, or parts of proteins, evolve more slowly than functionally less important ones.

VARIATION IN THE EVOLUTION OF DNA SEQUENCES

Variation in the evolutionary rate is also seen when DNA sequences are examined. The DNA sequences in pseudogenes—duplicated genes that do not encode functional products because they have sustained one or more lesions such as frameshifting or nonsense mutations—have the highest evolutionary rates. For example, the evolutionary rate of the $\psi \alpha 1$ pseudogene of α -globin is 5.1 nucleotide substitutions per site every billion years. By contrast, nucleotides in the first or second positions of codons in a functional α-globin gene evolve at the rate of 0.7 nucleotide substitutions per site every billion years. This sevenfold difference in the evolutionary rate can be explained by the concept of functional constraint. The nucleotides in a pseudogene are not constrained by selection because the function of the pseudogene has already been destroyed. However, the nucleotides in the first and second positions of a codon in a functional gene are constrained because changing them will almost always change the amino acid specified by that codon. Some of these changes will be conservative in the sense that the new amino acid will be structurally and functionally like the original amino acid. For example, if the first nucleotide in the codon CTT mutates to A, the amino acid specified by this codon will change from leucine to isoleucine. These two amino acids have similar properties. However, other substitutions in this codon may cause a nonconservative change in the amino acid sequence. For instance, if CTT mutates to TTT, the amino acid specified by the codon will change from leucine to phenylalanine, which has very different chemical properties.

Nucleotides in the third position of codons within functional genes present a special—and interesting—case. These nucleotides evolve much faster than nucleotides in either the first or the second position. This more rapid evolution is due to the degeneracy of the genetic code. Many amino acids are specified by more than one codon. For example, proline is specified by four different codons: CCT, CCC, CCA, and CCG. As long as the first two nucleotides in a codon are both C, any nucleotide can be present in the third position and the codon will specify proline—that is, the third nucleotide position is fourfold degenerate. Changing the last nucleotide in a proline codon—for example, changing the T in CCT to C to create the codon CCC—should therefore be inconsequential for the structure and function of the polypeptide encoded by a gene. However, changing either the first or second nucleotide in CCT to any other nucleotide will change the amino acid specified by the codon. The first two positions in the CCT codon are therefore more constrained than the third position.



About half of all codons are fourfold degenerate in the third nucleotide position. A majority of all the other codons are twofold degenerate in this position—that is, either of two nucleotides in the third position will specify the same amino acid. This high level of degeneracy accounts for the faster evolutionary rate of third position nucleotides.

A nucleotide substitution that does not change the amino acid specified by a codon is called a *synonymous* substitution. A nucleotide substitution that does change the amino acid specified by a codon is called a *nonsynonymous* substitution. A wealth of DNA sequence data has now established that synonymous substitutions occur more frequently than nonsynonymous substitutions in evolving lineages.

We also see variation in the evolutionary rates of nucleotides in the noncoding portions of genes (**■** Figure 24.8). Nucleotides in introns evolve more rapidly than nucleotides in 5' and 3' untranslated regions. The different evolutionary rates observed for these types of noncoding sequences presumably reflect variation in the functional constraints on them. In general, these types of sequences do not evolve as fast as pseudogenes, nor do they evolve as slowly as nucleotides in the first or second positions of codons. Rather, they show intermediate evolutionary rates.

THE NEUTRAL THEORY OF MOLECULAR EVOLUTION

Evolutionary geneticists have developed a theory—called the Neutral Theory—to explain the evolution of DNA and protein sequences. It focuses on three processes: mutation, purifying selection, and random genetic drift.

Mutation is at the root of all nucleotide and amino acid substitutions that occur during evolution. Without mutation, DNA and protein molecules could not evolve. Experimentally determined mutation rates are on the order of 10^{-9} – 10^{-8} events per nucleotide each generation. These rates reflect the effects of polymerase errors and chemical damage to DNA. They would surely be higher if cells were not equipped with an assortment of mechanisms to prevent replication errors and to repair damaged DNA.

Some of the mutations that occur spontaneously improve the fitness of organisms—that is, they are beneficial mutations that might, over time, spread through a population and become fixed. Other mutations depress fitness and are eliminated from a population by the force of *purifying selection*. Because each gene is already the end result of a long evolutionary process, it is improbable that very many new mutations will improve a gene's function. Many mutations, like random changes in a piece of complex machinery, are likely to impair function. However, some mutations may have little or no effect on fitness. Geneticists say that such mutations are *selectively neutral*. We could easily imagine that synonymous nucleotide substitutions in the third positions of codons might be selectively neutral, as might any type of nucleotide substitution in a pseudogene, which has already been impaired by a previous mutation. Conservative amino acid substitutions in proteins might also be selectively neutral.

■ **FIGURE 24.8** Variation in evolutionary rates among different parts of genes.

Solve It!

Evolution by Mutation and Genetic Drift

Suppose that the rate at which new, neutral mutations of a gene occur in a population is *u* and that size of the population, *N*, remains constant over time. Formulate an argument to show that the rate at which neutral mutations are fixed by random genetic drift is simply *u*, the mutation rate.

To see the solution to this problem, visit the Student Companion site. The fate of a selectively neutral mutation depends completely on *random genetic drift*. Most selectively neutral mutations are lost from a population shortly after they first appear. A very small fraction of them survive for a few generations, and an even smaller fraction ultimately spread throughout the population and become fixed. When evolution occurs by random genetic drift, the rate of fixation is the rate at which genes mutate to selectively neutral alleles. Solve It: Evolution by Mutation and Genetic Drift challenges you to construct an argument to justify this statement.

In the Neutral Theory, the rate of evolution does not depend on population size, efficiency of selection, or peculiarities of the mating system. It simply depends on the neutral mutation rate, which is expected to be more or less constant in different lineages over time. Thus, the Neutral Theory explains why amino acid and nucleotide substitutions seem to occur in clocklike fashion.

The Neutral Theory, however, does not require that all polypeptide and DNA sequences evolve at the same rate. For some positions within a sequence, all or nearly all mutations will be selectively neutral—for example, the nucleotides in a pseudogene or in the third position of a codon that is fourfold degenerate. For other positions, a smaller fraction of all mutations will be selectively neutral, and for some positions, almost no mutations will be selectively neutral. Thus, the Neutral Theory explains the variation in evolutionary rates that is observed among proteins and DNA regions by invoking differences in functional constraints. The highest rates are observed in molecules or in portions of molecules that are not constrained by selection to preserve a function—that is, in molecules in which mutational changes have little or no effect on function. The lowest evolutionary rates are observed in molecules where selection pressure is strongest.

The Neutral Theory has had an enormous impact on the study of evolution at the molecular level. We discuss its intellectual roots in A Milestone in Genetics: The Neutral Theory of Molecular Evolution on the Student Companion site.

MOLECULAR EVOLUTION AND PHENOTYPIC EVOLUTION

By definition, the Neutral Theory has nothing to say about the evolution of traits that are adaptive. The giraffe's long neck, the elephant's trunk, and the camel's hump are all adaptations that enhance fitness. So, too, is the large, highly convoluted brain of a human being. Darwin emphasized that adaptations such as these evolve because natural selection favors them. In the classic Darwinian sense, then, evolution implies positive selection for something, not just negative selection against deleterious mutants, or, as the Neutral Theory assumes, no selection at all. In addition, Darwin recognized a connection between the evolution of adaptations and the diversification of organisms. As organisms adapt to what Darwin called "the conditions of life," they become different from one another. The phenotypic changes that occur during this process produce new varieties and eventually new species.

The evolution of adaptations and the diversification of organisms must ultimately be due to change at the molecular level. However, change at the molecular level is not a guarantee that phenotypic evolution will occur. Crocodiles, sharks, and horseshoe crabs (**■** Figure 24.9) have all accumulated amino acid and nucleotide changes at rates



(b) Great white shark (Carcharodon carcharias)





(a) Nile crocodile (Crocodylus niloticus)

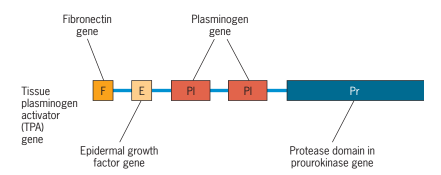
FIGURE 24.9 Some organisms considered to be "living fossils."

similar to highly diversified groups of animals such as birds, mammals, and insects. Yet, to judge from the fossil record, these types of organisms have changed very little in phenotype since they first appeared hundreds of millions of years ago. "Living fossils" therefore seem to have roughly the same rate of molecular evolution as organisms that have diverged extensively at the phenotypic level. This observation suggests that many nucleotide and amino acid substitutions have little to do with phenotypic evolution.

What sorts of genetic changes might be responsible for the evolution of novel phenotypes? Some possible answers are coming from the genome sequencing projects and from ongoing studies in developmental genetics. From these investigations we know that *gene duplication* is an important evolutionary event. The classic example comes from the study of the globin genes in animals (**■** Figure 24.10). Today we find two classes of globin genes—those encoding components of hemoglobin, which carries oxygen in the blood, and those encoding myoglobin, which stores oxygen in muscle. These functionally different classes of genes are derived from a primordial globin gene, which was duplicated about 800 mya, long before the diversification of the animals at the start of the Paleozoic Era.

The hemoglobin genes have, in turn, been duplicated several times during the evolution of the vertebrates. As best we can tell, the α - and β -globin genes were created by a duplication more than 450 mya in the evolutionary line that produced the jawed fishes. Jawless fish-the lampreys and their kin-have only one kind of hemoglobin gene (α); sharks and bony fish have at least two. About 300 to 350 mya, the α - and β -globin genes were separated from each other and took up residence on different chromosomes. Each of these genes subsequently underwent several duplication events to produce clusters of α - and β -globin genes. In humans, for example, seven α -globin genes are clustered together on chromosome 16, and six β -globin genes are clustered together on chromosome 11. Three of the α -globin genes and one of the β -globin genes in humans are nonfunctional pseudogenes. The other globin genes in these clusters encode different, but related, polypeptides that carry oxygen in the blood at different times during life. Some of these polypeptides function only in the embryo, others only in the fetus, and still others only in the adult (see Chapter 19). Thus, these families of hemoglobin genes indicate that duplicated genes can acquire different functions.

Another phenomenon that might help to explain phenotypic evolution is that portions of genes may be duplicated and recombined with other genes. Eukaryotic genes are segmented into exons and introns. Shortly after this segmented structure was discovered, Walter Gilbert speculated that each exon in a gene encodes a separate functional domain in the gene's polypeptide product. He further speculated that exons from one gene could be combined with exons from another gene to create a coding sequence that would specify a protein with some of the properties of each of the original gene products. Thus, he proposed that novel proteins could be created by combining exons in modular fashion—a process now called *exon shuffling* (**— Figure 24.11**).



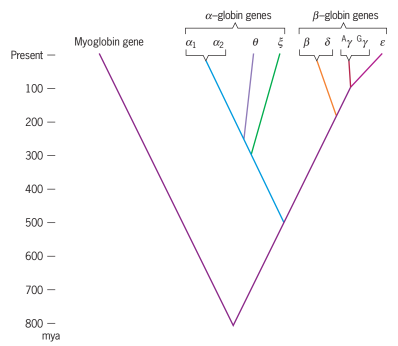
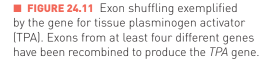


FIGURE 24.10 Role of gene duplication in the evolution of the globin genes.



DNA sequencing studies have provided evidence for Gilbert's hypothesis. For example, tissue plasminogen activator (TPA), a protein involved in the breakup of blood clots, is encoded by a gene that seems to have acquired exons from several different sources. One exon comes from the gene for fibronectin, another from the gene for epidermal growth factor, two exons come from the gene for plasminogen, and one comes from a gene that encodes a protease. Altogether, then, at least four genes have contributed exons to the formation of the *TPA* gene. The recombination of evolutionarily proven exons provides almost limitless possibilities to form mosaic proteins. Mixing and matching exons, and the polypeptide domains they encode, may be an important process in evolution, and it may partly explain why eukaryotes are anatomically, physiologically, and behaviorally so diverse.

In addition to gene duplication and exon shuffling, evolutionary diversification seems to have involved spatial and temporal changes in the expression of genes, especially those whose products regulate the expression of other genes. For example, the homeobox genes play important roles in the formation of animal bodies along an anterior–posterior axis; these genes encode transcription factors. Changing the time or place in which specific homeobox genes are expressed may profoundly change the appearance of the animal. In *Drosophila*, where the homeobox genes have been studied thoroughly, it is clear that altering the pattern of expression of one or a few of these genes can produce a fly with four wings instead of two, or a fly with extra appendages on either the head or the thorax. With these kinds of observations in the laboratory, it is not hard to imagine that similar kinds of changes might have occurred in nature during the course of evolution.

- **KEY POINTS** *Phylogenetic trees based on the comparison of DNA and protein sequences show the evolutionary relationships among organisms.*
 - The rate of molecular evolution can be determined by calculating the average number of amino acid or nucleotide changes that have occurred per site in a molecule since two or more evolving lineages diverged from a common ancestor.
 - The near uniformity of the rate of molecular evolution in different lineages is metaphorically described as a "molecular clock."
 - The rate of evolution varies among different protein and DNA sequences and appears to depend on the extent to which these sequences are constrained by natural selection to preserve their function.
 - Selectively neutral mutations are fixed in a population at a rate equal to the neutral mutation rate.
 - Gene duplication and exon shuffling have played important roles in evolution.
 - Changes in the spatial and temporal aspects of gene regulation may have contributed to the rapid evolution of some types of organisms.

Speciation

Species arise when a population of organisms splits into genetically distinct groups that can no longer interbreed with each other. Biologists have named and described a large number of plant, animal, and microbial species. Many more species have yet to be identified. Where did all this diversity come from? How is it maintained? Why are different species distinct from one another? What factors contribute to the formation of species? Charles Darwin

raised these kinds of questions more than 150 years ago when he wrote *The Origin of Species*. Today, biologists continue to grapple with them as they address the central problem of evolutionary genetics—the problem of speciation.

WHAT IS A SPECIES?

The term *species* is usually applied to a group of organisms that share certain characteristics. However, species have been defined in different ways. In classical taxonomy, a species is defined exclusively on the basis of phenotypic characteristics. If the characteristics of two groups of organisms are sufficiently different, then the groups are considered to be separate species. This approach to defining a species relies on careful observation of the organisms, either as specimens in a zoo, arboretum, herbarium, or museum collection, or, better still, as the inhabitants of a natural environment where their behavior as well as their morphology can be studied. This approach to defining a species also relies on the expertise of the taxonomist, who must decide if groups of organisms are sufficiently different to warrant their classification as separate species. Thus, it is a subjective approach that may lead to different classifications in the hands of different people.

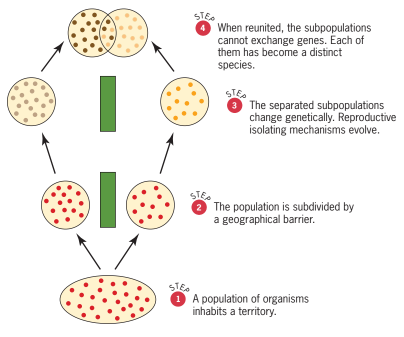
In evolutionary genetics, a species is defined on the basis of a shared gene pool. A group of interbreeding, or potentially interbreeding, organisms that does not exchange genes with other such groups is considered to be a species. Evolutionary geneticists say that each species is *reproductively isolated* from every other species. This approach to defining a species relies on a researcher's ability to determine whether groups of organisms exchange genes in nature. If they do, they are classified as a single species; if they do not, they are classified as separate species. The genetic approach to defining species therefore involves an objective assessment of whether or not groups of organisms are reproductively isolated from each other.

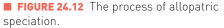
These two ways of defining species are not always in agreement. Organisms may be reproductively isolated, but they may not be distinguished by easily recognized phenotypic characteristics. In taxonomy, such organisms would be regarded as a single species, whereas in evolutionary genetics, they would be regarded as separate species. Conversely, organisms may have different phenotypic characteristics, but they may not be reproductively isolated. A taxonomist would regard such organisms as separate species, whereas an evolutionary geneticist would regard them as a single species. When it is possible to determine whether different organisms are reproductively isolated, we can apply the genetic definition of species. However, when such determinations are not possible—as, for example, with fossilized organisms—we are limited to the taxonomic definition of species.

Reproductive isolation is the key to the genetic definition of a species. Groups of organisms that inhabit the same territory can be reproductively isolated from each other by different mechanisms. *Prezygotic isolating mechanisms* prevent the members of different groups from producing hybrid offspring. *Postzygotic isolating mechanisms* prevent any hybrid offspring that are produced from passing on their genes to subsequent generations.

Prezygotic isolating mechanisms operate by preventing matings between individuals from different populations of organisms, or by preventing the gametes of these individuals from uniting to form zygotes. For example, two populations of organisms that inhabit the same area might seek out different habitats within that area. If the habitat preference is strong, the two populations will have little or no contact with each other. Ecological isolation based on habitat preference can therefore prevent the populations from producing hybrid zygotes. Temporal or behavioral factors can also bring about reproductive isolation between populations of organisms. For instance, the organisms might become sexually mature at different times, or they might have different courting rituals. If ecological, temporal, and behavioral isolating mechanisms fail to prevent mating between different organisms, then anatomical or chemical incompatibilities in their reproductive organs or gametes might prevent them from producing hybrid zygotes. The organisms might be unable to copulate successfully, or to exchange pollen, or their sperm or pollen might die in the reproductive tissues of their mates. Any of these prezygotic isolating mechanisms could prevent genes from being exchanged between populations occupying the same territory.

Postzygotic isolating mechanisms operate after hybrid zygotes have been formed, either by reducing hybrid viability or by impairing hybrid fertility. The zygotes from matings between different organisms might not survive, or they might not reach sexual maturity. If they do reach sexual maturity, they might not produce functional gametes. Any of these circumstances could prevent populations of organisms that live in the same territory from exchanging genes.





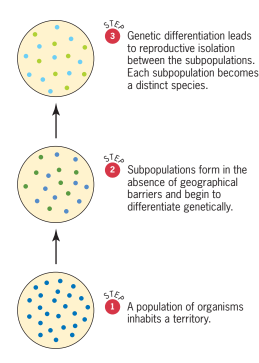


FIGURE 24.13 The process of sympatric speciation.

MODES OF SPECIATION

The key event in speciation is the splitting of a population of organisms into one or more subpopulations that become reproductively isolated from each other. The most straightforward way for this event to happen is for the subpopulations to become geographically separated so that they evolve independently-that is, geographical barriers keep the subpopulations apart so that they accumulate their own sets of genetic changes over time (**Figure 24.12**). Then, if the subpopulations are reunited by the disappearance of the geographical barriers, the genetic changes they have accumulated may make them reproductively isolated from each other. For example, one subpopulation may have evolved a preference for a particular food source, and another subpopulation may have evolved a preference for a different food source. When the two subpopulations are rejoined in the same territory, their distinctive food preferences may limit contact between them to such an extent that interpopulational matings never occur. Another possibility is that during the time the subpopulations

were separated, they may have evolved different physiological processes or mating habits. When the subpopulations are reunited, they may not be able to mate with each other, or if they can mate with each other, their hybrids may not be viable or fertile. The process whereby subpopulations evolve reproductive isolation while they are geographically separated is called **allopatric speciation** (from Greek roots meaning "in other villages").

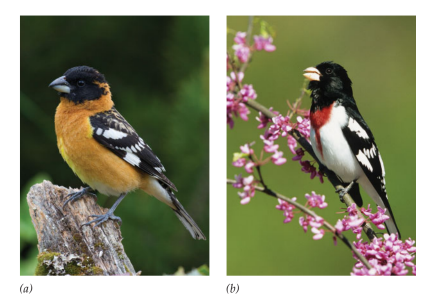
It is conceivable that subpopulations might evolve reproductive isolation without being separated geographically (**Figure 24.13**). Perhaps the subpopulations become ecologically specialized so that they evolve more or less independently, or perhaps their members mate only with individuals like themselves so that there is little or no genetic exchange between the subpopulations. The process of evolving reproductive isolation between subpopulations that exist in the same territory is called **sympatric speciation** (from Greek roots meaning "in the same villages").

Because the evolution of reproductive isolation may require hundreds of thousands of years, it is not easily studied. Most investigations of speciation are done *post factum*— that is after the species have already formed. Based on data collected from the species, researchers attempt to determine how and why they became reproductively isolated from each other.

One issue in these studies is whether the species evolved allopatrically or sympatrically. Did they develop reproductive isolating mechanisms while they were geographically separated, or did they develop these mechanisms while they inhabited the same territory? Usually this question cannot be answered with certainty. However, most evolutionary geneticists are inclined toward the view that allopatric speciation is more prevalent than sympatric speciation, if only because allopatric speciation is a more straightforward process. For example, imagine that a small number of organisms migrate to a remote oceanic island where they found a population that evolves independently of the main population on the nearest continent. The island population may change significantly over time and eventually become reproductively isolated from its closest relatives on the continent. This scenario-which is allopatric speciation pure and simple-may have played out many times on oceanic islands (Figure 24.14). Indeed, Darwin proposed it as an explanation for the species of plants and animals he observed on the Galapagos Islands off the west coast of South America. It is not too hard to imagine other types of geographic separation that would permit allopatric speciation to occur. Deserts and mountain ranges can subdivide continents; reductions in rainfall can isolate lakes and river systems; land masses can rise up to separate oceans. Populations that are subdivided by these kinds



■ FIGURE 24.14 Four species of *Drosophila* from the Hawaiian Islands. Starting at the upper left and moving clockwise: *D. heteroneura*, *D. grimshawi*, *D. ornata*, and *D. differens*. These and hundreds of other *Drosophila* species have evolved during the last few million years on the Hawaiian Islands, which are far removed from other land masses in and around the Pacific Ocean.



■ FIGURE 24.15 Species of grosbeaks that may have arisen by allopatric speciation. (a) The black-headed grosbeak (*Pheucticus melanocephalus*) found in the western United States. (b) The rose-breasted grosbeak (*P. ludoviciannus*) found in the eastern United States.

of barriers have the potential to evolve into distinct, reproductively isolated species (**Figure 24.15**).

Although allopatric speciation may have been the prevalent mode in creating the species that exist today, there is evidence that sympatric speciation has also contributed to species diversity. The strongest case for sympatric speciation comes from the study of cichlid fish in two small crater lakes located in west central Africa. Today these lakes are isolated from other significant bodies of water. However, in the relatively recent past they were apparently colonized by cichlids from the surrounding river systems. These colonists then evolved into the groups of species now present in the lakes. Analysis of mitochondrial DNA sequences indicates that the cichlid species within each lake are derived from a common ancestor and that they are more closely related to each other than to the cichlid species found in the surrounding river systems. There are no obvious geographic barriers within these lakes. Their shorelines are regular, and they do not seem to have been subdivided during their history. Thus, it appears that the crater-lake cichlids evolved into different species sympatrically.

Cichlid fish inhabit many of the lakes and rivers in tropical Africa, especially the East African Great Lakes—Lake Victoria, Lake Malawi, and Lake Tanganyika— where over 1500 cichlid species have been identified. The apparent sympatric speciation of cichlids in the small crater lakes of west central Africa raises the possibility that some of the species in these large lakes may also have originated sympatrically. More research is needed to determine how the Great Lake cichlids evolved.

- In evolutionary genetics, a species is a group of populations that share a common gene pool.
- The development of reproductive isolation between populations is the key event in the speciation process.
- Speciation may occur when the populations are geographically separated (allopatric) or when they coexist in the same territory (sympatric).

KEY POINTS

Human Evolution

Fossil evidence and DNA sequence analysis have provided information about the origin of modern humans. When Darwin proposed his theory of evolution in 1859, and later, when he suggested that human beings had evolved from more primitive organisms, he provoked a great controversy. The idea that organisms evolve, and more specifically, that humans evolve, has troubled many people. In the ensuing 150 years, much has been learned about the course of human evolu-

tion. Paleontologists have analyzed the fossilized remains of organisms that are likely to have been the ancestors of modern humans, and geneticists have analyzed DNA sequence data in order to study the relationships among humans and their closest nonhuman relatives, the great apes. In the following sections, we discuss some of these analyses.

HUMANS AND THE GREAT APES

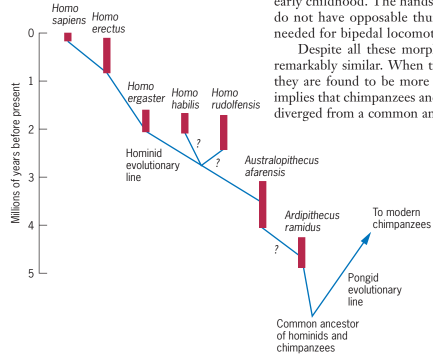
Several morphological features distinguish human beings from chimpanzees and gorillas. The apes have larger canine and incisor teeth than modern humans, and their jaws are larger and heavier. Ape brains are smaller than human brains, and the point where the ape brain attaches to the spinal cord is placed farther to the back of the skull than it is in humans. The shape and proportions of an ape's body are also different from those of a human. In an ape, the body's trunk widens toward the base, whereas in a human, it tends to have the same width from the shoulders to the waist. The legs of an ape are proportionately shorter than those of a human, and the pelvis is not constructed to accommodate a regular upright stance. Although apes can walk upright on two legs, they cannot do so for long periods of time. By contrast, humans are exclusively bipedal—except, of course, in early childhood. The hands and feet of apes also differ from those of humans. Apes do not have opposable thumbs, and their feet do not provide the support that is needed for bipedal locomotion.

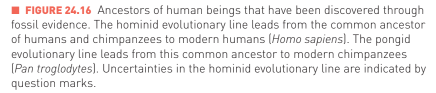
Despite all these morphological differences, the DNA of apes and humans is remarkably similar. When the genomes of chimpanzees and humans are compared, they are found to be more than 99 percent identical. This high degree of identity implies that chimpanzees and humans are quite closely related, and suggests that they diverged from a common ancestor rather recently in evolutionary time, perhaps 5 to

6 million years ago. The other great ape species, the gorilla, appears to be less closely related to humans than the chimpanzee is.

HUMAN EVOLUTION IN THE FOSSIL RECORD

Though rare, fossils have provided important information about human evolution (**Figure 24.16**). The oldest fossils that appear to be strictly within the human evolutionary line come from East Africa where they were formed 4 to 5 million years ago. These first humanlike—that is, *hominin*—creatures have been given the name *Ardipithecus ramidus*. Later in the fossil record, 3 to 4 million years ago, another hominin creature appeared. This organism, known as *Australopithecus afarensis*, probably stood 1 to 1.5 m tall and walked upright, at least for short distances. The fossil known as Lucy is a specimen of *Australopithecus afarensis*.





The first organisms to be classified in the same genus as *Homo sapiens* appeared 2 to 2.5 million years ago. Two species have been named, *H. rudolfensis* and *H. habilis*. Both of these "early *Homo*" species have many apelike features; however, compared to *Australopithecus*, the opening for the spinal cord is closer to the middle of the skull, and the skull itself is reduced in length and increased in width—all hominin characteristics. Nevertheless, many paleontologists have questioned the inclusion of these two species within the genus *Homo*, and there is some sentiment to reclassify them in the genus *Australopithecus*.

Between 1.9 and 1.5 million years ago, another hominin appeared in the fossil record. This creature, called *Homo ergaster*; had a body shape and limb proportions like those of modern humans, and its teeth and jaws were also human in structure. Thus, *H. ergaster* is the first hominin that can confidently be placed within the genus *Homo*.

All the early hominin fossils come from Africa. The first hominin species to produce fossils outside of Africa was *Homo erectus*. These fossils, formed about 1 million years ago, have been found in China and Indonesia. Thus, *H. erectus* was widespread and probably gave rise to archaic populations of humans in Europe, Asia, and Africa. The best known of the archaic humans were the Neanderthals, a species that evolved in Europe and the Near East several hundred thousand years ago. Ultimately, however, they lost out in competition with the ancestors of the modern human species, *H. sapiens*, and became extinct.

Modern humans may have evolved simultaneously in Europe, Asia, and Africa from the archaic human populations that existed on each of those continents, or they may have evolved on one continent—for instance, Africa—and subsequently spread to the others. Fossil evidence cannot discriminate between these two hypotheses. However, genetic evidence obtained by studying DNA sequences in living human beings has provided ways of testing them.

DNA SEQUENCE VARIATION AND HUMAN ORIGINS

Genetic data allow researchers to study human evolution by investigating the relationships among extant human populations. Populations that are closely related share genetic properties that distantly related populations do not. Thus, by analyzing variation in genes, gene products, and DNA sequences, it is possible to determine the relatedness of different racial and ethnic groups, and to arrange them in a phylogenetic tree. Genetic analysis also permits researchers to decipher key events in human evolutionary history.

Many types of genetic variation have been used to study human evolution: blood group and protein polymorphisms, and variation in the composition of DNA sequences themselves. Both nuclear and mitochondrial genetic variation has been investigated. The nuclear genome contains the preponderance of human polymorphisms, but the mitochondrial genome has the unique property of being transmitted exclusively through females. Variation in mitochondrial DNA therefore provides a way of tracing maternal lineages in human evolutionary history.

Compared to other species, the human species is genetically rather uniform. At the nucleotide level, humans have about one-fourth the genetic variation of chimpanzees and about one-tenth that of *Drosophila*. Furthermore, most of the genetic variation in the human species—perhaps 85 to 95 percent of it—is within rather than between populations.

The relative absence of genetic variation in human populations implies that during its evolutionary history, the genetically effective size of the human population was small—between 10,000 and 100,000 individuals. The census size may have been larger (today, it certainly is), but the mating system, various constraints on reproduction, and bottlenecks in size caused by famine, disease, or weather-related catastrophes apparently conspired to keep the effective population size under 100,000. In such a population, random genetic drift dominates over mutation to

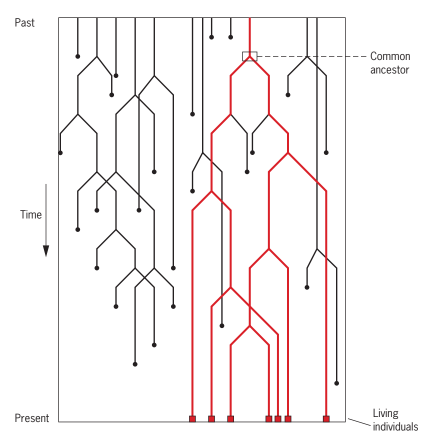


FIGURE 24.17 A coalescence process. If the lineages of the DNA sequences found in living individuals are traced back into the past, they coalesce in a common ancestor. These lineages are highlighted in red in the time line. Other DNA sequences from the past are not represented in living individuals; the time each became extinct is indicated by a dot.

determine the equilibrium level of variability for selectively neutral alleles (see Chapter 23).

When different human populations are analyzed for genetic variation, those in Africa are found to have more variation than those in other continents. The greater accumulation of genetic variation in African populations suggests that these populations are the oldest-an idea that is consistent with the hypothesis that humans originated in Africa and then spread to other continents. Fairly strong evidence for this hypothesis has come from studies of mitochondrial DNA sequences from different human populations. By analyzing sequences from living individuals, it is possible to work back to the ancestral sequence from which all the existing sequences could have sprung. This ancestral sequence represents the point at which the lineages of the living individuals coalesce into one individual, the common ancestor of them all (**Figure 24.17**). Then, by counting the number of mutations that occurred between the ancestral DNA sequence and the current sequences, and by dividing this number by the known mutation rate, it is possible to calculate the time that has elapsed since the common ancestor existed.

When this type of analysis is performed on mitochondrial DNA sequences, the elapsed time between the present and the time when the common ancestor lived is estimated to be 100,000 to 200,000 years. Analyses of DNA sequences on the Y chromosome, which is transmitted exclusively through males, yield a similar estimate. Thus, the coalescent principle suggests that all modern humans are descended from maternal and paternal common ancestors who lived

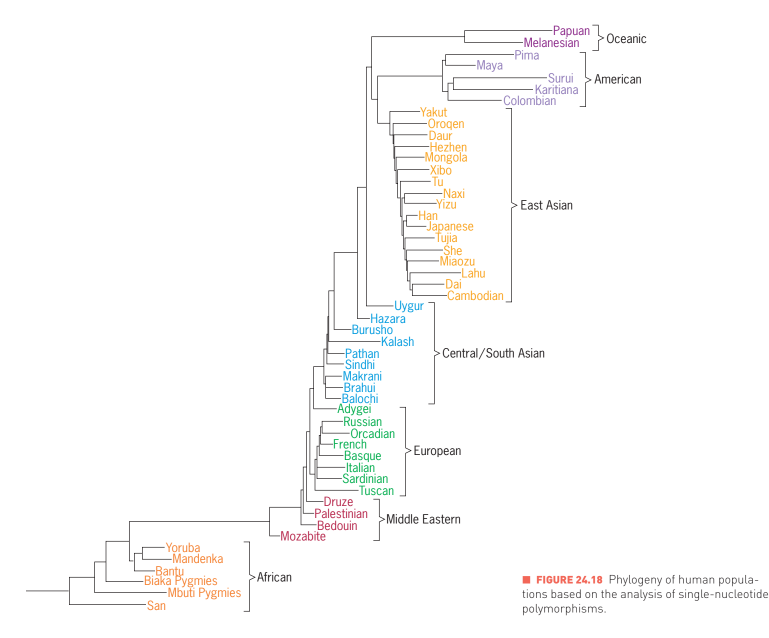
between 100,000 and 200,000 years ago. This result does not imply, however, that these common ancestors were the only two people alive at that remote time. Certainly many others were alive too. Their genetic lineages—mitochondrial in the case of females and Y chromosomal in the case of males—simply became extinct. With the coalescent method, current DNA sequences can be traced back to the individuals whose mitochondrial or Y chromosomal lineages were lucky enough to survive and spread through the species, modified, of course, by the random process of mutation.

These analyses of mitochondrial and Y-linked DNA sequences have now been supplemented with analyses of autosomal DNA. One recent study analyzed single-nucleotide polymorphisms in more than 900 human genomes from 51 different populations all over the world (**■** Figure 24.18). The results indicate that the modern human species is relatively young and that it originated in the archaic human populations of Africa. From Africa, humans migrated to Asia and Europe, and later to Australia and the Americas, ultimately becoming the dominant species on the Earth.

KEY POINTS

• Fossil evidence indicates that the remote ancestors of human beings evolved in Africa, beginning about 4 to 5 million years ago.

• Genetic evidence indicates that modern human populations may have emerged from Africa about 100,000 to 200,000 years ago and subsequently spread to other continents.



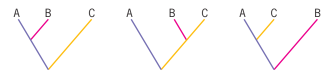
Basic Exercises

Illustrate Basic Genetic Analysis

1. Nevin Aspinwall investigated the frequencies of electrophoretically distinguishable alleles of the gene encoding alphaglycerophosphate dehydrogenase (α -GPDH) in the pink salmon (*Onchorbynchus gorbuscha*) in rivers along the northwest coast of North America, from Alaska to Washington State (1974, *Evolution* 28: 295–305). Fast, slow, and hybrid forms of α -GPDH were detected in this study; the fast and slow forms were each encoded by different alleles of the gene, and the hybrid form was produced in fish heterozy-gous for these alleles. In the sample from Dungeness River, Washington, Aspinwall observed 32 fish with the slow form, 6 with the hybrid form, and 1 with the fast form. What are the frequencies of the "fast" and "slow" alleles of the α -*GPDH* gene in the sample from this locality?

- **Answer:** In the Dungeness River sample of 39 fish, each with two copies of the α -*GPDH* gene, the frequency of the fast allele is $(2 \times 1 + 6)/(2 \times 39) = 0.10$, and the frequency of the slow allele is 1 0.10 = 0.90.
- 2. How many distinct rooted, bifurcating phylogenetic trees could show the evolutionary relationships among three different organisms?

Answer: If we denote the organisms as A, B, and C, three distinct rooted, bifurcating phylogenetic trees could show the evolutionary relationships among them:



- 3. Human and horse α -globin polypeptides differ in 18 of 141 amino acid positions. On average, how many amino acid substitutions have occurred per site in this polypeptide since the human and horse lineages diverged from a common ancestor? If the evolutionary rate for α -globin among mammals has been 0.74 substitutions per site every billion years, how much time has elapsed since the common ancestor of humans and horses existed?
- **Answer:** Human and horse α -globin differ in 18/141 = 0.128 of their amino acids. To obtain the average number of amino acid substitutions that have occurred per site since the human and the horse α -globins began evolving independently, we use the Poisson correction (see Appendix E:

Testing Your Knowledge

Integrate Different Concepts and Techniques

- 1. In his study of protein polymorphism in populations of pink salmon in rivers from Alaska to Washington State, Nevin Aspinwall collected data from mature salmon captured in 1969, 1970, and 1971. Salmon are born in rivers, and after about nine months, they migrate into the ocean, where they increase in size. When they reach two years of age, the salmon return to the river of their birth to spawn, and then they die. Because of this two-year life cycle, Pacific salmon are split into odd- and even-year populations that do not interbreed. Aspinwall found that among the salmon captured in odd years, 870 were homozygous for the slow allele of α -GPDH, 17 were homozygous for the fast allele, and 231 were heterozygous. Among the salmon captured in the even year, 649 were homozygous for the slow allele, 45 were homozygous for the fast allele, and 309 were heterozygous. What is interesting about these data?
- **Answer:** From Aspinwall's summary data, we can calculate the frequencies of the fast allele of the α -*GPDH* gene in the odd- and even-year populations. In the odd-year population, the frequency is $(2 \times 17 + 231)/(2 \times 1118) = 0.119$, and in the even-year population, it is $(2 \times 45 + 309)/(2 \times 1003) = 0.199$. Thus, the frequency of the fast allele in the even-year population is almost twice the corresponding frequency in the odd-year population. Because the two salmon populations inhabit the same territory, they are presumably subject to the same selection pressures. Thus, the observed difference in allele frequency between these populations suggests that they have diverged by random genetic drift.

Evolutionary Rates): $-\ln(1 - 0.128) = 0.136$ amino acid substitutions per site. Then, to calculate the total time that has elapsed since the common ancestor of humans and horses, we divide 0.136 amino acid substitutions per site by the estimated evolutionary rate for mammals (0.74 amino acid substitutions per site every billion years): 0.136/0.74 = 184 million years. This span of time must be divided equally between the human and horse lineages to obtain the time since their common ancestor existed: 184 million years/2 = 92 million years.

- 4. Under the Neutral Theory of Molecular Evolution, what is the rate at which selectively neutral mutations are fixed in a population by random genetic drift?
- **Answer:** The rate of fixation of selectively neutral mutations is simply the rate at which these mutations occur.
- 5. What is the genetic definition of a species?
- **Answer:** A species is a population that is reproductively isolated from all other populations—that is, it cannot exchange genes with other populations.

2.	The	following	table	shows	the	number	of	amino	acid
	diffe	rences amo	ng mo	olecules	of cy	ytochrom	e c.		

	Tuna	Silkworm	Wheat
Human	20	26	35
Tuna		27	40
Silkworm			37

If the number of amino acid sites that can be compared among these molecules is 110, what is the average number of amino acid substitutions that have occurred per site during the evolution of each pair of organisms? What is the rate at which cytochrome c has evolved among the vertebrates? If the evolutionary rate among the vertebrates can be applied to other branches of cytochrome c's phylogeny, how long ago did the insect and fish lineages diverge from a common ancestor? How long ago did the animal and plant lineages diverge from a common ancestor?

Answer: To estimate the average number of amino acid substitutions per site, we first compute the proportion of amino acid differences for each pair of organisms by dividing the observed number of differences by 110, which is the total number of sites in the cytochrome c molecule. Then we use the Poisson correction (see Appendix E: Evolutionary Rates) to calculate the average number of amino acid substitutions per site. If *d* is the proportion of amino acid differences between the cytochrome c molecules of two organisms, then the average number of substitutions per site is obtained from the formula $-\ln(1 - d)$. In the following table, the proportion

of amino acid differences is given in black and the average number of amino acid substitutions per site is given in red:

	Tuna	Silkworm	Wheat
Human	0.18	0.24	0.32
	0.20	0.27	0.38
Tuna		0.24	0.36
		0.28	0.45
Silkworm			0.34
			0.42

To calculate the rate of evolution among the vertebrates, we focus on the comparison between human and tuna cytochrome c molecules. The observed proportion of amino acid differences is 0.18, and the estimated average number of substitutions per site is slightly higher, 0.20. From the fossil record, the fish (represented by the tuna) and the tetrapod (represented by the human being) lineages are estimated to have split about 440 million years ago (mya). The total elapsed evolutionary time in these lineages is therefore 2×440 my = 880 my. We obtain the rate of amino acid substitution per site in cytochrome c by dividing the average number of amino acid substitutions per site by the total elapsed evolutionary time: 0.20 amino acid substitutions per site/880 my = 0.23 amino acid substitutions per site every billion years.

If we assume that this rate holds throughout the phylogeny of tetrapods, fish, insects, and plants—that is, if we assume a molecular clock—then we can calculate the time that has passed since the fish and insect lineages and the animal and plant lineages split from common ancestors. For the common ancestor of fish and insects, we focus on the comparison between tuna and silkworm cytochrome c molecules. The observed proportion of amino acid differences is 0.24, and the estimated average number of amino acid substitutions per site is 0.28. Dividing this

average by the estimated rate of evolution of cytochrome c (0.23 amino acid substitutions per site every billion years), we obtain the total elapsed evolutionary time: 0.28 substitutions per site/0.23 substitutions per site every billion years = 1.2 billion years. We must apportion this time equally between the fish and insect lineages. Thus, the time since they diverged from a common ancestor is calculated to be 600 million years. For the ancestor of animals and plants, we focus on the comparison between silkworm and wheat cytochrome c molecules. The observed proportion of amino acid differences is 0.34, and the estimated average number of amino acid substitutions per site is 0.42. Dividing this average by the assumed rate of evolution of cytochrome c, we estimate the total elapsed evolutionary time to be 1.82 billion years. The time since these two lineages diverged from a common ancestor is therefore 910 million years.

- **3.** In an extensive analysis of single-nucleotide polymorphisms (SNPs) among three groups of Americans, David Hinds and his collaborators (2005, *Science* 307: 1072–1079) found that among 1,586,383 SNPs examined by microarray technology, 93.5 percent were segregating in a sample of 23 African Americans, 81.1 percent were segregating in a sample of 24 European Americans, and 73.6 percent were segregating in a sample of 24 Han Chinese Americans. What do these data indicate about genetic diversity among these three groups, and how do they fit with current ideas about human evolutionary history?
- **Answer:** If we use the percentage of SNPs segregating in a population as an indicator of its genetic diversity, then clearly the African American group is the most diverse of the three groups studied. The fact that African Americans are the most diverse also fits with the idea that modern humans originated in Africa. African populations, being the oldest among all populations of modern humans, have had the longest time to accumulate genetic variants.

Questions and Problems

Enhance Understanding and Develop Analytical Skills

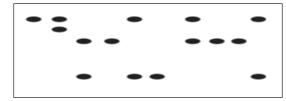
- 24.1 What was some of the evidence that led Charles Darwin to argue that species change over time?
- **24.2** Darwin stressed that species evolve by natural selection. What was the main gap in his theory?
- **24.3** Using the data in Table 24.1, and assuming that mating is random with respect to the blood type, predict the frequencies of the three genotypes of the Duffy blood-type locus in a South African and an English population.
- 24.4 Theodosius Dobzahnsky and his collaborators studied chromosomal polymorphisms in *Drosophila pseudoobscura* and its sister species in the western United States. In one study of polymorphisms in chromosome III of *D. pseudoobscura* sampled from populations at different locations in the Yosemite region of the Sierra Nevada, Dobzhansky (1948, *Genetics* 33: 158–176) recorded the

following frequencies of the Standard (ST) banding pattern:

Location	Frequency ST	Elevation (in feet)
Jacksonville	0.46	850
Lost Claim	0.41	3,000
Mather	0.32	4,600
Aspen	0.26	6,200
Porcupine	0.14	8,000
Tuolumne	0.11	8,600
Timberline	0.10	9,900
Lyell Base	0.10	10,500

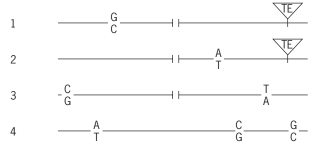
What is interesting about these data?

- **24.5** In a survey of electrophoretically detectable genetic variation in the alcohol dehydrogenase gene of *Drosophila melanogaster*; a researcher found two forms, denoted F (fast) and S (slow) in a population; 32 individuals were homozygous for the *F* allele of the gene, 22 were homozygous for the *S* allele, and 46 were heterozygous for the *F* and *S* alleles. Are the observed frequencies of the three genotypes consistent with the assumption that the population is in Hardy–Weinberg equilibrium?
- **24.6** A researcher has been studying genetic variation in fish populations by using PCR to amplify microsatellite repeats at a particular site on a chromosome (see Chapter 16). The following diagram shows the gel-fractionated products of amplifications with DNA samples from 10 different fish. How many distinct alleles of this microsatellite locus are evident in the gel?

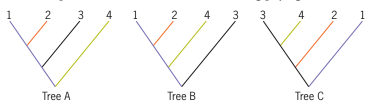


- 24.7 Within the coding region of a gene, where would you most likely find silent polymorphisms?
- **24.8** Why are the nucleotide sequences of introns more polymorphic than the nucleotide sequences of exons?
- **24.9** DNA and protein molecules are "documents of evolutionary history." Why aren't complex carbohydrate molecules such as starch, cellulose, and glycogen considered "documents of evolutionary history"?
- **24.10** A geneticist analyzed the sequences of a gene cloned from four different individuals. The four clones were identical except for a few base-pair differences, a deletion (gap), and a transposable element (TE) insertion:

Sequences



Using this information, compute the minimum number of mutations required to explain the derivation of the four sequences (1, 2, 3, and 4) in the following phylogenetic trees:



Which of these trees provides the most parsimonious explanation for the evolutionary history of the four DNA sequences?

- **24.11** The heme group in hemoglobin is held in place by histidines in the globin polypeptides. All vertebrate globins possess these histidines. Explain this observation in terms of the Neutral Theory of Molecular Evolution.
- 24.12 \bigcirc During the early evolutionary history of the vertebrates, a primordial globin gene was duplicated to form the α and β -globin genes. The rate of evolution of the polypeptides encoded by these duplicate genes has been estimated to be about 0.9 amino acid substitutions per site every billion years. By comparing the human α and β -globins, the average number of amino acid substitutions per site has been estimated to be 0.800. From this estimate, calculate when the duplication event that produced the α and β -globin genes must have occurred.
- **24.13** Ribonuclease, a protein that degrades RNA, is 124 amino acids long. A comparison between the amino acid sequences of cow and rat ribonucleases reveals 40 differences. What is the average number of amino acid substitutions that have occurred per site in these two evolutionary lineages? If the cow and the rat lineages diverged from a common ancestor 80 million years ago, what is the rate of ribonuclease evolution?
- 24.14 If a randomly mating population is segregating *n* selectively neutral alleles of a gene and each allele has the same frequency, what is the frequency of all the homozygotes in the population?
- **24.15** If the evolutionary rate of amino acid substitution in a protein is *K*, what is the average length of time between successive amino acid substitutions in this protein?
- 24.16 The coding sequence of the alcohol dehydrogenase (*Adb*) gene of *Drosophila melanogaster* consists of 765 nucleotides (255 codons); 192 of these nucleotides are functionally silent—that is, they can be changed without changing an amino acid in the Adh polypeptide. In a study of genetic variation in the *Adh* gene, Martin Kreitman observed that 13 of the 192 silent nucleotides were polymorphic. If the same level of polymorphisms existed among the nonsilent nucleotides of the *Adh* gene, how many amino acid polymorphisms would Kreitman have observed in the populations he studied?
- **24.17** How might you explain the thousandfold difference in the evolutionary rates of fibrinopeptide and histone 3?
- 24.18 A geneticist has studied the sequence of a gene in each of three species, A, B, and C. Species A and species B are sister species; species C is more distantly related. The geneticist has calculated the ratio of nonsynonymous (NS) to synonymous (S) nucleotide substitutions in the coding region of the gene in two ways—first, by comparing the gene sequences of species A and C, and second, by comparing the gene sequences of species B and C. The NS:S ratio for the comparison of species B and C is five

times greater than it is for the comparison of species A and C. What might this difference in the NS:S ratios suggest?

- 24.19 Dispersed, repetitive sequences such as transposable elements may have played a role in duplicating short regions in a genome. Can you suggest a mechanism? (*Hint:* see Chapter 17.)
- **24.20** Exon shuffling is a mechanism that combines exons from different sources into a coherent sequence that can encode a composite protein—one that contains peptides from each of the contributing exons. Alternate splicing is a mechanism that allows exons to be deleted during the expression of a gene; the mRNAs produced by alternate splicing may encode different, but related, polypeptides (see Chapter 19). What bearing do these two mechanisms have on the number of genes in a eukaryotic genome? Do these mechanisms help to explain why the gene number in the nematode *Caenorabditis elegans* is not too different from the gene number in *Homo sapiens*?
- 24.21 Drosophila mauritiana inhabits the island of Mauritius in the Indian Ocean. Drosophila simulans, a close relative, is widely distributed throughout the world. What experimental tests would you perform to determine if D. mauritiana and D. simulans are genetically different species?
- **24.22** Distinguish between allopatric and sympatric modes of speciation.

- 24.23 The *prune* gene (symbol *pn*) is X-linked in *Drosophila melanogaster*. Mutant alleles of this gene cause the eyes to be brown instead of red. A dominant mutant allele of another gene located on a large autosome causes hemizygous or homozygous *pn* flies to die; this dominant mutant allele is therefore called *Killer of prune* (symbol *Kpn*). How could mutants such as these play a role in the evolution of reproductive isolation between populations?
- 24.24 A segment of DNA in an individual may differ at several nucleotide positions from a corresponding DNA segment in another individual. For instance, one individual may have the sequence ... A ... G ... C ... and another individual may have the sequence ... T ... A ... A ... These two DNA segments differ in three nucleotide positions. Because the nucleotides within each segment are tightly linked, they will tend to be inherited together as a unit, that is, without being scrambled by recombination. We call such heritable units DNA haplotypes. Through sampling and DNA sequencing, researchers can determine which DNA haplotypes are present in a particular population. When this kind of analysis is performed on human populations by sequencing, for example, a segment of mitochondrial DNA, it is found that samples from Africa exhibit more haplotype diversity than samples from other continents. What does this observation tell us about human evolution?

Genomics on the Web at http://www.ncbi.nlm.nih.gov

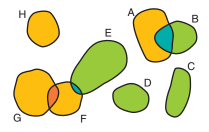
- 1. Search GenBank for AY149291, a 357-bp fragment of mitochondrial DNA (mtDNA) obtained from a Neanderthal fossil found in Germany. Use the BLAST tool to find the homologous DNA sequence in the mtDNA of modern humans. What are the coordinates of the modern human DNA sequence? How similar is the Neanderthal sequence to the modern human sequence?
- **2.** Now use the BLAST tool to find the homologous DNA sequence in the mtDNA of chimpanzee (*Pan troglodytes*). Click on the first item in the list of results to see the comparison of the Neanderthal and chimpanzee mtDNA sequences. How similar are these two sequences?
- **3.** Now search GenBank for AF347015, the complete sequence of the mtDNA of a modern human. When this sequence appears, copy the part of it that corresponds to the 357-bp fragment of Neanderthal mtDNA into a text file, delete the numbers and spaces from the copied text, and then use the resulting sequence in BLAST to compare this region in modern human mtDNA to the homologous region in chimpanzee mtDNA. How similar are these two sequences?
- **4.** From this exercise, can you draw a phylogenetic tree that shows the relationships among modern human, Neanderthal, and chimpanzee mtDNAs?

This page intentionally left blank

Appendix A The Rules of Probability

Probability theory accounts for the frequency of events—for example, the chance of getting a head on a coin toss, drawing an ace from a deck of cards, or obtaining a dominant homozygote from a mating between two heterozygotes. In each case, the event is the outcome of a process—tossing a coin, drawing a card, producing an off-spring. To determine the probability of a particular event, we must consider all possible outcomes of the process. The collection of all events is called the *sample space*. For a coin toss, the sample space contains two events, head and tail; for drawing a card, it contains 52, one for each card; and for heterozygotes producing an offspring, it contains three, *GG*, *Gg*, and *gg*. *The probability of an event is the frequency of that event in the sample space*. For example, the probabilities associated with each of the progeny from a mating between two heterozygotes are 1/4 (for *GG*), 1/2 (for *Gg*), and 1/4 (for *gg*).

Two kinds of questions often arise in problems involving probabilities: (1) What is the probability that two events, A and B, will occur together? (2) What is the probability that at least one of two events, A or B, will occur at all? The first question specifies the joint occurrence of two events—A *and* B must occur together to satisfy this question. The second question is less stringent—if *either* A *or* B occurs, the question will be satisfied. A simple diagram can help to explain the different meanings of these two questions.



The shapes in the diagram represent events in the sample space, and the sizes of the shapes reflect their relative frequencies. Overlaps between shapes indicate the joint occurrence of two events. If the events do not overlap, then they can never occur together. The first question seeks the probability that both A and B will occur; this probability is represented by the size of the overlap between the two events. The second question seeks the probability that either A or B will occur; this probability is represented by the combined shapes of the two events, including, of course, the overlap between them.

The Multiplicative Rule: If the events A and B are independent, the probability that they occur together, denoted P(A and B), is $P(A) \times P(B)$.

Here P(A) and P(B) are the probabilities of the individual events. Note that independent does not mean that they do not overlap in the sample space. In fact, nonoverlapping, or disjoint, events are not independent, for if one occurs, then the other cannot. In probability theory, independent means that one event provides no information about the other. For example, if a card drawn from a deck turns out to be an ace, we have no clue about the card's suit. Thus, drawing the ace of hearts represents the joint occurrence of two independent events—the card is an ace (A) and it is a heart (H). According to the Multiplicative Rule, $P(A \text{ and } H) = P(A) \times P(H)$, and because P(A) = 4/52 and P(H) = 1/4, $P(A \text{ and } H) = (4/52) \times (1/4) = 1/52$.

The Additive Rule: If the events A and B are independent, the probability that at least one of them occurs, denoted P(A or B), is $P(A) + P(B) - [P(A) \times P(B)]$.

Here the term P(A) × P(B), which is the probability that A and B occur together, is subtracted from the sum of the probabilities, P(A) + P(B), because the straight sum includes this term twice. As an example, suppose we seek the probability that a card drawn from a deck is either an ace or a heart. According to the Additive Rule, P(A or H) = P(A) + P(H) - [P(A) × P(H)] = (4/52) + (1/4) - [(4/52) × (1/4)] = 16/52.

If the two events do not overlap in the sample space, the Additive Rule reduces to a simpler expression: P(A or B) = P(A) + P(B). For example, suppose we seek the probability that a card drawn from a deck is either an ace or a king (K). These two events do not overlap in the sample space—that is, they are mutually exclusive. Thus, P(A or K) = P(A) + P(K) = (4/52) + (4/52) = 8/52.

Appendix B Binomial Probabilities

The progeny of crosses sometimes segregate into two distinct classes—for example, male or female, healthy or diseased, normal or mutant, dominant phenotype or recessive phenotype. To be general, we can refer to these two kinds of progeny as P and Q, and note that for any individual offspring, the probability of being P is p and the probability of being Q is q. Because there are only two classes, q = 1 - p. Suppose that the total number of progeny is n and that each one is produced independently. We can calculate the **binomial probability** that exactly x of the progeny will fall into one class and y into the other:

Probability of x in class P and y in class Q =
$$\left[\frac{n!}{x! y!}\right] p^x q^y$$

The bracketed term contains three factorial functions (n!, x!, and y!), each of which is computed as a descending series of products. For example, $n! = n(n - 1)(n - 2)(n - 3) \dots (3)(2)(1)$. If 0! is needed, it is defined as one. In the formula, the bracketed term, often called the **binomial coefficient**, counts the different ways, or orders, in which *n* offspring can be segregated so that *x* fall in the P class and *y* fall in the Q class. The other term, p^xq^y , gives the probability of obtaining a particular way or order. Because each of the orders is equally likely, multiplying this term by the bracketed term gives the probability of obtaining *x* progeny in the P class and *y* in the Q class, regardless of the order of occurrence.

If, for fixed values of *n*, *p*, and *q*, we systematically vary *x* and *y*, we can calculate a whole set of probabilities. This set constitutes a *binomial probability distribution*. With the distribution, we can answer questions such as "What is the probability that *x* will exceed a particular value?" or "What is the probability that *x* will lie between two particular values?" For example, let's consider a family with six children. What is the probability that at least four will be girls? To answer this question, we note that for any given child, the probability that it will be a girl (*p*) is 1/2 and the probability that it will be girls (and two will be boys) is therefore $[(6!)/(4! 2!)](1/2)^4 (1/2)^2 = 15/64$, which is one of the terms in the binomial distribution. However, the probability that at least four will be girls (and that no more than two will be boys) is the sum of three terms from this distribution:

Event	Binomial Formula	Probability
4 girls and 2 boys	$[(6!)/(4! 2!)] \times (1/2)^4 (1/2)^2 =$	15/64
5 girls and 1 boy	$[(6!)/(5! 1!)] \times (1/2)^5 (1/2)^1 =$	6/64
6 girls and 0 boys	$[(6!)/(6! \ 0!)] \times (1/2)^6 \ (1/2)^0 =$	1/64

Therefore, the answer is (15/64) + (6/64) + (1/64) = 22/64.

EventBinomial FormulaProbability1 girl and 5 boys $[(6!)/(1! 5!)] \times (1/2)^1 (1/2)^5 =$ 6/642 girls and 4 boys $[(6!)/(2! 4!)] \times (1/2)^2 (1/2)^4 =$ 15/643 girls and 3 boys $[(6!)/(3! 3!)] \times (1/2)^3 (1/2)^3 =$ 20/644 girls and 2 boys $[(6!)/(4! 2!)] \times (1/2)^4 (1/2)^2 =$ 15/64

The binomial distribution also provides answers to other kinds of questions. For example, what is the probability that at least one but no more than four of the children will be girls? Here the answer is the sum of four terms:

Summing up, we find that the answer is 56/64.

Let's now consider the example discussed in Chapter 3. A man and a woman, who are both heterozygous for the recessive mutant allele that causes cystic fibrosis, plan to have four children. What is the chance that one of these children will have cystic fibrosis and the other three will not? We have already seen by enumeration that the answer to this question is 108/256 (see Figure 3.14). However, this answer could also be obtained by using the binomial formula. The probability that a particular child will be affected is p = 1/4, and the probability that it will not be affected is q = 3/4. The total number of children is n = 4, the number of affected children is x = 1, and the number of unaffected children is y = 3. Putting all this together, we can calculate the probability that exactly one of the couple's four children will have cystic fibrosis as

$$[4!/1! 3!] (1/4)^1 (3/4)^3 = 4 \times (1/4) \times (27/64) = 108/256$$

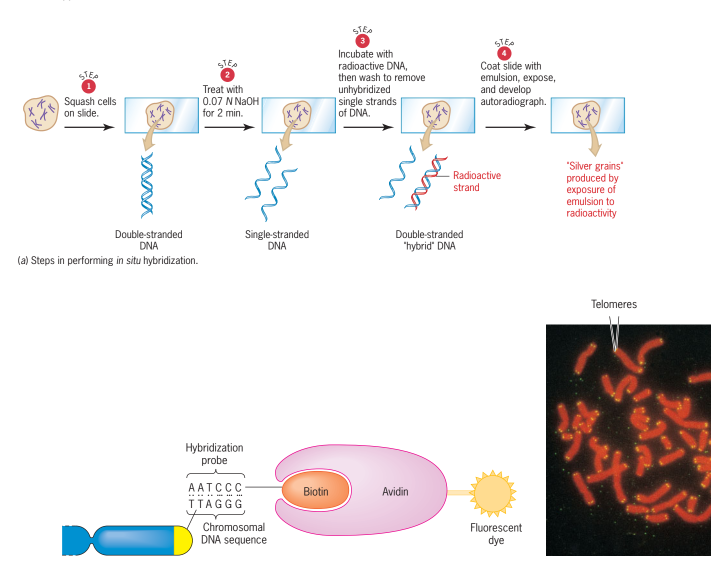
Appendix C *In Situ* Hybridization

In 1969, Mary Lou Pardue and Joseph Gall developed a procedure by which they could hybridize radioactive single strands of DNA with complementary strands of DNA in chromosomes on glass slides. By using this procedure, called *in situ* **hybridization**, Pardue and Gall were able to determine the chromosomal locations of repetitive DNA sequences. (The Latin term *in situ* means "in its original place"; hybridization is the formation of "hybrid" duplex molecules by the base pairing of complementary or partially complementary strands of DNA or RNA.) Classical *in situ* hybridization involved spreading mitotic chromosomes on a glass slide (see Figure 6.1), denaturing the DNA in the chromosomes by exposure to alkali (0.07 N NaOH) for a few minutes, rinsing with buffer to remove the alkaline solution, incubating the slide in hybridization solution containing radioactive copies of the nucleotide sequence of interest, washing off the radioactive strands that have not hybridized with complementary sequences in the chromosomes, exposing the slide to a photographic emulsion that is sensitive to low-energy radioactivity, developing the autoradiograph, and superimposing the autoradiograph on a photograph of the chromosomes (Figure 1*a*).

One of the first *in situ* hybridization experiments that Pardue and Gall performed demonstrated that the satellite DNA sequence of the mouse is located in heterochromatic regions that flank the centromeres of the mouse chromosomes. The mouse genome contains about 10⁶ copies of this satellite DNA sequence, which is about 400 nucleotide pairs long and makes up about 10 percent of the mouse genome. Similar studies have subsequently been done with the satellite DNAs of several other species, and these repetitive DNA sequences are usually located in centromeric heterochromatin or adjacent to telomeres.

A repetitive DNA sequence can be identified as satellite DNA only if the sequence has a base composition sufficiently different from that of main-band DNA to produce a distinct band during density-gradient centrifugation. Therefore, centrifugation cannot be used to identify all repetitive DNA sequences. Satellite DNA sequences usually are not expressed; that is, they do not encode RNA or protein products.

Today, *in situ* hybridization experiments are often done by using hybridization probes that are linked to fluorescent dyes or antibodies tagged with fluorescent compounds (Figure 1*b* and 1*c*). In one protocol, DNA or RNA hybridization probes are linked to the vitamin biotin, which is bound with high affinity by the egg protein avidin (Figure 1*b*). By using avidin covalently linked to a fluorescent dye, the chromosomal location of the hybridized probe can be detected by the fluorescence of the dye. This procedure, called **FISH** (Fluorescent *In Situ* Hybridization), has been used to demonstrate the presence of the repetitive sequence TTAGGG in the telomeres of human chromosomes (Figure 1*c*). The FISH procedure is very sensitive and can be used to detect the locations of single-copy sequences in human mitotic and interphase chromosomes.



1 μm

(b) Visualization of human telomeres by using fluorescent dyes and in situ hybridization.

(c) Human telomeres visualized using fluorescent probes and *in situ* hybridization.

FIGURE 1 Localization of repeated DNA sequences in chromosomes by *in situ* hybridization performed with radioactive probes (*a*) or fluorescent probes (*b* and *c*). The *in situ* hybridization procedure developed by Pardue and Gall is shown in (*a*). The use of fluorescent dyes to localize the TTAGGG repeat sequence to the telomeres of human chromosomes is illustrated in (*b*), and a photomicrograph demonstrating its telomeric location is shown in (*c*).

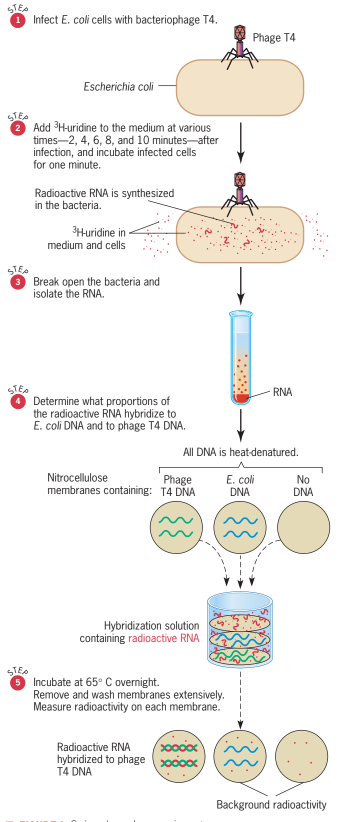
Appendix D Evidence for an Unstable Messenger RNA

The first evidence for the existence of an RNA intermediary in protein synthesis came from studies by Elliot Volkin and Lawrence Astrachan on bacteria infected with bacterial viruses. Their results, published in 1956, suggested that the synthesis of viral proteins in infected bacteria involved unstable RNA molecules specified by viral DNA. Volkin and Astrachan observed a burst of RNA synthesis after infecting *E. coli* cells with bacteriophage T2. By labeling RNA with the radioactive isotope ³²P, they demonstrated that the newly synthesized RNA molecules were unstable, turning over with half-lives of only a few minutes. In addition, they showed that the nucleotide composition of the unstable RNAs was similar to the composition of T2 DNA and unlike that of *E. coli* DNA. Their results were soon extended by studies in other laboratories.

In 1961, Sol Spiegelman and coworkers reported that the unstable RNAs synthesized in phage T4-infected cells could form RNA–DNA duplexes with denatured T4 DNA, but not with denatured *E. coli* DNA. They pulse-labeled bacteria with ³H-uridine at various times after infection with T4 phage, isolated total RNA from these cells, and determined whether the radioactive RNA molecules hybridized with *E. coli* DNA or phage T4 DNA. Their experiment is diagrammed in Figure 1.

Their results (Figure 2) demonstrated that most of the short-lived RNA molecules synthesized after infection were complementary to single strands of phage T4 DNA and not complementary to single strands of *E. coli* DNA. This finding indicated that they were produced from phage T4 DNA templates, not from *E. coli* DNA templates.

In the same year that Spiegelman and colleagues published their results, Sydney Brenner, François Jacob, and Matthew Meselson demonstrated that phage T4 proteins were synthesized on *E. coli* ribosomes. Thus, the amino acid sequences of T4 proteins were not controlled by components of the ribosomes. Instead, the ribosomes provided the workbenches on which protein synthesis occurred, but did not provide the specifications for individual proteins. These results strengthened the idea, first formally proposed by François Jacob and Jacques Monod in 1961, that unstable RNA molecules carried the specifications for the amino acid sequences of individual gene products from the genes to the ribosomes. Subsequent research firmly established the role of these unstable RNAs, now called messenger RNAs or mRNAs, in the transfer of genetic information from genes to the sites of protein synthesis in the cytoplasm.





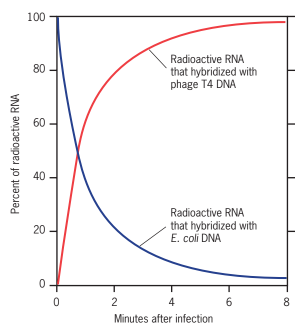


FIGURE 2 Rapid switch from the transcription of *E. coli* genes to phage T4 genes in T4-infected bacteria.

Appendix E Evolutionary Rates

Nucleotide and amino acid sequences are the fundamental data for the study of molecular evolution. Once homologous sequences from different organisms have been aligned, we can ascertain how many positions in the molecules are the same or different; then, with the help of fossil data on the history of the organisms, we can estimate the rate of molecular evolution.

The simplest case is when we compare the amino acid sequences of two homologous polypeptides. Consider, for example, the two polypeptides shown in Figure 1. In three of the four positions in these two polypeptides, the amino acids are identical; in the remaining position, they are different—glycine in one polypeptide and serine in the other. This single amino acid difference indicates that at least one amino acid substitution occurred during the evolution of the two polypeptides. The ancestral amino acid might have been serine, in which case the glycine in one polypeptide represents a substitution event, or the ancestral amino acid might have been glycine, in which case the serine in the other polypeptide represents a substitution event.

However, the history of these polypeptides might have been more complicated. The ancestral amino acid at the variable position might have been something other than serine or glycine—say, for example, arginine. In this case, both of the descendant polypeptides must have sustained amino acid substitutions during their evolution. Thus, the minimum number of amino acid substitutions would be two. We say "minimum" because multiple substitutions might have occurred at the variable position in either of the descendant polypeptides during their evolution. Thus, by focusing on amino acid differences at corresponding positions in homologous polypeptides, we cannot count the actual number of amino acid substitutions that have taken place. All we can say is that *at least* one such substitution has occurred. This uncertainty poses a problem for estimating the rate of molecular evolution, which, after all, is the total number of amino acid substitutions that have been evolving.

To get around this problem we focus—paradoxically—on the amino acids that are the same in the two polypeptides. These amino acids have presumably not changed in either polypeptide since the two evolving lineages diverged from a common ancestor. Thus, they provide information about the probability that an amino acid substitution does *not* occur during the course of evolution. If we can estimate this probability, then

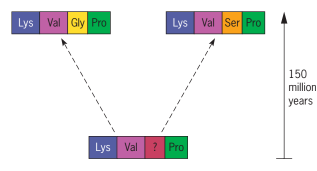


FIGURE 1 Comparison of two homologous polypeptides that have evolved independently for 150 million years.

we can turn the situation around and estimate the probability that a substitution does occur, and from it we can obtain the evolutionary rate.

Suppose that S is the proportion of amino acids that are the *same* in two polypeptides—in our example, S = 0.75—and suppose that v is the probability that an amino acid substitution occurs at a site in either polypeptide during one year of evolutionary time—that is, v is the yearly rate of amino acid substitution per site in these polypeptides. By defining v in this way, 1 - v is the probability that an amino acid substitution does not occur at a site in any one year of evolutionary time.

From the fossil record we can determine when the two lineages carrying these polypeptides diverged from a common ancestor. For the polypeptides in Figure 1, this divergence occurred 150 million years ago. In general, if the time since divergence from the common ancestor is T years, then the total evolutionary time for the two lineages is T + T = 2T years. This sum represents the total number of yearly opportunities for an amino acid substitution to occur at a particular site in the evolving polypeptides. It also represents the total number of yearly opportunities for a substitution *not* to occur at this site. Thus, at the end of the evolutionary process, the probability that an amino acid substitution has not occurred at a particular site in either of the polypeptides is the product of all the individual, independent chances for it not to occur, which equals $(1 - v)^{2T}$. To say it another way, the probability that corresponding amino acids in the two polypeptides have remained the same during the evolutionary process is the probability that neither of them has changed in any one year, which is $(1 - v)^{2T}$. We can estimate this probability by the proportion of amino acids that are currently the same in the two polypeptides—that is, by *S*. Thus,

$$S = (1 - v)^{27}$$

To solve for v, the yearly rate of amino acid substitution per site, we take the natural logarithm of both sides of the equation.

$$\ln S = \ln(1 - v)^{2T}$$
$$\ln S = 2T \ln(1 - v)$$

Because v is a very small number—in fact quite close to zero— $\ln(1 - v)$ is approximately equal to -v (the logarithm curve is nearly linear when the argument of the logarithm function is close to 1). Thus,

$$\ln S = -2Tv$$

which implies that

$$v = (-\ln S)/2T$$

With this formula we can estimate the rate of molecular evolution of two homologous polypeptides by (1) calculating the proportion of sites in them that are the same, (2) taking the natural logarithm of this proportion, and then (3) dividing by the total elapsed evolutionary time. In our example, S = 0.75 and 2T = 300 million years; thus, v is $[-\ln(0.75)]/300 = 0.97$ amino acid substitutions per site every billion years.

As discussed above, some of the amino acid sites that are different in two polypeptides have changed once, others have changed twice, and still others have changed multiple times during the evolutionary process. The quantity 2Tv is the average number of amino acid substitutions that have occurred per site during the evolution of the polypeptides. If we assume that amino acid substitutions occur randomly and independently throughout time, then we can use this average to calculate the probability that a site has changed a specified number of times. The calculation uses the formula for a probability distribution that is widely used by scientists. It is called the *Poisson probability distribution*. In the context of molecular evolution, the Poisson formula is

Probability of *n* changes occurring at an amino acid site $= e^{-2Tv}(2Tv)^n/n!$

The average number of amino acid substitutions that have occurred per site (2Tv) appears twice in this formula—as the exponent of the first term and as the argument of the power function in the second term. Thus, it is the key parameter of the Poisson formula.

In our example, 2Tv is estimated from $-\ln S = -\ln(0.75)$ to be 0.29 amino acid substitutions per site. This estimate is slightly greater than the proportion of amino acids that are different in the two polypeptides (1 - S = 0.25) because it takes into account the possibility that multiple substitutions may have occurred at individual amino acid sites. We say that 2Tv is the Poisson-corrected number of amino acid differences between the two polypeptides.

With an estimate of 2Tv, we can use the Poisson formula to calculate the probability that a particular amino acid site has changed exactly once, twice, and so on.

Probability of 1 change =
$$e^{-2Tv}(2Tv) = 0.22$$

Probability of 2 changes = $e^{-2Tv}(2Tv)^2/2 = 0.03$

The probability that no changes have occurred is

Probability of 0 changes = $e^{-2Tv} = 0.75$

In this example, the probability that more than two changes have occurred is negligible. However, if the Poisson parameter 2Tv were greater, multiple changes would have some chance of occurring. For example, if 2Tv = 0.7, the probability for three changes at a site is 0.03, and the probability for four changes is 0.005.

Statistical procedures analogous to the Poisson correction have been developed to estimate evolutionary rates from comparisons of homologous DNA sequences. However, these procedures are more complicated because the identity of a nucleotide in two DNA sequences does not necessarily imply that this nucleotide remained unchanged during the evolution of these sequences. Methods to deal with this issue can be found in specialized texts on the subject of molecular evolution. This page intentionally left blank

Answers to Odd-Numbered Questions and Problems

CHAPTER 1

- 1.1 Mendel postulated transmissible factors—genes—to explain the inheritance of traits. He discovered that genes exist in different forms, which we now call alleles. Each organism carries two copies of each gene. During reproduction, one of the gene copies is randomly incorporated into each gamete. When the male and female gametes unite at fertilization, the gene copy number is restored to two. Different alleles may coexist in an organism. During the production of gametes, they separate from each other without having been altered by coexistence.
- **1.3** The bases present in DNA are adenine, thymine, guanine, and cytosine; the bases present in RNA are adenine, uracil, guanine, and cytosine. The sugar in DNA is deoxyribose; the sugar in RNA is ribose.
- 1.5 TAACGGCAG.
- 1.7 GAACGGUCT.
- **1.9** Sometimes DNA is synthesized from RNA in a process called reverse transcription. This process plays an important role in the life cycles of some viruses.
- **1.11** The two mutant forms of the β -globin gene are properly described as alleles. Because neither of the mutant alleles can specify a "normal" polypeptide, an individual who carries each of them would probably suffer from anemia.

CHAPTER 2

- **2.1** Sugars combine to form carbohydrates; amino acids combine to form proteins.
- **2.3** In a eukaryotic cell the many chromosomes are contained within a membrane-bounded structure called the nucleus; the chromosomes of prokaryotic cells are not contained within a special subcellular compartment. Eukaryotic cells usually possess a well-developed internal system of membranes, and they also have membranebounded subcellular organelles such as mitochondria and chloroplasts; prokaryotic cells do not typically have a system of internal membranes (although some do), nor do they possess membrane-bounded organelles.
- **2.5** Prokaryotic chromosomes are typically (but not always) smaller than eukaryotic chromosomes; in addition, pro-karyotic chromosomes are circular, whereas eukaryotic

chromosomes are linear. For example, the circular chromosome of *E. coli*, a prokaryote, is about 1.4 mm in circumference. By contrast, a linear human chromosome may be 10 to 30 cm long. Prokaryotic chromosomes also have a comparatively simple composition: DNA, some RNA, and some protein. Eukaryotic chromosomes are more complex: DNA, some RNA, and lots of protein.

- **2.7** Interphase typically lasts longer than M phase. During interphase, DNA must be synthesized to replicate all the chromosomes. Other materials must also be synthesized to prepare for the upcoming cell division.
- **2.9** (1) Anaphase: (f), (h); (2) metaphase: (e), (i); (3) prophase: (b), (c), (d); (4) telophase: (a), (g).
- **2.11** Chromosomes 11 and 16 would not be expected to pair with each other during meiosis; these chromosomes are heterologues, not homologues.
- **2.13** Crossing over occurs *after* chromosomes have duplicated in cells going through meiosis.
- **2.15** Chromosome disjunction occurs during anaphase I. Chromatid disjunction occurs during anaphase II.
- **2.17** Among eukaryotes, there doesn't seem to be a clear relationship between genome size and gene number. For example, humans, with 3.2 billion base pairs of genomic DNA, have about 20,500 genes, and *Arabidopsis* plants, with about 150 million base pairs of genomic DNA, have roughly the same number of genes as humans. However, among prokaryotes, gene number is rather tightly correlated with genome size, probably because there is so little nongenic DNA.
- **2.19** It is a bit surprising that yeast chromosomes are, on average, smaller than *E. coli* chromosomes because, as a rule, eukaryotic chromosomes are larger than prokaryotic chromosomes. Yeast is an exception because its genome—not quite three times the size of the *E. coli* genome—is distributed over 16 separate chromosomes.
- **2.21** One of the pollen nuclei fuses with the egg nucleus in the female gametophyte to form the zygote, which then develops into an embryo and ultimately into a sporophyte. The other genetically functional pollen nucleus fuses with two nuclei in the female gametophyte to form a triploid nucleus, which then develops into a triploid

tissue, the endosperm; this tissue nourishes the developing plant embryo.

2.23 (a) 5, (b) 5, (c) 15, (d) 10.

CHAPTER 3

- **3.1** (a) All tall; (b) 3/4 tall, 1/4 dwarf; (c) all tall; (d) 1/2 tall, 1/2 dwarf.
- 3.3 The data suggest that coat color is controlled by a single gene with two alleles, C (gray) and c (albino), and that C is dominant over c. On this hypothesis, the crosses are: gray (CC) × albino (cc) → F₁ gray (Cc); F₁ × F₁ → 3/4 gray (1 CC: 2 Cc), 1/4 albino (cc). The expected results in the F₂ are 203 gray, 67 albino. To compare the observed and expected results, compute χ² with one degree of freedom: (198 203)²/203 + (72 67)²/67 = 0.496, which is not significant at the 5 percent level. Thus, the results are consistent with the hypothesis.
- 3.5 (a) Checkered, red (CC BB) × plain, brown (cc bb) → F₁ all checkered, red (Cc Bb); (b) F₂ progeny: 9/16 checkered, red (C- B-), 3/16 plain, red (cc B-), 3/16 checkered, brown (C- bb), 1/16 plain, brown (cc bb).
- **3.7** Among the F₂ progeny with long, black fur, the genotypic ratio is 1 *BB RR*: 2 *BB Rr*: 2 *Bb RR*: 4 *Bb Rr*; thus, 1/9 of the rabbits with long, black fur are homozygous for both genes.
- 3.9

F ₁ Gametes	F ₂ Genotypes	F ₂ Phenotypes
(a) 2	3	2
(b) $2 \times 2 = 4$	$3 \times 3 = 9$	$2 \times 2 = 4$
(c) $2 \times 2 \times 2 = 8$	$3 \times 3 \times 3 = 27$	$2 \times 2 \times 2 = 8$
(d) 2^n	3 ⁿ	2^n , where <i>n</i> is the number of genes

3.11 (a) 1, reject; (b) 2, reject; (c) 3, accept; (d) 3, accept.

- **3.13** $\chi^2 = (30 25)^2/25 + (20 25)^2/25 = 2$, which is less than 3.84, the 5 percent critical value for a chi-square statistic with one degree of freedom; consequently, the observed segregation ratio is consistent with the expected ratio of 1:1.
- **3.15** Half the children from $Aa \times aa$ matings would have albinism. In a family of three children, the chance that one will be unaffected and two affected is $3 \times (1/2)^1 \times (1/2)^2 = 3/8$.
- **3.17** Man (*Ccff*) × woman (*ccFf*). (a) *ccff*, $(1/2) \times (1/2) = 1/4$; (b) *Cc ff*, $(1/2) \times (1/2) = 1/4$; (c) *cc Ff*, $(1/2) \times (1/2) = 1/4$; (d) *Cc Ff*, $(1/2) \times (1/2) = 1/4$.
- **3.19** $(1/2)^3 = 1/8$.
- **3.21** (20/64) + (15/64) + (6/64) + (1/64) = 42/64.
- **3.23** (a) $(1/2) \times (1/4) = 1/8$; (b) $(1/2) \times (1/2) \times (1/4) = 1/16$; (c) $(2/3) \times (1/4) = 1/6$; (d) $(2/3) \times (1/2) \times (1/2) \times (1/4) = 1/24$.
- **3.25** For III-1 \times III-2, the chance of an affected child is 1/2. For IV-2 \times IV-3, the chance is zero.

3.27 1/2.

3.29 The researcher has obtained what appears to be a non-Mendelian ratio because he has been studying only families in which at least one child shows albinism. In these families, both parents are heterozygous for the mutant allele that causes albinism. However, other couples in the population might also be heterozygous for this allele but, simply due to chance, have failed to produce a child with albinism. If a man and a woman are both heterozygous carriers of the mutant allele, the chance that a child they produce will not have albinism is 3/4. The chance that four children they produce will not have albinism is therefore $(3/4)^4 = 0.316$. In the entire population of families in which two heterozygous parents have produced a total of four children, the average number of affected children is 1. Among families in which two heterozygous parents have produced at least one affected child among a total of four children, the average must be greater than 1. To calculate this *conditional average*, let's denote the number of children with albinism by x, and the probability that exactly x of the four children have albinism by P(x). The average number of affected children among families in which at least one of the four children is affected-that is, the conditional average-is therefore $\sum xP(x)/(1 - P(0))$, where the sum starts at x = 1 and ends at x = 4. We start the sum at x = 1 because we must exclude those cases in which none of the four children is affected. The divisor (1 - P(0)) is the probability that the couple has had at least one affected child among their four children. Now P(0) = 0.316 and $\sum xP(x) = 1$. Therefore, the average we seek is simply 1/(1 - 0.316) = 1.46. If, in the subset of families with at least one affected child, the average number of affected children is 1.46, then the average number of unaffected children is 4 - 1.46 = 2.54. Thus the expected ratio of unaffected to affected children in these families is 2.54:1.56, or 1.74:1, which is what the researcher has observed.

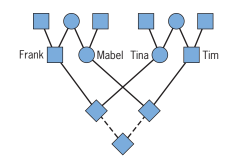
CHAPTER 4

4.1 M and MN.

4.3

	Parents	Offspring
(a)	yellow \times yellow	2 yellow: 1 light belly
(b)	yellow \times light belly	2 yellow: 1 light belly: 1 black and tan
(c)	black and tan $ imes$ yellow	2 yellow: 1 black and tan: 1 black
(d)	light belly $ imes$ light belly	all light belly
(e)	light belly $ imes$ yellow	1 yellow: 1 light belly
(f)	agouti $ imes$ black and tan	1 agouti: 1 black and tan
(g)	black and tan $ imes$ black	1 black and tan: 1 black
(h)	yellow $ imes$ agouti	1 yellow: 1 light belly
(i)	yellow \times yellow	2 yellow: 1 light belly

- **4.5** (a) all AB; (b) 1 A: 1 B; (c) 1 A: 1 B: 1 AB: 1 O; (d) 1 A: 1 O.
- 4.7 No. The woman is I^AI^B. One man could be either I^AI^A or I^Ai; the other could be either I^BI^B or I^Bi. Given the uncertainty in the genotype of each man, either could be the father of the child.
- **4.9** The woman is *ii L^ML^M*; the man is *I^AI^B L^ML^N*; the blood types of the children will be A and M, A and MN, B and M, and B and MN, all equally likely.
- **4.11** The individuals III-4 and III-5 must be homozygous for recessive mutations in different genes; that is, one is *aa BB* and the other is *AA bb*; none of their children is deaf because all of them are heterozygous for both genes (*Aa Bb*).
- **4.13** No. The test for allelism cannot be performed with dominant mutations.
- **4.15** The mother is *Bb* and the father is *bb*. The chance that a daughter is *Bb* is 1/2. (a) The chance that the daughter will have a bald son is $(1/2) \times (1/2) = 1/4$. (b) The chance that the daughter will have a bald daughter is zero.
- **4.17** (a) 3/4 walnut, 1/4 rose; (b) 1/2 walnut, 1/2 pea; (c) 3/8 walnut, 3/8 rose, 1/8 pea, 1/8 single; (d) 1/2 rose, 1/2 single.
- **4.19** 12/16 white, 3/16 yellow, 1/16 green.
- **4.21** 9/16 dark red (wild-type), 3/16 brownish-purple, 3/16 bright red, 1/16 white.
- **4.23** 9 black: 39 gray: 16 white.
- **4.25** (a) purple × red; (b) proportion white (aa) = 1/4; (c) proportion red (A B C dd) = (3/4)(3/4)(3/4)(1/2) = 27/128, proportion white (aa) = 1/4 = 32/128, proportion blue (A B cc Dd) = (3/4)(3/4)(1/4)(1/2) = 9/128.
- **4.27** (a) Because the F_2 segregation is approximately 9 red: 7 white, flower color is due to epistasis between two independently assorting genes: red = A- B- and white = aa B-, A- bb, or aa bb. (b) colorless precursor—A→ colorless product—B→ red pigment.
- **4.29** $F_A = (1/2)^5 = 1/32$; $F_B = 2 \times (1/2)^6 = 1/32$; $F_C = 2 \times (1/2)^7 = 1/64$.
- **4.31** The pedigree is as follows.



The coefficient of relationship between the offspring of the two couples is obtained by calculating the inbreeding coefficient of the imaginary child from a mating

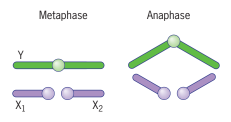
between these offspring and multiplying by 2: $[(1/2)^5 \times 2] \times 2 = 1/8$.

4.33 The mean ear length for randomly mated maize is 24 cm, and that for maize from one generation of self-fertilization is 20 cm. The inbreeding coefficient of the offspring of one generation of self-fertilization is 1/2, and the inbreeding coefficient of the offspring of two generations of self-fertilization is (1/2)(1 + 1/2) = 3/4. Mean ear length (*Y*) is expected to decline linearly with inbreeding according to the equation $Y = 24 - b F_1$ where *b* is the slope of the line. The value of *b* can be determined from the two values of *Y* that are given. The difference between these two values (4 cm) corresponds to an increase in *F* from 0 to 1/2. Thus, b = 4/(1/2) = 8 cm, and for F = 3/4, the predicted mean ear length is $Y = 24 - 8 \times (3/4) = 18$ cm.

CHAPTER 5

- **5.1** The male-determining sperm carries a Y chromosome; the female-determining sperm carries an X chromosome.
- 5.3 All the daughters will be green and all the sons will be rosy.
- **5.5** XX is female, XY is male, XXY is female, XXX is female (but barely viable), XO is male (but sterile).
- **5.7** No. Defective color vision is caused by an X-linked mutation. The son's X chromosome came from his mother, not his father.
- **5.9** The risk for the child is P(mother is C/c) × P(mother transmits c) × P(child is male) = $(1/2) \times (1/2) \times (1/2) = 1/8$; if the couple has already had a child with color blindness, P(mother is C/c) = 1, and the risk for each subsequent child is 1/4.
- **5.11** Each of the rare vermilion daughters must have resulted from the union of an X(v) X(v) egg with a Y-bearing sperm. The diplo-X eggs must have originated through nondisjunction of the X chromosomes during oogenesis in the mother. However, we cannot determine if the nondisjunction occurred in the first or the second meiotic division.
- **5.13** Each of the rare white-eyed daughters must have resulted from the union of an X(w) X(w) egg with a Y-bearing sperm. The rare diplo-X eggs must have originated through nondisjunction of the X chromosomes during the second meiotic division in the mother.
- 5.15 Female.
- 5.17 Male.
- **5.19** (a) Female; (b) intersex; (c) intersex; (d) male: (e) female; (f) male.
- **5.21** *Drosophila* does not achieve dosage compensation by inactivating one of the X chromosomes in females.
- **5.23** Because the centromere is at the end of each small X chromosome but in the middle of the larger Y, X₁ and X₂ both pair at the centromere of the Y during metaphase.

Then during anaphase, the two X chromosomes disjoin together and segregate from the Y chromosome.

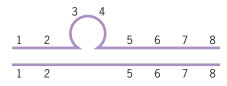


5.25 Eye color in canaries is due to a gene on the Z chromosome, which is present in two copies in males and one copy in females. The allele for pink color at hatching (*p*) is recessive to the allele for black color at hatching (P). There is no eve color gene on the other sex chromosome (W), which is present in one copy in females and absent in males. The parental birds were genotypically p/W (cinnamon females) and P/P (green males). Their F_1 sons were genotypically p/P (with black eyes at hatching). When these sons were crossed to green females (genotype P/W), they produced F, progeny that sorted into three categories: males with black eyes at hatching (P/-, half the total progeny), females with black eyes at hatching (P/W), a fourth of the total progeny), and females with pink eyes at hatching (p/W, a)fourth of the total progeny). When these sons were crossed to cinnamon females (genotype p/W), they produced F₂ progeny that sorted into four equally frequent categories: males with black eyes at hatching (genotype P/p), males with pink eyes at hatching (genotype p/p), females with black eyes at hatching (genotype P/W), and females with pink eyes at hatching (genotype p/W).

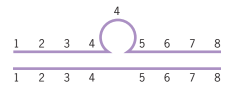
CHAPTER 6

- 6.1 Use one of the banding techniques.
- **6.3** In allotetraploids, each member of the different sets of chromosomes can pair with a homologous partner during prophase I and then disjoin during anaphase I. In triploids, disjunction is irregular because homologous chromosomes associate during prophase I by forming either bivalents and univalents or by forming trivalents.
- **6.5** The fertile plant is an allotetraploid with 7 pairs of chromosomes from species A and 9 pairs of chromosomes from species B; the total number of chromosomes is $(2 \times 7) + (2 \times 9) = 32$.
- **6.7** XX is female, XY is male, XO is female (but sterile), XXX is female, XXY is male (but sterile), XYY is male.
- **6.9** The fly is a gynandromorph, that is, a sexual mosaic. The yellow tissue is X(y)/O and the gray tissue is X(y)/X(+). This mosaicism must have arisen through loss of the X chromosome that carried the wild-type allele, presumably during one of the early embryonic cleavage divisions.

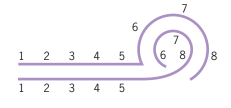
- **6.11** Nondisjunction must have occurred in the mother. The color-blind woman with Turner syndrome was produced by the union of an X-bearing sperm, which carried the mutant allele for color blindness, and a nullo-X egg.
- **6.13** XYY men would produce more children with sex chromosome abnormalities because their three sex chromosomes will disjoin irregularly during meiosis. This irregular disjunction will produce a variety of aneuploid gametes, including the XY, YY, XYY, and nullo sex chromosome constitutions.
- **6.15** (a) Deletion:



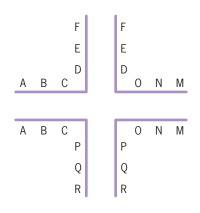
(b) Duplication:



(c) A terminal inversion:

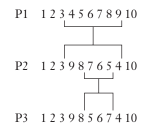


6.17



6.19 The boy carries a translocation between chromosome 21 and another chromosome, say chromosome 14. He also carries a normal chromosome 21 and a normal chromosome 14. The boy's sister carries the translocation, one normal chromosome 14, and two normal copies of chromosome 21.

- **6.21** All the daughters will be yellow-bodied, and all the sons will be white-eyed.
- 6.23 The three populations are related by a series of inversions:



- **6.25** The mother is heterozygous for a reciprocal translocation between the long arms of the large and small chromosomes; a piece from the long arm of the large chromosome has been broken off and attached to the long arm of the short chromosome. The child has inherited the rearranged large chromosome and the normal small chromosome from the mother. Thus, because the rearranged large chromosome is deficient for some of its genes, the child is hypoploid.
- **6.27** The sons will have bright red eyes because they will inherit the Y chromosome with the bw^+ allele from their father. The daughters will have white eyes because they will inherit an X chromosome from their father.
- **6.29** XX zygotes will develop into males because one of their X chromosomes carries the *SRY* gene that was translocated from the Y chromosome. XY zygotes will develop into females because their Y chromosome has lost the *SRY* gene.

CHAPTER 7

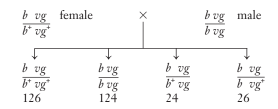
- 7.1 If Mendel had known of the existence of chromosomes, he would have realized that the number of factors determining traits exceeds the number of chromosomes, and he would have concluded that some factors must be linked on the same chromosome. Thus, Mendel would have revised the Principle of Independent Assortment to say that factors on different chromosomes (or far apart on the same chromosome) are inherited independently.
- **7.3** No. The genes *a* and *d* could be very far apart on the same chromosome—so far apart that they recombine freely, that is, 50 percent of the time.
- 7.5 Yes, if they are very far apart.
- 7.7 (a) Cross: a⁺ b⁺/a⁺ b⁺ × a b/a b. Gametes: a⁺ b⁺ from one parent, a b from the other. F₁: a⁺ b⁺/a b. (b) 40% a⁺ b⁺, 40% a b, 10% a⁺ b, 10% a b⁺. (c) F₂ from testcross: 40% a⁺ b⁺/a b, 40% a b/a b, 10% a⁺ b/a b, 10% a b⁺/a b. (d) Coupling linkage phase. (e) F₂ from intercross:

		Sperm			
		40% <i>a</i> ⁺ <i>b</i> ⁺	40% a b	$10\% a^+ b$	$10\% \ a \ b^+$
	$40\% a^+ b^+$	16%	16%	4%	4%
		$a^+ b^+/a^+ b^+$	a ⁺ b ⁺ /a b	$a^{+}b^{+}/a^{+}b$	$a^{+}b^{+}/ab^{+}$
	40% a b	16%	16%	4%	4%
Eggs		$a b/a^{+} b^{+}$	a b/a b	a b/a ⁺ b	$a b/a b^+$
	10% a ⁺ b	4%	4%	1%	1%
		$a^{+} b/a^{+} b^{+}$	a ⁺ b/a b	$a^+ b/a^+ b$	$a^+ b/a b^+$
	10% <i>a b</i> ⁺	4%	4%	1%	1%
		$a b^{+}/a^{+} b^{+}$	a b ⁺ /a b	$a b^+/a^+ b$	$a b^+/a b^+$

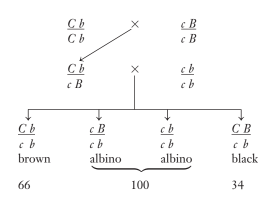
Summary of phenotypes:

a^+ and b^+	66%	a and b^+	9%
a^+ and b	9%	a and b	16%

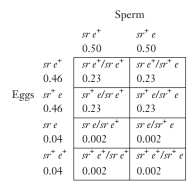
- 7.9 Coupling heterozygotes a⁺ b⁺/a b would produce the following gametes: 30% a⁺ b⁺, 30% a b, 20% a⁺ b, 20% a b⁺; repulsion heterozygotes a⁺ b/a b⁺ would produce the following gametes: 30% a⁺ b, 30% a b⁺, 20% a⁺ b⁺, 20% a b. In each case, the frequencies of the testcross progeny would correspond to the frequencies of the gametes.
- 7.11 Yes. Recombination frequency = (24 + 26)/(126 + 24 + 26 + 124) = 0.167. Cross:



7.13 Yes. Recombination frequency is estimated by the frequency of black offspring among the colored offspring: 34/(66 + 34) = 0.34. Cross:



7.15 (a) The F₁ females, which are sr e⁺/sr⁺ e, produce four types of gametes: 46% sr e⁺, 46% sr⁺ e, 4% sr e, 4% sr⁺ e⁺. (b) The F₁ males, which have the same genotype as the F₁ females, produce two types of gametes: 50% sr e⁺, 50% sr⁺ e; remember, there is no crossing over in *Drosophila* males. (c) 46% striped, gray; 46% unstriped, ebony; 4% striped, ebony; 4% unstriped, gray. (d) The offspring from the intercross can be obtained from the following table.



- 7.17 (a) The F_1 females, which are $cn vg^+/cn^+ vg$, produce four types of gametes: 45% $cn vg^+$, 45% $cn^+ vg$, 5% $cn^+ vg^+$, 5% cn vg. (b) 45% cinnabar eyes, normal wings; 45% reddish-brown eyes, vestigial wings; 5% reddish-brown eyes, normal wings; 5% cinnabar eyes, vestigial wings.
- **7.19** In the enumeration below, classes 1 and 2 are parental types, classes 3 and 4 result from a single crossover between *Pl* and *Sm*, classes 5 and 6 result from a single crossover between *Sm* and *Py*, and classes 7 and 8 result from a double crossover, with one of the exchanges between *Pl* and *Sm* and the other between *Sm* and *Py*.

Class	Phenotypes	(a) Fre- quency with no Inter- ference	(b) Fre- quency with Complete Inter- ference
1	purple, salmon, pigmy	0.405	0.40
2	green, yellow, normal	0.405	0.40
3	purple, yellow, normal	0.045	0.05
4	green, salmon, pigmy	0.045	0.05
5	purple, salmon, normal	0.045	0.05
6	green, yellow, pigmy	0.045	0.05
7	purple, yellow, pigmy	0.005	0
8	green, salmon, normal	0.005	0

- 7.21 The double crossover classes, which are the two that were not observed, establish that the gene order is y-w-ec. Thus, the F_1 females had the genotype y w ec/+ + +. The distance between y and w is estimated by the frequency of recombination between these two genes: (8 + 7)/1000 = 0.015; similarly, the distance between w and ec is (18 + 23)/1000 = 0.041. Thus, the genetic map for this segment of the X chromosome is y-1.5 cM-w-4.1 cM-ec.
- 7.23 (a) Two of the classes (the parental types) vastly outnumber the other six classes (recombinant types);
 (b) st + +/+ ss e; (c) st—ss—e; (d) [(145 + 122) × 1 +

 $(18) \times 2$]/1000 = 30.3 cM; (e) (122 + 18)/1000 = 14.0 cM; (f) (0.018)/(0.163 × 0.140) = 0.789. (g) *st* + +/+ *ss e* females × *st ss e*/*st ss e* males \rightarrow 2 parental classes and 6 recombinant classes.

- **7.25** The F₁ females are genotypically pn + l + g. Among their sons, 40 percent will be recombinant for the two X-linked genes, and half of the recombinants will have the wild-type alleles of these genes. Thus the frequency of sons with dark red eyes will be $1/2 \times 40\% = 20\%$.
- **7.27** $(P/2)^2$.
- **7.29** From the parental classes, + c and a b +, the heterozygous females must have had the genotype + c/a b +. The missing classes, + b + and a + c, which would represent double crossovers, establish that the gene order is b-a-c. The distance between b and a is (96 + 110)/1000 = 20.6 cM, and that between a and c is (65 + 75)/1000 = 14.0 cM. Thus, the genetic map is b-20.6 cM-a-14.0 cM-c.
- **7.31** II-1 has the genotype C h/c H; that is, she is a repulsion heterozygote for the alleles for color blindness (*c*) and hemophilia (*h*). None of her children are recombinant for these alleles.
- **7.33** The woman is a repulsion heterozygote for the alleles for color blindness and hemophilia—that is, she is C h/c H. If the woman has a boy, the chance that he will have hemophilia is 0.5 and the chance that he will have color blindness is 0.5. If we specify that the boy have only one of these two conditions, then the chance that he will have color blindness is 0.45. The reason is that the boy will inherit a nonrecombinant X chromosome with a probability of 0.9, and half the nonrecombinant X chromosomes will carry the mutant allele for color blindness and the other half will carry the mutant allele for hemophilia. The chance that the boy will have both conditions is 0.05, and the chance that he will have neither condition is 0.05. The reason is that the boy will inherit a recombinant X chromosome with a probability of 0.1, and half the recombinant X chromosomes will carry both mutant alleles and the other half will carry neither mutant allele.
- **7.35** A two-strand double crossover within the inversion; the exchange points of the double crossover must lie between the genetic markers and the inversion breakpoints.

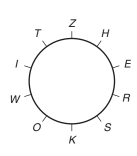
CHAPTER 8

- **8.1** Viruses reproduce and transmit their genes to progeny viruses. They utilize energy provided by host cells and respond to environmental and cellular signals like other living organisms. However, viruses are obligate parasites; they can reproduce only in appropriate host cells.
- **8.3** Bacteriophage T4 is a virulent phage. When it infects a host cell, it reproduces and kills the host cell in the process. Bacteriophage lambda can reproduce and kill the host bacterium—the lytic response—just like phage T4, or it can insert its chromosome into the chromosome of

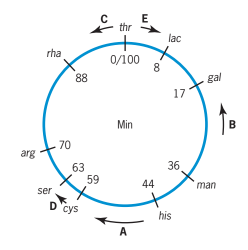
the host and remain there in a dormant state—the lysogenic response.

- 8.5 The insertion of the phage λ chromosome into the host chromosome is a site-specific recombination process catalyzed by an enzyme that recognizes specific sequences in the λ and *E. coli* chromosomes. Crossing over between homologous chromosomes is not sequence-specific. It can occur at many sites along the two chromosomes.
- 8.7 The *a*, *b*, and *c* mutations are closely linked and in the order b-a-c on the chromosome.
- **8.9** Perform two experiments: (1) determine whether the process is sensitive to DNase, and (2) determine whether cell contact is required for the process to take place. The cell contact requirement can be tested by a U-tube experiment (see Figure 8.9). If the process is sensitive to DNase, it is similar to transformation. If cell contact is required, it is similar to conjugation. If it is neither sensitive to DNase nor requires cell contact, it is similar to transduction.
- **8.11** (a) F' factors are useful for genetic analyses where two copies of a gene must be present in the same cell, for example, in determining dominance relationships. (b) F' factors are formed by abnormal excision of F factors from Hfr chromosomes (see Figure 8.21). (c) By the conjugative transfer of an F' factor from a donor cell to a recipient (F⁻) cell.
- **8.13** IS elements (or insertion sequences) are short (800–1400 nucleotide pairs) DNA sequences that are transposable—that is, capable of moving from one position in a chromosome to another position or from one chromosome to another chromosome. IS elements mediate recombination between nonhomologous DNA molecules—for example, between F factors and bacterial chromosomes.
- **8.15** Cotransduction refers to the simultaneous transduction of two different genetic markers to a single recipient cell. Since bacteriophage particles can package only 1/100 to 1/50 of the total bacterial chromosome, only markers that are relatively closely linked can be cotransduced. The frequency of cotransduction of any two markers will be an inverse function of the distance between them on the chromosome. As such, this frequency can be used as an estimate of the linkage distance. Specific cotransduction-linkage functions must be prepared for each phage–host system studied.









CHAPTER 9

- 9.1 (a) Griffith's *in vivo* experiments demonstrated the occurrence of transformation in pneumococcus. They provided no indication as to the molecular basis of the transformation phenomenon. Avery and colleagues carried out *in vitro* experiments, employing biochemical analyses to demonstrate that transformation was mediated by DNA. (b) Griffith showed that a transforming substance existed; Avery et al. defined it as DNA. (c) Griffith's experiments did not include any attempt to characterize the substance responsible for transformation. Avery et al. isolated DNA in "pure" form and demonstrated that it could mediate transformation.
- **9.3** Purified DNA from Type III cells was shown to be sufficient to transform Type II cells. This occurred in the absence of any dead Type III cells.
- **9.5** (a) The objective was to determine whether the genetic material was DNA or protein. (b) By labeling phosphorus, a constituent of DNA, and sulfur, a constituent of protein, in a virus, it was possible to demonstrate that only the labeled phosphorus was introduced into the host cell during the viral reproductive cycle. The DNA was enough to produce new phages. (c) Therefore DNA, not protein, is the genetic material.
- **9.7** (a) The ladderlike pattern was known from X-ray diffraction studies. Chemical analyses had shown that a 1:1 relationship existed between the organic bases adenine and thymine and between cytosine and guanine. Physical data concerning the length of each spiral and the stacking of bases were also available. (b) Watson and Crick developed the model of a double helix, with the rigid strands of sugar and phosphorus forming spirals around an axis, and hydrogen bonds connecting the complementary bases in nucleotide pairs.
- **9.9** (a) 400,000; (b) 20,000; (c) 400,000; (d) 68,000 nm.
- **9.11** No. TMV RNA is single-stranded. Thus the base-pair stoichiometry of DNA does not apply.

8.19 pro—pur—bis.

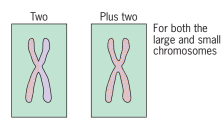
9.13 3'-CAGTACTG-5'.

- **9.15** (a) double-stranded DNA; (b) single-stranded DNA; (c) single-stranded RNA.
- **9.17** The value of $T_{\rm m}$ increases with the GC content because GC base pairs, connected by three hydrogen bonds, are stronger than AT base pairs connected by two hydrogen bonds.
- **9.19** (1) The nucleosome level; the core containing an octamer of histones plus 146 nucleotide pairs of DNA arranged as $1\frac{3}{4}$ turns of a supercoil (see Figure 9.18), yielding an approximately 11-nm diameter spherical body; or juxtaposed, a roughly 11-nm diameter fiber. (2) The 30-nm fiber observed in condensed mitotic and meiotic chromosomes; it appears to be formed by coiling or folding the 11-nm nucleosome fiber. (3) The highly condensed mitotic and meiotic chromosomes); the tight folding or coiling maintained by a "scaffold" composed of nonhistone chromosomal proteins (see Figure 9.22).
- **9.21** (a) 89.5° C. (b) about 39 percent.
- **9.23** The satellite DNA fragments would renature much more rapidly than the main-band DNA fragments. In *D. virilus* satellite DNAs, all three have repeating heptanucleotide-pair sequences. Thus essentially every 40 nucleotide-long (average) single-stranded fragment from one strand will have a sequence complementary (in part) with every single-stranded fragment from the complementary strand. Many of the nucleotide-pair sequences in main-band DNA will be unique sequences (present only once in the genome).
- **9.25** Interphase. Chromosomes are for the most part metabolically inactive (exhibiting little transcription) during the various stages of condensation in mitosis and meiosis.
- 9.27 (a) Histories have been highly conserved throughout the evolution of eukaryotes. A major function of histones is to package DNA into nucleosomes and chromatin fibers. Since DNA is composed of the same four nucleotides and has the same basic structure in all eukaryotes, one might expect that the proteins that play a structural role in packaging this DNA would be similarly conserved. (b) The nonhistone chromosomal proteins exhibit the greater heterogeneity in chromatin from different tissues and cell types of an organism. The histone composition is largely the same in all cell types within a given species-consistent with the role of histones in packaging DNA into nucleosomes. The nonhistone chromosomal proteins include proteins that regulate gene expression. Because different sets of genes are transcribed in different cell types, one would expect heterogeneity in some of the nonhistone chromosomal proteins of different tissues.

CHAPTER 10

- 10.1 (a) Both $3' \rightarrow 5'$ and $5' \rightarrow 3'$ exonuclease activities. (b) The $3' \rightarrow 5'$ exonuclease "proofreads" the nascent DNA strand during its synthesis. If a mismatched base pair occurs at the 3'-OH end of the primer, the $3' \rightarrow 5'$ exonuclease removes the incorrect terminal nucleotide before polymerization proceeds again. The 5' \rightarrow 3' exonuclease is responsible for the removal of RNA primers during DNA replication and functions in pathways involved in the repair of damaged DNA (see Chapter 13). (c) Yes, both exonuclease activities appear to be very important. Without the $3' \rightarrow 5'$ proofreading activity during replication, an intolerable mutation frequency would occur. The $5' \rightarrow 3'$ exonuclease activity is essential to the survival of the cell. Conditional mutations that alter the 5' \rightarrow 3' exonuclease activity of DNA polymerase I are lethal to the cell under conditions where the exonuclease is nonfunctional.
- 10.3 ¹⁵Nitrogen contains eight neutrons instead of the seven neutrons in the normal isotope of nitrogen, ¹⁴N. Therefore, ¹⁵N has an atomic mass of about 15, whereas ¹⁴N has a mass of about 14. This difference means that purines and pyrimidines containing ¹⁵N have a greater density (weight per unit volume) than those containing ¹⁴N. Equilibrium density-gradient centrifugation in 6M CsCl separates DNAs or other macromolecules based on their densities, and *E. coli* DNA, for example, that contains ¹⁵N has a density of 1.724 g/cm², whereas *E. coli* DNA that contains ¹⁴N has a density of 1.710 g/cm².
- **10.5** If nascent DNA is labeled by exposure to ³H-thymidine for very short periods of time, continuous replication predicts that the label would be incorporated into chromosome-sized DNA molecules, whereas discontinuous replication predicts that the label would first appear in small pieces of nascent DNA (prior to covalent joining, catalyzed by DNA ligase).

10.7



10.9 That DNA replication was unidirectional rather than bidirectional. As the intracellular pools of radioactive ³H-thymidine are gradually diluted after transfer to nonradioactive medium, less and less ³H-thymidine will be incorporated into DNA at each replicating fork. This will produce autoradiograms with tails of decreasing grain density at each growing point. Since such tails appear at only one end of each track, replication must be unidirectional. Bidirectional replication would

produce such tails at both ends of an autoradiographic track (see Figure 10.31).

- **10.11** Current evidence suggests that polymerases α , δ , and/or ε are required for the replication of nuclear DNA. Polymerase δ and/or ε are thought to catalyze the continuous synthesis of the leading strand, and polymerase α is believed to function as a primase in the discontinuous synthesis of the lagging strand. Polymerase γ catalyzes replication of organellar chromosomes. Polymerases β , ζ , η , θ , ι , κ , λ , μ , σ , ϕ , and Rev1 function in various DNA repair pathways (see Chapter 13).
- 10.13 No DNA will band at the "light" position; 12.5 percent (2 of 16 DNA molecules) will band at the "hybrid" density; and 87.5 percent (14 of 16 DNA molecules) will band at the "heavy" position.
- 10.15 (a) DNA gyrase; (b) primase; (c) the $5' \rightarrow 3'$ exonuclease activity of DNA polymerase I; (d) the $5' \rightarrow 3'$ polymerase activity of DNA polymerase III; (e) the $3' \rightarrow 5'$ exonuclease activity of DNA polymerase III.
- **10.17** In eukaryotes, the rate of DNA synthesis at each replication fork is about 2500 to 3000 nucleotide pairs per minute. Large eukaryotic chromosomes often contain 10^7 to 10^8 nucleotide pairs. A single replication fork could not replicate the giant DNA in one of these large chromosomes fast enough to permit the observed cell generation times.
- **10.19** No. *E. coli* strains carrying *polA* mutations that eliminate the $3' \rightarrow 5'$ exonuclease activity of DNA polymerase I will exhibit unusually high mutation rates.
- **10.21** (a) Rolling-circle replication begins when an endonuclease cleaves one strand of a circular DNA double helix. This cleavage produces a free 3'-OH on one end of the cut strand, allowing it to function as a primer. (b) The discontinuous synthesis of the lagging strand requires the *de novo* initiation of each Okazaki fragment, which requires DNA primase activity.
- **10.23** DNA helicase unwinds the DNA double helix, and single-strand DNA-binding protein coats the unwound strands, keeping them in an extended state. DNA gyrase catalyzes the formation of negative supercoiling in *E. coli* DNA, and this negative supercoiling behind the replication forks is thought to drive the unwinding process because superhelical tension is reduced by unwinding the complementary strands.
- **10.25** DnaA protein initiates the formation of the replication bubble by binding to the 9-bp repeats of *OriC*. DnaA protein is known to be required for the initiation process because bacteria with temperature-sensitive mutations in the *dnaA* gene cannot initiate DNA replication at restrictive temperatures.
- **10.27** Nucleosomes and replisomes are both large macromolecular structures, and the packaging of eukaryotic DNA into nucleosomes raises the question of how a replisome can move past a nucleosome and replicate

the DNA in the nucleosome in the process. The most obvious solution to this problem would be to completely or partially disassemble the nucleosome to allow the replisome to pass. The nucleosome would then reassemble after the replisome had passed. One popular model has the nucleosome partially disassembling, allowing the replisome to move past it (see Figure 10.33*b*).

- 10.29 (1) DNA replication usually occurs continuously in rapidly growing prokaryotic cells but is restricted to the S phase of the cell cycle in eukaryotes. (2) Most eukaryotic chromosomes contain multiple origins of replication, whereas most prokaryotic chromosomes contain a single origin of replication. (3) Prokaryotes utilize two catalytic complexes that contain the same DNA polymerase to replicate the leading and lagging strands, whereas eukaryotes utilize two or three distinct DNA polymerases for leading and lagging strand synthesis. (4) Replication of eukaryotic chromosomes requires the partial disassembly and reassembly of nucleosomes as replisomes move along parental DNA molecules. In prokaryotes, replication probably involves a similar partial disassembly/reassembly of nucleosome-like structures. (5) Most prokaryotic chromosomes are circular and thus have no ends. Most eukaryotic chromosomes are linear and have unique termini called telomeres that are added to replicating DNA molecules by a unique, RNA-containing enzyme called telomerase.
- **10.31** The chromosomes of haploid yeast cells that carry the *est1* mutation become shorter during each cell division. Eventually, chromosome instability results from the complete loss of telomeres, and cell death occurs because of the deletion of essential genes near the ends of chromosomes.

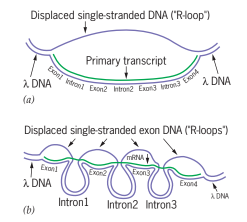
CHAPTER 11

11.1 (a) RNA contains the sugar ribose, which has an hydroxyl (OH) group on the 2-carbon; DNA contains the sugar 2-deoxyribose, with only hydrogens on the 2-carbon. RNA usually contains the base uracil at positions where thymine is present in DNA. However, some DNAs contain uracil, and some RNAs contain thymine. DNA exists most frequently as a double helix (double-stranded molecule); RNA exists more frequently as a single-stranded molecule; but some DNAs are single-stranded and some RNAs are double-stranded. (b) The main function of DNA is to store genetic information and to transmit that information from cell to cell and from generation to generation. RNA stores and transmits genetic information in some viruses that contain no DNA. In cells with both DNA and RNA: (1) mRNA acts as an intermediary in protein synthesis, carrying the information from DNA in the chromosomes to the ribosomes (sites at which proteins are synthesized). (2) tRNAs carry amino acids to the

ribosomes and function in codon recognition during the synthesis of polypeptides. (3) rRNA molecules are essential components of the ribosomes. (4) snRNAs are important components of spliceosomes, and (5) miRNAs play key roles in regulating gene expression (see Chapter 19). (c) DNA is located primarily in the chromosomes (with some in cytoplasmic organelles, such as mitochondria and chloroplasts), whereas RNA is located throughout cells.

- 11.3 3'—GACTA—5'.
- **11.5** Protein synthesis occurs on ribosomes. In eukaryotes, most of the ribosomes are located in the cytoplasm and are attached to the extensive membranous network of endoplasmic reticulum. Some protein synthesis also occurs in cytoplasmic organelles such as chloroplasts and mitochondria.
- 11.7 Both prokaryotic and eukaryotic organisms contain messenger RNAs, transfer RNAs, and ribosomal RNAs. In addition, eukaryotes contain small nuclear RNAs and micro RNAs. Messenger RNA molecules carry genetic information from the chromosomes (where the information is stored) to the ribosomes in the cytoplasm (where the information is expressed during protein synthesis). The linear sequence of triplet codons in an mRNA molecule specifies the linear sequence of amino acids in the polypeptides produced during translation of that mRNA. Transfer RNA molecules are small (about 80 nucleotides long) molecules that carry amino acids to the ribosomes and provide the codon-recognition specificity during translation. Ribosomal RNA molecules provide part of the structure and function of ribosomes; they represent an important part of the machinery required for the synthesis of polypeptides. Small nuclear RNAs are structural components of spliceosomes, which excise noncoding intron sequences from nuclear gene transcripts. Micro RNAs are involved in the regulation of gene expression.
- **11.9** "Self-splicing" of RNA precursors demonstrates that RNA molecules can also contain catalytic sites; this property is not restricted to proteins.
- **11.11** The introns of protein-encoding nuclear genes of higher eukaryotes almost invariably begin (5') with GT and end (3') with AG. In addition, the 3' subterminal A in the "TACTAAC box" is completely conserved; this A is involved in bond formation during intron excision.
- 11.13 (a) Sequence 5. It contains the conserved intron sequences:
 a 5' GU, a 3' AG, and a UACUAAC internal sequence providing a potential bonding site for intron excision. Sequence 4 has a 5' GU and a 3' AG, but contains no internal A for the bonding site during intron excision. (b) 5'—UAGUCUCAA—3'; the putative intron from the 5' GU through the 3' AG has been removed.

11.15



- 11.17 Assuming that there is a -35 sequence upstream from the consensus -10 sequence in this segment of the DNA molecule, the nucleotide sequence of the transcript will be 5'-ACCCGACAUAGCUACGAUGACGAUAAGC-GACAUAGC-3'.
- **11.19** Assuming that there is a CAAT box located upstream from the TATA box shown in this segment of DNA, the nucleotide sequence of the transcript will be 5'-ACCC-GACAUAGCUACGAUGACGAUA-3'.
- **11.21** According to the central dogma, genetic information is stored in DNA and is transferred from DNA to RNA to protein during gene expression. RNA tumor viruses store their genetic information in RNA, and that information is copied into DNA by the enzyme reverse transcriptase after a virus infects a host cell. Thus the discovery of RNA tumor viruses or retroviruses—retro for backwards flow of genetic information—provided an exception to the central dogma.
- **11.23** DNA, RNA, and protein synthesis all involve the synthesis of long chains of repeating subunits. All three processes can be divided into three stages: chain initiation, chain elongation, and chain termination.
- **11.25** The primary transcripts of eukaryotes undergo more extensive posttranscriptional processing than those of prokaryotes. Thus the largest differences between mRNAs and primary transcripts occur in eukaryotes. Transcript processing is usually restricted to the excision of terminal sequences in prokaryotes. In contrast, eukaryotic transcripts are usually modified by (1) the excision of intron sequences; (2) the addition of 7-methyl guanosine caps to the 5' termini; (3) the addition of poly(A) tails to the 3' termini. In addition, the sequences of some eukaryotic transcripts are modified by RNA editing processes.
- 11.27 In eukaryotes, the genetic information is stored in DNA in the nucleus, whereas proteins are synthesized on ribosomes in the cytoplasm. How could the genes, which are separated from the sites of protein synthesis by a double membrane—the nuclear envelope—direct the synthesis

of polypeptides without some kind of intermediary to carry the specifications for the polypeptides from the nucleus to the cytoplasm? Researchers first used labeled RNA and protein precursors and autoradiography to demonstrate that RNA synthesis and protein synthesis occurred in the nucleus and the cytoplasm, respectively.

- 11.29 A simple pulse- and pulse/chase-labeling experiment will demonstrate that RNA is synthesized in the nucleus and is subsequently transported to the cytoplasm. This experiment has two parts: (1) Pulse-label eukaryotic culture cells by growing them in ³H-uridine for a few minutes, and localize the incorporated radioactivity by autoradiography. (2) Repeat the experiment, but this time add a large excess of nonradioactive uridine to the medium in which the cells are growing after the labeling period, and allow the cells to grow in the nonradioactive medium for about an hour. Then localize the incorporated radioactivity will be located in the nucleus when the culture cells are pulse-labeled with ³H-uridine and in the cytoplasm on ribosomes in the pulse-chase experiment.
- **11.31** The first preparation of RNA polymerase is probably lacking the sigma subunit and, as a result, initiates the synthesis of RNA chains at random sites along both strands of the *argH* DNA. The second preparation probably contains the sigma subunit and initiates RNA chains only at the site used *in vivo*, which is governed by the position of the -10 and -35 sequences of the promoter.
- 11.33 TATA and CAAT boxes. The TATA and CAAT boxes are usually centered at positions -30 and -80, respectively, relative to the startpoint (+1) of transcription. The TATA box is responsible for positioning the transcription startpoint; it is the binding site for the first basal transcription factor that interacts with the promoter. The CAAT box enhances the efficiency of transcriptional initiation.
- **11.35** RNA editing sometimes leads to the synthesis of two or more distinct polypeptides from a single mRNA.
- **11.37** This zygote will probably be nonviable because the gene product is essential and the elimination of the 5' splice site will almost certainly result in the production of a nonfunctional gene product.

CHAPTER 12

12.1 Proteins are long chainlike molecules made up of amino acids linked together by peptide bonds. Proteins are composed of carbon, hydrogen, nitrogen, oxygen, and usually sulfur. They provide the enzymatic capacity and much of the structure of living organisms. DNA is composed of phosphate, the pentose sugar 2-deoxyribose, and four nitrogen-containing organic bases (adenine, cytosine, guanine, and thymine). DNA stores and transmits the genetic information in most living organisms. Protein synthesis is of particular interest to geneticists because proteins are the primary gene products—the key

intermediates through which genes control the phenotypes of living organisms.

- 12.3 It depends on how you define alleles. If every variation in nucleotide sequence is considered to be a different allele, even if the gene product and the phenotype of the organism carrying the mutation are unchanged, then the number of alleles will be directly related to gene size. However, if the nucleotide sequence change must produce an altered gene product or phenotype before it is considered a distinct allele, then there will be a positive correlation, but not a direct relationship, between the number of alleles of a gene and its size in nucleotide pairs. The relationship is more likely to occur in prokaryotes where most genes lack introns. In eukaryotic genes, nucleotide sequence changes within introns usually are neutral; that is, they do not affect the activity of the gene product or the phenotype of the organism. Thus, in the case of eukaryotic genes with introns, there may be no correlation between gene size and number of alleles producing altered phenotypes.
- 12.5 (a) Singlet and doublet codes provide a maximum of 4 and (4)² or 16 codons, respectively. Thus neither code would be able to specify all 20 amino acids. (b) 20. (c) (20)¹⁴⁶.
- **12.7** (a) The genetic code is degenerate in that all but 2 of the 20 amino acids are specified by two or more codons. Some amino acids are specified by six different codons. The degeneracy occurs largely at the third or 3' base of the codons. "Partial degeneracy" occurs where the third base of the codon may be either of the two purines or either of the two pyrimidines and the codon still specifies the same amino acid. "Complete degeneracy" occurs where the third base of the codon may be any one of the four bases and the codon still specifies the same amino acid. (b) The code is ordered in the sense that related codons (codons that differ by a single base change) specify chemically similar amino acids. For example, the codons CUU, AUU, and GUU specify the structurally related amino acids, leucine, isoleucine, and valine, respectively. (c) The code appears to be almost completely universal. Known exceptions to universality include strains carrying suppressor mutations that alter the reading of certain codons (with low efficiencies in most cases) and the use of UGA as a tryptophan codon in yeast and human mitochondria.
- **12.9** His \rightarrow Arg results from a transition; His \rightarrow Pro would require a transversion (not induced by 5-bromouracil).
- **12.11** Ribosomes are from 10 to 20 nm in diameter. They are located primarily in the cytoplasm of cells. In bacteria, they are largely free in the cytoplasm. In eukaryotes, many of the ribosomes are attached to the endoplasmic reticulum. Ribosomes are complex structures composed of over 50 different polypeptides and three to five different RNA molecules.

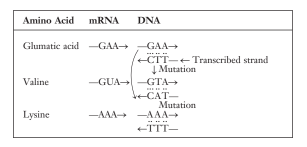
- **12.13** Messenger RNA molecules carry genetic information from the chromosomes (where the information is stored) to the ribosomes in the cytoplasm (where the information is expressed during protein synthesis). The linear sequence of triplet codons in an mRNA molecule specifies the linear sequence of amino acids in the polypeptide(s) produced during translation of that mRNA. Transfer RNA molecules are small (about 80 nucleotides long) molecules that carry amino acids to the ribosomes and provide the codon-recognition specificity during translation. Ribosomal RNA molecules provide part of the structure and function of ribosomes; they represent an important part of the machinery required for the synthesis of polypeptides.
- **12.15** A specific aminoacyl-tRNA synthetase catalyzes the formation of an amino acid-AMP complex from the appropriate amino acid and ATP (with the release of pyrophosphate). The same enzyme then catalyzes the formation of the aminoacyl-tRNA complex, with the release of AMP. The amino acid-AMP and aminoacyl-tRNA linkages are both high-energy phosphate bonds.
- **12.17** Crick's wobble hypothesis explains how the anticodon of a given tRNA can base-pair with two or three different mRNA codons. Crick proposed that the base-pairing between the 5' base of the anticodon in tRNA and the 3' base of the codon in mRNA was less stringent than normal and thus allowed some "wobble" at this site. As a result, a single tRNA often recognizes two or three of the related codons specifying a given amino acid (see Table 12.2).
- 12.19 (a) Inosine. (b) Two.
- 12.21 Translation occurs by very similar mechanisms in prokaryotes and eukaryotes; however, there are some differences. (1) In prokaryotes, the initiation of translation involves base-pairing between a conserved sequence (AGGAGG)-the Shine-Dalgarno box-in mRNA and a complementary sequence near the 3' end of the 16S rRNA. In eukaryotes, the initiation complex forms at the 5' end of the transcript when a cap-binding protein interacts with the 7-methyl guanosine on the mRNA. The complex then scans the mRNA processively and initiates translation (with a few exceptions) at the AUG closest to the 5' terminus. (2) In prokaryotes, the amino group of the initiator methionyl-tRNA^{Met} is formylated; in eukaryotes, the amino group of methionyl-tRNA_i^{Met} is not formylated. (3) In prokaryotes, two soluble protein release factors (RFs) are required for chain termination. RF-1 terminates polypeptides in response to UAA and UAG condons; RF-2 terminates chains in response to UAA and UGA codons. In eukaryotes, one release factor responds to all three termination codons.
- **12.23** Assuming 0.34 nm per nucleotide pair in B-DNA, a gene 68 nm long would contain 200 nucleotide pairs. Given the triplet code, this gene would contain 200/3 = 66.7 triplets, one of which must specify chain termination. Disregarding the partial triplet, this gene could encode a maximum of 65 amino acids.

- **12.25** 426 nucleotides— $3 \times 141 = 423$ specifying amino acids plus three (one codon) specifying chain termination.
- **12.27** (a) Related codons often specify the same or very similar amino acids. As a result, single base-pair substitutions frequently result in the synthesis of identical proteins (degeneracy) or proteins with amino acid substitutions involving very similar amino acids. (b) Leucine and valine have very similar structures and chemical properties; both have nonpolar side groups and fold into essentially the same three-dimensional structures when present in polypeptides. Thus, substitutions of leucine for valine or valine for leucine seldom alter the function of a protein.
- 12.29 (a) Ribosomes and spliceosomes both play essential roles in gene expression, and both are complex macromolecular structures composed of RNA and protein molecules. (b) Ribosomes are located in the cytoplasm; spliceosomes in the nucleus. Ribosomes are larger and more complex than spliceosomes.
- 12.31 Met-Ser-Ile-Cys-Leu-Phe-Gln-Ser-Leu-Ala-Ala-Gln-Asp-Arg-Pro-Gly.
- **12.33** (UAG). This is the only nonsense codon that is related to tryptophan, serine, tyrosine, leucine, glutamic acid, glutamine, and lysine codons by a single base-pair substitution in each case.

- **13.1** (a) Transition, (b) transition, (c) transversion, (d) transversion, (e) frameshift, (f) transition.
- **13.3** (a) *ClB* method, (b) attached-X method (see Chapter 6).
- **13.5** Probably not. A human is larger than a bacterium, with more cells and a longer lifespan. If mutation frequencies are calculated in terms of cell generations, the rates for human cells and bacterial cells are similar.
- **13.7** The X-linked gene is carried by mothers, and the disease is expressed in half of their sons. Such a disease is difficult to follow in pedigree studies because of the recessive nature of the gene, the tendency for the expression to skip generations in a family line, and the loss of the males who carry the gene. One explanation for the sporadic occurrence and tendency for the gene to persist is that, by mutation, new defective genes are constantly being added to the load already present in the population.
- **13.9** The sheep with short legs could be mated to unrelated animals with long legs. If the trait is expressed in the first generation, it could be presumed to be inherited and to depend on a dominant gene. On the other hand, if it does not appear in the first generation, F_1 sheep could be crossed back to the short-legged parent. If the trait is expressed in one-half of the backcross progeny, it is probably inherited as a simple recessive. If two short-legged sheep of different sex could be obtained, they could be mated repeatedly to test the hypothesis of

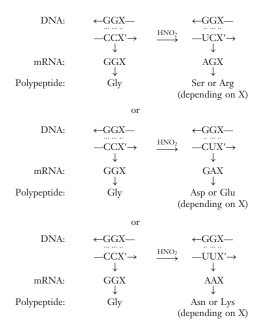
- dominance. In the event that the trait is not transmitted to the progeny that result from these matings, it might be considered to be environmental or dependent on some complex genetic mechanism that could not be identified by the simple test used in the experiments.
- **13.11** If both mutators and antimutators operate in the same living system, an optimum mutation rate for a particular organism in a given environment may result from natural selection.
- **13.13** (a) Yes. (b) A block would result in the accumulation of phenylalanine and a decrease in the amount of tyrosine, which would be expected to result in several different phenotypic expressions.

13.15



- 13.17 Mutations: transitions, transversions, and frameshifts.
- 13.19 3%; 4%; 6%.
- **13.21** Radioactive iodine is concentrated by living organisms and food chains.
- 13.23 $(x^+ m^+ z) (x^+ m^+ z^+) (x m^+ z) (x m z^+)$ or equivalent.
- 13.25 Transitions.
- 13.27 Nitrous acid acts as a mutagen on either replicating or nonreplicating DNA and produces transitions from A to G or C to T, whereas 5-bromouracil does not affect nonreplicating DNA but acts during the replication process causing GC ↔ AT transitions. 5-Bromouracil must be incorporated into DNA during the replication process in order to induce mispairing of bases and thus mutations.
- 13.29 5-BU causes GC ↔ AT transitions. 5-BU can, therefore, revert almost all of the mutations that it induces by enhancing the transition event that is the reverse of the one that produced the mutation. In contrast, the spontaneous mutations will include transversions, frameshifts, deletions, and other types of mutations, including transitions. Only the spontaneous transitions will show enhanced reversion after treatment with 5-BU.
- 13.31 (a) Frameshift due to the insertion of C at the 9th, 10th, or 11th nucleotide from the 5' end. (b) Normal: 5'-AUGCCGUACUGCCAGCUAACUGCU-AAAGAACAAUUA-3'. Mutant: 5'-AUGCCCGUA-CUGCCAGCUAACUGCUAAAGAACAAUUA-3'. (c) Normal: NH₂-Met-Pro-Tyr-Cys-Gln-Leu-Thr-Ala-Lys-Glu-Gln-Leu. Mutant: NH₂-Met-Pro-Val-Leu-Pro-Ala-Asn-Cys.

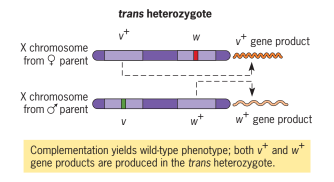
- 13.33 No. Leucine → proline would occur more frequently. Leu (CUA) -5-BU→ Pro (CCA) occurs by a single base-pair transition, whereas Leu (CUA) -5-BU→ Ser (UCA) requires two base-pair transitions. Recall that 5-bromouracil (5-BU) induces only transitions (see Figure 13.7).
- 13.35 Yes:



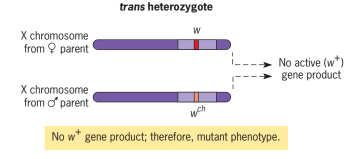
Note: The X at the third position in each codon in mRNA and in each triplet of base pairs in DNA refers to the fact that there is complete degeneracy at the third base in the glycine codon. Any base may be present in the codon, and it will still specify glycine.

- **13.37** Tyr \rightarrow Cys substitutions; Tyr to Cys requires a transition, which is induced by nitrous acid. Tyr to Ser would require a transversion, and nitrous acid is not expected to induce transversions.
- **13.39** 5'-UGG-UGG-UGG-AUG-CGA or AGA-GAA or GAG-UGG-AUG-3'.
- **13.41** Two genes; mutations 1, 2, 3, 4, 5, 6, and 8 are in one gene; mutation 7 is in a second gene.
- **13.43** The complementation test for allelism involves placing mutations pairwise in a common protoplasm in the *trans* configuration and determining whether the resulting *trans* heterozygotes have wild-type or mutant phenotypes. If the two mutations are in different genes, the two mutations will complement each other, because the wild-type copies of each gene will produce functional gene products (see Figure 13.23*a*). However, if the two mutations are in the same gene, both copies of the gene in the *trans* heterozygote will produce defective gene products, resulting in a mutant phenotype (see Figure 13.23*b*). When complementation occurs, the *trans* heterozygote will have the wild-type phenotype. Thus, the complementation test allows one to determine

whether any two recessive mutations are located in the same gene or in different genes. Because the mutations of interest are sex-linked, all the male progeny will have the same phenotype as the female parent. They are hemizygous, with one X chromosome obtained from their mother. In contrast, the female progeny are *trans* heterozygotes. In the cross between the white-eyed female and the vermilion-eyed male, the female progeny have red eyes, the wild-type phenotype. Thus, the *white* and *vermilion* mutations are in different genes, as illustrated in the following diagram:



In the cross between a white-eyed female and a white cherry-eyed male, the female progeny have light cherrycolored eyes (a mutant phenotype), not wild-type red eyes as in the first cross. Since the *trans* heterozygote has a mutant phenotype, the two mutations, *white* and *white cherry*, are in the same gene:



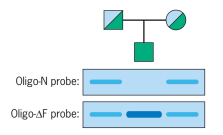
CHAPTER 14

14.1 (a) Both introduce new genetic variability into the cell. In both cases, only one gene or a small segment of DNA representing a small fraction of the total genome is changed or added to the genome. The vast majority of the genes of the organism remain the same. (b) The introduction of recombinant DNA molecules, if they come from a very different species, is more likely to result in a novel, functional gene product in the cell, if the introduced gene (or genes) is capable of being expressed in the foreign protoplasm. The introduction of recombinant DNA molecules is more analogous to duplication mutations (see Chapter 6) than to other types of mutations.

- **14.3** (a) $(1/4)^4 = 1/256$; (b) $(1/4)^6 = 1/4096$.
- 14.5 Recombinant DNA and gene-cloning techniques allow geneticists to isolate essentially any gene or DNA sequence of interest and to characterize it structurally and functionally. Large quantities of a given gene can be obtained in pure form, which permits one to determine its nucleotide-pair sequence (to "sequence it" in common lab jargon). From the nucleotide sequence and our knowledge of the genetic code, geneticists can predict the amino acid sequence of any polypeptide encoded by the gene. By using an appropriate subclone of the gene as a hybridization probe in northern blot analyses, geneticists can identify the tissues in which the gene is expressed. Based on the predicted amino acid sequence of a polypeptide encoded by a gene, geneticists can synthesize oligopeptides and use these to raise antibodies that, in turn, can be used to identify the actual product of the gene and localize it within cells or tissues of the organism. Thus, recombinant DNA and gene-cloning technologies provide very powerful tools with which to study the genetic control of essentially all biological processes. These tools have played major roles in the explosive progress in the field of biology during the last three decades.
- 14.7 Restriction endonucleases are believed to provide a kind of primitive immune system to the microorganisms that produce them—protecting their genetic material from "invasion" by foreign DNAs from viruses or other pathogens or just DNA in the environment that might be taken up by the microorganism. Obviously, these microorganisms do not have a sophisticated immune system like that of higher animals (Chapter 20).
- **14.9** A foreign DNA cloned using an enzyme that produces single-stranded complementary ends can always be excised from the cloning vector by cleavage with the same restriction enzyme that was originally used to clone it. If a *Hin*dIII fragment containing your favorite gene was cloned into *Hin*dIII-cleaved Bluescript vector DNA, it will be flanked in the recombinant Bluescript clone by *Hin*dIII cleavage sites. Therefore, you can excise that *Hin*dIII fragment by digestion of the Bluescript clone with endonuclease *Hin*dIII.
- 14.11 Most genes of higher plants and animals contain noncoding intron sequences. These intron sequences will be present in genomic clones, but not in cDNA clones, because cDNAs are synthesized using mRNA templates and intron sequences are removed during the processing of the primary transcripts to produce mature mRNAs.
- 14.13 The maize *gln2* gene contains many introns, and one of the introns contains a *Hind*III cleavage site. The intron sequences (and thus the *Hind*III cleavage site) are not present in mRNA sequences and thus are also not present in full-length *gln2* cDNA clones.
- 14.15 (a) Southern, northern, and western blot procedures all share one common step, namely, the transfer of

macromolecules (DNAs, RNAs, and proteins, respectively) that have been separated by gel electrophoresis to a solid support—usually a nitrocellulose or nylon membrane—for further analysis. (b) The major difference between these techniques is the class of macromolecules that are separated during the electrophoresis step: DNA for Southern blots, RNA for northern blots, and protein for western blots.

- 14.17 All modern cloning vectors contain a "polycloning site" or "multiple cloning site" (MCS)—a cluster of unique cleavage sites for a number of different restriction endonucleases in a nonessential region of the vector into which the foreign DNA can be inserted. In general, the greater the complexity of the MCS—that is, the more restriction endonuclease cleavage sites that are present—the greater the utility of the vector for cloning a wide variety of different restriction fragments. For example, see the MCS present in plasmid Bluescript II shown in Figure 14.3.
- 14.19 Because the nucleotide-pair sequences of both the normal CF gene and the $CF \Delta 508$ mutant gene are known, labeled oligonucleotides can be synthesized and used as hybridization probes to detect the presence of each allele (normal and $\Delta 508$). Under high-stringency hybridization conditions, each probe will hybridize only with the CF allele that exhibits perfect complementarity to itself. Since the sequences of the CF gene flanking the $\Delta 508$ site are known, oligonucleotide PCR primers can be synthesized and used to amplify this segment of the DNA obtained from small tissue explants of putative CF patients and their relatives by PCR. The amplified DNAs can then be separated by agarose gel electrophoresis, transferred to nylon membranes, and hybridized to the respective labeled oligonucleotide probes, and the presence of each CF allele can be detected by autoradiography. For a demonstration of the utility of this procedure, see Focus on Detection of a Mutant Gene Causing Cystic Fibrosis. In the procedure described there, two synthetic oligonucleotide probesoligo-N = 3'-CTTTTATAGTAGAAACCAC-5' and oligo- $\Delta F = 3'$ -TTCTTTTATAGTA—ACCACAA-5' (the dash indicates the deleted nucleotides in the $CF\Delta 508$ mutant allele) were used to analyze the DNA of CF patients and their parents. For confirmed CF families, the results of these Southern blot hybridizations with the oligo-N (normal) and oligo- ΔF (CF Δ 508) labeled probes were often as follows:



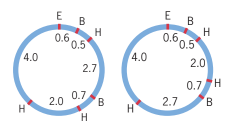
rare recessive trait, and the CF patient was homozygous for the $CF \Delta 508$ allele. In such families, one-fourth of the children would be expected to be homozygous for the $\Delta 508$ mutant allele and exhibit the symptoms of CF, whereas three-fourths would be normal (not have CF). However, two-thirds of these normal children would be expected to be heterozygous and transmit the allele to their children. Only one-fourth of the children of this family would be homozygous for the normal *CF* allele and have no chance of transmitting the mutant *CF* gene to their offspring. Note that the screening procedure described here can be used to determine which of the normal children are carriers of the *CF* $\Delta 508$ allele: that is, the mutant gene can be detected in heterozygotes as well as homozygotes.

14.21 Genetic selection is the most efficient approach to cloning genes of this type. Prepare a genomic library in an expression vector such as Bluescript (see Figure 14.3) using DNA from the kanamycin-resistant strain of *Shigella dysenteriae*. Then, screen the library for the kanamycin-resistance gene by transforming kanamycin-sensitive *E. coli* cells with the clones in the library and plating the transformed cells on medium containing kanamycin. Only cells that are transformed with the kanamycin-resistance gene will produce colonies in the presence of kanamycin.

14.23

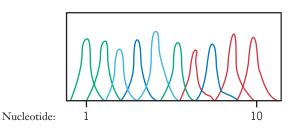


14.25 There are two possible restriction maps for these data as shown below:



Restriction enzyme cleavage sites for *Bam*HI, *Eco*RI, and *Hin*dIII are denoted by B, E, and H, respectively. The numbers give distances in kilobase pairs.

14.27



- **15.1** Genetic map distances are determined by crossover frequencies. Cytogenetic maps are based on chromosome morphology or physical features of chromosomes. Physical maps are based on actual physical distances—the number of nucleotide pairs (0.34 nm per bp)—separating genetic markers. If a gene or other DNA sequence of interest is shown to be located near a mutant gene, a specific band on a chromosome, or a particular DNA restriction fragment, that genetic or physical marker (mutation, band, or restriction fragment) can be used to initiate a chromosome walk to the gene of interest (see Figure 15.7).
- 15.3 A contig (contiguous clones) is a physical map of a chromosome or part of a chromosome prepared from a set of overlapping genomic DNA clones. An RFLP (restriction fragment length polymorphism) is a variation in the length of a specific restriction fragment excised from a chromosome by digestion with one or more restriction endonucleases. A VNTR (variable number tandem repeat) is a short DNA sequence that is present in the genome as tandem repeats and in highly variable copy number. An STS (sequence tagged site) is a unique DNA sequence that has been mapped to a specific site on a chromosome. An EST (expressed sequence tag) is a cDNA sequence-a genomic sequence that is transcribed. Contig maps permit researchers to obtain clones harboring genes of interest directly from DNA Stock Centers-to "clone by phone." RFLPs are used to construct the high-density genetic maps that are needed for positional cloning. VNTRs are especially valuable RFLPs that are used to identify multiple sites in genomes. STSs and ESTs provide molecular probes that can be used to initiate chromosome walks to nearby genes of interest.
- 15.5 (a)

 10 cM
 25 cM
 15 cM
 1 cM
 14 cM

 STS1
 STS5
 STS3
 C
 STS4
 STS2

(b) $3.3 \times 10^{\circ}$ bp/ 3.3×10^{3} cM = 1×10^{6} bp/cM. The total map length is 65 cM, which equates to about 65×10^{6} or 65 million bp.

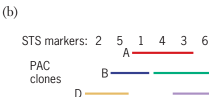
(c) The cancer gene (*C*) and STS4 are separated by 1 cM or about one million base pairs.

- **15.7** With a clone of the gene available, fluorescent *in situ* hybridization (FISH) can be used to determine which human chromosome carries the gene and to localize the gene on the chromosome. Single-stranded copies of the clone are coupled to a fluorescent probe and hybridized to denatured DNA in chromosomes spread on a slide. After hybridization, free probe is removed by washing, and the location of the fluorescent probe is determined by photography using a fluorescence microscope (see Appendix C: *In Situ* Hybridization).
- **15.9** Variable number tandem repeats (VNTRs) are composed of repeated sequences 10 to 80 nucleotide pairs

long, and short tandem repeats (STRs) are composed of repeated sequences 2 to 10 nucleotide pairs long.

- **15.11** The resolution of radiation hybrid mapping is higher than that of standard somatic-cell hybrid mapping because the frequency of recombination is greatly increased in radiation hybrids by using X rays to fragment the human chromosomes prior to cell fusion. The rationale of radiation hybrid mapping is that the probability of breaking a DNA molecule in the region between the two genes and thus separating them is directly proportional to the physical distance (number of base pairs) between them.
- 15.13 The goals of the Human Genome Project were to prepare genetic and physical maps showing the locations of all the genes in the human genome and to determine the nucleotide sequences of all 24 chromosomes in the human genome. These maps and nucleotide sequences of the human chromosomes helped scientists identify mutant genes that result in inherited diseases. Hopefully, the identification of these mutant disease genes will lead to successful treatments, including gene therapies, for at least some of these diseases in the future. Potential misuses of these data include invasions of privacy by governments and businesses-especially employment agencies and insurance companies. Individuals must not be denied educational opportunities, employment, or insurance because of inherited diseases or mutant genes that result in a predisposition to mental or physical abnormalities.
- **15.15** An EST is more likely than an RFLP to occur in a disease-causing human gene. ESTs all correspond to expressed sequences in a genome. RFLPs occur throughout a genome, in both expressed and unexpressed sequences. Because less than 2 percent of the human genome encodes proteins, most RFLPs occur in noncoding DNA.
- **15.17** (a) Segment 5; (b) segment 4; (c) segment 1, 6, or 10.
- **15.19** The major advantage of gene chips as a microarray hybridization tool is that a single gene chip can be used to quantify thousands of distinct nucleotide sequences simultaneously. The gene-chip technology allows researchers to investigate the levels of expression of large numbers of genes more efficiently than was possible using earlier microarray procedures.
- **15.21** The DNA sequences in human chromosome-specific cDNA libraries can be coupled to fluorescent dyes and hybridized *in situ* to the chromosomes of other primates. The hybridization patterns can be used to detect changes in genome structure that have occurred during the evolution of the various species of primates from common ancestors. Such comparisons are especially effective in detecting new linkage relationships resulting from translocations and centric fusions.





15.25 All of the sequences identified by the megablast search encode histone H2a proteins. The query sequence is identical to the coding sequence of the *Drosophila melanogaster* histone H2aV gene (a member of the gene family encoding histone H2a proteins). The query sequence encodes a *Drosophila* histone H2a polypeptide designated variant V. The same databank sequences are identified when one-half or one-fourth of the given nucleotide sequence is used as the query in the megablast search. Query sequences as short as 15 to 20 nucleotides can be used to identify the *Drosophila* gene encoding the histone H2a variant. However, the results will vary depending on the specific nucleotide sequence used as the query sequence.

С

– F

15.27 Reading frame $5' \rightarrow 3'$ number 1 has a large open reading frame with a methionine codon near the 5' end. You can verify that this is the correct reading frame by using the predicted translation product as a query to search one of the protein databases (see Question 15.26).

CHAPTER 16

- **16.1** CpG islands are clusters of cytosines and guanines that are often located just upstream (5') from the coding regions of human genes. Their presence in nucleotide sequences can provide hints as to the location of genes in human chromosomes.
- **16.3** The *CF* gene was identified by map position-based cloning, and the nucleotide sequences of *CF* cDNAs were used to predict the amino acid sequence of the *CF* gene product. A computer search of the protein data banks revealed that the *CF* gene product was similar to several ion channel proteins. This result focused the attention of scientists studying cystic fibrosis on proteins involved in the transport of salts between cells and led to the discovery that the *CF* gene product was a transmembrane conductance regulator—now called the CFTR protein.
- **16.5** Oligonucleotide primers complementary to DNA sequences on both sides (upstream and downstream) of the CAG repeat region in the *MD* gene can be synthesized and used to amplify the repeat region by PCR. One primer must be complementary to an upstream region of the template strand, and the other primer must be complementary to a downstream region of the nontemplate strand. After amplification, the size(s) of the CAG repeat regions can be determined by gel electrophoresis (see Figure 16.2). Trinucleotide repeat lengths can be measured by including repeat regions of known length

on the gel. If less than 30 copies of the trinucleotide repeat are present on each chromosome, the newborn, fetus, or pre-embryo is homozygous for a wild-type *MD* allele or heterozygous for two different wild-type *MD* alleles. If more than 50 copies of the repeat are present on each of the homologous chromosomes, the individual, fetus, or cell is homozygous for a dominant mutant *MD* allele or heterozygous for two different mutant alleles. If one chromosome contains fewer than 30 copies of the CAG repeat and the homologous chromosome contains more than 50 copies, the newborn, fetus, or pre-embryo is heterozygous, carrying one wild-type *MD* allele and one mutant *MD* allele.

- **16.7** The transcription initiation and termination and translation initiation signals or eukaryotes differ from those of prokaryotes such as *E. coli*. Therefore, to produce a human protein in *E. coli*, the coding sequence of the human gene must be joined to appropriate *E. coli* regulatory signals—promoter, transcription terminator, and translation initiator sequences. Moreover, if the gene contains introns, they must be removed or the coding sequence of a cDNA must be used, because *E. coli* does not possess the spliceosomes required for the excision of introns from nuclear gene transcripts. In addition, many eukaryotic proteins undergo posttranslational processing events that are not carried out in prokaryotic cells. Such proteins are more easily produced in transgenic eukaryotic cells growing in culture.
- **16.9** Eleven, ranging in multiples of 3, from 15 to 45 nucleotides long.
- **16.11** DNA profiles are the specific patterns (1) of peaks present in electropherograms of chromosomal STRs or VNTRs amplified by PCR using primers tagged with fluorescent dyes and separated by capillary gel electrophoresis (see Figures 16.11 and 16.12) or (2) of bands on Southern blots of genomic DNAs that have been digested with specific restriction enzymes and hybridized to appropriate STR or VNTR sequences (see Figure 16.10). DNA profiles, like epidermal fingerprints, are used as evidence for identity or nonidentity in forensic cases. Geneticists have expressed concerns about the statistical uses of DNA profile data. In particular, they have questioned some of the methods used to calculate the probability that DNA from someone other than the suspect could have produced an observed profile. These concerns have been based in part on the lack of adequate databases for various human subpopulations and the lack of precise information about the amount of variability in DNA profiles for individuals of different ethnic backgrounds. These concerns have been addressed by the acquisition of data on profile frequencies in different populations and ethnic groups from throughout the world.
- **16.13** Contamination of blood samples would introduce more variability into DNA profiles. This would lead to a lack of allelic matching of profiles obtained from the blood

samples and from the defendant. Mixing errors would be expected to lead to the acquittal of a guilty person and not to the conviction of an innocent person. Only the mislabeling of samples could implicate someone who is innocent.

- **16.15** Probing Southern blots of restriction enzyme-digested DNA of the transgenic plants with ³²P-labeled transgene may provide evidence of multiple insertions, but would not reveal the genomic location of the inserts. Fluorescence *in situ* hybridization (FISH) is a powerful procedure for determining the genomic location of gene inserts. FISH is used to visualize the location of transgenes in chromosomes (see Appendix C).
- **16.17** Transgenic mice are usually produced by microinjecting the genes of interest into pronuclei of fertilized eggs or by infecting pre-implantation embryos with retroviral vectors containing the genes of interest. Transgenic mice provide invaluable tools for studies of gene expression, mammalian development, and the immune system of mammals. Transgenic mice are of major importance in medicine; they provide the model system most closely related to humans. They have been, and undoubtedly will continue to be, of great value in developing the tools and technology that will be used for human gene therapy in the future.
- 16.19 Posttranslationally modified proteins can be produced in transgenic eukaryotic cells growing in culture or in transgenic plants and animals. Indeed, transgenic sheep have been produced that secrete human blood-clotting factor IX and α 1-antitrypsin in their milk. These sheep were produced by fusing the coding sequences of the respective genes to a DNA sequence that encodes the signal peptide required for secretion, and introducing this chimeric gene into fertilized eggs that were then implanted and allowed to develop into transgenic animals. In principle, this approach could be used to produce any protein of interest.
- **16.21** The vector described contains the HGH gene; however, it does not contain a mammalian HGH-promoter that will regulate the expression of the transgene in the appropriate tissues. Construction of vectors containing a properly positioned mammalian HGH-promoter sequence should result in transgenic mice in which HGH synthesis is restricted to the pituitary gland.
- **16.23** RNAi involves the use of double-stranded RNAs, where one strand is complementary to the mRNA and the other strand is equivalent to the mRNA, to silence the expression of target genes. RNAi makes use of the RNA-induced silencing complex (RISC) to block gene expression (see Figure 16.23).
- **16.25** Plants have an advantage over animals in that once insertional mutations are induced they can be stored for long periods of time and distributed to researchers as dormant seeds.
- **16.27** (a) You would first want to check the Salk Institute's Genome Analysis Laboratory web site to see if a T-DNA

or transposon insertion has already been identified in this gene (see Question 16.28). If so, you can simply order seeds of the transgenic line from the Arabidopsis Biological Resource Center at Ohio State University. If no insertion is available in the gene, you can determine where it maps in the genome and use transposons that preferentially jump to nearby sites to identify a new insertional mutation (see http://www.arabidopsis.org/ abrc/ima/jsp). (b) You can construct a gene that has sense and antisense sequences transcribed to a single mRNA molecule (see Figure 16.23b), introduce it into Arabidopsis plants by A. tumefaciens-mediated transformation, and study its effect(s) on the expression of the gene and the phenotype of transgenic plants. The transcript will form a partially base-paired hairpin that will enter the RISC silencing pathway and block the expression of the gene (see Figure 16.23b).

CHAPTER 17

- **17.1** The pair in (d) are inverted repeats and could therefore qualify.
- **17.3** Resistance for the second antibiotic was acquired by conjugative gene transfer between the two types of cells.
- 17.5 In the first strain, the F factor integrated into the chromosome by recombination with the IS element between genes C and D. In the second strain, it integrated by recombination with the IS element between genes Dand E. The two strains transfer their genes in different orders because the two chromosomal IS elements are in opposite orientation.
- 17.7 No. IS1 and IS2 are mobilized by different transposases.
- **17.9** The *tnpA* mutation: no; the *tnpR* mutation: yes.
- 17.11 Many bacterial transposons carry genes for antibiotic resistance, and it is relatively simple for these genes to move from one DNA molecule to another. DNA molecules that acquire resistance genes can be passed to other cells in a bacterial population, both vertically (by descent) and horizontally (by conjugative transfer). Over time, continued exposure to an antibiotic will select for cells that have acquired a gene for resistance to that antibiotic. The antibiotic will therefore no longer be useful in combating these bacteria.
- **17.13** The c^m mutation is due to a *Ds* or an *Ac* insertion.
- **17.15** The paternally inherited *Bz* allele was inactivated by a transposable element insertion.
- **17.17** Cross dysgenic (highly mutable) males carrying a wildtype X chromosome to females homozygous for a balancer X chromosome; then cross the heterozygous F_1 daughters individually to their brothers and screen the F_2 males that lack the balancer chromosome for mutant phenotypes, including failure to survive (lethality). Mutations identified in this screen are probably due to *P* element insertions in X-linked genes.

- **17.19** Factors made by the fly's genome are required for transposition; other insects apparently lack the ability to provide these factors.
- **17.21** Through crossing over between the LTRs of a Ty1 element.
- **17.23** *In situ* hybridization to polytene chromosomes using a *TART* probe (see Appendix C).
- **17.25** *TART* and *HeT-A* replenish the ends of *Drosophila* chromosomes.
- 17.27 The Sleeping Beauty element could be used as a transformation vector in vertebrates much like the P element has been used in Drosophila. The gfp gene could be inserted between the ends of the Sleeping Beauty element and injected into eggs or embryos along with an intact Sleeping Beauty element capable of encoding the element's transposase. If the transposase that is produced in the injected egg or embryo acts on the element that contains the gfp gene, it might cause the latter to be inserted into genomic DNA. Then, if the egg or embryo develops into an adult, that adult can be bred to determine if a Sleeping Beauty/gfp transgene is transmitted to the next generation. In this way, it would be possible to obtain strains of mice or zebra fish that express the gfp gene.

18.1 By studying the synthesis or lack of synthesis of the enzyme in cells grown on chemically defined media. If the enzyme is synthesized only in the presence of a certain metabolite or a particular set of metabolites, it is probably inducible. If it is synthesized in the absence but not in the presence of a particular metabolite or group of metabolites, it is probably repressible.

18.3

Gene or Regulatory Element	Function	
(a) Regulator gene	Codes for repressor	
(b) Operator	Binding site of repressor	
(c) Promoter	Binding site of RNA polymerase and CAP-cAMP complex	
(d) Structural gene Z	Encodes β-galactosidase	
(e) Structural gene Y	Encodes β-galactoside permease	

18.5 (a) 1, 2, 3, and 5; (b) 2, 3, and 5.

18.7 The O^c mutant prevents the repressor from binding to the operator. The I^s mutant repressor cannot bind to O^c . The I^s mutant protein has a defect in the allosteric site that binds allolactose, but has a normal operator binding site.

18.9 (a)
$$\frac{I^+ O^c Z^+ Y^-}{I^+ O^+ Z^- Y^+};$$

(b) $\frac{I^s O^c Z^+ Y^-}{I^s O^+ Z^- Y^+}$

- **18.11** (a) The O^c mutations map very close to the Z structural gene; I^- mutations map slightly farther from the structural gene (but still very close by; see Figure 18.5). (b) An $I^+O^+Z^+Y^+/I^+O^cZ^+Y^+$ partial diploid would exhibit constitutive synthesis of β -galactosidase and β -galactoside permease, whereas an $I^+O^+Z^+Y^+/I^-O^+Z^+Y^+$ partial diploid would be inducible for the synthesis of these enzymes. (c) The O^c mutation is *cis*-dominant; the I^- mutation is *trans*-recessive.
- **18.13** Catabolite repression has evolved to assure the use of glucose as a carbon source when this carbohydrate is available, rather than less efficient energy sources.
- **18.15** Positive regulation; the CAP-cAMP complex has a positive effect on the expression of the *lac* operon. It functions in turning on the transcription of the structural genes in the operon.
- **18.17** Negative regulatory mechanisms such as that involving the repressor in the lactose operon block the transcription of the structural genes of the operon, whereas positive mechanisms such as the CAP-cAMP complex in the *lac* operon promote the transcription of the structural genes of the operon.
- **18.19** Repression/derepression of the *trp* operon occurs at the level of transcription initiation, modulating the frequency at which RNA polymerase initiates transcription from the *trp* operon promoters. Attenuation modulates *trp* transcript levels by altering the frequency of termination of transcription within the *trp* operon leader region (*trpL*).
- 18.21 First, remember that transcription and translation are coupled in prokaryotes. When tryptophan is present in cells, tryptophan-charged tRNA^{Trp} is produced. This allows translation of the *trp* leader sequence through the two UGG Trp codons to the *trp* leader sequence UGA termination codon. This translation of the *trp* leader region prevents base-pairing between the partially complementary mRNA leader sequences 75–83 and 110–121 (see Figure 18.15*b*), which in turn permits formation of the transcription–termination "hairpin" involving leader sequences 110–121 and 126–134 (see Figure 18.15*c*).
- **18.23** Both *trp* attenuation and the lysine riboswitch turn off gene expression by terminating transcription upstream from the coding regions of the regulated genes. Both involve the formation of alternative mRNA secondary structures—switching between the formation of antiterminator and transcription–terminator hairpins—in response to the presence or absence of a specific metabolite (compare Figure 18.15 and Figure 2 in On the Cutting Edge: The Lysine Riboswitch).

- **19.1** In multicellular eukaryotes, the environment of an individual cell is relatively stable. There is no need to respond quickly to changes in the external environment. In addition, the development of a multicellular organism involves complex regulatory hierarchies composed of hundreds of different genes. The expression of these genes is regulated spatially and temporally, often through intricate intercellular signaling processes.
- **19.3** Activity of the *dystrophin* gene could be assessed by blotting RNA extracted from the different types of cells and hybridizing it with a probe from the gene (northern blotting); or the RNA could be reverse-transcribed into cDNA using one or more primers specific to the *dystrophin* gene and the resulting cDNA could be amplified by the polymerase chain reaction (RT-PCR). Another technique would be to hybridize *dystrophin* RNA *in situ*—that is, in the cells themselves—with a probe from the gene. It would also be possible to check each cell type for production of dystrophin protein by using anti-dystrophin antibodies to analyze proteins from the different cell types on western blots, or to analyze the proteins in the cells themselves—that is, *in situ*.
- **19.5** One procedure would be to provide larvae with radioactively labeled UTP, a building block of RNA, under different conditions—with and without heat shock. Then prepare samples of polytene cells from these larvae for autoradiography. If the heat shock-induced puffs contain genes that are vigorously transcribed, the radioactive signal should be abundant in the puffs.
- 19.7 By alternate splicing of the transcript.
- **19.9** Northern blotting of RNA extracted from plants grown with and without light, or PCR amplification of cDNA made by reverse-transcribing these same RNA extracts.
- **19.11** That enhancers can function in either orientation.
- **19.13** Probably not unless the promoter of the *gfp* gene is recognized and transcribed by the *Drosophila* RNA polymerase independently of the heat-shock response elements.
- **19.15** The mutation is likely to be lethal in homozygous condition because the transcription factor controls so many different genes and a frameshift mutation in the coding sequence will almost certainly destroy the transcription factor's function.
- **19.17** Exon 3 contains an in-frame stop codon. Thus, the protein translated from the *Sxl* mRNA in males will be shorter than the protein translated from the shorter *Sxl* mRNA in females.
- **19.19** The intron could be placed in a GUS expression vector, which could then be inserted into *Arabidopsis* plants. If the intron contains an enhancer that drives gene expression in root tips, transgenic plants should show GUS expression in their root tips. See the Problem-Solving Skills feature in Chapter 19 for an example of this type of analysis.

- **19.21** Yes. The diffuse, bloated appearance indicates that the genes on this chromosome are being transcribed vigorously—the chromatin is "open for business."
- **19.23** Short interfering RNAs target messenger RNA molecules, which are devoid of introns. Thus, if siRNA were made from double-stranded RNA derived from an intron, it would be ineffective against an mRNA target.
- **19.25** The paternally contributed allele (*b*) will be expressed in the F_1 progeny.
- **19.27** RNA could be isolated from liver and brain tissue. Northern blotting or RT-PCR with this RNA could then establish which of the genes (*A* or *B*) is transcribed in which tissue. For northern blotting, the RNA samples would be fractionated in a denaturing gel and blotted to a membrane, and then the RNA on the membrane would be hybridized with gene-specific probes, first for one gene, then for the other (or the researcher could prepare two separate blots and hybridize each one with a different probe). For RT-PCR, the RNA samples would be reverse-transcribed into cDNA using primers specific for each gene; then the cDNA molecules would be amplified by standard PCR, and the products of the amplifications would be fractionated by gel electrophoresis to determine which gene's RNA was present in the original samples.
- 19.29 The *msl* gene is not functional in females.
- **19.31** HP1, the protein encoded by the wild-type allele of the suppressor gene, is involved in chromatin organization. Perhaps this heterochromatic protein spreads from the region near the inversion breakpoint in the chromosome that carries the *white mottled* allele and brings about the "heterochromatization" of the *white* locus. When HP1 is depleted by knocking out one copy of the gene encoding it—that is, by putting the suppressor mutation into the fly's genotype, the "heterochromatization" of the *white* locus would be less likely to occur, and perhaps not occur at all. The white locus would then function fully in all eye cells, producing a uniform red eye color.

CHAPTER 20

- **20.1** Unequal division of the cytoplasm during the meiotic divisions; transport of substances into the oocyte from surrounding cells such as the nurse cells in *Drosophila*.
- **20.3** Collect mutations with diagnostic phenotypes; map the mutations and test them for allelism with one another; perform epistasis tests with mutations in different genes; clone individual genes and analyze their function at the molecular level.
- 20.5 Imaginal discs.
- **20.7** In homozygous condition, the mutation that causes phenylketonuria has a maternal effect. Women homozygous for this mutation influence the development of their children *in utero*.

- **20.9** Female sterility. Females affected by the mutations will lay abnormal eggs that will not develop into viable embryos.
- **20.11** The somatic cells surrounding a developing *Drosophila* egg in the ovary determine where the spätzle protein, which is the ligand for the Toll receptor protein, will be cleaved. This cleavage will eventually occur on the ventral side of the developing embryo.
- **20.13** $ey \rightarrow boss \rightarrow sev \rightarrow R7$ differentiation
- **20.15** Because the *Pax6* gene gave the same phenotype in flies as overexpression of the *eyeless* gene, the genes must be functionally homologous, as well as structurally homologous. Therefore, expect extra mouse eyes or eye primorida when expressing *eyeless* in the mouse.
- **20.17** Northern blotting of RNA extracted from the tissues at different times during development. Hybridize the blot with gene-specific probes.
- **20.19** Reproductive cloning of mammals such as sheep, mice, and cats indicates that somatic-cell nuclei have all the genetic information to direct the development of a complete, viable organism. It also shows that epigenetic modifications of chromatin, such as X chromosome inactivation, can be reset.
- **20.21** If each antibody consists of one kind of light chain and one kind of heavy chain, and if light and heavy chains can combine freely, the potential to produce 100 million different antibodies implies the existence of 10,000 light chain genes and 10,000 heavy chain genes (10,000 \times 10,000 = 100 million). If each light chain is 220 amino acids long, each light chain gene must comprise $3 \times 220 =$ 660 nucleotides because each amino acid is specified by a triplet of nucleotides; similarly, each heavy chain gene must comprise $3 \times 450 = 1350$ nucleotides. Therefore, the genome must contain $10,000 \times 660 = 6.6$ million nucleotides devoted to light chain production and $10,000 \times 1,350 = 13.5$ million nucleotides devoted to heavy chain production. Altogether, then, the genome must contain 19.5 million nucleotides dedicated to encoding the amino acids of the various antibody chains.

- **21.1** Cancer has been called a genetic disease because it results from mutations of genes that regulate cell growth and division. Nonhereditary forms of cancer result from mutations in somatic cells. These mutations, however, can be induced by environmental factors including tobacco smoke, chemical pollutants, ionizing radiation, and UV light. Hereditary forms of cancer also frequently involve the occurrence of environmentally induced somatic mutations.
- **21.3** Aneuploidy might involve the loss of functional copies of tumor suppressor genes, or it might involve the inappropriate duplication of proto-oncogenes. Loss of tumor suppressor genes would remove natural brakes on cell division, and duplication of proto-oncogenes would increase the abundance of factors that promote cell division.

- 21.5 They possess introns.
- **21.7** The products of these genes play important roles in cell activities.
- **21.9** The cultured NIH 3T3 cells probably carry other mutations that predispose them to become cancerous; transfection of such cells with a mutant *c*-*H*-*ras* oncogene may be the last step in the process of transforming the cells into cancer cells. Cultured embryonic cells probably do not carry the predisposing mutations needed for them to become cancerous; thus, when they are transfected with the mutant *c*-*H*-*ras* oncogene, they continue to divide normally.
- **21.11** Retinoblastoma results from homozygosity for a loss-of-function (recessive) allele. The sporadic occurrence of retinoblastoma requires two mutations of this gene in the same cell or cell lineage. Therefore retinoblastoma is rare among individuals who, at conception, are homozygous for the wild-type allele of the RB gene. For such individuals, we would expect the frequency of tumors in both eyes to be the square of the frequency of tumors in one eye. Individuals who are heterozygous for a mutant RB allele require only one somatic mutation to occur for them to develop retinoblastoma. Because there are millions of cells in each retina, there is a high probability that this somatic mutation will occur in at least one cell in each eye, causing both eyes to develop tumors.
- **21.13** At the cellular level, loss-of-function mutations in the RB gene are recessive; a cell that is heterozygous for such a mutation divides normally. However, when a second mutation occurs, that cell becomes cancerous. If the first RB mutation was inherited, there is a high probability that the individual carrying this mutation will develop retinoblastoma because a second mutation can occur any time during the formation of the retinas in either eye. Thus, the individual is predisposed to develop retinoblastoma, and it is this predisposition that shows a dominant pattern of inheritance.
- **21.15** By binding to E2F transcription factors, pRB prevents those transcription factors from activating their target genes—which encode proteins involved in progression of the cell cycle; pRB is therefore a negative regulator of transcription factors that stimulate cell division.
- **21.17** Cells homozygous for a loss-of-function mutation in the p16 gene might be expected to divide in an uncontrolled fashion because the p16 protein would not be able to inhibit cyclin-CDK activity during the cell cycle. The p16 gene would therefore be classified as a tumor suppressor gene.
- **21.19** Cells homozygous for a loss-of-function mutation in the *BAX* gene would be unable to prevent repression of the programmed cell death pathway by the BCL-2 gene product. Consequently, these cells would be unable to execute that pathway in response to DNA damage induced by radiation treatment. Such cells would continue to divide and accumulate mutations; ultimately, they would have a good

chance of becoming cancerous. The *BAX* gene would therefore be classified as a tumor suppressor gene.

- **21.21** If a cell were heterozygous for a mutation that caused p53 to bind tightly and constitutively to the DNA of its target genes, its growth and division might be retarded, or it might be induced to undergo apoptosis. Such a cell would be expected to be more sensitive to the effects of ionizing radiation because radiation increases the expression of p53, and in this case, the p53 would be predisposed to activate its target genes, causing the cell to respond vigorously to the radiation treatment.
- 21.23 They would probably decrease the ability of pAPC to bind β -catenin.
- **21.25** The increased irritation to the intestinal epithelium caused by a fiber-poor, fat-rich diet would be expected to increase the need for cell division in this tissue (to replace the cells that were lost because of the irritation), with a corresponding increase in the opportunity for the occurrence of cancer-causing mutations.
- **21.27** No. Apparently there is another pathway—one not mediated by p53—that leads to the activation of the *p21* gene.

CHAPTER 22

- **22.1** Some of the genes implicated in heart disease are listed in Table 22.2. Environmental factors might include diet, amount of exercise, and whether or not the person smokes.
- **22.3** The concordance for monozygotic twins is almost twice as great as that for dizygotic twins. Monozygotic twins share twice as many genes as dizygotic twins. The data strongly suggest that alcoholism has a genetic basis.
- **22.5** Because 8/2012 is approximately $1/256 = (1/4)^4$, it appears that four size-determining genes were segregating in the crosses.
- **22.7** Because $\Sigma(X_i \text{mean}) = 0$.
- **22.9** 3.17/6.08 = 0.52.
- **22.11** V_e is estimated by the average of the variances of the inbreds: 9.4 cm². V_g is estimated by the difference between the variances of the randomly pollinated population and the inbreds: $(26.4 9.4) = 17.0 \text{ cm}^2$. The broad-sense heritability is $H^2 = V_e/V_T = 17.0/26.4 = 0.64$.
- **22.13** Broad-sense heritability must be greater than narrowsense heritability because $H^2 = V_a/V_T > V_a/V_T = b^2$.
- **22.15** (15 12)(0.3) + 12 = 12.9 bristles.
- **22.17** $b^2 = R/S = (12.5 10)/(15 10) = 0.5$; selection for increased growth rate should be effective.
- **22.19** Half-siblings share 25 percent of their genes. The maximum value for b^2 is therefore 0.14/0.25 = 0.56.
- **22.21** The correlations for MZT are not much different from those for MZA. Evidently, for these personality traits, the environmentality (C^2 in Table 22.3) is negligible.

CHAPTER 23

- **23.1** Frequency of L^M in Central American population: $p = (2 \times 53 + 29)/(2 \times 86) = 0.78$; q = 0.22. Frequency of L^M in North American population: $p = (2 \times 78 + 61)/(2 \times 278) = 0.39$; q = 0.61.
- **23.3** $q^2 = 0.0004; q = 0.02.$
- **23.5** Frequency of tasters (genotypes *TT* and *Tt*): $(0.4)^2 + 2(0.4)(0.6) = 0.64$. Frequency of *TT* tasters among all tasters: $(0.4)^2/(0.64) = 0.25$.
- **23.7** $(0.00025)^2 = 6.25 \times 10^{-8}$.
- **23.9** In females, the frequency of the dominant phenotype is 0.36. The frequency of the recessive phenotype is $0.64 = q^2$; thus, q = 0.8 and p = 0.2. The frequency of the dominant phenotype in males is therefore p = 0.2.
- **23.11** Frequency of heterozygotes = H = 2pq = 2p(1 p). Using calculus, take the derivative of H and set the result to zero to solve for the value of p that maximizes H: dH/dp = 2 - 4p = 0 implies that p = 2/4 = 0.5.
- **23.13** Under the assumption that the population is in Hardy–Weinberg equilibrium, the frequency of the allele for light coloration is the square root of the frequency of recessive homozygotes. Thus, $q = \sqrt{0.49} = 0.7$, and the frequency of the allele for dark color is 1 q = p = 0.3. From $p^2 = 0.09$, we estimate that $0.09 \times 100 = 9$ of the dark moths in the sample are homozygous for the dominant allele.
- **23.15** Ultimate frequency of *GG* is 0.2; ultimate frequency of *gg* is 0.8.
- 23.17 (a) Frequency of A in merged population is 0.5, and that of a is also 0.5; (b) 0.25 (AA), 0.50 (Aa), and 0.25 (aa); (c) frequencies in (b) will persist.
- **23.19** The relative fitnesses can be obtained by dividing each of the survival probabilities by the largest probability (0.92). Thus, the relative fitnesses are 1 for *GG*, 0.98 = 1 0.02 for *Gg*, and 0.61 = 1 0.39 for *gg*. The selection coefficients are $s_1 = 0.02$ for *Gg* and $s_2 = 0.39$ for *gg*.
- **23.21** (a) Use the following scheme:

Genotype	CC	Cc	сс
Hardy–Weinberg frequency	$(0.98)^2 = 0.9604$	2(0.98)(0.02) = 0.0392	$(0.02)^2 = 0.0004$
Relative fitness	1	1	0
Relative contribution to next generation	(0.9604) × 1	(0.0392) × 1	0
Proportional contribution	0.9604/0.9996 = 0.9608	$\begin{array}{l} 0.0392/0.9996 \\ = 0.0392 \end{array}$	0

The new frequency of the allele for cystic fibrosis is (0.5)(0.0392) = 0.0196; thus, the incidence of the disease

- **23.23** $q^2 = 4 \times 10^{-5}$; thus $q = 6.3 \times 10^{-3}$ and 2pq = 0.0126.
- **23.25** Probability of ultimate fixation of A_2 is 0.5; probability of ultimate loss of A_3 is 1 0.3 = 0.7.
- **23.27** p = 0.2; at equilibrium, p = t/(s + t). Because s = 1, we can solve for t; t = 0.25.
- **23.29** At mutation-selection equilibrium $q = \sqrt{u/s} = \sqrt{10^{-6}/1} = 0.001$.

- 24.1 Among other things, Darwin observed species on islands that were different from each other and from continental species, but that were still similar enough to indicate that they were related. He also observed variation within species, especially within domesticated breeds, and saw how the characteristics of an organism could be changed by selective breeding. His observations of fossilized organisms indicated that some species have become extinct.
- **24.3** The frequency of the *a* allele is 0.06 in the South African population and 0.42 in the English population. The predicted genotype frequencies under the assumption of random mating are:

Genotype	South Africa	England
aa	$(0.06)^2 = 0.004$	$(0.42)^2 = 0.18$
ab	2(0.06)(0.94) = 0.11	2(0.42)(0.58) = 0.49
bb	$(0.94)^2 = 0.88$	$(0.58)^2 = 0.33$

24.5 In the sample, the frequency of the *F* allele is $(2 \times 32 + 46)/(2 \times 100) = 0.55$ and the frequency of the *S* allele is 1 - 0.55 = 0.45. The predicted and observed genotype frequencies are:

Genotype	Observed	Hardy-Weinberg Predicted
FF	32	$100 \times (0.55)^2 = 30.25$
FS	46	$100 \times 2(0.55)(0.45) = 49.5$
SS	22	$100 \times (0.45)^2 = 20.25$

To test for agreement between the observed and predicted values, we compute a chi-square statistic with 1 degree of freedom: $\chi^2 = \Sigma$ (obs. – pred.)²/pred. = 0.50, which is not significant at the 5 percent level. Thus, the population appears to be in Hardy–Weinberg equilibrium for the alcohol dehydrogenase locus.

- **24.7** In the third position of some of the codons. Due to the degeneracy of the genetic code, different codons can specify the same amino acid. The degeneracy is most pronounced in the third position of many codons, where different nucleotides can be present without changing the amino acid that is specified.
- **24.9** Complex carbohydrates are not "documents of evolutionary history" because, although they are polymers, they are typically made of one subunit incorporated repetitiously into a chain. Such a polymer has little or no "information content." Thus, there is little or no opportunity to distinguish a complex carbohydrate obtained from two different organisms. Moreover, complex carbohydrates are not part of the genetic machinery; their formation is ultimately specified by the action of enzymes, which are gene products, but they themselves are not genetic material or the products of the genetic material.
- **24.11** The histidines are rigorously conserved because they perform an important function—anchoring the heme group in hemoglobin. Because these amino acids are strongly constrained by natural selection, they do not evolve by mutation and random genetic drift.
- **24.13** Estimate the average number of substitutions per site in the ribonuclease molecule as $-\ln(S)$, where S = (124 - 40)/124 = 0.68, the proportion of amino acids that are the same in the rat and cow molecules. The average number of substitutions per site since the cow and rat lineages diverged from a common ancestor is therefore 0.39. The evolutionary rate in the cow and rat lineages is 0.39/(2 × 80 million years) = 2.4 substitutions per site every billion years.
- **24.15** The reciprocal of the rate, that is, 1/*K*.
- **24.17** The protein with the higher evolutionary rate is not as constrained by natural selection as the protein with the lower evolutionary rate.
- **24.19** Repetitive sequences that are near each other can mediate displaced pairing during meiosis. Exchange involving the displaced sequences can duplicate the region between them.
- **24.21** Cross *D. mauritiana* with *D. simulans* and determine if these two species are reproductively isolated. For instance, can they produce offspring? If they can, are the offspring fertile?
- **24.23** The *Kpn-pn* interaction is an example of the kind of negative epistasis that might prevent populations that have evolved separately from merging into one panmictic population. The *Kpn* mutation would have evolved in one population and the *pn* mutation in another, geographically separate population. When the populations merge, the two mutations can be brought into the same fly by interbreeding. If the combination of these mutations is lethal, then the previously separate populations will not be able to exchange genes; that is, they will be reproductively isolated.

Glossary

This glossary provides an introduction to some basic and recurring terms in the text. Names of chemical compounds, definitions of specialized terms, and variants of basic names have been omitted from the glossary but are given in the index. Please locate terms that are not in the glossary by referring to the index.

A

Abscissa. The horizontal scale on a graph.

Acentric chromosome. Chromosome fragment lacking a centromere.

Acquired immune deficiency syndrome. See AIDS.

- Acridine dyes. A class of positively charged polycyclic molecules that intercalate into DNA and induce frameshift mutations.
- **Acrocentric.** A modifying term for a chromosome or chromatid that has its centromere near the end.
- Activator (of gene expression). Regulator gene products that turn on, or activate, the expression of other genes.
- Activator (Ac). A transposable element in maize that encodes a transacting transposase capable of catalyzing the movement of Acelements and other members of the Ac/Ds family.
- Adaptation. Adjustment of an organism or a population to an environment.
- ADA⁻ SCID (adenosine deaminase-deficient severe combined immunodeficiency disease). An autosomal recessive disorder in humans caused by a lack of the enzyme adenosine deaminase, which catalyzes the breakdown of deoxyadenosine. In the absence of this enzyme, toxic derivatives of this nucleoside accumulate and kill cells required for normal immune responses to infections.
- Additive allelic effects. Genetic factors that raise or lower the value of a phenotype on a linear scale of measurement.
- Additive genetic variance. The portion of the total phenotypic variance in a quantitive trait that is due to the additive effects of alleles.
- Adenine (A). A purine base found in RNA and DNA.
- **A-DNA.** A right-handed DNA double helix that has 11 base pairs per turn. DNA exists in this form when partially dehydrated.
- *Agrobacterium tumefaciens*-mediated transformation. A naturally occurring process of DNA transfer from the bacterium *A. tumefaciens* to plants.
- **AIDS (acquired immunodeficiency syndrome).** The usually fatal human disease in which the immune system is destroyed by the human immunodeficiency virus (HIV).
- **Albinism.** Absence of pigment in skin, hair, and eyes of an animal. Absence of chlorophyll in plants.
- Aleurone. The outermost layer of the endosperm in a seed.
- Alkaptonuria. An inherited metabolic disorder. Alkaptonurics excrete excessive amounts of homogentisic acid (alkapton) in the urine.
- **Alkylating agents.** Chemicals that transfer alkyl (methyl, ethyl, and so on) groups to the bases in DNA.

- Allele (allelomorph; *adj.*, allelic, allelomorphic). One of a pair, or series, of alternative forms of a gene that occur at a given locus in a chromosome. Alleles are symbolized with the same basic symbol (for example, *D* for tall peas and *d* for dwarf). (See also **Multiple alleles**.)
- **Allele frequency.** The proportion of one allele relative to all alleles at a locus in a population.
- Allopatric speciation. Speciation occurring at least in part because of geographic isolation.
- **Allopolyploid.** A polyploid having chromosome sets from different species; a polyploid containing genetically different chromosome sets derived from two or more species.
- **Allosteric transition.** A reversible interaction of a small molecule with a protein molecule that causes a change in the shape of the protein and a consequent alteration of the interaction of that protein with a third molecule.
- Allotetraploid. An organism with four genomes derived from hybridization of different species. Usually, in forms that become established, two of the four genomes are from one species and two are from another species.
- Allozyme. A variant of an enzyme detected by electrophoresis.
- **Amino acid.** Any one of a class of organic compounds containing an amino (NH₂) group and a carboxyl (COOH) group. Amino acids are the building blocks of proteins. Alanine, proline, threonine, histidine, lysine, glutamine, phenylalanine, tryptophan, valine, arginine, tyrosine, and leucine are among the common amino acids.
- **Aminoacyl** (*A*) site. The ribosome binding site that contains the incoming aminoacyl-tRNA.
- **Aminoacyl-tRNA synthetases.** Enzymes that catalyze the formation of high energy bonds between amino acids and tRNA molecules.
- **Amniocentesis.** A procedure for obtaining amniotic fluid from a pregnant woman. Chemical contents of the fluid are studied directly for the diagnosis of some diseases. Cells are cultured, and metaphase chromosomes are examined for irregularities (for example, trisomy).
- **Amnion.** The thin membrane that lines the fluid-filled sac in which the embryo develops in higher vertebrates.
- **Amniotic fluid.** Liquid contents of the amniotic sac of higher vertebrates containing cells of the embryo (not of the mother). Both fluid and cells are used for diagnosis of genetic abnormalities of the embryo or fetus.
- **Amorphic.** A term applied to a mutant allele that completely abolishes gene expression. Such a mutant allele is called an amorph.

- **Amphidiploid.** A species or type of plant derived from doubling the chromosomes in the F_1 hybrid of two species; an allopolyploid. In an amphidiploid the two species are known, whereas in other allopolyploids they may not be known.
- **Amplification (recombinant DNA molecules).** The production of many copies of a newly constructed recombinant DNA molecule.
- **Anabolic pathway.** A pathway by which a metabolite is synthesized; a biosynthetic pathway.
- **Anaphase.** The stage of mitosis or meiosis during which the daughter chromosomes pass from the equatorial plate to opposite poles of the cell (toward the ends of the spindle). Anaphase follows metaphase and precedes telophase.
- **Anaphase I.** The stage during the first meiotic division when duplicated homologous chromosomes separate from each other and begin moving to opposite poles of the cell.
- **Anaphase II.** The stage during the second meiotic division when sister chromatids of a duplicated chromosome separate from each other and begin moving to opposite poles of the cell.
- **Anchor gene.** A gene that has been positioned on both the physical map and the genetic map of a chromosome.
- **Androgen.** A male hormone that controls sexual activity in vertebrate animals.
- **Anemia.** Abnormal condition characterized by pallor, weakness, and breathlessness, resulting from a deficiency of hemoglobin or a reduced number of red blood cells.
- **Aneuploid.** An organism or cell having a chromosome number that is not an exact multiple of the monoploid (*n*) with one genome, that is, hyperploid, higher (for example, 2n + 1), or hypoploid, lower (for example, 2n 1). Also applied to cases where part of a chromosome is duplicated or deficient.
- Anther. The organ in flowers that produces pollen.
- **Antibody.** Substance in a tissue or fluid of the body that acts in antagonism to a foreign substance (antigen).
- **Anticodon.** Three bases in a transfer RNA molecule that are complementary to the three bases of a specific codon in messenger RNA.
- **Antigen.** A substance, usually a protein, that is bound by an antibody or a T-cell receptor when introduced into a vertebrate organism.
- **Antisense RNA.** RNA that is complementary to the pre-mRNA or mRNA produced from a gene.
- **Apomixis.** An asexual method of reproduction involving the production of unreduced (usually diploid) eggs, which then develop without fertilization.
- **Apoptosis.** A phenomenon in which eukaryotic cells die because of genetically programmed events within those cells.
- Aptamer domain. The metabolite-binding region of a riboswitch.
- **Artificial selection.** The practice of choosing individuals from a population for reproduction, usually because these individuals possess one or more desirable traits.
- **Ascospore.** One of the spores contained in the ascus of certain fungi such as *Neurospora*.
- **Ascus** (*pl.*, **asci**). Reproductive sac in the sexual stage of a type of fungi (Ascomycetes) in which ascospores are produced.
- **Asexual reproduction.** Any process of reproduction that does not involve the formation and union of gametes from the different sexes or mating types.
- **Assortative mating.** Mating in which the partners are chosen because they are phenotypically similar.

- **Asynapsis.** The failure or partial failure in the pairing of homologous chromosomes during the meiotic prophase.
- **ATP.** Adenosine triphosphate: an energy-rich compound that promotes certain activities in the cell.
- **Attenuation.** A mechanism for controlling gene expression in prokaryotes that involves premature termination of transcription.
- **Attenuator.** A nucleotide sequence in the 5' region of a prokaryotic gene (or in its RNA) that causes premature termination of transcription, possibly by forming a secondary structure.
- **Autocatalytic reaction.** A reaction catalyzed by a substrate without the involvement of any other catalytic agent.
- Autoimmune diseases. Disorders in which the immune systems of affected individuals produce antibodies against self antigens— antigens synthesized in their own cells.
- **Autonomous.** A term applied to any biological unit that can function on its own, that is, without the help of another unit. For example, a transposable element that encodes an enzyme for its own transposition (cf. **Nonautonomous**).
- **Autopolyploid.** A polyploid that has multiple and identical or nearly identical sets of chromosomes (genomes). A polyploid species with genomes derived from the same original species.
- **Autoradiograph.** A record or photograph prepared by labeling a substance such as DNA with a radioactive material such as tritiated thymidine and allowing the image produced by radioactive decay to develop on a film over a period of time.
- Autosome. Any chromosome that is not a sex chromosome.
- **Auxotroph.** A mutant microorganism (for example, bacterium or yeast) that will not grow on a minimal medium but that requires the addition of some compound such as an amino acid or a vitamin.

В

- **Backcross.** The cross of an F_1 hybrid to one of the parental types. The offspring of such a cross are referred to as the backcross generation or backcross progeny. (See also **Testcross**.)
- **Back mutation.** A second mutation at the same site in a gene as the original mutation, which restores the wild-type nucleotide sequence.
- **BACs (bacterial artificial chromosomes).** Cloning vectors constructed from bacterial fertility (F) factors; like YAC vectors, they accept large inserts of size 200 to 500 kb.
- **Bacteriophage.** A virus that attacks bacteria. Such viruses are called bacteriophages because they destroy their bacterial hosts.
- **Balanced lethal.** Lethal mutations in different genes on the same pair of chromosomes that remain in repulsion because of close linkage or crossover suppression. In a closed population, only the *trans*-heterozygotes $(l_1 + l_2 + l_3)$ for the lethal mutations survive.
- **Balanced polymorphism.** Two or more types of individuals maintained in the same breeding population by a selection mechanism.
- **Balancer chromosome.** In *Drosophila* genetics, a dominantly marked, multiply-inverted chromosome that suppresses recombination with a homologous chromosome that is structurally normal.
- **Barr body.** A condensed mass of chromatin found in the nuclei of placental mammals that contains one or more X chromosomes; named for its discoverer, Murray Barr.
- Basal body. Small granule to which a cilium or flagellum is attached.
- **Basal transcription factors.** Proteins required for the initiation of transcription in eukaryotes.

- **Base analogs.** Unnatural purine or pyrimidine bases that differ slightly from the normal bases and that can be incorporated into nucleic acids. They are often mutagenic.
- **Base excision repair.** The removal of abnormal or chemically modified bases from DNA.
- **Base substitution.** A single base change in a DNA molecule. (See also **Transition; Transversion.**)
- **B-DNA.** Double-stranded DNA that exists as a right-handed helix with 10.4 base pairs per turn; the conformation of DNA when present in aqueous solutions containing low salt concentrations.
- **Binomial coefficient.** The term that gives the number of ways of obtaining the two possible outcomes in an experiment in which only two outcomes are possible.
- **Binomial expansion.** Exponential multiplication of an expression consisting of two terms connected by a plus (+) or minus (-) sign, such as $(a + b)^n$.
- **Binomial probability.** The frequency associated with the occurrence of an outcome in an experiment that has only two possible outcomes, such as head or tail in coin tossing.
- **Bioinformatics.** The study of genetic and other biological information using computer and statistical techniques.
- **Biometry.** Application of statistical methods to the study of biological problems.
- **Bivalent.** A pair of synapsed or associated homologous chromosomes that have undergone the duplication process to form a group of four chromatids.
- **Blastomere.** Any one of the cells formed from the first few cleavages in animal development.
- **Blastula.** In animals, an early embryo form that follows the morula stage; typically, a single-layered sheet or ball of cells.
- **B lymphocytes (B cells).** An important class of cells that mature in bone marrow and are largely responsible for the antibodymediated or humoral immune response; they give rise to the antibody-producing plasma cells and some other cells of the immune system.
- **Broad-sense heritability.** In quantitative genetics, the proportion of the total phenotypic variance that is due to genetic factors.

С

- **CAAT box.** A conserved nucleotide sequence in eukaryotic promoters involved in the initiation of transcription.
- **Carbohydrate.** A molecule consisting of carbon, hydrogen, and oxygen in the proportions 1:2:1; a molecule of sugar or a macromolecule composed of sugar subunits.
- **5' cap (mRNA).** The 7-methy guanosine cap that is added to most eukaryotic mRNAs posttranscriptionally.
- Carcinogen. An agent capable of inducing cancer in an organism.
- **Carrier.** An individual who carries a recessive allele that is not expressed (that is, is obscured by a dominant allele).
- **Catabolic pathway.** A pathway by which an organic molecule is degraded in order to obtain energy for growth and other cellular processes; degradative pathway.
- **Catabolite activator protein (CAP).** A positive regulatory protein that in the presence of cyclic AMP (cAMP) binds to the promoter regions of operons and stimulates their transcription. CAP/cAMP assures that glucose is used as a carbon source when present rather than less efficient energy sources such as lactose, arabinose, and

other sugars. When glucose is present, it prevents the synthesis of cAMP and thus the activation of transcription by CAP/cAMP.

- **Catabolite repression.** Glucose-mediated reduction in the rates of transcription of operons that specify enzymes involved in catabolic pathways (such as the *lac* operon).
- **cDNA (complementary DNA).** A DNA molecule synthesized *in vitro* from an RNA template.
- **cDNA library.** A collection of cDNA clones containing copies of the RNAs isolated from an organism or a specific tissue or cell type of an organism.
- **Cell cycle.** The cyclical events that occur during the divisions of mitotic cells. The cell cycle oscillates between mitosis and the interphase, which is divided into G_1 , S, and G_2 .

CentiMorgan. See Crossover unit.

- **Centriole.** An organelle in many animal cells that appears to be involved in the formation of the spindle during mitosis.
- Centromere. Spindle-fiber attachment region of a chromosome.
- **Centrosome.** A barrel-shaped organelle associated with the mitotic spindle in animal cells.
- **Chain-termination codon.** A codon that specifies polypeptide chain termination rather than the incorporation of an amino acid. There are three such codons (UAA, UAG, and UGA), and they are recognized by protein release factors rather than tRNAs.
- **Chaperone.** A protein that helps nascent polypeptides fold into their proper three-dimensional structures.
- **Character** *(contraction of the word* **characteristic)**. One of the many details of structure, form, substance, or function that make up an individual organism.
- **Checkpoint.** A mechanism that halts progression through the eukaryotic cell cycle.
- **Chemotaxis.** Attraction or repulsion of organisms by a diffusing substance.
- **Chiasma** (*pl.*, **Chiasmata**). A visible change of partners in two of a group of four chromatids during the first meiotic prophase. In the diplotene stage of meiosis, the four chromatids of a bivalent are associated in pairs, but in such a way that one part of two chromatids is exchanged. This point of "change of partner" is the chiasma.
- **Chimera (animal).** Individual derived from two embryos by experimental intervention.
- **Chimera (plant).** Part of a plant with a genetically different constitution as compared with other parts of the same plant. It may result from different zygotes that grow together or from artificial fusion (grafting); it may either be pernical, with parallel layers of genetically different tissues, or sectorial.
- **Chimeric selectable marker gene.** A gene constructed using DNA sequences from two or more sources that allows a cell or organism to survive under conditions where it would otherwise die.
- **Chi-square.** A statistic used to test the goodness of fit of data to the predictions of an hypothesis.
- **Chloroplast.** A green organelle in the cytoplasm of plants that contains chlorophyll and in which starch is synthesized. A mode of cytoplasmic inheritance, independent of nuclear genes, has been associated with these cytoplasmic organelles.

Chloroplast DNA. See cpDNA.

Chorionic biopsy. A procedure in which cells are taken from an embryo for the purpose of genetic testing.

- **Chromatid.** In mitosis or meiosis, one of the two identical strands resulting from self-duplication of a chromosome.
- **Chromatin.** The complex of DNA and proteins in eukaryotic chromosomes; originally named because of the readiness with which it stains with certain dyes.
- **Chromatin fiber.** A basic organizational unit of eukaryotic chromosomes that consists of DNA and associated proteins assembled into a strand of average diameter 30 nm.
- **Chromatin remodeling.** The alteration of the structure of DNA and its associated protein molecules, especially histones, by a protein complex; this remodeling often involves the chemical modification of the histones.
- **Chromatography.** A method for separating and identifying the components from mixtures of molecules having similar chemical and physical properties.
- **Chromocenter.** Body produced by fusion of the heterochromatic regions of the chromosomes in the polytene tissues (for example, the salivary glands) of certain *Diptera*.
- **Chromomeres.** Small bodies that are identified by their characteristic size and linear arrangement along a chromosome.
- **Chromonema** (*pl.*, **chromonemata**). An optically single thread forming an axial structure within each chromosome.
- **Chromosome aberration.** Abnormal structure or number of chromosomes; includes deficiency, duplication, inversion, translocation, aneuploidy, polyploidy, or any other change from the normal pattern.
- **Chromosome banding.** Staining of chromosomes in such a way that light and dark areas occur along the length of the chromosomes. Lateral comparisons identify pairs. Each human chromosome can be identified by its banding pattern.
- **Chromosome jumping.** A procedure that uses large DNA fragments to move discontinuously along a chromosome from one site to another site. (See also **Positional cloning.**)
- **Chromosome painting.** The study of the organization and evolution of chromosomes by *in situ* hybridization using DNA probes labeled with fluorescent dyes that emit light at different wavelengths.
- **Chromosomes.** Darkly staining nucleoprotein bodies that are observed in cells during division. Each chromosome carries a linear array of genes.
- **Chromosome Theory of Heredity.** The theory that chromosomes carry the genetic information and that their behavior during meiosis provides the physical basis for the segregation and independent assortment of genes.
- **Chromosome walking.** A procedure that uses overlapping clones to move sequentially down a chromosome from one site to another site. (See also **Positional cloning.**)
- **Cilium** (*pl.*, **cilia**; *adj.*, **ciliate**). Hairlike locomotor structure on certain cells; a locomotor structure on a ciliate protozoan.
- *cis*-acting sequence. A nucleotide sequence that only affects the expression of genes located on the same chromosome, that is, *cis* to itself.
- cis configuration. See Coupling.
- *cis* heterozygote. A heterozygote that contains two mutations arranged in the *cis* configuration—for example, $a^+ b^+ / a b$.
- *cis-trans* **position effect.** The occurrence of different phenotypes when two mutations are present in *cis-* and *trans*-heterozygotes.

- *cis-trans* test. The construction and analysis of *cis* and *trans* heterozygotes of pairs of mutations to determine whether the mutations are in the same gene or in two different genes. For the test to be informative, the *cis* heterozygote must have the wild-type phenotype. If this condition is met, the two mutations are in the same gene if the *trans* heterozygote has the mutant phenotype, and they are in two different genes if the *trans* heterozygote has the wildtype phenotype.
- *ClB* chromosome. An X chromosome in *Drosophila* that carries a mutation causing bar-shaped eyes and a recessive lethal mutation within a large inversion.
- *CIB* method. The use of a special X chromosome in *Drosophila* that carries a mutation causing bar-shaped eyes and a recessive lethal mutation within a long inversion to detect new recessive X-linked lethal mutations. H. J. Muller used this chromosome to demonstrate that X rays are mutagenic. See also *CIB* chromosome.
- **Clone.** All the individuals derived by vegetative propagation from a single original individual. In molecular biology, a population of identical DNA molecules all carrying a particular DNA sequence from an organism.
- **Cloning (gene).** The production of many copies of a gene or specific DNA sequence.
- **Cloning vector.** A small, self-replicating DNA molecule—usually a plasmid or viral chromosome—into which foreign DNAs are inserted in the process of cloning genes or other DNA sequences of interest.
- **Codominant alleles.** Alleles that produce independent effects when heterozygous.
- **Codon.** A set of three adjacent nucleotides in an mRNA molecule that specifies the incorporation of an amino acid into a polypeptide chain or that signals the end of polypeptide synthesis. Codons with the latter function are called termination codons.
- **Coefficient.** A number expressing the amount of some change or effect under certain conditions (for example, the coefficient of inbreeding).
- **Coefficient of coincidence.** The ratio of the observed frequency of double crossovers to the expected frequency, which is calculated on the assumption that crossovers in adjacent segments of the chromosome occur independently.
- **Coefficient of relationship.** The fraction of genes two individuals share by virtue of common ancestry.
- **Coenzyme.** A substance necessary for the activity of an enzyme.
- **Coincidence.** The ratio of the observed frequency of double crossovers to the expected frequency, where the expected frequency is calculated by assuming that the two crossover events occur independently of each other.
- **Cointegrate.** A DNA molecule formed by the fusion of two different DNA molecules, usually mediated by a transposable element.
- **Colchicine.** An alkaloid derived from the autumn crocus that is used as an agent to arrest spindle formation and interrupt mitosis.
- **Colinearity** (*adj.*, **colinear**). A relationship in which the units in one molecule occur in the same sequence as the units in another molecule which they specify; for example, the nucleotides in a gene are colinear with the amino acids in the polypeptide encoded by that gene.
- **Colony.** A compact collection of cells produced by the division of a single progenitor cell.

Comparative genomics. The branch of genomics that compares the structure and function of the genomes of different species.

- **Competence** (*adj.*, **competent**). Ability of a bacterial cell to incorporate DNA and become genetically transformed.
- **Competence (Com) proteins.** Proteins that mediate the process of transformation in bacteria. Their synthesis is induced by small peptides called competence pheromones.
- **Complementarity.** The relationship between the two strands of a double helix of DNA. Thymine in one strand pairs with adenine in the other strand, and cytosine in one strand pairs with guanine in the other strand.
- **Complementation screening.** Screening expression libraries for cDNA or genomic clones based on their ability to rescue mutant host cells.
- **Complementation test (***trans* **test).** Introduction of two recessive mutations into the same cell to determine whether they are alleles of the same gene, that is, whether they affect the same genetic function. If the mutations are allelic, the genotype $m_1 + / + m_2$ will exhibit a mutant phenotype, whereas if they are nonallelic, it will exhibit the wild phenotype.
- **Composite transposon.** A transposable element formed when two identical or nearly identical transposons insert on either side of a nontransposable segment of DNA—for example, the bacterial transposon Tn5.
- **Compound chromosome.** A chromosome formed by the union of two separate chromosomes from the same pair, as in attached-X chromosomes or attached X-Y chromosomes.
- **Concordance rate.** Among pairs of items identified because one member of the pair has a particular trait, the frequency with which the other member of the pair has the same trait.
- **Conditional lethal mutation.** A mutation that is lethal under one set of environmental conditions—the restrictive conditions—but is viable under another set of environmental conditions—the permissive conditions.
- **Conidium** (*pl.*, **conidia**). An asexual spore produced by a specialized hypha in certain fungi.
- **Conjugation.** Union of sex cells (gametes) or unicellular organisms during fertilization; in *Escherichia coli*, a one-way transfer of genetic material from a donor ("male" cell) to a recipient ("female" cell).
- **Conjugative R plasmid.** A circular DNA molecule that can be transferred from one bacterium to another during conjugation.
- **Consanguineous mating.** A mating between relatives.
- **Consanguinity.** Relationship due to descent from a common ancestor.
- **Consensus sequence.** The nucleotide sequence that is present in the majority of genetic signals or elements that perform a specific function.
- **Constitutive enzyme.** An enzyme that is synthesized continually regardless of growth conditions (cf. **Inducible enzyme** and **Repressible enzyme**).
- **Constitutive gene.** A gene that is continually expressed in all cells of an organism.
- **Contig.** A set of overlapping clones that provide a physical map of a portion of a chromosome.
- **Continuous replication.** The synthesis of a nascent strand of DNA by the sequential addition of nucleotides to the 3'-OH terminus of the strand. Characteristic of the synthesis of the leading strand—the strand being extended in the overall $5' \rightarrow 3'$ direction.

- **Continuous variation.** Variation not represented by distinct classes. Individuals grade into each other, and measurement data are required for analysis (cf. **Discontinuous variation**). Multiple genes are usually responsible for this type of variation.
- **Controlling element.** In maize, a transposable element such as Ac or Ds that is capable of influencing the expression of a nearby gene.
- **Coordinate repression.** Correlated regulation of the structural genes in an operon by a molecule that interacts with the operator sequence.
- **Copolymers.** Mixtures consisting of more than one monomer; for example, polymers of two kinds of organic bases such as uracil and cytosine (poly-UC) have been combined for studies of the genetic code.
- **Co-repressor.** An effector molecule that forms a complex with a repressor and turns off the expression of a gene or set of genes.
- Correlation. A statistical association between variables.
- **Cosmids.** Cloning vectors that are hybrids between phage λ chromosomes and plasmids; they contain λ *cos* sites and plasmid origins of replication.
- **Coupling (***cis* **configuration).** The condition in which a double heterozygote has received two linked mutations from one parent and their wild-type alleles from the other parent (for example, $a b/a b \times + +/+ + produces a b/+ + (cf. Repulsion)$).
- **Covalent bond.** A bond in which an electron pair is equally shared by protons in two adjacent atoms.
- Covariance. A measure of the statistical association between variables.
- **cpDNA.** The DNA of plant plastids, including chloroplasts.
- **CpG islands.** Clusters of cytosines and guanines that often occur upstream of human genes.
- **Cri-du-chat syndrome.** A condition produced when a small region in the short arm of one human chromosome 5 is deleted.
- **Critical value.** The threshold value of a statistic that marks off a fraction of the statistic's frequency distribution. A sample statistic greater than this critical value warrants rejection of the hypothesis being tested.
- **Crossbreeding.** Mating between members of different races or species.
- **Crossing over.** A process in which chromosomes exchange material through the breakage and reunion of their DNA molecules. (See also **Recombination**.)
- **Crossover unit.** A measure of distance on genetic maps that is based on the average number of crossing-over events that take place during meiosis. A map interval that is one crossover unit in length (sometimes called a centiMorgan) implies that only one in every hundred chromatids recovered from meiosis will have undergone a crossing-over event in this interval.
- **Cut-and-paste transposon.** A transposable element that is excised from one position in the genome and inserted into another position through the action of a transposon-encoded enzyme called the transposase.
- **Cyclic AMP.** Adenosine-3', 5'-monophosphate, a small molecule that must be bound by the catabolite activator protein (CAP) in order for the complex (CAP/cAMP) to bind to the promoters of operons and stimulate transcription.
- Cystic fibrosis (CF). An autosomal recessive disorder in humans characterized by clogging of the lungs, pancreas, and liver with

mucus and, as a result, chronic infections. The average life expectancy of an individual with cystic fibrosis is about 35 years.

- **Cytogenetics.** Area of biology concerned with chromosomes and their implications in genetics.
- **Cytokinesis.** Cytoplasmic division and other changes exclusive of nuclear division that are a part of mitosis or meiosis.
- **Cytological map.** A diagram of a chromosome based on differential staining—the "banding pattern"—along its length.
- Cytology. The study of the structure and function of cells.
- **Cytoplasm.** The protoplasm of a cell outside the nucleus in which cell organelles (mitochondria, plastids, and the like) reside; all living parts of the cell except the nucleus.
- **Cytoplasmic inheritance.** Hereditary transmission dependent on the cytoplasm or structures in the cytoplasm rather than the nuclear genes; extrachromosomal inheritance. Example: Plastid characteristics in plants may be inherited by a mechanism independent of nuclear genes.
- **Cytosine (C).** A pyrimidine base found in RNA and DNA.
- **Cytoskeleton.** A complex system of fibers and filaments that provides support for cells and that is involved in moving the components of cells throughout the cytoplasm.

D

Dalton. The mass of a hydrogen atom.

Daughter cell. A product of cell division.

- **Deficiency (deletion).** Absence of a segment of a chromosome, reducing the number of loci.
- **Degeneracy (of the genetic code).** The specification of an amino acid by more than one codon.
- **Degrees of freedom.** An index associated with the frequency distribution of a test statistic calculated from sample data.
- **Denaturation.** Loss of native configuration of a macromolecule, usually accompanied by loss of biological activity. Denatured proteins often unfold their polypeptide chains and express changed properties of solubility.
- de novo. Arising anew, afresh, once more.
- **Deoxyribonuclease (DNase).** Any enzyme that hydrolyzes DNA.
- Deoxyribonucleic acid. See DNA.
- **Derepression.** The process of turning on the expression of a gene or set of genes whose expression has been repressed (turned off).
- **Determination.** Process by which undifferentiated cells in an embryo become committed to develop into specific cell types, such as neuron, fibroblast, and muscle cell.
- Deviation. As used in statistics, a departure from an expected value.
- **Diakinesis.** A stage of meiosis just before metaphase I in which the bivalents are shortened and thickened.
- Dicentric chromosome. One chromosome having two centromeres.

Dicot. A plant with two cotyledons, or seed leaves.

- 2',3'-Dideoxyribonucleoside triphosphates (ddNTPs). Chainterminating DNA precursors (nucleoside triphosphates) with a hydrogen (H) linked to the 3' carbon in place of the hydroxyl (OH) group in normal DNA precursors (2'-deoxyribonucleotide triphosphates); ddNTPs are used in DNA sequencing reactions.
- **Differentiation.** A process in which unspecialized cells develop characteristic structures and functions.

- **Dihybrid, Dihybrid cross.** An individual that is heterozygous for two pairs of alleles; the progeny of a cross between homozygous parents differing in two respects.
- **Dimer.** A compound having the same percentage composition as another but twice the molecular weight; one formed by polymerization.
- **Dimorphism.** Two different forms in a group as determined by such characteristics as sex, size, or coloration.
- **Diploid.** An organism or cell with two sets of chromosomes (2n) or two genomes. Somatic tissues of higher plants and animals are ordinarily diploid in chromosome constitution in contrast with the haploid (monoploid) gametes.
- **Diplonema** (*adj.*, **diplotene**). That stage in prophase of meiosis I following the pachytene stage, but preceding diakinesis, in which the chromosomes of bivalents separate from each other at and around their centromeres.
- **Discontinuous replication.** The synthesis of a nascent strand of DNA by the formation of short segments of DNA (Okazaki fragments) that are subsequently joined by DNA ligase. Characteristic of the synthesis of the lagging strand—the strand being extended in the overall $3' \rightarrow 5'$ direction.
- **Discontinuous variation.** Phenotypic variability involving distinct classes such as red versus white, tall versus dwarf (cf. **Continuous variation**).
- **Discordant.** Members of a pair showing different, rather than similar, characteristics.
- **Disjunction.** Separation of homologous chromosomes during anaphase of mitotic or meiotic divisions. (See also **Nondisjunction**.)
- **Dissociation** (Ds). A transposable element in maize, originally detected as an agent that mediates chromosome breakage in response to the effect of *Activator* (*Ac*), another transposable element.

Dizygotic (DZ) twins. Two-egg or fraternal twins.

- **DNA.** Deoxyribonucleic acid; the information-carrying genetic material that comprises the genes. DNA is a macromolecule composed of a long chain of deoxyribonucleotides joined by phosphodiester linkages. Each deoxyribonucleotide contains a phosphate group, the five-carbon sugar 2-deoxyribose, and a nitrogen-containing base.
- DNA chip. See Gene chip.
- DNA fingerprint. See DNA profile.
- **DNA gyrase.** An enzyme in bacteria that catalyzes the formation of negative supercoils in DNA.
- **DNA helicase.** An enzyme that catalyzes the unwinding of the complementary strands of a DNA double helix.
- **DNA ligase.** An enzyme that catalyzes covalent closure of nicks in DNA double helices.
- **DNA photolyase.** An enzyme that uses energy from blue light to cleave ultraviolet light-induced covalent cross-linkages in thymine, cytosine, and cytosine-thymine dimers in DNA.
- **DNA polymerase.** An enzyme that catalyzes the synthesis of DNA.
- **DNA primase.** An enzyme that catalyzes the synthesis of short strands of RNA that initiate the synthesis of DNA strands.
- **DNA profile (DNA print).** A recorded pattern of DNA polymorphisms.
- **DNA profiling (DNA fingerprinting).** The use of DNA sequence data—especially highly polymorphic short tandem repeats (STRs) and variable number tandem repeats (VNTRs)—in personal identity cases.

- **DNA repair enzymes.** Enzymes that catalyze the repair of damaged DNA.
- **DNA topoisomerase.** An enzyme that catalyzes the introduction or removal of supercoils from DNA.
- **Dominant.** A term applied to an allele that is manifested to the exclusion of a different allele in a heterozygote.
- **Dominant-negative mutation.** A mutant allele of a gene that interferes with the function of a wild-type allele so that individuals heterozygous for the mutant and wild-type alleles have a mutant phenotype.
- **Dominant selectable marker gene.** A gene that allows the host cell to survive under conditions where it would otherwise die.
- **Donor cell.** A bacterium that donates DNA to another (recipient) cell during recombination in bacteria (cf. **Recipient cell**).
- **Dosage compensation.** A phenomenon in which the activity of a gene is increased or decreased according to the number of copies of that gene in the cell.
- **Double helix.** A DNA molecule composed of two complementary strands.
- **Downstream sequence.** A sequence in a unit of transcription that follows (is located 3' to) the transcription start site. The nucleotide pair in DNA corresponding to the nucleotide at the 5' end of the transcript (RNA) is designated +1. The following nucleotide pair is designated +2. All of the following (+) nucleotide sequences are downstream sequences (cf. **Upstream sequence**).
- **Down syndrome.** The phenotype due to the presence of an extra chromosome 21 in humans.
- Drift. See Random genetic drift.
- **Duplication.** The occurrence of a segment more than once in the same chromosome or genome; also, the multiplication of cells.

Ε

- Ecdysone. A hormone that influences development in insects.
- Eclosion. Emergence of an adult insect from the pupal stage.
- **Ecotype.** A population or strain of organisms that is adapted to a particular habitat.
- **Ectopic.** A term used to describe a phenomenon that occurs in an abnormal place.
- **Effector molecule.** A molecule that influences the behavior of a regulatory molecule, such as a repressor protein, thereby influencing gene expression.
- **Egg (ovum).** A germ cell produced by a female organism.
- **Electrophoresis.** The migration of suspended particles in an electric field.
- **Electroporation.** A process whereby cell membranes are made permeable to DNA by applying an intense electric current.
- **Elongation (of DNA, RNA, or protein synthesis).** The incorporation of the second and subsequent subunits (nucleotides or amino acids) during the synthesis of a macromolecule (DNA, RNA, or polypeptide).
- **Elongation factors.** Soluble proteins that are required for polypeptide chain elongation.
- **Embryo.** An organism in the early stages of development; in humans, the first two months in the uterus.
- **Embryoid bodies.** Masses of differentiated and undifferentiated cells derived from embryonic stem cells.

- **Embryonic stem cells (ES cells).** Cells present in embryos that can differentiate into many different types of tissues and/or organs.
- **Embryo sac.** A large thin-walled space within the ovule of the seed plant in which the egg and, after fertilization, the embryo develop; the mature female gametophyte in higher plants.
- **Endomitosis.** Duplication of chromosomes without division of the nucleus, resulting in increased chromosome number within a cell. Chromosome strands separate, but the cell does not divide.
- **Endonuclease.** An enzyme that breaks strands of DNA at internal positions; some are involved in recombination of DNA.
- **Endoplasmic reticulum.** Network of membranes in the cytoplasm to which ribosomes adhere.
- **Endopolyploidy.** A state in which the cells of a diploid organism contain multiples of the diploid chromosome number (that is, 4n, 8n, and so on).
- **Endosperm.** Nutritive tissue that develops in the embryo sac of most angiosperms. It usually forms after the fertilization of the two fused primary endosperm nuclei of the embryo sac with one of the two male gamete nuclei. In most diploid plants, the endosperm is triploid (3n).
- **Endosymbiosis.** A mutually beneficial relationship in which one organism lives inside another organism.
- End-product inhibition. See Feedback inhibition.
- **Enhancer.** A substance or an object that increases a chemical activity or a physiological process; a major or modifier gene that increases a physiological process; a DNA sequence that influences transcription of a nearby gene.
- **Environment.** The aggregate of all the external conditions and influences affecting the life and development of an organism.
- **Environmentality.** The proportion of the total phenotypic variance in a quantitative trait that is due to the effects of a shared environment.
- **Enzyme.** A protein that accelerates a specific chemical reaction in a living system.
- **Epigenetic.** A term referring to the nongenetic causes of a phenotype.
- **Episome.** A genetic element that may be present or absent in different cells and that may be inserted in a chromosome or independent in the cytoplasm (for example, the fertility factor (F) in *Escherichia coli*).
- **Epistasis.** Interactions between products of nonallelic genes. Genes suppressed are said to be hypostatic. Dominance is associated with members of allelic pairs, whereas epistasis results from interactions of the products of nonalleles.
- **Equational division.** Mitotic-type division that is usually the second division in the meiotic sequence; somatic mitosis and the nonreductional division of meiosis.
- **Equatorial plate.** The figure formed by the chromosomes in the center (equatorial plane) of the spindle in mitosis.
- **Equilibrium.** A state of dynamical systems in which there is no net change.
- **Equilibrium density-gradient centrifugation.** A procedure used to separate macromolecules based on their density (mass per unit volume).
- Estrogen. Female hormone or estrus-producing compound.
- **ESTs (expressed sequence tags).** Short cDNA sequences that are used to link physical maps and genetic (RFLP) maps.

- **Euchromatin.** Genetic material that is not stained so intensely by certain dyes during interphase and that comprises many different kinds of genes (cf. **Heterochromatin**).
- **Eugenics.** The application of the principles of genetics to the improvement of humankind.
- **Eukaryote.** A member of the large group of organisms that have nuclei enclosed by a membrane within their cells (cf. **Prokaryote**).
- **Eukaryotic cells.** The cells of organisms classified as eukaryotes. These cells are characterized by having a membrane-bound nucleus that contains the chromosomal DNA.
- **Euploid.** An organism or cell having a chromosome number that is an exact multiple of the monoploid (*n*) or haploid number. Terms used to identify different levels in an euploid series are diploid, triploid, tetraploid, and so on (cf. **Aneuploid**).
- **Excinuclease.** The endonuclease-containing protein complex that excises a segment of damaged DNA during excision repair.
- **Excision repair.** DNA repair processes that involve the removal of the damaged segment of DNA and its replacement by the synthesis of a new strand using the complementary strand of DNA as template.
- **Exit** (*E*) site. The ribosome binding site that contains the free tRNA prior to its release.
- **Exon amplification.** A procedure that is used to identify coding regions (exons) that are flanked by 5' and 3' intron splice sites.
- **Exons.** The segments of a eukaryotic gene that correspond to the sequences in the final processed RNA transcript of that gene.
- **Exonuclease.** An enzyme that digests DNA or RNA, beginning at the ends of strands.
- **Expression domain.** The region of a riboswitch that can fold into two conformations, one facilitating gene expression and the other blocking gene expression.
- **Extrachromosomal.** Structures that are not part of the chromosomes; DNA units in the cytoplasm that control cytoplasmic inheritance.

F

- \mathbf{F}_{1} . The first filial generation; the first generation of descent from a given mating.
- F_2 . The second filial generation produced by crossing *inter se* or by self-pollinating the F_1 . The inbred "grandchildren" of a given mating, but in controlled genetic experimentation, self-fertilization of the F_1 (or equivalent) is implied.
- F⁺ cell. A bacterium that contains an autonomous fertility (F) factor.See F factor.
- **F factor.** A bacterial episome that confers the ability to function as a genetic donor ("male") in conjugation; the fertility factor in bacteria.
- **Feedback inhibition (or end-product inhibition).** The accumulated end product of a biochemical pathway stops synthesis of that product. A late metabolite of a synthetic pathway regulates synthesis at an earlier step of the pathway.
- **Female gametophyte.** A large thin-walled space within the ovule of the seed plant that contains the eight identical haploid nuclei derived by mitosis from the megaspore that was produced by meiosis.
- **Fertilization.** The fusion of a male gamete (sperm) with a female gamete (egg) to form a zygote.
- **Fetus.** Prenatal stage of a viviparous animal between the embryonic stage and the time of birth; in humans, the final seven months before birth.

Filial. See \mathbf{F}_1 and \mathbf{F}_2 .

- **Fission.** A mode of cell division among the prokaryotes in which the genetic material of the mother cell is first duplicated and then apportioned equally to the two daughter cells.
- **Fitness.** The number of offspring left by an individual, often compared with the average of the population or with some other standard, such as the number left by a particular genotype.
- **Fixation.** An event that occurs when all the alleles at a locus except one are eliminated from a population. The remaining allele, with frequency 100 percent, is said to have been fixed.
- **Flagellum** (*pl.*, **flagella**; *adj.* **flagellate**). A whiplike organelle of locomotion in certain cells; locomotor structures in flagellate protozoa.
- Fluorescence *in situ* hybridization (FISH). *In situ* hybridization performed using a DNA or RNA probe coupled to a fluorescent dye.
- **Folded genome.** The condensed intracellular state of the DNA in the nucleoid of a bacterium. The DNA is segregated into domains, and each domain is independently negatively supercoiled.
- **Founder principle.** The possibility that a new, small, isolated population may diverge genetically because the founding individuals are a random sample from a large, main population.
- **Frameshift mutation.** A mutation that changes the reading frame of an mRNA, either by inserting or deleting nucleotides.
- **Frequency distribution.** A graph showing either the relative or absolute incidence of classes in a population. The classes may be defined by either a discrete or a continuous variable; in the latter case, each class represents a different interval on the scale of measurement.
- **Fusion protein.** A polypeptide made from a recombinant gene that contains portions of two or more different genes. The different genes are joined so that their coding sequences are in the same reading frame.

G

- **Gain-of-function mutation.** A mutation that endows a gene product with a new function.
- Gall. A tumorous growth in plants.
- Gamete. A mature male or female reproductive cell (sperm or egg).
- Gametogenesis. The formation of gametes.
- **Gametophyte.** That phase of the plant life cycle that bears the gametes; the cells have *n* chromosomes.
- **Gametophytic incompatibility.** A botanical phenomenon controlled by the complex *S* locus in which a pollen grain cannot fertilize an ovule produced by a plant that carries the same *S* allele as the pollen grain. For example, S_1 pollen cannot fertilize an ovule made by an S_1/S_2 plant.
- **Gap gene.** A gene that controls the formation of adjacent segments in the body of *Drosophila*.
- **Gastrula.** An early animal embryo consisting of two layers of cells; an embryological stage following the blastula.
- **GenBank.** The DNA sequence databank maintained by the National Center for Biotechnology Information at the National Institutes of Health in the United States. Similar databanks are maintained in Europe (the European Molecular Biology Laboratory Data Library) and Japan (the DNA DataBank of Japan).
- Gene. A hereditary determinant of a specific biological function; a unit of inheritance (DNA) located in a fixed position on a

728 Glossary

chromosome; a segment of DNA encoding one polypeptide and defined operationally by the *cis-trans* or complementation test.

Gene addition. The addition of a functional copy of a gene to the genome of an organism.

Gene amplification. A phenomenon whereby the DNA of a specific gene or set of genes is replicated independently of the rest of the genome to increase the number of gene copies.

- **Gene chip.** A small silicon wafer or other solid support containing a large number of oligonucleotide or cDNA hybridization probes arranged on its surface in a specific pattern, or microarray.
- **Gene cloning.** The incorporation of a gene of interest into a self-replicating DNA molecule and the amplification of the resulting recombinant DNA molecule in an appropriate host cell.
- **Gene conversion.** A process, often associated with recombination, during which one allele is replicated at the expense of another, leading to non-Mendelian segregation ratios. In whole tetrads, for example, the ratio may be 6:2 or 5:3 instead of the expected 4:4.
- **Gene expression.** The process by which genes produce RNAs and proteins and exert their effects on the phenotype of an organism.
- **Gene flow.** The spread of genes from one breeding population to another by migration, possibly leading to allele frequency changes.
- **Gene pool.** The sum total of all different alleles in the breeding members of a population at a given time.
- **Generalized transduction.** Recombination in bacteria mediated by a bacteriophage that can transfer any bacterial gene of the donor cell to a recipient cell (cf. **Specialized transduction**).
- **Gene replacement.** The incorporation of a transgene into a chromosome at its normal location by homologous recombination, thus replacing the copy of the gene originally present at the locus.
- **Gene therapy.** The treatment of inherited diseases by introducing wild-type copies of the defective gene causing the disorder into the cells of affected individuals. If reproductive cells are modified, the procedure is called *germ-line* or *heritable gene therapy*. If cells other than reproductive cells are modified, the procedure is called *somatic-cell* or *noninheritable gene therapy*.
- **Genetic code.** The set of 64 nucleotide triplets that specify the 20 amino acids and polypeptide chain initiation and termination.
- Genetic drift. See Random genetic drift.
- **Genetic equilibrium.** Condition in a group of interbreeding organisms in which the allele frequencies remain constant over time.
- **Genetic map.** A diagram of a chromosome with distances based on recombination frequencies—centiMorgans.
- Genetics. The science of heredity and variation.
- **Genetic selection.** The exposure of a cell or an organism to environmental conditions in which it can survive only if it carries a specific gene or genetic element.
- **Genome.** A complete set (*n*) of chromosomes (hence, of genes) inherited as a unit from one parent.
- **Genomic DNA library.** A collection of clones containing the genomic DNA sequences of an organism.
- Genomics. The study of the structure and function of entire genomes.
- **Genotype.** The genetic constitution (gene makeup) of an organism (cf. **Phenotype**).
- **Germ cell.** A reproductive cell capable when mature of being fertilized and reproducing an entire organism (cf. **Somatic cell**).

- **Germinal mutation.** A mutation that occurs in the reproductive cells (germ-line cells) of the body and is transmitted to progeny (cf. **Somatic mutation**).
- Germ line. The tissue that ultimately produces the gametes.
- Germ-line (heritable) gene therapy. Treatment of an inherited disorder by adding functional (wild-type) copies of a gene to reproductive (germ-line) cells of an individual carrying defective copies of that gene (cf. Somatic-cell [nonheritable] gene therapy).
- **Germ plasm.** The hereditary material transmitted to the offspring through the germ cells.
- **Globulins.** Common proteins in the blood that are insoluble in water and soluble in salt solutions. Alpha, beta, and gamma globulins can be distinguished in human blood serum. Gamma globulins are important in developing immunity to diseases.
- **Glucocorticoid.** A steroid hormone that regulates gene expression in higher animals.
- **Golgi complex.** A membranous system within cells that is involved in the secretion of cellular substances.
- **Gonad.** A sexual organ (that is, ovary or testis) that produces gametes.
- **Green fluorescent protein (GFP).** A naturally occurring fluorescent protein synthesized by the jellyfish *Aequorea victoria*.
- Guanine (G). A purine base found in DNA and RNA.
- **Guide RNAs.** RNA molecules that contain sequences that function as templates during RNA editing.
- **Gynandromorph.** An individual in which one part of the body is female and another part is male; a sex mosaic.

Н

- **Haploid (monoploid).** An organism or cell having only one complete set (*n*) of chromosomes or one genome.
- **Haplotype.** A set of linked genetic variants, especially single-nucleotide polymorphisms (SNPs), on a chromosome.
- Haptoglobin. A serum protein, alpha globulin, in the blood.
- Hardy–Weinberg Principle. Mathematical relationship that allows the frequencies of genotypes in a population to be predicted from their constituent allele frequencies; a consequence of random mating.
- **Helix.** Any structure with a spiral shape. The Watson and Crick model of DNA is in the form of a double helix.
- **Helper T cells.** T cells that respond to an antigen displayed by a macrophage by stimulating B and T lymphocytes to develop into antibody-producing plasma cells and killer T cells, respectively.
- **Hemizygote.** An individual that carries one copy of a chromosome or gene, as in sex linkage or as a result of deletion.
- **Hemoglobin.** Conjugated protein compound containing iron, located in erythrocytes of vertebrates; important in the transportation of oxygen to the cells of the body.
- **Hemolymph.** The mixture of blood and other fluids in the body cavity of an invertebrate.
- **Hemophilia.** A bleeder's disease; tendency to bleed freely from even a slight wound; hereditary condition dependent on a sex-linked recessive gene.
- **Heredity.** Resemblance among individuals related by descent; transmission of traits from parents to offspring.

- Heritability. Degree to which a given trait is controlled by inheritance. (See also Broad-sense heritability and Narrow-sense heritability.)
- **Hermaphrodite.** An individual with both male and female reproductive organs.
- **Heteroalleles.** Mutations that are functionally allelic but structurally nonallelic; mutations at different sites in a gene.
- **Heterochromatin.** Chromatin staining darkly even during interphase, often containing repetitive DNA with few genes.
- **Heteroduplex.** A double-stranded nucleic acid containing one or more mismatched (noncomplementary) base pairs.
- **Heterogametic sex.** Producing unlike gametes with regard to the sex chromosomes. In humans, the XY male is heterogametic, and the XX female is homogametic.
- **Heterogeneous nuclear RNA (hnRNA).** The population of primary transcripts in the nucleus of a eukaryotic cell.
- **Heterologous chromosome.** A chromosome that contains a different set of genes than the chromosome to which it is compared.
- **Heterosis.** Superiority of heterozygous genotypes in respect to one or more traits in comparison with corresponding homozygotes.
- **Heterozygosity**. The proportion of heterozygous individuals in a population; used as a measure of genetic variability.
- **Heterozygote** (*adj.*, **heterozygous**). An organism with unlike members of any given pair or series of alleles that consequently produces unlike gametes.
- **Hfr.** High-frequency recombination strain of *Escherichia coli*; in such strains, the F episome is integrated into the bacterial chromosome.
- **Histones.** Group of proteins rich in basic amino acids. They function in the coiling of DNA in chromosomes and in the regulation of gene activity.
- **HIV (human immunodeficiency virus).** The retrovirus that causes AIDS in humans.
- **Holoenzyme.** The form of a multimeric enzyme in which all of the component polypeptides are present.
- **Homeobox.** A DNA sequence found in several genes that are involved in the specification of organs in different body parts in animals; characteristic of genes that influence segmentation in animals. The homeobox corresponds to an amino acid sequence in the polypeptide encoded by these genes; this sequence is called the homeodomain.
- Homeodomain. See Homeobox.
- **Homeotic genes.** A group of genes whose products control formation of the body of an embryo by regulating the expression of other genes in segmental regions along the anterior-posterior axis.
- **Homeotic mutation.** A mutation that causes a body part to develop in an inappropriate position in an organism; for example, a mutation in *Drosophila* that causes legs to develop on the head in the place of antennae.
- **Homoalleles.** Mutations that are both functionally and structurally allelic; mutations at the same site in the same gene.
- **Homogametic sex.** Producing like gametes with regard to the sex chromosomes (cf. **Heterogametic sex**).
- **Homologous chromosomes.** Chromosomes that occur in pairs and are generally similar in size and shape, one having come from the male parent and the other from the female parent. Such chromosomes contain the same array of genes.

- Homologous genes. Genes that have evolved from a common ancestral gene (cf. Orthologous genes; Paralogous genes).
- Homologues. See Homologous chromosomes; Homologous genes.
- **Homozygote** (*adj.*, **homozygous**). An individual in which the two copies of a gene are the same allele.
- **Hormone.** An organic product of cells of one part of the body that is transported by the body fluids to another part where it influences activity or serves as a coordinating agent.
- **Human Genome Organization (HUGO).** An international group of scientists formed to coordinate the sequencing and mapping of the human genome.
- **Human Genome Project.** A huge international effort to map and sequence the entire human genome.
- **Human growth hormone (HGH).** A signaling polypeptide required for normal growth in humans; it is deficient in individuals with certain types of dwarfism.
- **Human immunodeficiency virus (HIV).** The retrovirus that causes acquired immune deficiency syndrome (AIDS) in humans.
- Huntington's disease (HD). A late-onset (age 30 to 50 years) neurodegenerative disorder in humans caused by an autosomal dominant mutation. The genetic defect is an expanded $(CAG)_n$ trinucleotide repeat that encodes an abnormally long polyglutamine region near the amino terminus of the *huntingtin* gene product.
- **Hybrid.** An offspring of homozygous parents differing in one or more genes; more generally, an offspring of a cross between unrelated strains.
- **Hybrid dysgenesis.** In *Drosophila*, a syndrome of abnormal germline traits, including mutation, chromosome breakage, and sterility, which results from transposable element activity.
- **Hybridization.** Interbreeding of species, races, varieties, and so on, among plants or animals; a process of forming a hybrid by cross pollination of plants or by mating animals of different types.
- **Hybrid vigor (heterosis).** Unusual growth, strength, and health of heterozygous hybrids derived from two less vigorous homozygous parents.
- **Hydrogen bonds.** Weak interactions between electronegative atoms and hydrogen atoms (electropositive) that are linked to other electronegative atoms.
- **Hydrophobic interactions.** Association of nonpolar groups with each other when present in aqueous solutions because of their insolubility in water.
- **Hydroxylating agent.** A chemical—such as the mutagen hydroxylamine—that transfers hydroxyl groups to other molecules.
- **Hyperploid.** A genetic condition in which a chromosome or a segment of a chromosome is overrepresented in the genotype (cf. **Hypoploid**).
- **Hypersensitive sites.** Regions in the DNA that are highly susceptible to digestion with endonucleases.
- **Hypomorphic.** A term applied to a mutant allele that has less expression than a wild-type allele but that does not completely abolish expression. Such a mutant allele is called a hypomorph.
- **Hypoploid.** A genetic condition in which a chromosome or segment of a chromosome is underrepresented in the genotype (cf. **Hyperploid**).
- **Hypothesis.** In science, a statement about how a phenomenon can be explained.

Imaginal disc. A mass of cells in the larvae of *Drosophila* and other holometabolous insects that gives rise to a particular adult organ such as an antenna, eye, or wing.

Immunoglobulin. See Globulins.

- **Imprinting.** A process that alters the state of a gene without altering its nucleotide sequence; often associated with methylation of specific nucleotides in the gene. The altered state is established in the germ line and is transmitted to the offspring where it may persist throughout the offspring's life. A gene that has been altered in this way is said to have been imprinted.
- *in situ.* From the Latin, meaning in the natural place; refers to experimental treatments performed on cells or tissue rather than on extracts from them.
- *in situ* colony or plaque hybridization. A procedure for screening colonies or plaques growing on plates or membranes for the presence of specific DNA sequences by the hybridization of nucleic acid probes to the DNA molecules present in these colonies or plaques.
- *in situ* hybridization. A method for determining the location of specific DNA sequences in chromosomes by hybridizing labeled DNA or RNA to denatured DNA in chromosome preparations and visualizing the hybridized probe by autoradiography or fluorescence microscopy.
- **Intein.** A short amino acid sequence in a primary translation product that can excise itself from the polypeptide.
- *in vitro.* From the Latin meaning "within glass"; biological processes made to occur experimentally outside the organism in a test tube or other container.
- in vivo. From the Latin meaning "within the living organism."
- **Inbred line.** A strain produced by many generations of systematic inbreeding, for example, by repeated self-fertilization or by repeated full-sib mating.
- Inbreeding. Matings between related individuals.
- **Inbreeding coefficient.** The probability that two alleles in an individual are identical to each other by descent from a common ancestor.
- **Inbreeding depression.** The observation that inbred lines are weaker than noninbred lines.
- **Incomplete dominance.** Expression of two alleles in a heterozygote that allows the heterozygote to be distinguished from either of its homozygous parents.
- **Independent assortment.** The random distribution of alleles to the gametes that occurs when genes are located in different chromosomes. The distribution of one pair of alleles is independent of other genes located in nonhomologous chromosomes.
- **Induced mutation.** A mutation that results from the exposure of an organism to a chemical or physical agent that causes changes in the structure of DNA or RNA (cf. **Spontaneous mutation**).
- **Inducer.** A substance of low molecular weight that is bound by a repressor to produce a complex that can no longer bind to the operator; thus, the presence of the inducer turns on the expression of the gene(s) controlled by the operator.
- **Inducible enzyme.** An enzyme that is synthesized only in the presence of the substrate that acts as an inducer.
- **Inducible gene.** A gene that is expressed only in the presence of a specific metabolite, the inducer.

- **Induction.** The process of turning on the expression of a gene or set of genes by an inducer.
- **Inhibitor.** Any substance or object that retards a chemical reaction; a major or modifier gene that interferes with a reaction.
- **Initiation (of DNA, RNA, or protein synthesis).** The incorporation of the first subunit (nucleotide or amino acid) during the synthesis of a macromolecule (DNA, RNA, or polypeptide).
- **Initiation codon.** A sequence of three nucleotides in mRNA usually AUG, sometimes GUG—that signals the initiation of a new polypeptide during translation.
- **Initiation factors.** Soluble proteins required for the initiation of translation.
- Insertion Sequence. See IS element.
- **Insertional mutation.** A mutation caused by the insertion of foreign DNA such as a transposable element or the T-DNA of the Ti plasmid of *Agrobacterium tumefaciens*.
- **Interaction.** In statistics, an effect that cannot be explained by the additive action of contributing factors; a departure from strict additivity.
- **Intercalating agent.** A chemical capable of inserting between adjacent base pairs in a DNA molecule.
- **Intercross.** A cross between the F_1 hybrids derived from a cross between two parental strains.
- **Interference.** Crossing over at one point that reduces the chance of another crossover nearby; detected by studying the pattern of crossing over with three or more linked genes.
- **Interphase.** The stage in the cell cycle when the cell is not dividing; the metabolic stage during which DNA replication occurs; the stage following telophase of one division and extending to the beginning of prophase in the next division.
- **Intersex.** An organism displaying secondary sexual characters intermediate between male and female; a type that shows some phenotypic characteristics of both males and females.
- **Introns.** Intervening sequences of DNA bases within eukaryotic genes that are not represented in the mature RNA transcript because they are spliced out of the primary RNA transcript.
- **Invariant.** Constant, unchanging, usually referring to the portion of a molecule that is the same across species.
- **Inversion.** A rearrangement that reverses the order of a linear array of genes in a chromosome.
- **Inverted repeat.** A sequence present twice in a DNA molecule but in reverse orientation.
- Ionic bonds. Attractions between oppositely charged chemical groups.
- **Ionizing radiation.** The portion of the electromagnetic spectrum that results in the production of positive and negative charges (ion pairs) in molecules. X rays and gamma rays are examples of ionizing radiation (cf. **Nonionizing radiation**).
- **IS element (insertion sequence).** A short (800–1400 nucleotide pairs) DNA sequence found in bacteria that is capable of transposing to a new genomic location; other DNA sequences that are bounded by IS elements may also be transposed.
- **Isoalleles.** Different forms of a gene that produce the same phenotype or very similar phenotypes.
- **Isochromosome.** A chromosome with two identical arms and identical genes. The arms are mirror images of each other.
- **Isoform.** A member of a family of closely related proteins—proteins that have some amino acid sequences in common and some different.

Κ

- Kappa chain. One of two classes of antibody light chains (cf. Lambda chain).
- **Karyotype.** The chromosome constitution of a cell or an individual; chromosomes arranged in order of length and according to position of centromere; also, the abbreviated formula for the chromosome constitution, such as 47, XX + 21 for human trisomy-21.
- **Kinetics.** A dynamic process involving motion.
- **Kinetochore.** A proteinaceous structure associated with the centromere of a chromosome during eukaryotic cell division; the point at which microtubules attach to move the chromosome through the division process.
- **Klinefelter syndrome.** A condition produced when two X chromosomes and one Y chromosome are present in the human karyotype.
- **Knockout mutation.** A mutation that completely abolishes a gene's function.
- **Kozak's rules.** The sequence requirements—5'-GCC(A or G) CCAUGG-3'—for optimal initiation of translation at the first (5') AUG in eukaryotic mRNAs (named after Marilyn Kozak who first proposed them).

L

- **Lagging strand.** The strand of DNA that is synthesized discontinuously during replication.
- Lambda chain. One of two classes of antibody light chains (cf. Kappa chain).
- **Lamella.** A double-membrane structure, plate, or vesicle that is formed by two membranes lying parallel to each other.
- **Leader sequence.** The segment of an mRNA molecule from the 5' terminus to the translation initiation codon.
- **Leading strand.** The strand of DNA that is synthesized continuously during replication.
- Leptonema (*adj.*, leptotene). Stage in meiosis immediately preceding synapsis in which the chromosomes appear as single, fine, threadlike structures (but they are really double because DNA replication has already taken place).
- Ligand. A molecule that can bind to another molecule in or on cells.
- Ligase. An enzyme that joins the ends of two strands of nucleic acid.
- Ligation. The joining of two or more DNA molecules by covalent bonds.
- **LINEs (long interspersed nuclear elements).** Families of long (average length = 6500 bp) moderately repetitive transposable elements in eukaryotes.
- **Linkage.** The tendency of different genes to be inherited together because they are located on the same chromosome.
- **Linkage equilibrium.** A state in which the alleles of linked loci are randomized with respect to each other on the chromosomes of a population.
- **Linkage map.** A linear or circular diagram that shows the relative positions of genes on a chromosome as determined by genetic analysis.
- **Linkage phase.** The arrangement of linked genetic markers in a heterozygote. The markers can be in the coupling (A B/a b) or in the repulsion (A b/a B) phase.
- **Linker (DNA).** The unprotected DNA double helix that connects adjacent nucleosomes.
- Lipid. A molecule composed of fatty acids and triglycerides.

- **Locus** (*pl.*, **loci**). A fixed position on a chromosome that is occupied by a given gene or one of its alleles.
- **Long terminal repeats.** Identical or nearly identical DNA sequences at opposite ends of an integrated retrovirus or a retroviruslike element. Typically these sequences are a least 300 base pairs in length. Abbreviation: LTRs.
- **Loss-of-function mutation.** A mutation that impairs or abolishes gene expression or the function of a gene product.
- **Lymphocyte.** A general class of white blood cells that are important components of the immune system of vertebrate animals.
- **Lysine riboswitch.** An mRNA in bacteria that undergoes a change from an active (transcribed) conformation to an inactive (nontranscribed) structure when it binds lysine.
- **Lysis.** Bursting of a cell by the destruction of the cell membrane following infection by a virus.
- Lysogenic bacteria. Those harboring temperate bacteriophages.
- **Lysosome.** A small, membrane-bound cellular organelle that contains enzymes dedicated to the degradation of macromolecules.

Lytic phage. See Virulent phage.

Μ

- **Macromolecule.** A large molecule; term used to identify molecules of proteins and nucleic acids.
- **Male gametophyte.** The three identical haploid nuclei within a pollen grain.
- Map unit. See Crossover unit.
- **Mass selection.** As practiced in plant and animal breeding, the choosing of individuals for reproduction from the entire population on the basis of the individual's phenotypes rather than the phenotypes of their relatives.
- **Maternal effect.** Trait controlled by a gene of the mother but expressed in the progeny.
- **Maternal-effect gene.** A gene whose product acts in the offspring of the female who carries the gene.
- **Maternal-effect mutation.** A mutation that causes a mutant phenotype in the offspring of a female that carries the mutation; however, the female herself may not show the mutant phenotype.
- **Maternal inheritance.** Inheritance controlled by extrachromosomal (that is, cytoplasmic) factors that are transmitted through the egg.
- **Mean.** The arithmetic average; the sum of all measurements or values in a sample divided by the sample size.
- **Median.** In a set of measurements, the central value above and below which there are an equal number of measurements.
- **Megaspore.** The single large cell produced at the end of meiosis in the female reproductive tissues of plants.
- **Meiosis.** The process by which the chromosome number of a reproductive cell becomes reduced to half the diploid (2n) or somatic number; results in the formation of gametes in animals or of spores in plants; important source of variability through recombination.
- Melanin. Brown or black pigment.
- **Membrane.** A macromolecular structure composed of lipids and proteins that surrounds a cell or certain of the organelles within a cell, such as the mitochondria and chloroplasts; also, a component of the endoplasmic reticulum within cells.
- **Mendelian population.** A natural interbreeding unit of sexually reproducing plants or animals sharing a common gene pool.

- **Mesoderm.** The middle germ layer that forms in the early animal embryo and gives rise to such parts as bone and connective tissue.
- **Messenger RNA (mRNA).** RNA that carries information necessary for protein synthesis from the DNA to the ribosomes.
- **Metabolism.** Sum total of all chemical processes in living cells by which energy is provided and used.
- **Metacentric chromosome.** A chromosome with the centromere near the middle and two arms of about equal length.
- **Metafemale (superfemale).** In *Drosophila*, abnormal female, usually sterile, with an excess of X chromosomes compared with sets of autosomes (for example, XXX; AA).
- **Metaphase.** That stage of cell division in which the chromosomes are most discrete and arranged in an equatorial plate; stage following prophase and preceding anaphase.
- **Metaphase I.** The stage during the first meiotic division when duplicated homologous chromosomes that have paired condense and gather at the equatorial plane of the cell.
- **Metaphase II.** The stage during the second meiotic division when duplicated chromosomes gather at the equatorial plane of the cell.
- **Metaphase plate.** The equatorial plane where duplicated chromosomes gather in a cell during the metaphase of mitosis.
- Metastasis. The spread of cancer cells to previously unaffected organs.
- **Methylation (of DNA and RNA).** The addition of a methyl $(-CH_3)$ group(s) to one or more of the nucleotides in a nucleic acid.
- **Microarray.** A membrane or other solid support containing thousands of oligonucleotides or nucleic acid hybridization probes for use in detecting complementary DNAs or RNAs.
- **Microprojectile bombardment.** A procedure for transforming plant cells by shooting DNA-coated tungsten or gold particles into the cells.
- MicroRNA. See Short interfering RNA.

Microsatellite See Short tandem repeat (STR).

- **Microspore.** One of the four end products of meiosis in the male reproductive tissues of plants.
- **Microtubule Organizing Center (MTOC).** A region in a eukaryotic cell that generates the microtubules used during cell division. In animal cells, the MTOC is associated with distinct organelles called centrosomes.
- **Microtubules.** Hollow filaments in the cytoplasm making up a part of the locomotor apparatus of a motile cell; component of the mitotic spindle.
- **Midparent value.** In quantitative genetics, the average of the phenotypes of two mates.
- **Mismatch repair.** DNA repair processes that correct base pairs that are not properly hydrogen-bonded.
- **Missense mutation.** A mutation that changes a codon specifying an amino acid to a codon specifying a different amino acid.
- **Mitochondria.** Organelles in the cytoplasm of plant and animal cells where oxidative phosphorylation takes place to produce ATP.
- Mitochondrial DNA. See mtDNA.
- **Mitosis.** Disjunction of duplicated chromosomes and division of the cytoplasm to produce two genetically identical daughter cells.
- **Modal class.** In a frequency distribution, the class having the greatest frequency.
- Model. A mathematical description of a biological phenomenon.

- **Model organisms.** Plants, animals, and microbes that are routinely used in genetic analysis.
- **Modifier (modifying gene).** A gene that affects the expression of some other gene.
- **Monohybrid.** An offspring of two homozygous parents that differ from one another by the alleles present at only one gene locus.
- **Monohybrid cross.** A cross between parents differing in only one trait or in which only one trait is being considered.
- **Monomer.** A single molecular entity that may combine with others to form more complex structures.
- **Monoploid.** Organism or cell having a single set of chromosomes or one genome (chromosome number *n*).
- **Monosomic.** A diploid cell or organism lacking one chromosome of its proper complement (chromosome formula 2n 1). A specific case of this condition is called a monosomy (*pl.* monosomies).
- Monozygotic twins. One-egg or identical twins.
- **Morphogen.** A substance that stimulates the development of form or structure in an organism.
- **Morphology.** Study of the form of an organism; developmental history of visible structures and the comparative relation of similar structures in different organisms.
- **Mosaic.** An organism or part of an organism that is composed of cells of different genotypes.
- **Mother cell.** A cell that is prepared to divide mitotically or meiotically.
- **Motility.** Cell movement, usually accomplished through the action of specialized structures such as cilia and flagella.
- mtDNA. The DNA of mitochondria.
- **Multifactorial trait**. A trait determined by a combination of several genetic and environmental factors.
- **Multigene family.** A group of genes that are similar in nucleotide sequence or that produce polypeptides with similar amino acid sequences.
- **Multiple alleles.** A condition in which a particular gene occurs in three or more allelic forms in a population of organisms.
- **Multiple Factor Hypothesis.** A theory advanced by R. A. Fisher and others to explain variation in complex phenotypes such as height, weight, and disease susceptibility.
- Mutable genes. Genes with an unusually high mutation rate.
- **Mutagen.** An environmental agent, either physical or chemical, that is capable of inducing mutations.
- **Mutant.** A cell or individual organism that shows a change brought about by a mutation; a changed gene.
- **Mutation.** A change in the DNA at a particular locus in an organism. The term is used loosely to include point mutations involving a single gene change as well as a chromosomal change.
- **Mutation pressure.** A constant mutation rate that adds mutant genes to a population; repeated occurrences of mutations in a population.
- **Mycelium** (*pl.*, **mycelia**). Threadlike filament making up the vegetative portion of thallus fungi.

Ν

Narrow-sense heritability. In quantitative genetics, the proportion of the phenotypic variance that is due to the additive effects of alleles.

- **Natural selection.** Differential survival and reproduction in nature that favors individuals that are better adapted to their environment; elimination of less fit organisms.
- **Negative control mechanism.** A mechanism in which the regulatory protein(s) is required to turn off gene expression.
- **Negative supercoiling.** The formation of coiled tertiary structures in double-stranded DNA molecules with fixed (not free to rotate) ends when the molecules are underwound.
- **Neutral mutation.** A mutation that changes the nucleotide sequence of a gene but has no effect on the fitness of the organism.
- **Neutral theory.** The theory that the evolution of traits with little or no effect on fitness is a random process involving mutation and genetic drift.
- **Nitrous acid.** HNO₂, a potent chemical mutagen.
- **Nonautonomous.** A term referring to biological units that cannot function by themselves; such units require the assistance of another unit, or "helper" (cf. **Autonomous**).
- **Nondisjunction.** Failure of disjunction or separation of homologous chromosomes in mitosis or meiosis, resulting in too many chromosomes in some daughter cells and too few in others. Examples: In meiosis, both members of a pair of chromosomes go to one pole so that the other pole does not receive either of them; in mitosis, both sister chromatids go to the same pole.
- **Nonhistone chromosomal proteins.** All of the proteins in chromosomes except the histones.
- **Nonionizing radiation.** The portion of the electromagnetic spectrum that does not lead to the production of positive and negative charges (ion pairs) in molecules. Visible and ultraviolet light are examples of nonionizing radiation (cf. **Ionizing radiation**).
- **Nonpolypoid Colorectal Cancer.** A form of cancer found in the lower digestive tract, sometimes inherited as a dominant condition.
- **Nonsense mutation.** A mutation that changes a codon specifying an amino acid to a termination codon.
- **Nonsynonymous substitution.** A base-pair change in a codon that alters the amino acid specified by the codon.
- **Nontemplate strand.** In transcription, the nontranscribed strand of DNA. It will have the same sequence as the RNA transcript, except that T is present at positions where U is present in the RNA transcript.
- **Northern blot.** The transfer of RNA molecules from an electrophoretic gel to a cellulose or nylon membrane by capillary action.
- Nuclease. An enzyme that catalyzes the degradation of nucleic acids.
- **Nucleic acid.** A macromolecule composed of phosphoric acid, pentose sugar, and organic bases; DNA and RNA.
- **Nucleolar Organizer (NO).** A chromosomal segment containing genes that control the synthesis of ribosomal RNA, located at the secondary constriction of some chromosomes.
- **Nucleolus.** An RNA-rich, spherical sack in the nucleus of metabolic cells; associated with the nucleolar organizer; storage place for ribosomes and ribosome precursors.
- **Nucleoprotein.** Conjugated protein composed of nucleic acid and protein; the material of which the chromosomes are made.
- **Nucleoside.** An organic compound consisting of a base covalently linked to ribose or deoxyribose.
- **Nucleosome, nucleosome core.** The nucleose-resistant subunit of chromatin that consists of about 146 nucleotides of DNA wrapped

as 1.65 turns of negative superhelix around an octamer of histones—two molecules each of histones H2a, H2b, H3, and H4.

- **Nucleotide.** A subunit of DNA and RNA molecules containing a phosphate group, a sugar, and a nitrogen-containing organic base.
- **Nucleotide excision repair.** The removal of relatively large defects such as thymine dimers in DNA via the excision of a segment of the DNA strand spanning the defect and repair synthesis by a DNA polymerase using the complementary strand as template.
- **Nucleus.** The part of a eukaryotic cell that contains the chromosomes; separated from the cytoplasm by a membrane.
- **Null allele.** A mutant form of a gene that either produces no product or produces a totally nonfunctional product.
- **Nullisomic.** An otherwise diploid cell or organism lacking both members of a chromosome pair (chromosome formula 2n 2).
- **Null mutation.** A mutation that abolishes the expression of a gene. (See also **Amorphic.**)

0

- **Octoploid.** Cell or organism with eight genomes or sets of chromosomes (chromosome number *8n*).
- **Oncogene.** A gene that can cause cancerous transformation in animal cells growing in culture and tumor formation in animals themselves; a gene that promotes cell division.
- **Oocyte.** The egg-mother cell; the cell that undergoes two meiotic divisions (oogenesis) to form the egg cell. Primary oocyte—before completion of the first meiotic division; secondary oocyte—after completion of the first meiotic division.
- **Oogenesis.** The formation of the egg or ovum in animals.
- **Oogonium** (*pl.*, **oogonia**). A germ cell of the female animal before meiosis begins.
- **Open reading frame (ORF).** A DNA segment containing the sequences required to encode a polypeptide. The RNA transcript of an ORF begins with a translation start codon, followed by a sequence of codons specifying amino acids, and ending with a translation stop codon. An ORF is presumed, but not known, to encode a polypeptide.
- **Operator.** A part of an operon that controls the expression of one or more structural genes by serving as the binding site for one or more regulatory proteins.
- **Operon.** A group of genes making up a regulatory or control unit. The unit includes an operator, a promoter, and structural genes.
- **Operon model.** The negative control mechanism proposed by Jacob and Monod in 1961 to explain the coordinate regulation of cotranscribed sets of structural genes. The mechanism involves a regulator gene encoding a repressor that controls transcription of the set of genes by binding to an operator region and blocking transcription by RNA polymerase.
- **Order (in the genetic code).** There are two types of order in the genetic code: (1) multiple codons for a given amino acid usually differ only at the third position, and (2) the codons for amino acids with similar chemical properties are closely related.
- **Ordinate.** The vertical axis in a graph.
- **Organelle.** Specialized part of a cell with a particular function or functions (for example, the cilium of a protozoan).
- **Organizer.** An inductor; a chemical substance in a living system that determines the fate in development of certain cells or groups of cells.

Origin of replication. The site or nucleotide sequence on a chromosome or DNA molecule at which replication is initiated.

Orthologous genes. Homologous genes present in different species (cf. **Homologous genes**).

Orthologues. See Orthologous genes.

Outbreeding. Mating of unrelated individuals.

- **Ovary.** The swollen part of the pistil of a plant flower that contains the ovules; the female reproductive organ or gonad in animals.
- **Overdominance.** A condition in which heterozygotes are superior (on some scale of measurement) to either of the associated homo-zygotes.
- **Ovule.** The macrosporangium of a flowering plant that becomes the seed. It includes the nucellus and the integuments.

Ρ

- P. Symbol for the parental generation or parents of a given individual.
- **Pachynema** (*adj.*, **pachytene**). A mid-prophase stage in meiosis immediately following zygonema and preceding diplonema. In favorable microscopic preparations, the chromosomes are visible as long, paired threads. Rarely, four chromatids are detectable.
- **Pair-rule gene.** A gene that influences the formation of body segments in *Drosophila*.
- **Palindrome.** A segment of DNA in which the base-pair sequence reads the same in both directions from a central point of symmetry.
- **Panmictic population.** A population in which mating occurs at random.
- Panmixis. Random mating in a population.

Paracentric inversion. An inversion that is entirely within one arm of a chromosome and does not include the centromere.

- Paralogues. See Paralogous genes.
- **Paralogous genes.** Homologous genes present within a species (cf. **Homologous genes**).
- **Parameter.** A value or constant based on an entire population (cf. **Statistic**).
- **Parental.** Pertaining to the founding strains used in a cross; having the characteristics of these strains. In a series of crosses, the parental generation is symbolized as **P**.
- **Parthenogenesis.** The development of a new individual from an egg without fertilization.
- **Paternal.** Pertaining to the father.
- Pathogen. An organism that causes a disease.
- **Pattern baldness.** A hereditary form of baldness in which the thinning of the hair begins on the crown of the head.

PCR. See Polymerase chain reaction.

- **Pedigree.** A table, chart, or diagram representing the ancestry of an individual.
- **Pelement.** A transposable element in *Drosophila* that, when activated, causes hybrid dysgenesis.
- **Penetrance.** The percentage of individuals that show a particular phenotype among those capable of showing it.
- **Peptide.** A compound containing amino acids; a breakdown or buildup unit in protein metabolism.
- **Peptide bond.** A chemical bond holding amino acid subunits together in proteins.

- **Peptidyl** (*P*) **site.** The ribosome binding site that contains the tRNA to which the growing polypeptide chain is attached.
- **Peptidyl transferase.** An enzyme activity—built into the large subunit of the ribosome—that catalyzes the formation of peptide bonds between amino acids during translation.
- **Pericentric inversion.** An inversion including the centromere, hence involving both arms of a chromosome.
- **Peroxisome.** A subcellular organelle that contains enzymes involved in the degradation of fatty acids and amino acids.

Phage. See Bacteriophage.

- **Phagemids.** Cloning vectors that contain components derived from both phage chromosomes and plasmids.
- **Phenocopy.** An organism whose phenotype (but not genotype) has been changed by the environment to resemble the phenotype of a different (mutant) organism.
- **Phenotype.** The observable characteristics of an organism.

Phenylalanine. See Amino acid.

- **Phenylketonuria.** Metabolic disorder resulting in mental retardation; transmitted as a Mendelian recessive and treated in early childhood by special diet.
- Photoreactivation. A DNA repair process that is light-dependent.
- **Phylogeny.** A diagram showing the evolutionary relationships among a group of organisms; an evolutionary tree.
- **Physical map.** A diagram of a chromosome or DNA molecule with distances given in base pairs, kilobases, or megabases.
- Pistil. The centrally located organ in flowers that contains the ovary.
- **Plasma cells.** Antibody-producing white blood cells devived from B lymphocytes.
- **Plasmid.** An extrachromosomal hereditary determinant that exists in an autonomous state and is transferred independently of chromosomes.
- **Plastid.** A cytoplasmic body found in the cells of plants and some protozoa. Chloroplasts, for example, produce chlorophyll that is involved in photosynthesis.
- **Pleiotropy** (*adj.*, **Pleiotropic**). Condition in which a single gene influences more than one trait.
- **Pluripotent.** An adjective applied to cells that have the potential to differentiate into many different types.
- **Point mutations.** Changes that occur at specific sites in genes. They include nucleotide-pair substitutions and the insertion or deletion of one or a few nucleotide pairs.
- **Polar bodies.** In female animals, the smaller cells produced at meiosis that do not develop into egg cells. The first polar body is produced at division I and may not go through division II. The second polar body is produced at division II.
- **Pole cells.** A group of cells in the posterior of *Drosophila* embryos that are precursors to the adult germ line.
- **Pollen grain.** The male gametophyte in higher plants.
- **Polyadenylation.** The addition of poly(A) tails to eukaryotic gene transcripts (RNAs).
- **Poly(A) polymerase.** An enzyme that adds the poly(A) tails to the 3' termini of eukaryotic gene transcripts (RNAs).
- **Poly(A) tail (mRNA).** A polyadenosine tract 20 to 200 nucleotides long that is added to the 3' ends of most eukaryotic mRNAs postranscriptionally.

- **Polydactyly.** The occurrence of more than the usual number of fingers or toes.
- **Polygene** (*adj.*, **polygenic**). One of many genes involved in quantitative inheritance.
- **Polylinker (multiple cloning site).** A segment of DNA that contains a set of unique restriction enzyme cleavage sites.
- **Polymer.** A compound composed of many smaller subunits; results from the process of polymerization.
- **Polymerase.** An enzyme that catalyzes the formation of DNA or RNA.
- **Polymerase chain reaction (PCR).** A procedure involving multiple cycles of denaturation, hybridization to oligonucleotide primers, and polynucleotide synthesis that amplifies a particular DNA sequence.
- **Polymerization.** Chemical union of two or more molecules of the same kind to form a new compound having the same elements in the same proportions but a higher molecular weight and different physical properties.
- **Polymorphism.** The existence of two or more variants in a population of individuals, with at least two of the variants having frequencies greater than 1 percent.
- **Polynucleotide.** A linear sequence of joined nucleotides in DNA or RNA.
- **Polypeptide.** A linear molecule with two or more amino acids and one or more peptide groups. They are called dipeptides, tripeptides, and so on, according to the number of amino acids present.
- **Polyploid.** An organism with more than two sets of chromosomes (2*n* diploid) or genomes—for example, triploid (3*n*), tetraploid (4*n*), pentaploid (5*n*), hexaploid (6*n*), heptaploid (7*n*), octoploid (8*n*).
- **Polysaccharide capsules.** Carbohydrate coverings with antigenic specificity that are present on some types of bacteria.
- **Polytene chromosomes.** Giant chromosomes produced by interphase replication without division and consisting of many identical chromatids arranged side by side in a cablelike pattern.
- **Population.** Entire group of organisms of one kind; an interbreeding group of plants or animals; the extensive group from which a sample might be taken.
- Population (effective). Breeding members of the population.
- **Population genetics.** The branch of genetics that deals with frequencies of alleles and genotypes in breeding populations.
- **Positional cloning.** The isolation of a clone of a gene or other DNA sequence based on its map position in the genome.
- **Position effect variegation.** Phenotypic variation within an individual that is due to a change in the genomic position of a gene. Usually this type of variation is seen when a gene naturally located in euchromatin is moved by a chromosome rearrangement to a heterochromatic region of the genome.
- **Positive control mechanism.** A mechanism in which the regulatory protein(s) is required to turn on gene expression.
- **Postreplication repair.** A recombination-dependent mechanism for repairing damaged DNA.
- **Pre-mRNA.** The primary transcript of a eukaryotic gene prior to processing to produce an mRNA.
- **Primary transcript.** The RNA molecule produced by transcription prior to any posttranscriptional modifications; also called a pre-mRNA in eukaryotes.

- **Primer.** A short nucleotide sequence with a reactive 3' OH that can initiate DNA synthesis along a template.
- **Primosome.** A protein replication complex that catalyzes the initiation of Okazaki fragments during discontinuous DNA synthesis. It contains DNA primase and DNA helicase activities.
- Probability. The frequency of occurrence of an event.
- **Proband.** The individual in a family in whom an inherited trait is first identified.
- Progerias. Inherited diseases characterized by premature aging.
- **Prokaryote.** A member of a large group of organisms (including bacteria and bluegreen algae) that lack true nuclei in their cells and that do not undergo mitosis.
- **Prokaryotic cells.** The cells of organisms classified as prokaryotes. These cells are characterized by not having a membrane-bound nucleus that contains the chromosomal DNA.
- **Promoter.** A nucleotide sequence to which RNA polymerase binds and initiates transcription; also, a chemical substance that enhances the transformation of benign cells into cancerous cells.
- **Proofreading.** The enzymatic scanning of DNA for structural defects such as mismatched base pairs.
- **Prophage (provirus).** The genome of a temperate bacteriophage integrated into the chromosome of a lysogenic bacterium and replicated along with the host chromosome.
- **Prophase.** The stage of mitosis between interphase and metaphase. During this phase, the centriole divides and the two daughter centrioles move apart. Each sister DNA strand from interphase replication becomes coiled, and the chromosome is longitudinally double except in the region of the centromere. Each partially separated chromosome is called a chromatid. The two chromatids of a chromosome are sister chromatids.
- **Prophase I.** The stage during the first meiotic division when duplicated chromosomes condense and pair with their homologues.
- **Prophase II.** The stage during the second meiotic division when duplicated chromosomes condense and prepare to move to the equatorial plane of the cell.
- **Protamines.** Small basic proteins that replace the histones in the chromosomes of some sperm cells.
- **Protease.** Any enzyme that hydrolyzes proteins.
- **Protein.** A macromolecule composed of one to several polypeptides. Each polypeptide consists of a chain of amino acids linked together by peptide bonds.
- **Proteome.** The complete set of proteins encoded by a genome.
- **Proteomics.** The science focused on determining the structures and functions of all the proteins produced by living organisms.
- **Proto-oncogene.** A normal cellular gene that can be changed to an oncogene by mutation.
- **Protoplast.** A plant or bacterial cell from which the wall has been removed.
- **Prototroph.** An organism such as a bacterium that will grow on a minimal medium.
- **Provirus.** A viral chromosome that has integrated into a host—either prokaryotic or eukaryotic—genome (cf. **Prophage**).
- **Pseudoautosomal gene.** A gene located on both the X and Y chromosomes.
- **Pseudogene.** An inactive but stable component of a genome resembling a gene; apparently derived from active genes by mutation.

- **Purine.** A double-ring nitrogen-containing base present in nucleic acids; adenine and guanine are the two purines present in most DNA and RNA molecules.
- **Pyrimidine.** A single-ring nitrogen-containing base present in nucleic acids; cytosine and thymine are commonly present in DNA, whereas uracil usually replaces thymine in RNA.

Q

Quantitative inheritance. Inheritance of measurable traits (height, weight, color intensity) that depend on the cumulative action of many genes, each producing a small effect on the phenotype.

- **Quantitative trait loci (QTL).** Two or more genes that affect a single quantitative trait.
- **Quantitative traits.** Phenotypes that can be measured, such as height, weight, and growth rate.

R

- Race. A distinguishable group of organisms of a particular species.
- **Radiation hybrid mapping.** The use of human–rodent hybrid cells containing fragments of human chromosomes (produced by irradiation) fused to rodent chromosomes to determine the linkage relationships of human genes.
- **Radioactive isotope.** An unstable isotope (form of an atom) that emits ionizing radiation.
- **Random genetic drift.** Changes in allele frequency in small breeding populations due to chance fluctuations.
- **Reading frame.** The series of nucleotide triplets that are sequentially positioned in the *A* site of the ribosome during translation of an mRNA; also, the sequence of nucleotide-pair triplets in DNA that correspond to these codons in mRNA.
- **Receptor.** A molecule that can accept the binding of a ligand.
- **Recessive.** A term applied to one member of an allelic pair lacking the ability to manifest itself when the other or dominant member is present.
- **Recessive lethal mutation.** A mutant form of a gene that results in the death of an organism that is homozygous for it.
- **Recipient cell.** A bacterium that receives DNA from another (donor) cell during recombination in bacteria (cf. **Donor cell**).
- **Reciprocal crosses.** Crosses between different strains with the sexes reversed; for example, female $A \times$ male B and male $A \times$ female B are reciprocal crosses.
- **Recognition sequence (-35 sequence).** A nucleotide sequence (consensus TTGACA) in prokaryotic promoters to which the sigma factor of RNA polymerase binds during the initiation of transcription.
- **Recombinant DNA molecule.** A DNA molecule constructed *in vitro* by joining all or parts of two different DNA molecules.
- **Recombination.** The production of gene combinations not found in the parents by the assortment of nonhomologous chromosomes and crossing over between homologous chromosomes during meiosis. For linked genes, the frequency of recombination can be used to estimate the genetic map distance; however, high frequencies (approaching 50 percent) do not yield accurate estimates.
- **Reduction division.** Phase of meiosis in which the maternal and paternal chromosomes of the bivalent separate (cf. **Equational division**).

- **Regulator gene.** A gene that controls the rate of expression of another gene or genes. Example: The *lacI* gene produces a protein that controls the expression of the structural genes of the *lac* operon in *Escherichia coli*.
- **Relative fitness.** The survival and reproductive ability of a genotype in a population in comparison to the survival and reproductive ability of another genotype in that population.
- **Release factors (RF).** Soluble proteins that recognize termination codons in mRNAs and terminate translation in response to these codons.
- **Renaturation.** The restoration of a molecule to its native form. In nucleic acid biochemistry, this term usually refers to the formation of a double-stranded helix from complementary single-stranded molecules.
- **Repetitive DNA.** DNA sequences that are present in a genome in multiple copies—sometimes a million times or more.
- **Replica plating.** A procedure for duplicating the bacterial colonies growing on agar medium in one petri plate to agar medium in another petri plate.
- **Replication.** A duplication process that is accomplished by copying from a template (for example, reproduction at the level of DNA).
- **Replication bubble.** The localized region of complementary strand separation that occurs at the origin of replication during the initiation of DNA replication.
- **Replication fork.** The Y-shaped structure where the two parental strands of a DNA double helix are unwound and are being used as templates for the synthesis of new complementary strands.
- **Replicative transposon.** A transposable element that is replicated during the transposition process. Tn3 in *E. coli* is an example.
- **Replicon.** A unit of replication. In bacteria, replicons are associated with segments of the cell membrane that control replication and coordinate it with cell division.
- **Replisome.** The complete replication apparatus—present at a replication fork—that carries out the semiconservative replication of DNA.
- **Repressible enzyme.** An enzyme whose synthesis is diminished by a regulatory molecule.
- **Repression.** The process of turning off the expression of a gene or set of genes in response to some signal.
- **Repressor.** A protein that binds to DNA and turns off gene expression.
- **Repressor gene.** A gene that encodes a repressor.
- **Reproductive cloning.** A process in which the nucleus of an egg cell is replaced with the nucleus of a cell from a developed organism with the purpose of producing a new organism genetically identical to the donor.
- **Repulsion** (*trans* configuration). The condition in which a double heterozygote has received a mutant and a wild-type allele from each parent; for example, $a + / a + \times + b/+ b$ produces a + / + b (cf. Coupling).
- **Resistance factor.** A plasmid that confers antibiotic resistance to a bacterium.
- Restriction endonuclease. See Restriction enzyme.
- **Restriction enzyme.** An endonuclease that recognizes a specific short sequence in DNA and cleaves the DNA molecule at or near that site.
- **Restriction fragment.** A fragment of DNA produced by cleaving a DNA molecule with one or more restriction endonucleases.

- **Restriction fragment-length polymorphism (RFLP).** The existence of two or more genetic variants detectable by visualizing fragments of genomic DNA that were obtained by digesting the DNA with a restriction enzyme. Usually the DNA fragments are fractionated by electrophoresis, transferred to a membrane by Southern blotting, and then visualized by autoradiography after hybridization to a labeled DNA probe.
- **Restriction map.** A linear or circular physical map of a DNA molecule showing the sites that are cleaved by different restriction enzymes.
- **Restriction site.** A DNA sequence that is cleaved by a restriction enzyme.
- **Reticulocyte.** A young red blood cell.
- **Retroelement.** Any of the integrated retroviruses or the transposable elements that resemble them.
- **Retroposon.** A transposable element that creates new copies via reverse transcription of RNA into DNA but that lacks the long terminal repeat sequences.
- **Retrotransposon.** A transposable element that creates new copies by reverse transcription of RNA into DNA.
- **Retrovirus.** A virus that stores its genetic information in RNA and replicates by using reverse transcriptase to synthesize a DNA copy of its RNA genome.
- **Retroviruslike element.** A type of retrotransposon that resembles the integrated form of a retrovirus.
- **Reverse genetics.** Genetic approaches that use the nucleotide sequence of a gene to devise procedures for isolating mutations in the gene or shutting off its expression.
- **Reverse transcriptase.** An enzyme that catalyzes the synthesis of DNA using an RNA template.
- **Reversion (reverse mutation).** Restitution of a mutant gene to the wild-type condition, or at least to a form that gives the wild phenotype; more generally, the appearance of a trait expressed by a remote ancestor.
- RFLP. See Restriction fragment-length polymorphism.
- **Ribonuclease (RNase).** Any enzyme that hydrolyzes RNA.
- Ribonucleic acid. See RNA.
- **Ribosomal RNAs (rRNAs).** The RNA molecules that are structural components of ribosomes.
- **Ribosome.** Cytoplasmic organelle on which proteins are synthesized.
- **Riboswitch.** A mRNA molecule that can regulate gene expression transcription or translation—by undergoing a change in conformation upon binding a specific metabolite.
- **RNA-induced silencing complex (RISC).** A protein complex that uses double-stranded RNA to produce and target small interfering RNAs to complementary messenger RNAs within eukaryotic cells.
- **R-loops.** Single-stranded DNA regions in RNA-DNA hybrids formed *in vitro* under conditions where RNA-DNA duplexes are more stable than DNA-DNA duplexes.
- **RNA.** Ribonucleic acid; the information-carrying material in some viruses; more generally, a molecule derived from DNA by transcription that may carry information (messenger or mRNA), provide subcellular structure (ribosomal or rRNA), transport amino acids (transfer or tRNA), or facilitate the biochemical modification of itself or other RNA molecules.
- **RNA editing.** Posttranscriptional processes that alter the information encoded in gene transcripts (RNAs).

- **RNA interference (RNAi).** A phenomenon in which doublestranded RNA prevents the expression of a gene homologous to at least part of the RNA.
- **RNA polymerase.** An enzyme that catalyzes the synthesis of RNA.
- **RNA primer.** A short (10 to 60 nucleotides) segment of RNA that is used to initiate the synthesis of a new strand of DNA; synthesized by the enzyme DNA primase.
- **Robertsonian translocation.** A rearrangement in which the long arms of two nonhomologous chromosomes have been joined at or near their centromeres and the short arms of these chromosomes have been lost.
- Roentgen (r). Unit of ionizing radiation.
- **Rolling-circle replication.** A mechanism of replication of circular DNA molecules in which one parental strand of DNA is cleaved at the origin of replication while the other strand remains intact. The 5' terminus of the cleaved strand is unwound and replicated discontinuously while continuous replication of the other strand occurs at the 3' terminus with the intact circular strand as template.

S

Sample. A group of items selected to represent a large population.

- **Satellite band.** A band formed by DNA in a density gradient that is smaller than, and distinct from, main-band DNA. A satellite band contains repeated DNA sequences called satellite DNAs with lower or higher densities than main-band DNA.
- **Satellite DNA.** A component of the genome that can be isolated from the rest of the DNA by density-gradient centrifugation. Usually, it consists of short, highly repetitious sequences.
- **Scaffold.** The central core structure of condensed chromosomes. The scaffold is composed of nonhistone chromosomal proteins.
- **SCID** (severe combined immunodeficiency syndrome). A group of diseases characterized by the inability to mount an immune response, either humoral or cellular.
- Secondary oocyte. See Oocyte.
- Secondary spermatocyte. See Spermatocyte.
- **Segmentation genes.** A group of genes that control the early development of *Drosophila* embryos. Their products define segments along the anterior-posterior axis.
- **Segment-polarity genes.** A group of genes whose products define the anterior and posterior compartments in each of the segments that form along the anterior-posterior axis of *Drosophila* embryos.
- **Segregation** (*v.*, **segregate**). The separation of paternal and maternal chromosomes from each other at meiosis; the separation of alleles from each other in heterozygotes; the occurrence of different phenotypes among offspring, resulting from chromosome or allele separation in their heterozygous parents; Mendel's first principle of inheritance.
- **Selection.** Differential survival and reproduction among genotypes; the most important of the factors that change allele frequencies in large populations.
- **Selection coefficient.** A number that measures the fitness of a genotype relative to a standard.
- **Selection differential.** In plant and animal breeding, the difference between the mean of the individuals selected to be parents and the mean of the overall population.
- **Selection pressure.** Effectiveness of differential survival and reproduction in changing the frequency of alleles in a population.

- **Selection response.** In plant and animal breeding, the difference between the mean of the individuals selected to be parents and the mean of their offspring.
- **Selector gene.** A gene that influences the development of specific body segments in *Drosophila*; a homeotic gene.
- **Selenocysteine.** An amino acid that contains selenium (atomic number 34) in place of the sulfur group in cysteine.
- **Selenoprotein.** A protein that contains the amino acid selenocysteine.
- **Self-fertilization.** The process by which pollen of a given plant fertilizes the ovules of the same plant. Plants fertilized in this way are said to have been selfed. An analogous process occurs in some animals, such as nematodes and molluscs.
- **Semiconservative replication.** Replication of DNA by a mechanism in which the parental strands are conserved (remain intact) and serve as templates for the synthesis of new complementary strands.
- **Semidominant.** A term applied to alleles in which the phenotype of a heterozygote is midway between the phenotypes of the corresponding homozygotes.
- **Semisterility.** A condition of only partial fertility in plant zygotes (for example, maize); usually associated with translocations.
- **Sense RNA.** A primary transcript or mRNA that contains a coding region (contiguous sequence of codons) that is translated to produce a polypeptide.
- Sense strand (of RNA). See Sense RNA.
- -20 Sequence. See TATAAT sequence.
- -35 Sequence. See Recognition sequence.
- **Sex chromosomes.** Chromosomes that are connected with the determination of sex.
- **Sexduction.** The incorporation of bacterial genes into F factors and their subsequent transfer by conjugation to a recipient cell.
- **Sex factor.** A bacterial episome (for example, the F plasmid in *E. coli*) that enables the cell to be a donor of genetic material. The sex factor may be propagated in the cytoplasm, or it may be integrated into the bacterial chromosome.
- **Sex-influenced dominance.** A dominant expression that depends on the sex of the individual. For example, horns in some breeds of sheep are dominant in males and recessive in females.
- **Sex-limited.** Expression of a trait in only one sex. Examples: milk production in mammals; horns in Rambouillet sheep; egg production in chickens.
- **Sex linkage.** Association or linkage of a hereditary trait with sex; the gene is in a sex chromosome, usually the X; often used synonymously with X-linkage.
- Sex mosaic. See Gynandromorph.
- **Sexual reproduction.** Reproduction involving the formation of mature germ cells (that is, eggs and sperm).
- **Shelterin.** A protein complex that binds to telomeres and protects the DNA in them from degradation.
- **Shine-Dalgarno sequence.** A conserved sequence in prokaryotic mRNAs that is complementary to a sequence near the 5' terminus of the 16S ribosomal RNA and is involved in the initiation of translation.
- **Short interfering RNA (siRNA).** Double-stranded RNA molecules 21–28 base pairs long that mediate the phenomenon of RNA interference; also known as microRNA molecules.

- **Short tandem repeat (STR) (microsatellite).** A highly polymorphic tandem repeat of a sequence only two to five nucleotide pairs in length.
- Shuttle vector. A plasmid capable of replicating in two different organisms, such as yeast and *E. coli*.
- **Sib-mating (crossing of siblings).** Matings involving two individuals of the same parentage; brother–sister matings.
- **Sigma factor.** The subunit of prokaryotic RNA polymerases that is responsible for the initiation of transcription at specific initiation sequences.
- **Signal transduction.** The process whereby a molecular signal such as a hormone is passed internally within a cell by a system of molecules to effect a change in the cell's state.
- **Silencer.** A DNA sequence that helps to reduce or shut off the expression of a nearby gene.
- **Silent polymorphism.** A variant in DNA that does not alter the amino acid sequence of a protein.
- **Simple tandem repeat.** A tandemly repeated unit in DNA of only one to six nucleotides in length.
- **SINEs (short interspersed nuclear elements).** Families of short (150 to 300 bp), moderately repetitive transposable elements of eukaryotes. The best known SINE family is the Alu family in humans.
- **Single-nucleotide polymorphism (SNP).** A single base pair in the DNA that varies in a population.
- **Single-strand DNA-binding protein.** A protein that coats DNA single strands, keeping them in an extended state.
- Sister chromatid. One of the products of chromosome duplication.
- **Small nuclear ribonucleoproteins (snRNPs).** RNA-protein complexes that are components of spliceosomes.
- **Small nuclear RNAs (snRNAs).** Small RNA molecules that are located in the nuclei of eukaryotic cells; most snRNAs are components of the spliceosomes that excise introns from pre-mRNAs.
- **Somatic cell.** A cell that is a component of the body, in contrast with a germ cell that is capable, when fertilized, of reproducing the organism.
- **Somatic-cell (nonheritable) gene therapy.** Treatment of an inherited disorder by adding functional (wild-type) copies of a gene to nongerm-line cells of an individual carrying defective copies of that gene (cf. **Germ-line [heritable] gene therapy**).
- **Somatic hypermutation.** A high frequency of mutation that occurs in the gene segments encoding the variable regions of antibodies during the differentiation of B lymphocytes into antibodyproducing plasma cells.
- **Somatic mutation.** A mutation that occurs in the nonreproductive cells (somatic cells) of the body and is not transmitted to progeny (cf. **Germinal mutation**).
- **SOS response.** The synthesis of a whole set of DNA repair, recombination, and replication proteins in bacteria containing severely damaged DNA (for example, following exposure to UV light).
- **Southern blot.** The transfer of DNA fragments from an electrophoretic gel to a cellulose or nylon membrane by capillary action.
- **Specialized transduction.** Recombination in bacteria mediated by a bacteriophage that can only transfer genes in a small segment of the chromosome of the donor cell to a recipient cell (cf. **General-ized transduction**).
- **Species.** Interbreeding, natural populations that are reproductively isolated from other such groups.

- **Sperm (abbreviation of spermatozoon,** *pl.*, **spermatozoa).** A mature male germ cell.
- **Spermatids.** The four cells formed by the meiotic divisions in spermatogenesis. Spermatids become mature spermatozoa or sperm.
- **Spermatocyte (sperm mother cell).** The cell that undergoes two meiotic divisions (spermatogenesis) to form four spermatids; the *primary* spermatocyte before completion of the first meiotic division; the *secondary* spermatocyte after completion of the first meiotic division.
- **Spermatogenesis.** The process by which maturation of the gametes (sperm) of the male takes place.
- **Spermatogonium** (*pl.*, **spermatogonia**). Primordial male germ cell that may divide by mitosis to produce more spermatogonia. A spermatogonium may enter a growth phase and give rise to a primary spermatocyte.
- **Spermiogenesis.** Formation of sperm from spermatids; the part of spermatogenesis that follows the meiotic divisions of spermatocytes.
- **Spindle.** A system of microtubules that distributes duplicated chromosomes equally and exactly to each of the daughters of a dividing eukaryotic cell.
- **Spliceosome.** The RNA/protein complex that excises introns from primary transcripts of nuclear genes in eukaryotes.
- **Splicing.** The process that covalently joins exon sequences of RNA and eliminates the intervening intron sequences.
- **Spontaneous mutation.** A mutation that occurs without a known cause (cf. **Induced mutation**).
- **Sporophyte.** The diploid generation in the life cycle of a plant that produces haploid spores by meiosis.
- *SRY* (*Sex-determining region Y*). A Y-linked gene in humans and other mammals encoding a protein, the testis-determining factor, which plays a key role in male development.
- **Stamen.** The elongated structure that bears the anthers in flowering plants.
- **Standard deviation.** A measure of variability in a set of data; the square root of the variance.
- **Standard error.** A measure of variation among a population of means.
- **Statistic.** A value based on a sample or samples of a population from which estimates of a population value or parameter may be obtained.
- **Stem cell.** A cell with the ability to proliferate extensively and whose offspring can differentiate into specialized cell types.
- Sterility. Inability to produce offspring.
- Structural gene. A gene that specifies the synthesis of a polypeptide.
- **STSs (sequence-tagged sites).** Short, unique DNA sequences (usually 200 to 500 bp) that are amplified by PCR and used to link physical maps and genetic maps.
- **Subspecies.** One of two or more morphologically or geographically distinct but interbreeding populations of a species.
- **Supercoil.** A DNA molecule that contains extra twists as a result of overwinding (positive supercoils) or underwinding (negative supercoils).
- **Suppressor mutation.** A mutation that partially or completely cancels the phenotypic effect of another mutation.
- **Suppressor-sensitive mutant.** An organism that can grow when a second genetic factor—a suppressor—is present, but not in the absence of this factor.

- **Suppressor tRNA.** A mutant tRNA that recognizes one or more of the termination codons and inserts an amino acid at a site where translation termination would normally occur.
- **Symbiont.** An organism living in intimate association with another, dissimilar organism.
- **Sympatric speciation.** The formation of new species by populations that inhabit the same or overlapping geographic regions.
- **Synapsis.** The pairing of homologous chromosomes in the meiotic prophase.
- **Synaptinemal complex.** A ribbonlike structure formed between synapsed homologues at the end of the first meiotic prophase, binding the chromatids along their length and facilitating chromatid exchange.
- **Syndrome.** A group of symptoms that occur together and represent a particular disease.
- **Synonymous substitution.** A base-pair change in a codon that does not alter the amino acid specified by the codon.
- **Synteny.** The occurrence of two loci on the same chromosome, without regard to the distance between them.

Т

- *Taq* polymerase. A heat-stable DNA polymerase isolated from the thermophilic bacterium *Thermus aquaticus*.
- **Target site duplication.** A sequence of DNA that is duplicated when a transposable element inserts; usually found at each end of the insertion.
- **TATAAT sequence (-10 sequence).** An AT-rich sequence in prokaryotic promoters that facilitates the localized unwinding of DNA and the initiation of RNA synthesis.
- **TATA box.** A conserved promoter sequence that determines the transcription start site.
- **Tautomeric shift.** The transfer of a hydrogen atom from one position in an organic molecule to another position.
- **Tay-Sachs disease.** A lethal autosomal recessive disorder in humans characterized by neurological degeneration and death in early childhood. The disease is caused by the absence of an enzyme called hexosaminidase A.
- **T-cell receptor.** An antigen-binding protein that is located on the surfaces of killer T cells and mediates the cellular immune response of mammals. The genes that encode T-cell antigens are assembled from gene segments by somatic recombination processes that occur during T lymphocyte differentiation.
- **T-DNA.** The segment of DNA in the Ti plasmid of *Agrobacterium tumefaciens* that is transferred to plant cells and inserted into the chromosomes of the plant.
- **Telomerase.** An enzyme that adds telomere sequences to the ends of eukaryotic chromosomes.
- **Telomere.** The unique structure found at the end of eukaryotic chromosomes.
- **Telophase.** The last stage in each mitotic or meiotic division in which the chromosomes are assembled at the poles of the division spindle.
- **Telophase I.** The stage during the first meiotic division when duplicated chromosomes gather at the pole of a dividing cell and begin to decondense.
- **Telophase II.** The stage during the second meiotic division when the chromosomes gather at the pole of a dividing cell and begin to decondense.

- **Temperate phage.** A phage (virus) that invades but may not destroy (lyse) the host (bacterial cell) (cf. **Virulent phage**). However, it may subsequently enter the lytic cycle.
- **Temperature-sensitive mutant.** An organism that can grow at one temperature but not at another.
- **Template.** A pattern or mold. DNA stores coded information and acts as a model or template from which information is copied into complementary strands of DNA or transcribed into messenger RNA.
- **Template strand.** In transcription, the DNA strand that is copied to produce a complementary strand of RNA.
- **Terminal inverted repeat.** Identical or nearly identical DNA sequences at opposite ends of a cut-and-paste transposon. One sequence is the inverted mirror image of the other.
- **Terminalization.** Repelling movement of the centromeres of bivalents in the diplotene stages of the meiotic prophase that tends to move the visible chiasmata toward the ends of the bivalents.
- **Terminal transferase.** An enzyme that adds nucleotides to the 3' termini of DNA molecules.
- **Termination (of DNA, RNA, or protein synthesis).** The release of a complete macromolecule (DNA, RNA, or polypeptide) after the incorporation of the final subunit (nucleotide or amino acid).
- **Termination signal.** In transcription, a nucleotide sequence that specifies RNA chain termination.
- **Testcross.** Backcross to the recessive parental type, or a cross between genetically unknown individuals with a fully recessive tester to determine whether an individual in question is heterozygous or homozygous for a certain allele. It is also used as a test for linkage.
- **Testis-determining factor (TDF).** A protein produced early in the development of male mammals that stimulates the differentiation of the testes from the embryonic gonads.
- **Testosterone.** A steroid hormone that induces the development of male characteristics.
- **Tetrad.** The four cells arising from the second meiotic division in plants (pollen tetrads) or fungi (ascospores). The term is also used to identify the quadruple group of chromatids that is formed by the association of duplicated homologous chromosomes during meiosis.
- **Tetraploid.** An organism whose cells contain four haploid (4*n*) sets of chromosomes or genomes.
- **Tetrasomic** (*noun*, **tetrasome**). Pertaining to a nucleus or an organism with four members of one of its chromosomes, whereas the remainder of its chromosome complement is diploid. (Chromosome formula: 2n + 2).
- **TFIIX (Transcription Factor X for RNA polymerase II).** A protein required for the initiation of transcription by RNA polymerase II in eukaryotes; X represents any one of several different factors designated A through F.
- **Therapeutic cloning.** A process in which the nucleus of egg cell is replaced with the nucleus of a donor cell (possibly differentiated) to produce a population of stem cells that have the same genotype as the donor cell. These stem cells could then be used to replace lost cells in the donor organism.
- **Threshold trait.** A trait that is manifested discontinuously but that is a function of underlying continuous genetic and environmental variation.
- **Thymine (T).** A pyrimidine base found in DNA. The other three organic bases—adenine, cytosine, and guanine—are found in both RNA and DNA, but in RNA, thymine is replaced by uracil.

- **Ti plasmid.** The large plasmid in *Agrobacterium tumefaciens*. It is responsible for the induction of tumors in plants with crown gall disease and is an important vector for transferring genes into plants, especially dicots.
- **t-loop.** A loop of DNA formed by telomere repeat sequences at the end of a linear chromosome when a single strand at the 3' terminus invades an upstream repeat unit and pairs with the complementary strand, while displacing the equivalent strand.
- **T lymphocytes (T cells).** Cells that differentiate in the thymus gland and are primarily responsible for the T-cell-mediated or cellular immune response.
- **Topoisomerase.** An enzyme that introduces or removes supercoils from DNA.
- **Totipotent cell (or nucleus).** An undifferentiated cell (or nucleus) such as a blastomere that when isolated or suitably transplanted can develop into a complete embryo.
- **Trafficking.** The movement of materials through the cytoplasm of a cell, usually guided by membranes, vesicles, and components of the cytoskeleton.
- *trans*-acting. A term describing substances that are diffusable and that can affect spatially separated entities within cells.
- trans configuration. See Repulsion.
- Transcript. The RNA molecule produced by transcription of a gene.
- **Transcription.** Process through which RNA is formed along a DNA template. The enzyme RNA polymerase catalyzes the formation of RNA from ribonucleoside triphosphates.
- **Transcriptional antiterminator.** A protein that prevents RNA polymerase from terminating transcription at specific transcription-termination sequences.
- **Transcription bubble.** A locally unwound segment of DNA in which an RNA transcript is being synthesized.
- **Transcription factor.** A protein that regulates the transcription of genes.
- **Transcription unit.** A segment of DNA that contains transcription initiation and termination signals and is transcribed into one RNA molecule.
- **Transcriptome.** The complete set of RNAs transcribed from a genome.
- **Transduction (t).** Genetic recombination in bacteria mediated by bacteriophage. Abortive t: Bacterial DNA is injected by a phage into a bacterium, but it does not replicate. Generalized t: Any bacterial gene may be transferred by a phage to a recipient bacterium. Restricted t: Transfer of bacterial DNA by a temperate phage is restricted to only one site on the bacterial chromosome.
- **Transfection.** The uptake of DNA by a eukaryotic cell, followed by the incorporation of genetic markers present in the DNA into the cell's genome.
- **Transfer RNAs (tRNAs).** RNAs that transport amino acids to the ribosomes, where the amino acids are assembled into proteins.
- **Transformation (cancerous).** The conversion of eukaryotic cells growing in culture to a state of uncontrolled cell growth (similar to tumor cell growth).
- **Transformation (genetic).** Genetic alteration of an organism brought about by the incorporation of foreign DNA into cells.
- **Transgene.** A foreign or modified gene that has been introduced into an organism.

- **Transgenic.** A term applied to organisms that have been altered by introducing DNA molecules into them.
- **Transgressive variation.** The appearance in the F_2 (or later) generation of individuals showing more extreme development of a trait than either of the original parents.
- *trans* heterozygote. A heterozygote that contains two mutations arranged in the *trans* configuration—for example, $a b^+ / a^+ b$.
- **Transition.** A mutation caused by the substitution of one purine by another purine or one pyrimidine by another pyrimidine in DNA or RNA.
- **Translation.** Protein (polypeptide) synthesis directed by a specific messenger RNA; occurs on ribosomes.
- **Translocation.** Change in position of a segment of a chromosome to another part of the same chromosome or to a different chromosome.
- **Transposable genetic element.** A DNA element that can move from one location in the genome to another.
- **Transposase.** An enzyme that catalyzes the movement of a DNA sequence to a different site in a DNA molecule.
- **Transposons.** DNA elements that can move ("transpose") from one position in a DNA molecule to another.
- **Transposon tagging.** The insertion of a transposable element into or near a gene, thereby marking that gene with a known DNA sequence.
- **Transversion.** A mutation caused by the substitution of a purine for a pyrimidine or a pyrimidine for a purine in DNA or RNA.
- **Trihybrid.** The offspring from homozygous parents differing in three pairs of genes.
- **Trinucleotide repeats.** Tandem repeats of three nucleotides that are present in many human genes. In several cases, these trinucleotide repeats have undergone expansions in copy number that have resulted in inherited diseases.
- **Trisomic.** An otherwise diploid cell or organism that has an extra chromosome of one pair (chromosome formula: 2n + 1). A specific case of this condition is called a trisomy (pl. trisomies).
- Trivalent. An association between three chromosomes during meiosis.
- **tRNA**^{Met}. The methionine tRNA that responds to internal methionine codons rather than initiation codons (cf. **tRNA**^{Met}, **tRNA**ⁱ).
- **tRNA**_{*f*}^{Met}. The methionine tRNA that specifies the initiation of polypeptide chains in prokaryotes (cf. **tRNA**^{Met}, **tRNA**_{*f*}^{Met}).
- $tRNA_i^{Met}$. The methionine tRNA that specifies the initiation of polypeptide chains in eukaryotes (cf. $tRNA_f^{Met}$, $tRNA^{Met}$).
- **Tubulin.** The major protein component of the microtubules of eukaryotic cells.
- **Tumor suppressor gene.** A gene whose product is involved in the repression of cell division.
- **Turner syndrome.** The phenotype due to the XO genotype in humans.

U

Ultraviolet (UV) radiation. The portion of the electromagnetic spectrum—wavelengths from about 1 to 350 nm—between ionizing radiation and visible light. UV is absorbed by DNA and is

highly mutagenic to unicellular organisms and to the epidermal cells of multicellular organisms.

- **Unequal crossing over.** Crossing over between repeated DNA sequences that have paired out of register, creating duplicated and deficient products.
- Univalent. An unpaired chromosome at meiosis.
- **Universality (of the genetic code).** The codons have the same meaning, with minor exceptions, in virtually all species.
- **Upstream sequence.** A sequence in a unit of transcription that precedes (is located 5' to) the transcription start site. The nucleotide pair in DNA corresponding to the nucleotide at the 5' end of the transcript (RNA) is designated +1. The preceding nucleotide pair is designated -1. All preceding (-) nucleotide sequences are upstream sequences (cf. **Downstream sequence**).
- **Uracil (U).** A pyrimidine base found in RNA but not in DNA. In DNA, uracil is replaced by thymine.

V

- Van der Waals interactions. Weak attractions between atoms placed in close proximity.
- Variable number tandem repeat (VNTR) (minisatellite). A highly polymorphic tandem repeat of a sequence of 10 to 80 nucleotide pairs in length.
- **Variance.** A measure of variation in a population; the square of the standard deviation.
- Variation. In biology, the occurrence of differences among individuals.
- **Vector.** A plasmid or viral chromosome that may be used to construct recombinant DNA molecules for introduction into living cells.
- **Viability.** The capability to live and develop normally.
- *vir* region (of Ti plasmid). The region of the Ti plasmid of *Agrobacterium tumefaciens* that contains genes encoding products required for the transfer of the T-DNA from the bacterium to plant cells.
- **Virulent phage.** A phage (virus) that destroys the host (bacterial) cell (cf. **Temperate phage**).

VNTR. See Variable number tandem repeat.

W

- **Western blot.** The transfer of proteins from an electrophoretic gel to a cellulose or nylon membrane by means of an electric force.
- Whole-genome shotgun sequencing. An approach to sequencing genomes that involves randomly cleaving the entire genome into small fragments, sequencing the ends of these fragments, and using supercomputers to assemble the complete sequence by aligning overlapping sequences.
- Wild type. The customary phenotype or standard for comparison.
- **Wobble hypothesis.** Hypothesis to explain how one tRNA may recognize two codons. The first two bases of the mRNA codon and anticodon pair properly, but the third base in the anticodon has some play (or wobble) that permits it to pair with more than one base.

Х

X chromosome. A chromosome associated with sex determination. In most animals, the female has two, and the male has one X chromosome.

Y

- **YACs (yeast artificial chromosomes).** Linear cloning vectors constructed from essential elements of yeast chromosomes. They can accommodate foreign DNA inserts of 200 to 500 kb in size.
- Y chromosome. The partner of the X chromosome in the male of many animal species.

Ζ

- **Z-DNA.** A left-handed double helix that forms in GC-rich DNA molecules. The Z refers to the zig-zagged paths of the sugar-phosphate backbones in this form of DNA.
- **Zygonema** (*adj.*, **zygotene**). Stage in meiosis during which synapsis occurs; after the leptotene stage and before the pachytene stage in the meiotic prophase.
- **Zygote.** The cell produced by the union of two mature sex cells (gametes) in reproduction; also used in genetics to designate the individual developing from such a cell.

Photo Credits

Chapter 1 Opener: Science Photo Library/Getty Images, Inc. Fig. 1.1: James King-Holmes/Photo Researchers, Inc. Fig. 1.3: Perrin Pierre/ ©Corbis. Fig. 1.5: Scott Sinklier/AGStock USA/Photo Researchers, Inc. Fig. 1.9 (left): Omikron/Photo Researchers. Fig.1.9 (right): Omikron/Photo Researchers, Inc. Fig. Fig. 1.11(left): A. Humek/Alamy. Fig. 1.11(center left): Lynn Stone/AG Stock Images/ ©Corbis. Fig. 1.11(center right): J.L. Klein & M.L Hubert/Photolibrary. 1.11(right): Armin Floreth/ imagebroker/Alamy. Fig. 1.12(left): Courtesy John Doebley, Genetics, University Wisconsin. Fig. 1.12(right): Courtesy John Doebley, Genetics, University Wisconsin. Fig. 1.13a: Courtesy Mycogen. Fig 1.13b: Courtesy Mycogen.

Chapter 2 Opener (left): Najlah Feanny-Hicks/ ©Corbis SABA; opener inset: Getty Images. Fig. 2.2a: Dr. Gopal Murti/ Photo Researchers, Inc. Fig. 2.2b: Alfred Pasieka/Photo Researchers, Inc. Fig. 2.4a: Courtesy Conley L. Rieder. Fig. 2.4b: From Jerome B. Rattner and Stephanie G. Phillips, 7. Cell Biol. 57:363, 1973. Reproduced with permission of Rockefeller University Press. Fig. 2.5: Dr. Andrew S. Bajer, Professor Emeritus, University of Oregon. Fig. 2.6a: ©Dr. David Phillips/Visuals Unlimited/ ©Corbis. Fig. 2.6b: Wood/ CMSP/Getty Images, Inc. Fig. 2.7: L. Willatt/Photo Researchers, Inc. Fig. 2.9: Clare A. Hasenkampf/ Biological Photo Service. Fig. 2.10a: From M. Westergaard and D. von Wettstein, Annual Review of Genetics,

Vol. 6; ©1972 by Annual Reviews, Inc. Original photograph courtesy of D. von Wettstein. Fig. 2.11: From Bernard John, Meiosis, Cambridge University Press, 1990.

Chapter 3 Opener: Melanie Acevedo/Botanica/Getty Images, Inc. Fig. 3.3, 3.4, & 3.5: Martin Shields/Photo Researchers, Inc. Fig. 3.10b: Niall Benvie/ ©Corbis. Fig. 3.11: Martin Shields/ Photo Researchers, Inc.

- Chapter 4 Opener: John Glover/Oxford Scientific/ Getty Images, Inc. Fig. 4.10a: Biophoto Associates/Photo Researchers. Fig. 4.12a: Photolibrary. Fig. 4.12b: Lauren Bess Berley/ Photolibrary. Fig. 4.12c&d: iStockphoto. Fig. 4.15a: ©TH Foto-Werbung/Science Photo Library/Photo Researchers, Inc. Fig. 4.15 (inset top right): ©TH Foto/ Alamy. Fig. 4.18a& b: Photo by Leah Sandall, University of Nebraska-Lincoln, Plant and Soil Science Library, http:// plantandsoil.unl.edu.
- Chapter 5 Opener: Nigel Cattlin/Photo Researchers, Inc. Fig. 5.8: Corbis-Bettmann. Fig. 5.15: Flickr/Getty Images, Inc. Fig. 5.16: Michael Abbey/ Photo Researchers, Inc.

Chapter 6 Opener: Richard Boll/Photographer's Choice/ Getty Images, Inc. Fig. 6.2: From C.G. Vosa, 1971. "The quinacrine-flourescence patterns of the chromosomes of Albium carinatum, Fig. 1, *Chromosoma* 33:382–385. Fig. 6.3: from R.M Patterson & J.C. Petricciani, 1973. "A Comparison of Prophase & Metaphase G-bands in the Muntjak." *J. of Heredity* 64 (2): 80-82. Fig. 2A. Fig. 6.4: Health Protection Agency/Photo Researchers. Fig. 6.5: Phototake. Fig. 6.7a: C. G. Maxwell/ Photo Researchers, Inc. Fig. 6.7b: Clive Champion/ Photographer's Choice/Getty Images, Inc. Fig. 6.7c: Ken Wagner/Phototake. Fig. 6.7d: Hal Beral/Visuals Unlimited/ Getty Images, Inc. Fig. 6.11: Courtesy Todd R. Laverty, HHMI/Berkeley Drosophila Genome Project. Fig. 6.14a: Hattie Young/Photo Researchers, Inc. Fig. 6.14b: Courtesy of Robert M. Fineman, Dean, Health and Human Services, North Seattle Community College. Page 123: Yoav Levy/Phototake. Fig. 6.17: L. Willatt, East Anglian Regional Genetics Service/ Photo Researchers. Fig. 6.17 inset: Addenbrookes Hospital/ Photo Researchers. Fig. 6.18: Courtesy J. K. Lim.

Chapter 7 Opener: Kevin Summers/Photographer's Choice/Getty Images, Inc. Fig. 7.9: Courtesy Bernard John.

- Chapter 8 Opener: Medical RF/Phototake. Fig. 8.1b: A. K. Kleinschmidt, Biophys. Acta. 61, 1962, p. 861, Fig. 1, Elsevier Publishing Company. Fig. 8.3: Courtesy A. F. Howatson, From Gene Expression, Vol. 3. John Wiley & Sons, 1977. Fig. 8.6: MN Tremblay/Flickr/ Getty Images, Inc. Fig. 8.10: From O.T. Avery, C.M. MacLeod, and M. McCarty, 7. Exp. Med. 79:153, 1944; by copyright permission of Rockefeller University Press. Fig. 8.13: T. F. Anderson, E. Wollman, and F. Jacob, from Annals Institute Pasteur, Masson and Cie, Editors.
- Chapter 9 Opener: G. Murti/ Photo Researchers. Fig. 9.8 (top left): A. Barrington Brown/Photo Researchers, Inc.

Fig 9.8 (top right): Corbis-Bettmann. Fig. 9.8 (bottom left): Courtesy National Institute of Health. Fig. 9.8 (bottom right): A. Barrington Brown/Photo Researchers, Inc. Fig. 9.9: Omikron/Science Source/Photo Researchers, Inc. Fig. 9.13: Courtesy Dr. Svend Freytag. Fig. 9.17: Courtesy Victoria Foe. Fig. 9.19: Reprinted with permission from Karolyn Luger, et al, Nature 389:251m 1997, courtesy of Timothy J. Richmond. Fig. 9.20: Gunter F. Bahr/ Biological Photo Service. Fig. 9.21a: Courtesy Barbara Hamkalo. Fig. 9.21b: Reprinted from Bednar et al.. 1998 Proc. Natl. Acad. Sci. USA 95: 14173-14178 with permission. Photos courtesy Dr. Christopher L. Woodcock, University of Massachusetts, Amherst. Fig. 9.22: From J.R. Paulson & U.K. Laemmli, Cell 12: 817-828, 1977. Copyright 1977, MIT; published by MIT Press. Original photo courtesy U.K. Laemmli. Fig. 9.24: From Huntington F. Willard, Trends Genetics 6:414, 1990.

Chapter 10 Opener: Joel Sartore/NG Image Collection. Fig. 10.5: From J. H. Taylor, "The Replication and Organization of DNA in Chromosomes," Molecular Genetics, Par I, J. H. Taylor (ed), Academic Press, New York, 1963. Fig. 10.7a: J. Cairns, Imperial Cancer Research Fund, London, England. Reproduced with permission from Cold Spring Harbor Sympos. Quant. Biol. 28:43, 1963. Original photo courtesy John Cairns. Fig. 10.9c: Photo reproduced with permission from M. Schnos and R.B. Inman, 7. Mol. Biol. 51: 61-73, 1970. © 1970 by Academic Press, Inc. (London), Ltd./Elsevier

Limited. Fig. 10.11: Original micrographs courtesy of David Dressler, Harvard University. Fig. 10.24: From S. Doublie, S. Tabor, A.M. Long, C.C. Richardson, and T. Ellenberger, Nature 391:251-258. Fig. 10.25: From X.P. Kong et al., Cell 69:425-437, 1992. © Cell Press/Elsevier. Original photograph courtesy Dr. John Kuriyan, Howard Hughes Medical Institute, Rockefeller University. Fig. 10.29 (inset): Original micrograph courtesy of David Dressler, Harvard University. Fig. 10.31a&b: Reproduced with permission from J. A. Huberman and A.D. Riggs, Journal of Molecular *Biology* 32:327–341 © 1968 by Academic Press. Original photographs courtesy J. A. Huberman. Fig. 10.33a: Courtesy of Steven L. McKnight and Oscar L. Miller, Jr. Fig. 10.35: Gerald Herbert/ ©AP/Wide World Photos.

Chapter 11 Opener: Dr. Mark J. Winter/Photo Researchers. Fig. 11.11: From O.L. Miller, Jr., B.A. Hamkalo, and C.A. Thomas, Jr., Science 169:392-395, 1970. Copyright (c) 1970 by the American Association for the Advancement of Science. Original micrograph courtesy O. L. Miller, Jr. Fig. 11.15a: P. Cramer, D.A. Bushnell, & R.D. Kornberg, "A Structural Basis of Transcription," Science, Vol. 292: 1863-1876, Fig. 1b (2001). Fig.11.19b: From Tilghman et al., Proceedings of The National Academy of Sciences, U.S.A. 75:1309-1313. Fig. 11.19c: Courtesy Pierre Chambon.

Chapter 12 Opener: Eye of Science/Photo Researchers.
Fig. 12.13a: From S.H. Kim,
F.L. Suddath, G.J. Quiqley, A.
McPherson, J.L. Sussman,
A.H.J. WAng, N.C. Seeman,
and A. Rich, *Science* 185:
435-440, 1974 by the American
Association for the Advancement of Science. Original
photo courtesy S. H. Kim.
Fig. 12.14b: Courtesy Dr. Joachim Frank. From Frank, et al. 1995. Biochemistry and Cell Biology 73: 357. Fig. 12.15a: Reproduced with permission of M. Yusupov et al., *Science* 292: 823-826, 2001. Courtesy Albion Baucom and Harry Noller. Fig. 12.19: From S.L. McKnight, N.L. Sullivan, and O.L. Miller, Jr., *Prog. Nucleic Acid Res. Mol. Biol.* 19:313–318, 1976. Micrograph courtesy S.L. McKnight and O.L. Miller, Jr., University of Virginia.

Chapter 13 Opener: Sarah Leen/NG Image Collection. Fig. 13.3: Courtesy of H. Potter, University of South Florida and D. Dressler, Harvard University. Fig. 13.14: ©Corbis. Fig. 13.24: From B.N. Ames, J. McCann, and E. Yamasaki, Mutat. Res. 31:347, 1975. Photograph courtesy B.N. Ames. Fig. 13.28: ©Frederic Larson/San Francisco Chronicle/Corbis. Fig. 13.30 Courtesy of H. Potter, University of South Florida and D. Dressler, Harvard University

Chapter 14 Opener: ©H. Raguet/ Pr. Lavery, Laboratory of Biochemical Theory/ Phototake. Fig. 14.1: Based on an X-ray structure by John Rosenberg, U. of Pittsburgh. Fig. 14.4: Courtesy S. Kopczak and D.P. Snustad, University of Minnesota. Fig. 14.10: Courtesy D.P. Snustad. Fig. 14.12a: Photographs courtesy of S.R. Ludwig and D.P. Snustad, University of Minnesota. Fig. 14.12b: Prof. D. Peter Snustad. Fig. 14.13: Courtesy of S.R. Ludwig and D.P. Snustad, University of Minnesota. Page 383: From Kerem, et al. (1989), Science 245:1073-1080.

Chapter 15 Opener: ©AP/ Wide World Photos. Fig. 15.15 Courtesy CLONTECH Laboratories, Inc. Fig. 15.16a: Sam Ogden/ Photo Researchers. Fig. 15.16b: Deco/Alamy. Fig. 15.17b-d: Ludin & Matus, 1998. Trends in Cell Biology, 8:72, Elsevier.

Chapter 16 Opener: Stephen Voss/Redux Pictures. Fig. 16.2: From *Cell* 72:971-983, Fig. 7, March 26,1993, Copyright © 1993 Cell Press, Elsevier. Page 443: From Richards and Sutherland, *Trends in Genetics*, vol. 8 (7), p. 249, 1992. Photo courtesy Grant Sutherland. Fig. 16.9: Getty Images, Inc. Fig. 16.13: Orchid Cellmark. Fig. 16.18: Dr. R. L. Brinster, School of Veterinary Medicine, University of Pennsylvania.

Chapter 17 Opener: Gregory G. Dimijian, M.D./Photo Researchers. Fig. 17.11: NIBSC/Photo Researchers.

Chapter 18 Opener: CNRI/ Photo Researchers. Fig. 18.11b: Courtesy T.A. Steitz, Yale University. Fig. 18.12: Courtesy Ponzy Lu and Mitchell Lewis, University of Pennsylania. Lewis et al. 1966. *Science*, 271:1247–1254, Fig. 6.

Chapter 19 Opener: Dennis Kunkel Microscopy, Inc/ Phototake. Fig. 19.10: Courtesy of Joseph Fong, University of Minnesota. Fig. 19.16: from M. I. Kuroda et al, *Cell* 66:935–947, 1991, Fig. 6. Photograph courtesy of Dr. Mitzi Kuroda.

Chapter 20 Opener: Petit Format/Photo Researchers. Fig. 20.1: Courtesy Kevin Haley, University of Minnesota. Fig. 20.2: Daniel St. Johnston and Christiane Nusslein-Vohard, Cell 68:201-219, 1992. Photograph courtesy of Christiane Nusslein-Vohard. Fig. 20.9: Courtesy of Matthew Scott, Howard Hughes Medical Institute. Fig. 20.12: Courtesy Walter Gehring, Universitat Basel, Switzerland. Fig. 20.15: From Amit et al., Science 233:747, Copyright (c) 1986 the American Association for the Advancement of Science. Photograph courtesy of R. J.

Poljak.

Chapter 21 Opener: VEM/ Photo Researchers.

Chapter 22 Opener: Zephyr/ Photo Researchers. Fig. 22.2: Th Foto-Werbung/SPL/Photo Researchers, Inc. Fig. 22.9: iStockphoto.

Chapter 23 Opener: Danita Delimont/Alamy. Fig. 23.6a: Stephen Dalton/Minden Pictures, Inc. Fig. 23.6b: Martin Withers/Minden Pictures, Inc. Fig. 23.9: Eye of Science/Photo Researchers, Inc.

Chapter 24 Opener: Super-Stock. Fig. 24.1(top left): Herbert Spichtinger/ zefa/@Corbis. Fig. 24.1(top right): Gregg Stott/Masterfile. Fig. 24.1(bottom left): Ron Chapple Stock/Photolibrary. Fig. 24.1(bottom right): Kenneth H. Thomas/Photo Researchers. Fig. 24.2: Tui De Roy/Minden Pictures, Inc. Fig. 24.3(top left and top right): J. H. Robinson/Photo Researchers, Inc. Fig. 24.3(center left): Alvin E. Staffan/Photo Researchers, Inc. Fig. 24.3(center right): Gregory K. Scott/ Photo Researchers, Inc. Fig. 24.3 (bottom left): John Kaprielian/Photo Researchers, Inc. Fig. 24.3 (bottom right): Millard Sharp/Photo Researchers, Inc. Fig. 24.9a: Taxi/Getty Images. Fig. 24.9b: Mike Parry/Minden Pictures, Inc. Fig. 24.9c: Rich Reid/National Geographic Society. Fig. 24.14: Photo by K.Y. Kaneshiro and K. T. Kaneshiro, U. of Hawaii. Fig. 24.15a: Donald M. Jones/Minden Pictures, Inc. Fig. 24.15b: S & D & K Maslowski/Minden Pictures, Inc.

Appendices

Appendix C, p. 690: From J. Meyne, in R.P. Wagner, *Chromosomes: A Synthesis*, Copyright (c) 1993. Reprinted with permissions of Wiley-Liss, Inc., a subsidiary of John Wiley & Sons, Inc.

Illustration Credits

- Chapter 3 Fig. 3.15: H.T. Lynch, R. Fusaro, and J.F. Lynch. 1997. Cancer genetics in the new era of molecular biology. *NY Acad. Sci.* 833:1.
- Chapter 4 Fig. 4.10*b*: Principles of Human Genetics, 3/e by Curt Stern, © 1973 by W. H. Freeman and Company. Used with permission. Fig. 4.17: Nance, W. E., Jackson, C. E., and Witkop, C. J., Jr. 1970. American Journal of Human Genetics 22:579–586. Used with permission of the University of Chicago Press.
- **Chapter 8** Fig. 8.3*b*: After Pfashne *A Genetic Switch* 2e. Cell and BSP Press. Blackwell.
- Chapter 9 Fig. 9.23: After Figure 1 in The ENCODE Project Consortium. *Science* 306:636–640, Oct. 22, 2004.
- **Chapter 10** Fig. 10.28: Adapted from DNA Replication by Kornberg and Baker © 1992 by W. H. Freeman and Company. Used with permission.
- Chapter 11 Fig. 11.21: Reprinted with permission of *Nature* from Zang, N. J., Grabowski, P. J., and Cech, T. R. 1983. *Nature* 301: 578–583. Copyright 1983 Macmillan Magazines Limited.
- Chapter 12 Figs. 12.3: Alberts, B., et al., *Molecular Biology of the Cell, 3/e*, page 114. New York: Garland Publishing Inc., 1994. Fig. 12.4: Figure from *Biology*, Second Edition by Claude A. Villee, Eldra Pearl Solomon, Charles E. Martin, Diana W. Martin, Linda R. Berg, and P. William Davis, copyright

© 1989 by Saunders College Publishing, reproduced by permission of Harcourt Brace & Company. Fig. 12.12: Reprinted with permission from Holley, R.W., et al., 1965. *Science* 147: 1462–1465. Copyright 1965 American Association for the Advancement of Science.

- Chapter 14 Fig. 14.1*b*: X-ray drawing by John Rosenberg, Univ. of Pittsburgh.
- Chapter 15 Fig. 15.1: After Messing, J. W., and Llaca, V. 1998. Proceedings of the National Academy of Sciences 95: 2017. Copyright 1998 National Academy of Sciences, U.S.A. Fig. 15.4: Based on data from NIH/CEPH Collaboration Mapping Group, 1992. Science 258: 67-86. Fig. 15.8: From Stewart, E. A., et al. 1997. Genome Research 7:422-433. Figs. 15.9 and 15.10: Venter et al. (2000). Science 291: 1304-1351. Fig. 16.12: Data from the International Human Genome Sequencing Consortium. (2001). Nature 409:860-921. Fig. 15.11: Data from the International Human Genome Sequencing Consortium. (2001). Nature 409:860-921. Fig. 15.12: After Figure 1 in The ENCODE Project Consortium. Science 306:636-640, Oct. 22, 2004. Fig. 15.20: Creation of a Bacterial Cell Controlled by a Chemically Synthesized Genome; Science 329 (5987): 52-26. Fig. 15.23: After Moore, et al. 1995. Current Opinion Genetics &

Development 5:537 and Gale, M. D., and Devos, K. M. 1998. Proceedings of the National Academy of Sciences 95: 1971– 1973. Fig. 15.24: Chowdhary B. P., et al., 1998 Genome Research 8: 577–584.

Chapter 16 Fig. 16.1: From Huntington's Disease Collaborative Research Group. 1993. Cell 72:971-983. Copyright Cell Press. Fig. 16.3: From Marx, J. L. 1989. Science 245:923-925. Reprinted with permission of Dr. Lapchee Tsui, The Hospital for Sick Children, Toronto, Canada. Figs. 16.4, 16.5: Reprinted by permission from Collins, F. S., 1992. Science 256:774-779. Copyright 1992 American Association for Advancement of Science. Fig. 16.8: McCormack and Rabbitts. 2004. New England Journal of Medicine. 350:913-922 (Feb. 26, 2004 #9). Fig. 16.12a: Promega International Symposium on Human Identification Conference Proceedings. Fig. 16.22: José M. Alonso, Anna N. Stepanova, Thomas J. Leisse, Christopher J. Kim, Huaming Chen, Paul Shinn, Denise K. Stevenson, Justin Zimmerman, Pascual Barajas, Rosa Cheuk, Carmelita Gadrinab, Collen Heller, Albert Jeske, Eric Koesema, Cristina C. Meyers, Holly Parker, Lance Prednis, Yasser Ansari, Nathan Choy, Hashim Deen, Michael Geralt, Nisha Hazari, Emily Hom, Meagan Karnes, Celene Mulholland,

Ral Ndubaku, Ian Schmidt, Plinio Guzman, Laura Aguilar-Henonin, Markus Schmid, Detlef Weigel, David E. Carter, Trudy Marchand, Eddy Risseeuw, Debra Brogden, Albana Zeko, William L. Crosby, Charles C. Berry, and Joseph R. Ecker (2003) Genome-Wide Insertional Mutagenesis of Arabidopsis thaliana Science 301: 653–657.

- Chapter 17 Fig. 17.17 and 17.18: From Lim, J. K., and Simmons, M. J. 1994. *BioEssays* 16:269–275. © ICSN Press.
- Chapter 18 On the Cutting Edge Fig. 1: Wellcome Sanger Institute: http://rfam.sanger. ac.uk/family/varna/RF00168. On the Cutting Edge Fig. 2:
- **Chapter 19** Fig. 19.3: *Molecular Cell Biology* by Darnell, Lodish and Baltimore. © 1996, 1990, 1985 by Scientific American Books. Used with permission by W. H. Freeman and Company.
- Chapter 21 Fig. 22.10*a*: From Kinzler, K. W., and Vogelstein, B. 1996. *Cell* 87:159–170. Copyright Cell Press. Fig. 22.120: Based on Latil, A., and Lidereau, R. 1998. *Virchow's Archive* 432:389–406.
- Chapter 24 Fig. 24.16: Based on Wood, B. 1996. BioEssays 18:945–954. Copyright © 1996 John Wiley & Sons, Inc. Reprinted with permission of Wiley-Liss, Inc., a division of John Wiley & Sons, Inc.

Index

A

A, see Adenine Aberrant Ds elements, 484 Abortion, pollen, 129 ABRC (Arabidopsis Biological Resource Center), 470 Ac element, 483-485 Acentric fragment, 154 Acetylation, 269 Achondroplasia, 53 Acquired immunodeficiency syndrome (AIDS), 9, 488 Acquired traits, 334 Acridine dyes, 326-328 Activation, of whole chromosomes, 550-553 Activators: Ac element, 483-485 catabolite activator protein, 514, 517, 518 dominant activator, 589 tissue plasminogen activator, 671 Acute lymphoblastic leukemia, 270 Adaptation, 670 ADA- SCID (adenosine deaminase-deficient severe combined immunodeficiency disease), 452-453 Adelburg, Edward, 182 Adenine (A), 4, 198-200, 202 Adenomatous polyposis coli, 597 Adenosine deaminase-deficient severe combined immunodeficiency disease (ADA- SCID), 452-453 Adenoviruses, 451 Adjacent disjunction, 128 African sleeping sickness, 531 Agarose gel electrophoresis, 381-383, 386 Agglutination, 63 Aging in humans: and DNA repair defects, 352 and telomere length, 249 Agriculture, 12-14, 187

Agrobacterium tumefaciens, 465-466, 469 Agrobacterium tumefaciensmediated transformation, 464-465 AIDS (acquired immunodeficiency syndrome), 9, 488 Albinism, 53, 55-56, 77, 339-340 Alexandra, Czarina, 98, 99 Algal cells, 20 Alkaptonuria, 53, 339 Alkylating agents, 326, 327 Alleles: defined, 42 Mendel's study of, 3 multiple, 64-65 Allele frequency, 635–641 estimating, 635-636 in genetic counseling, 640-641 and genotype frequencies, 636 Hardy-Weinberg principle of, 636-640 random genetic drift and changes in, 645-646 theory of, 635-641 Allelic series, 65 Allelic variation, 63-70 Allelism, testing mutations for, 65-66 Allopatric speciation, 674–675 Allopolyploids, 117 Allosteric transitions, 509 Alpha (α) chains, 338 a-globin, 664-667 α helices, 288 α mating type, yeast, 33 ALS (amyotrophic lateral sclerosis), 415 Alteration of generations, 31–32 Alternate disjunction, 128 Alternate splicing, 533 Altman, Sidney, 278 ALV (avian leukosis virus), 464 Alzheimer's disease, 120, 628 α mating type, yeast, 33

American Board of Genetic Counseling, 56 Ames, Bruce, 346 Ames test, 346-348 Amino acids: abbreviations for, 23, 423 codons for, 309-310 functional constraint on, 666-667 polypeptides built from, 67, 286-287 selenocysteine as, 315 substitutions, 323 Amino acid metabolism, 14 Aminoacyl (A) site, 297, 298 Aminoacyl-tRNA synthetases, 294, 296 2-Aminopurine (2-AP), 326 Amish people, 77 Amniocentesis, 123 Amorphic alleles, 65 Amplification: in cloning vectors, 372-373 by PCR, 374-376 Amylases, 462 Amyotrophic lateral sclerosis (ALS), 415 Anabolic pathways, 507 Analogous DNA sequences, 664 Analysis of variance, 614 Anaphase, 25, 26, 28, 31 Anaphase I, 28, 31 Anaphase II, 29, 31 Anchor markers, 403 Androgen-independent prostate cancer, 601 Aneuploidy, 119–126 Belling's study of, 119 defined, 114, 119 deletions, 124, 125 duplication, 124, 125 monosomy, 121-124 tests for, 123 trisomy in human beings, 120-122 Angiogenesis, 602 Animals. See also specific types

cell division in, 23 as eukaryotes, 20 transgenic, 463-464 Animal cells, 21 cytokinesis in, 26 mitotic spindle, 24 Animal development, 558-578 genetic perspective on, 559-561 germ line during, 22 maternal gene activity, 561-565 and stem cell therapy, 558 in vertebrates, 571-578 zygotic gene activity, 565-571 Animal feed, 187 Animal growth-promoters, 187 Anopheles gambiae, 398 Anterior-posterior axis, in Drosophila, 562-565 Anthers, 33 Antibiotics: discovery of, 163-164 insertion sequences and resistance to, 481 market value of, 461 mutant genes resistant to, 171, 333-335 Antibiotic-resistant bacteria, 181.186-187 Antibodies, 575 Anticipation, 332 Anticodons, 294, 312, 313 Antigens, 63, 575 Antipodal cells, 33, 34 Antirrhinum majus, 63, 616 Antisense RNA, 261 APC gene, 600 Apes, 676 Apolipoproteins, 273, 274 Apomixis, 116 Apoptosis, 584 Apple, 332 Aptamer domain, riboswitch, 524-525 Arabidopsis Biological Resource Center (ABRC), 470

Arabidopsis thaliana: β-tubulin genes of, 423–424 genome sequencing of, 430, 431 insertional mutagenesis in, 469-471 as model organism, 32 tubulins, 539 Arber, Werner, 368 Ardipithecus, 676 Arginine, 207 Argonaute protein, 541 ARSs (autonomously replicating sequences), 225, 227 Artificial chromosomes: bacterial, 373, 374 as cloning vectors, 373 P1, 373, 374 yeast, 373, 374 Artificial selection, 618-620 Ascospores, 33 Ascus, 33 Asexual reproduction, 33, 116 Ashkenazi Jews, 340, 600 Assays of genome function, 415-419 expressed sequence, 416 green fluorescent protein, 419-420 microarrays/gene chips, 416-419 Aster, 24, 25 Ataxia telangiectasia, 53, 352, 353, 592 AT-rich denaturation sites, 229 Attached-X chromosome, 128-129 Attenuation, 520–523 Attenuator, 521-523 Auerbach, Charlotte, 326 AUG initiator, 301 Australopithecus, 676-677 Autoantibodies, 280 Autocatalytic splicing, 278–279 Automated DNA sequencing, 391 Autonomously replicating sequences (ARSs), 225, 227 Autonomous state, 176 Autopolyploids, 117 Autoradiography, 223, 227-228 Autosomes, 91, 97 Auxotrophs, 171, 341 Avery, Oswald, 174, 194, 195

Avian leukosis virus (ALV), 464

В

BACs (bacterial artificial chromosomes), 373, 374 Bacillus licheniformis, 462 Bacillus subtilis, 174–175 Bacillus thuringiensis (BT), 14 Back mutation, 336 Bacteria, 169-187. See also specific bacteria, e.g.: Escherichia coli antibiotic-resistant, 181, 186-187, 481 cell walls of, 19 chromosome in, 22 conjugation, 172, 175-180 DNA base composition in, 200 drug-resistant, 163-164 episomes in, 181 eukaryotic protein production in, 461-463 evolution of bacterial genomes, 186 genetic exchange in, 172-186 genetic importance of, 164-165 genetics of, 169-172 mapping genes in, 182 multi-drug-resistant TB, 163-164 mutant genes in, 170-171 plasmids in, 179-181 as prokaryotes, 20 sexduction, 181-182 synthetic bacterial genome, 425-426 transduction, 182-185 transformation of, 173-175 transposons in, 479-483 unidirectional gene transfer in, 171 Bacterial artificial chromosomes (BACs), 373, 374 Bacterial cells, 21 Bacteriophages, 165-169 in E. coli, 165-169 mapping genes in, 166 virulent vs. temperate, 165 Bacteriophage lambda, 166-169, 373 chromosome forms of, 228-229 DNA base composition in, 200 specialized transduction in, 183-185 Bacteriophage P1, 183

Bacteriophage P22, 183 Bacteriophage $\phi 80, 184$ Bacteriophage T2: DNA as carrier of genetic information in, 195-197 DNA base composition in, 200 Bacteriophage T4: amber mutations of, 344-345 as lytic phage, 165-166 morphogenesis in, 341-342 Bacteriophage T7, 230, 240 Bacteriophage vectors, 372 Baker's yeast, see Saccharomyces cerevisiae Balanced polymorphism, 648 Balancers, 154 Balancing selection, 648-649 Baldness, pattern, 71 Baltimore, David, 488 Banding patterns, 402, 403 Barr, Murray, 105 Barr body, 105 Basal transcription factors, 271, 537 Base analogs, 326 Base excision repair, 348-349 Base-pairing rules, 4, 6 Base-pair substitutions, 314 Basic HLH (bHLH) proteins, 540 Basic Local Alignment Search Tool (BLAST), 401, 421 Bateson, William, 41, 62, 67, 68, 72, 73, 136, 137, 566 BAX gene, 595–596 Beadle, George W., 67, 289-291 Beads-on-a-string (gene concept), 208-209 Behavioral trait inheritance, 628-630 Belling, John, 119 Berg, Paul, 371 β-catenin, 596, 598 Beta (β) chains, 338, 339 β-galactosidase, 373 β-globin, 8, 9, 274–275, 314, 323, 546 β sheets, 288 bHLH (basic HLH) proteins, 540 bicoid gene, 563, 564 Bidirectional replication, 228-230 Biessmann, Harald, 493 Bingham, Paul, 486

Binomial probabilities, 54 Bioinformatics, 400, 421-424, 431 Birds, 103. See also specific birds, e.g.: Grosbeaks Biston beularia, 643, 644 Bivalent, 30, 31 Blackburn, Elizabeth, 248 Bladder cancer, 587 Blakeslee, Albert, 119 BLAST (Basic Local Alignment Search Tool), 401, 421 Blastoderm, 559-560 Bligh, William, 634 Blindness, congenital, 439-440 Blood, see β-globin; Hemoglobin; Sickle-cell anemia Blood-clotting factors, 461 Blood lily, 25 Blood type, 63-65, 616, 635-638,660 Bloom's syndrome, 352, 353, 592 Bluescript II, 372, 373 Body segmentation, 565-568 homeotic genes, 565-566 segmentation genes, 566-568 Bone marrow cells, 220 Bone marrow stem cells, 452 BOSS proteins, 571 Brachydactyly, 53 Bracon hebetor, 103 BRCA1 gene, 14, 420, 599 BRCA2 gene, 420, 599 Bread mold, 289-291 Breast cancer, 14, 582, 592, 599 Bridges, Calvin, 89, 93-94, 102, 118-119, 125, 142-145, 149, 565 Broad bean, 223 Broad-sense heritability, 615-616, 626 5-Bromouracil (5-BU), 326-327 Bruce, David, 531 BT (Bacillus thuringiensis), 14 Budding, 33 Bulbs, 116 Burkitt's lymphoma, 471, 589, 590 Burns, Sarah, 182 Bursa bursa-pastoris, see Sweet pea

С

C, see Cytosine CAAT box, 270–271 *Caenorhabditis elegans:* development of, 559 Caenorhabditis elegans: (Continued) genome sequencing of, 430, 431 hypoactivation of X chromosomes in, 553 as model organism, 32 programmed cell death in, 584 RNA interference in, 471 transcription units in, 267 Cairns, John, 221, 224, 227, 231 cAMP (cyclic AMP), 517, 518 Campion, 49 Cancer, 581-603 and cell cycle, 583-584 and chromosome rearrangements, 589-590 and deregulation of cell division, 23 forms of, 582-583 genetic basis for, 584-585 and genetic counseling, 600 genetic pathways to, 600-603 inherited, 590-593 molecular family connection to, 581 oncogenes, 585-590 and programmed cell death, 584 tumor suppressor genes, 593-599 CAP, see Catabolite activator protein Cap-binding protein (CBP), 301-302 Capecchi, Mario, 455 Capsule, 21 Carbapenems, 186 Carbohydrates, 19, 506 Carcinogens, 346-347, 583 Cardiovascular disease, 607, 623, 624 Caspases, 584 Cats, 104, 105 Catabolic pathways, 506 Catabolite activator protein (CAP), 514, 517, 518 Catabolite repression, 514–515, 517 β-Catenin, 596, 598 Cattle, 13, 433-434 caudal gene, 563, 564 CBP (cap-binding protein), 301-302 CDKs (cyclin-dependent kinases), 583 cDNAs, see Complementary **DNAs**

cDNA clones, 416 cDNA libraries, see Complementary DNA libraries Cech, Thomas, 278 Celera Genomics, 410, 412 Cells: prokaryotic vs. eukaryotic, 20, 21 specification of type of, 569-570 Cell cycle: and cancer, 583-584 defined. 23 in eukaryotic chromosome replication, 245 Cell death, programmed, 584, 602 Cell division, 23-32 human chromosomes during, 22 meiosis, 27-32 mitosis, 24-26 process of, 23 Cell motility, 20 Cell plate, 26 Cellular blastoderm, 559-560 Cellular environment, 19 Cellular oncogenes, 587-589 Cellular reproduction, 18-36 cells and chromosomes, 19-24 life cycles of model organisms, 32-36 meiosis, 27-32 mitosis, 24-26 Cellulose, 19, 26 Cell walls, 19, 21, 26 CentiMorgans, 142, 166 Central dogma of molecular biology, 9, 257 Central element, 30 Centrifugation, 222, 223 Centrioles, 21, 24, 25 Centromeres, 545 and cell division, 22, 23 and chromosome "arms," 114 in eukaryotic chromosomes, 211-212 and kinetochores, 25 Centrosomes, 25 Cereal grasses, 431-432 Cesium chloride (CsCl), 222 CF, see Cystic fibrosis CF gene, 15 CFTR protein (cystic fibrosis transmembrane conductance regulator protein), 447-448 Chain cleavage, 272-273

Chain-termination codons, 304, 305 Channels, in cell walls, 19 Chaperones, 288 Chargaff, Erwin, 199 Chase, Martha, 195, 196 Checkpoint, 583, 584 Chemicals, Ames testing of, 346-348 Chemical-induced mutation. 326-328 Chiasma(ta): defined, 136 in meiosis, 30, 31 and time of crossing over, 140 Chickens: comb shapes in, 71-72 ovalbumin gene of, 274 transgenic, 464 Chi forms, 356 Chimeras, 464 Chimeric selectable marker genes, 466 Chimpanzees, 368 Chinese hamster, 245, 410 Chi-square (χ^2) , 50 Chi-square test, 50-52 Chlamydomonas reinhardti, 421, 422, 428 Chloramphenicol, 187 Chloroplasts: in cell division, 22, 23 defined, 20 function of, 532 genomes of, 426-429 illustration of, 21 Chloroplast DNA (cpDNA), 428-429 Chorion, 559, 560 Chorionic biopsy, 123 Chowdhary, Bhanu, 433 Christian, Fletcher, 634, 647 Chromatids, sister, see Sister chromatids Chromatid disjunction, 31 Chromatin, 24, 90, 207-209, 272 Chromatin fibers, 30 Chromatin organization, 544-550 chromatin remodeling, 546-547 DNA methylation, 547-548 euchromatin/heterochromatin, 545 imprinting, 548-550

of transcriptionally active DNA, 545-546 Chromatin remodeling, 268-270, 546-547 Chromocenters, 118–119 Chromosome(s): activation/inactivation of whole, 550-553 artificial chromosomes as cloning vectors, 373 in cells, 19, 20 combinations in sperm cells, 31 correlated maps of, 402-406 homologous, see Homologous chromosomes as location of genes, 20, 22-23 microscopic examination of, 22 replication of, see Replication, of DNA and chromosomes Chromosome 1, 410, 411 Chromosome 2, 433 Chromosome 3, 433-434 Chromosome 17, 433 Chromosome 21, 120-122, 409, 433-434, 628 Chromosome 22, 410 Chromosome disjunction, 31 Chromosome jumps, 408-409, 446 Chromosome mapping, 13, 141-148 crossing over, 141 first chromosome map, 135 genetic map distance, 146-147 with nucleotide sequences, 387-391 three-point testcross, 142-146 two-point testcross, 141-142 Chromosome number (n), 90 Chromosome painting, 112–113, 432-434 Chromosome pairing, 117 Chromosome structure, 192–216 DNA, 193-205 in eukaryotes, 207-216 nuclein discovery, 192 in prokaryotes, 205-206 rearrangement of, see Rearrangement, of chromosome structure RNA. 197-199 variation in, 660-661 in viruses, 205-206 Chromosome termini, 248-249

Chromosome theory of heredity, 92-98 experimental evidence, 92-94 nondisjunction as proof of, 93-94 Chromosome walks, 408, 446 Chronic myelogenous leukemia (CML), 590 Cichlid fish, 675 Cigarette smoke, 346 Cilia, 21 cinnamon gene, 562 cis configuration, 343 cis heterozygote, 343 cis test, 343 cis-trans position effect, 343 cis-trans test, 342-346 Civilization, cytogenetic variation and, 110 Classical genetics, 11 ClB chromosome, 325 ClB method, 325, 496 Cleavage, 354-356 Cleavage furrow, 26 Clones, 23, 372 Clone banks, 405–406 Cloning. See also Positional cloning defined, 367 of Dolly, 18 and gene sequencing, 87 large genes and segments of genomes, 374 reproductive, 574-575 therapeutic, 574 Cloning vectors: amplification of recombinant DNA in, 372-373 defined, 367 for large genes and segments of genomes, 374 CML (chronic myelogenous leukemia), 590 Cockayne syndrome, 352, 353 Codes, 256 Coding sequence, 292 CODIS, 456, 458-461 Codominance, 63-64 Codon(s): defined, 7 in gene transcription, 257 initiation/termination, 309-310 suppressor mutations with altered codon recognition, 313-314

three nucleotides per, 306-309 tRNA interactions with, 312-314 wobble hypothesis, 312–313 Coefficient of coincidence, 145 Coefficient of relationship, 82 Cohen, Stanley, 371 Coincidence, coefficient of, 145 Cointegrate, 482 Colchicine, 223 Colinearity, 291-292 Coliphage, 165, 166, 169 Collins, Francis, 249, 409, 410, 412, 446 Colon cancer, 584–585, 592 Colony, 23 Colony hybridization, 379-380 Color blindness, 98-100 Colorectal cancer, 55, 592, 597, 601 Col plasmids, 181 Common ancestor, 79-81 Comparative genomics, 402, 420-434 bioinformatics, 421-424 cereal grasses, 431-432 chloroplast genomes, 426-429 eukaryotic genomes, 429-431 mammals, 432-434 mitochondrial genomes, 427-429 prokaryotic genomes, 424-425 synthetic bacterial genome, 425-426 Competence (Com) proteins, 174 Competent bacteria, 174 Complementarity, 203 Complementary DNAs (cDNAs), 377, 384, 385 Complementary DNA (cDNA) libraries: construction of, 378 screening of, 378-380 Complementation screening, 378-379 Complementation (cis-trans) test, 342-346 Complete medium, 291 Complete P elements, 486–487 Complex traits, inheritance of, see Inheritance of complex traits Composite transposons, 481 Compound chromosome, 128-129 Compound inbreeding, 81 Concordance rate, 610

Condensation, of chromosome, 24, 25, 30 Conditional lethal mutation, 340-342 Conformation, 288 Congenital blindness, 439-440 Congenital night blindness, 53 Conjugation, in bacteria, 172, 175 - 180Conjugation channel, 175 Conjugative plasmids, 181, 481 Conjugative R plasmids, 480, 481 Connective tissue disorders, 53 Consanguineous mating, 77, 79-81,638 Consensus sequences, 264 Conservative replication, 222 Constant region, 576 Constitutive genes, 506 Contigs, 403, 406 Continuous synthesis, of DNA strand, 232, 233 Controlled transcription, 532-533 Controlling elements, 484 copia element, 492 Corces, Victor, 538 Co-repressors, 507 Corn, 13, 14, 200. See also Maize Correns, Carl, 41 Cortical granules, 35 Cosmids, 372-373 Cosmid vectors, 372-373 Cotterman, Charles, 79, 81 Cotyledons, 34 Coumermycin, 238 Coupling, 138, 343 Covalent bonds, 201 CpDNA (chloroplast DNA), 428-429 CpG islands, 269, 446, 548 Creighton, Harriet, 139 Crick, Francis, 3-4, 199-201, 221, 296, 306, 307, 312, 313, 321, 322, 409 Cri-du-chat syndrome, 124, 125 Critical value, 50 Cross-fertilization, 42 Crossing over, 136 chiasmata and time of, 140 defined, 30 double-strand break model of, 356 Holliday model of, 354-355

as measure of genetic distance, 141 during meiosis, 31, 32 outcomes of, 139 and recombination, 138-139 as recombination mechanism, 354 with transposons, 499 Crown gall disease, 465 Cryoelectron microscopy, 211 CsCl (cesium chloride), 222 CsCl equilibrium densitygradient centrifugation, 223 c-src gene, 587 C-terminal homo-oligomerization domain (OD), 595, 596 Cultivation, 116 Culturing: of bacteria, 164, 170 of viruses, 164 Cut-and-paste transposons, 478 Ac and Ds elements in maize, 483-485 in eukaryotes, 483–488 P elements and hybrid dysgenesis in Drosophila, 485-487 Cuttings, cultivation from, 116 Cyclic AMP (cAMP), 517, 518 Cyclins, 583 Cyclin-dependent kinases (CDKs), 583 Cysteine, 315 Cystic fibrosis (CF), 15 detection of mutant gene causing, 383 as recessive-trait inherited condition, 53 recombinant DNA technology with, 445-448 segregation example using, 54 Cystic fibrosis transmembrane conductance regulator (CFTR) protein, 447-448 Cytochrome b gene sequences, 10 Cytogenetics, 111 Cytogenetic mapping, 148-150 of chromosomes, 405 deletions/duplications, 148-149 distance, 149-150 Cytogenetic variation: about, 114 aneuploidy, 119-126 and civilization, 110

Cytogenetic variation: *(Continued)* polyploidy, 115–119 rearrangement, 126–129 Cytokinesis, 23, 25, 26, 29 Cytological analysis, 111–114 of human karyotype, 113–114 of mitotic chromosomes, 111–113 Cytological maps, 402, 403, 405 Cytology, 89 Cytoplasm, 19, 21 Cytoplasmic bridges, 35 Cytoplasmic control, 533–534 Cytosine (C), 4, 198–200, 202 Cytoskeleton, 20

D

Danio rerio, 32 "Dark matter," in genome, 412, 413 Darwin, Charles, 10, 641, 645, 656-658, 672, 676 Darwin's theory of evolution, 10,657-658 Databases, DNA, see DNA databases Datura stramonium, 119 Daughter cells, 23, 26, 27, 29, 31 Dawson, Martin, 173, 194 DBD (DNA-binding core domain), 595, 596 DDBJ (DNA DataBank of Japan), 400 Deacetylation, 547 Deaminating agent, 326–328 decapentaplegic gene, 562 Decondensation, of chromosomes, 24, 26, 31 Deficiency, 124 Degeneracy, 310–312 Degrees of freedom, 52 Deletions: chromosomal, 124, 125 mapping, 148-149 Delicious apple, 332 DeLucia, Paula, 231 Denaturation, 215, 384 Denaturation sites, 229 Denisovans, 398 Denmark, 187 de novo, 261 Dentatorubro-pallidoluysian atrophy, 442 Deoxyribonuclease (DNase), 194, 206. See also DNase I

Deoxyribonucleic acid, see DNA Derepression, 507 The Descent of Man (Charles Darwin), 656 Development, animal, see Animal development DeVries, Hugo, 41, 49-50, 52 D'Hérelle, Felix, 504 Diabetes, 14 Diakinesis, 28, 31 Dicentric chromatid bridge, 154 Dicer enzyme, 260, 541, 542 2',3'-Dideoxyribonucleoside triphosphates, 388, 389 Differentiation, in Arabidopsis lifecycle, 34 Dihybrid cross, 44-45 Dimers, 19 Dimorphism, 100-101 Diploid, 22, 27, 90 Diploid zygote, 34 Diplonema, 28, 30, 31 Discontinuous synthesis, of DNA strand, 232, 233 Disjoin, 93 Dispersive replication, 222 Dissociation (Ds) element, 483-485 Disulfide bridges, 288 Divergence times, 667 Dizygotic (DZ) twins, 610, 627, 629,630 DMD gene, 276 DNA (deoxyribonucleic acid), 193-205 in cells, 19, 20 chemical bonds important in, 201 chemical subunits of, 198-199 in chromosomes, 22 controlled transcription of, 532-533 double helix structure of, 4, 201-204 and E. coli lactose operon, 518-519 evolution of DNA sequences, 668-669 gene expression using information in, 7-9 genetic information in, 195 - 197as genetic material, 6-10 illustration of, 1 methylation, 547-548

molecular organization of transcriptionally active, 545-546 mutation of, 9-10 negative supercoils in, 204-205 replication of, see Replication, of DNA and chromosomes RNA vs., 3-4 in sperm cells, 27 transformation mediated by, 194-195 unwinding, 236-238 variation of, 216 Watson and Crick's model of. 3-4 DNA analysis, 380-384 DnaA protein, 234 DNA-binding core domain (DBD), 595, 596 DNA-binding protein, 237 DnaB protein, 234 DNA chains, initiation of, 234-236 DnaC protein, 234 DNA DataBank of Japan (DDBJ), 400 DNA databases, 5, 400-401 DNA fingerprinting, 376, 456 DNA gyrase, 238 DNA helicases, 236–237 **DNA** libraries: construction of, 377-378 screening of, 378-380 DNA ligase, 228, 232-233, 370 DNA methylation, 549 DNA methyl transferases (DNMTs), 547, 548 DNA packaging, 208-212 DNA photolyase, 348 DNA polymerases: in E. coli, 231, 232 in eukarvotic chromosome replication, 246-247 multiple, 238-242 proofreading activities of, 238-242 translesion, 231 DNA polymerase I, 231 DNA polymerase II, 240 DNA polymerase III, 240-241 DNA polymerase IV, 240 DNA polymerase V, 240 DNA polymerase eta (η) , 231 DNA primase, 234

DNA prints, 456 DNA profiling, 455-461 forensic applications, 459-461 paternity tests, 459 DNA recombination, see Recombination DNA repair, 348-351 excision, 348-349 inherited human diseases with defects in, 351-354 light-dependent, 348 mismatch, 349-350 SOS response, 351 DNA repair enzymes, 320 DNA repetitive, 214–215 DNA replication, see Replication, of DNA and chromosomes DNase, 194, 206 DNase I, 545, 546 DNase I hypersensitive sites, 546 DNA sequences: databases of, 400-401 human origins and variation of, 677–679 in molecular control of transcription, 537-538 repeated, 214-215 DNA sequencing, 4-5 developments in, 398-400 and evolution, 10-11 growth of, 400-401 of human genome, 410-414 and variation, 661-662 DNA synthesis, 23 DNA testing: for cystic fibrosis, 448, 450 for Huntington's disease, 442.445 for sickle-cell anemia, 448-450 for Tay-Sachs disease, 340 DNA topoisomerase, 237, 238 DnaT protein, 234 DNMTs (DNA methyl transferases), 547, 548 Dobzhansky, Theodosius, 660-661 DOE (U.S. Department of Energy), 400 Dogs, blindness in, 439 Dolly (sheep), 18, 574 Domains, chromosome, 206 Dominance (principle of inheritance), 44 co-dominance, 63-64 incomplete, 63

Dominant activators, 589 Dominant genes, 42 in pedigrees, 53 recessive vs., 68-70 Dominant-negative mutations, 69 Dominant selectable marker gene, 372 Donahue, R. P., 152 Donor cell, 171, 172 Dorsal-ventral axis, in Drosophila, 562-563 Dosage compensation, of X-linked genes, 104–105, 551 Double crossover, 147, 154 Double Ds elements, 484 Double helix structure: alternate forms of, 203-204 diagram of, 201 of DNA, 201-204 heteroduplex, 174 Double-strand break model, 356 Double-stranded RNA (dsRNA), 471, 544 Doublié, Sylvie, 240 Down, Langdon, 120 Downstream region, 263 Downstream sequences, 264 Down syndrome, 120-122, 628 Drift: balance of mutation and, 650 - 651random genetic, 645-647, 670 Drosophila: anterior-posterior axis in, 562-565 body segmentation in, 565-567 chromocenters in, 118-119 cytological mapping of, 150 deletions/duplications in, 125 development of, 559-560 dorsal-ventral axis in, 562-563 dosage compensation of X-linked genes in, 104 heat-shock proteins in, 535 homeotic transformation in, 540 homologous polytene chromosome pairs in, 118 hybrid dysgenesis in, 485-487 hyperactivation of X chromosomes in, 552 maternal gene activity in, 562-565

organ formation in, 567–569

P elements in, 485-487, 497-498 position-effect variegation in, 545 rearrangement of chromosome structure in, 125 sex determination in, 102 speciation of, 675 specification of cell types in, 569-570 tissue-specific enhancers in, 538 transposon studies with, 496-498 ultrabithorax gene of, 276 variation among, 660, 661 X-linked genes and autosomal dominance, 97 Drosophila melanogaster: chromosome combinations in, 31 chromosome map of, 136 and chromosome theory of heredity, 92-98 DNA base composition in, 200 eye color in, 65-66, 68-69, 72 - 73genome sequencing of, 430, 431 as model organism, 32 Morgan's study of, 89 Drug-resistance genes, 481 Drug-resistant bacteria, 163-164, 171, 186 Ds element, 483-485 dsRNA, see Double-stranded RNA Duchenne muscular dystrophy, 53, 276 Duffy blood-typing system, 660 Duplex structure, 4 Duplication(s): chromosomal, 124, 125 mapping, 148-149 of nucleosomes, 247-248 Dwarfism, 15, 53, 366 Dynamic equilibrium, 647 Dysentery, 187, 504 Dysplastic cells, 601 Dystrophin, 374 DZ twins, see Dizygotic twins

E

East, Edward M., 609, 610, 614, 615 *easter* gene, 562 Ecdysone, 535 Edgar, Robert, 341, 342 Edward syndrome, 122 EES (ethyl ethane sulfonate), 326 EF (elongation factor), 302, 303 Effector molecules, 509 Egg cell, 3, 18, 26, 33-35 Egg coat, 35 Ehler-Danlos syndrome, 53 Electromagnetic spectrum, 328, 330 Electron microscopy, 211 Electropherograms, 456, 458 Electroporation, 464 Elongation: of polypeptide chain translation, 302-304 of RNA chains, 263-265, 271-272 Elongation factor (EF), 302, 303 EMBL (European Molecular Biology Laboratory), 400 Embryogenesis, 34 Embryoid bodies, 573 Embryonic stem (ES) cells, 464, 573-574 Embryo sac, 33, 34 EMS (ethyl methane sulfonate), 326 Encyclopedia of DNA Elements (ENCODE), 412, 413, 430 Endomitosis, 117-118 Endonuclease, 278 Endoplasmic reticulum, 20, 21, 23, 532 Endosperm, 34 End-product inhibition, 526 Energy sources, 170 Enfield, Franklin, 619 Engels, William, 485-486 Enhanceosome, 269 Enhancers, 271, 538, 546 Entrez retrieval system, 400-401 Environmental influence: on gene action, 70-71 on natural selection, 642-644 Environmentality, 627 Enzyme(s): in cells, 19 functions of, 67 one gene and one, 289-291 Enzyme activation, 506 Enzyme synthesis, 506-507 Ephrussi, Boris, 289

Epigenetic regulation, of gene expression, 545, 549 Episomes, 169, 176, 181 Epistasis, 72-75 Epistatic variance, 616 Epitwin project, 549 Equilibrium density-gradient centrifugation, 222 Equilibrium density gradients, 222 eRF, 304 Error-prone repair system, 351 ES (embryonic stem) cells, 464, 573-574 Escherichia coli: antibiotic-resistant, 186 bacteriophages in, see under Bacteriophages cell division in, 23 chromosome of, 206 conjugation in, 175 DNA base composition in, 200 DNA polymerase III in, 240-241 DNA polymerase I in, 231 DNA repair in, 348-351 F' in, 182 gene expression in, 506-507 gene transfer in, 172 Hfrs in, 181 lactose operon in, 511–519 linkage map of, 179 lysine riboswitch, 524 number of genes in, 259 plasmids in, 181 semiconservative replication in, 221–223 sequence-based map of, 424 size/gene content of, 424 translation in, 299-305 tryptophan operon in, 519-523 tryptophan synthetase in, 291-292 E (exit) site, 297, 298 Estrogen, 534, 535, 537 ESTs, *see* Expressed-sequence tags Ethics, 413-414 Ethyl ethane sulfonate (EES), 326 Ethyl methane sulfonate (EMS), 326

Euchromatin, 90, 545

Eukaryotes: cell division in, 23 cells of, 20, 21 chromosomes in, 20, 22 DNA base composition in, 200 introns, 274-275 meiosis in, 27 transposons in, 483-488 Eukaryote transcription and RNA processing, 267-274 addition of 5' methyl guanosine caps in, 272 elongation of RNA chains in, 271-272 initiation of RNA chains, 270-271 and posttranscriptional processing, 267 RNA editing, 273–274 RNA polymerases in, 267-270 termination of RNA chains in, 272-273 Eukaryotic chromosome(s), 207-216 centromeres/telomeres, 211-214 chemical composition of, 207 - 208DNA packaging levels in, 208-212 large DNA molecules in, 208 microscopic examination of, 22-23 repeated DNA sequences in, 214-215 Eukaryotic chromosome replication, 244-250 cell cycle, 245 duplication of nucleosomes at forks, 247-248 multiple DNA polymerases at single fork, 246-247 multiple replicons per chromosome, 245-246 telomerase, 248-249 telomere length and aging, 249 Eukaryotic gene expression, 531-553 about, 532-534 activation/inactivation of whole chromosomes, 550-553 and African trypanosomes, 531 alternate splicing of RNA, 533

and chromatin organization, 544-550 controlled transcription of DNA, 532-533 cytoplasmic control of mRNA stability, 533-534 dimensions of regulation, 532 induction of transcriptional activity, 534-537 molecular control of transcription, 537-541 posttranscriptional regulation by RNA interference, 541-544 sequences required for, 539 Eukaryotic genomes, 429-431 Eukaryotic protein production, in bacteria, 461-463 of human growth hormone, 461-462 with industrial applications, 462-463 European Conditional Mouse Mutagenesis Project, 469 European Molecular Biology Laboratory (EMBL), 400 Even-skipped mutants, 567 Evolution: of bacterial genomes, 186 Darwin's theory of, 10, 657-658 and recombination, 153-155 significance of genetic exchange in bacteria, 186 Wallace's ideas about, 10 Evolutionary genetics, 656-679 chromosome structure, 660-661 Darwin's theory of evolution, 657-658 discipline of, 658 evolution of DNA sequences, 668-669 evolution of protein sequences, 667-668 genetic variation, 659-662 human evolution, 676-679 molecular clock, 667 molecular evolution, 662-672 molecular phylogenies, 664 neutral theory of molecular evolution, 669–670 nucleotide sequences, 661-662 phenotypic evolution, 670-672 protein structure, 661

speciation, 672-675

Excinuclease, 348 Excision DNA repair, 348-349 Excitation, 329-330 Exit (E) site, 297, 298 Exons, 257, 274 Exon shuffling, 671 "Experiments in Plant Hybridization" (Gregor Mendel), 40 Expressed sequences, 416 Expressed-sequence tags (ESTs), 403, 416, 418 Expression domain, riboswitch, 524, 525 Expressivity, 71–72 Extensively drug-resistant (XDR) bacteria, 163-164 Eye color mutation, 92–95 Eye development, 567-571 F F⁺ cells, 176, 177, 181 F⁻ cells, 176–179, 182 F factors, 176-179, 181-182 F' factors, 181–182

FACT protein complex, 272 Familial adenomatous polyposis (FAP), 592, 598 Familial breast cancer, 592 Familial melanoma, 592 Fanconi anemia, 352, 353 FAP (familial adenomatous polyposis), 592, 598 FASTA search engine, 421 Fatty acids, 19 Federal Bureau of Investigation (FBI), 456 Federoff, Nina, 483 Feedback inhibition, 526 Felis domesticus, 36 Female gametophyte, 33, 34 Female meiosis, 33, 35 Ferns, 90 Fertile polyploids, 116–117 Fertility, 93-94, 100 Fertilization, 34, 41, 42 Fibroin, 303-304 Filament, of flower, 33 Filial generation, 43 Fingerprinting, DNA, 456 Fire, Andrew, 471 Fischer, R. A., 645 FISH (fluorescent in situ hybridization), 405, 409 Fisher, R. A., 614, 658 Fission, 23

Fitness, 641-642 5' cap, 267 Flagellum, 21 Fleming, Alexander, 163 Fluorescent in situ hybridization (FISH), 405, 409 Folded genome, 206 Forensics, 459-461 Forked-line method, 46, 47 Forward mutation, 335 Fossils, 663 Fossil record, 676-677 454 sequencing, 391 F pili, 176 Fraenkel-Conrat, Heinz, 197 Fragile X syndrome: DNA testing for, 450 as neurodegenerative abnormality, 442 and trinucleotide repeats, 443-444 Frameshift mutations, 323 Franklin, Rosalind, 200, 201 Free amino group, 286 Free carboxyl group, 286 Free ribosomes, 21 Frequency distributions, 51, 611-612 Friedreich ataxia, 442 Fruit fly, see Drosophila Functional allelism, 344, 346 Functional constraint, 666-667 Functional genomics, 402, 417 Fungi, 20 Fur coloration, 104, 105 fushi tarazu mutants, 567

G

G, see Guanine G₁ phase, cell cycle, 23 G, phase, cell cycle, 23 Gain-of-function mutations, 69-70 GAL4 transcription factor, 539 Galactosemia, 53 β-galactosidase, 373 Galapagos Islands, 658, 675 Gametes, 3, 22 Gametogenesis, 35, 36 Gametophytes, 33 Gaps, in DNA, 233 Gap genes, 567 Garrod, Sir Archibald, 14, 67 Gates, in cell walls, 19

and recombination frequency,

Gauguin, Paul, 656 GC box, 270, 271 GCG (Genetics Computer Group), 421, 422 Gehring, Walter, 568 Gel electrophoresis, 381–383, 385-387,661 Gelsinger, Jesse, 452 GenBank, 400-401 Gene(s): calculating distances between, 143 - 145in chromosomes, 20, 22-23 cloning large, 374 colinearity, 291-292 complementation test, 342-346 drug-resistance, 481 function of, 67-68 interrupted, 274-276 Mendel's "factors" as, 42 Mendel's study of, 3 number of human, 412 one gene-one enzyme, 289-291 one gene-one polypeptide, 289-292 sex-linked, 98-100 structure, 388-392 Gene action, 70-76 environmental effects on gene expression, 70-71 environmental influence, 70 epistasis, 72-75 interactions, 72 penetrance/expressivity, 71-72 pleiotropy, 75-76 Gene additions, 455 Gene chips, 400, 416-419, 662 Gene conversion, 356–358 Gene density, 431 Gene expression: and chromatin remodeling, 269 - 270eukaryotic, see Eukaryotic gene expression as gene function, 193 information flow from, 8-9 Mendel's principles applied to, 71-72 mRNA intermediary, 259, 261 process of, 7-9, 259-263 prokaryotic, see Prokaryotic gene expression RNA synthesis, 261–262 stages of, 7

Gene expression cassette, 471, 472 Gene families, 423 GeneFinder, 423 Gene function: and allelic variation, 63-70 with polypeptides, 67-68 Gene interactions, 72 Gene level, natural selection at. 642-644 Gene mapping: of bacteriophages, 166 of chromosomes, 141-148 with conjugation data, 180 cytogenetic, 148-150, 405 of human genome, 409-410 with nucleotide sequences, 387-391 with partial diploids, 183 physical, 405-406 and positional cloning, 402 RFLP/STR, 403-405 GeneMark, 423 Gene order, 142-143 Generalized transduction, 183 Gene Recognition and Analysis Internet Link (GRAIL), 423 Gene replacements, 455 GeneScan, 423 Gene structure, 388–392 colinearity between coding sequence/polypeptide product, 291-292 recombination between adjacent nucleotide pairs, 389-391 Gene therapy, 15, 439-440, 450-455 Genetics, 10-15 in agriculture, 12–14 classical, 11 and cytology, 89 and evolution, 10-11 evolutionary, see Evolutionary genetics in medicine, 14-15 molecular, 11-12 population, see Population genetics in society, 15 Genetically effective population sizes, 651 Genetically modified (GM) crops, 466 Genetically modified (GM) foods, 466

Genetically modified organisms (GMOs), 13-14, 466 Genetic analysis: of inbreeding, 78-82 levels of, 11-12 Genetic code, 306–312 deciphering, 307-309 degeneracy of, 310-312 in gene transcription, 257 initiation/termination codons, 309-310 nearly universal, 312 ordered, 311-312 properties of, 306 three nucleotides per codon, 306-309 Genetic control, of recombination, 155 Genetic counseling: allele frequencies in, 640-641 and cancer, 600 Mendel's principles applied to, 54-56 Genetic counselors, 14 Genetic distance, see Genetic map distance Genetic drift, random, 645-647, 670 Genetic elements, transposable, see Transposons Genetic equilibrium, 647–651 balancing selection, 648-649 mutation-drift balance, 650-651 mutation-selection balance, 649-650 Genetic exchange mechanisms, in bacteria, 172-186 conjugation, 175-180 evolutionary significance of, 186 and plasmids/episomes, 179-181 sexduction, 181-182 transduction, 172, 182-185 transformation, 172-175 Genetic information: in DNA, 195-197 in RNA, 197 transfer of, 257 Genetic Information Nondiscrimination Act (GINA), 414 Genetic map distance, 143-145 calculating, 143-145 and physical distance, 149-150

146-147 Genetic material, functions of, 193 Genetic mosaics, 104, 105, 463 Genetic pathways, to cancer, 600-603 Genetic relationships, measuring, 82 Genetics Computer Group (GCG), 421, 422 Genetic selection, of DNA libraries, 378–379 Genetic symbols, 68 Genetic variability, 321 Genetic variation, 659-662 in chromosome structure, 660-661 in nucleotide sequences, 661-662 in phenotypes, 659-660 in protein structure, 661 Gene transfer, unidirectional, 171 Genome(s): chloroplast, 426-429 cloning segments of, 374 defined, 1, 4 eukaryotic, 429-431 evolution of, 186 mitochondrial, 427-429 Neanderthal, 397–398 prokaryotic, 424-425 synthetic bacterial, 425-426 transposons and organization of. 498-499 Genome function, assays of, 415-419 Genomics: about, 402 assays of genome function, 415-419 chromosome jumps, 408-409 chromosome walks/jumps, 408-409 comparative, see Comparative genomics correlated maps of chromosomes, 402-406 cytogenetic maps, 405 defined, 5 developments in, 398-400 physical maps/clone banks, 405-406 position-based cloning maps, 407-409 RFLP maps, 403-405

Genomic DNA libraries: construction of, 377 screening of, 378-380 Genotypes: defined. 43 nomenclature for, 170-171 Genotype frequencies, 636 Germinal mutation, 332-333 Germ line, 22 Germ-line gene therapy, 451 Gever, Pamela, 538 GFP (green fluorescent protein), 419-420 Giemsa, Gustav, 112 Giemsa stain, 112 Gilbert, Walter, 388, 671-672 GINA (Genetic Information Nondiscrimination Act), 414 α-Globin, 664–667 β-Globin, 8, 9, 274–275, 314, 323, 546 Globin genes, 338-339, 671 Glucocorticoids, 535 Glucose, 19, 506 Glucose effect, 514 Glutamic acid, 9 Glutamines, 526 Glycogen storage disease, 53 GM (genetically modified) crops, 466 GM (genetically modified) foods, 466 GMOs (genetically modified organisms), 13-14, 466 Goad, Walter, 400 Golgi complex, 20, 21, 23 Grafts, 116 GRAIL (Gene Recognition and Analysis Internet Link), 423 Grasses, 431-432 Grasshoppers, 90, 91, 140 Great apes, 676 Green fluorescent protein (GFP), 419-420 Greider, Carol, 248 Griffith, Frederick, 173, 174, 194 Grosbeaks, 675 Groudine, Mark, 545 Growth, in Arabidopsis lifecycle, 34 Growth promoters, 187 Guanine (G), 4, 198-200, 202, 213 Guide RNAs, 274

Gusella, James, 440 Gynandromorphs, 122 gypsy element, 492–493

Н

Haas, Corey, 439, 440 Haemanthus, 25 Haemophilus influenzae, 174, 424 Hairpin structures, 237, 265-266, 444, 471, 472, 521-522 Haldane, J. B. S., 658 Hamkalo, Barbara, 266, 303 Hanahan, Douglas, 602 Haplo-diplo system, of sex determination, 102-103 Haploid, 22, 27, 31, 90, 205 Haploid genome, 90 Haplopappus gracilis, 249 Haplotypes, 414-415 HapMap Project, 414-415 Hardy, G. H., 636 Hardy-Weinberg principle, 636-640 applications of, 636-638 exceptions to, 638-640 Harris, H., 661 HATs (histone acetyl transferases), 547 Haw River syndrome, 442 Haves, William, 177 HBB human gene, 8 HD, see Huntington's disease HDACs (histone deacetylases), 547 Heat-shock proteins, 535 Heat-shock response elements (HSEs), 535 Heavy chains, 576 Helix-loop-helix motif, 540 Helix-turn-helix motif, 540 Hemizygote, 92 Hemoglobin, 338-339, 664-667,671 Hemophilia, 98-100 Hereditary nonpolyposis colorectal cancer (HNPCC), 592.598-599 Heredity, chromosome theory of, 92-98 Heritability: broad-sense, 615-616, 626 narrow-sense, 616-618, 626 Heritable gene therapy, 451 Hermaphrodites, 551

Herpes simplex virus (HSV), 200, 467, 468 Herrick, James, 285 Hershey, Alfred, 195, 196 Heteroalleles, 346 Heterochromatin, 90, 545 Heteroduplex, 57, 174 Heterogametic sex, 102-103 Heterogeneous nuclear RNA (hnRNA), 267 Heterologous chromosomes, 128 Heterologues, 27 Heterosis, 78 Heterozygosity, 646, 651 Heterozygous genes, 42 Hfr cell, 177-179, 181-182 hGH (human growth hormone), 14-15, 461-462 Hiatt, Jessie, 332 Histidine operon, 522 Histones, 207-211, 272, 544 Histone acetylation, 549 Histone acetyl transferases (HATs), 547 Histone deacetylases (HDACs), 547 Histone methyl transferases (HMTs), 547 HIV, see Human immunodeficiency virus H.M.S. Beagle, 657-658 H.M.S. Bounty, 634, 647 HMTs (histone methyl transferases), 547 HNPCC (hereditary nonpolyposis colorectal cancer), 592, 598-599 hnRNA (heterogeneous nuclear RNA), 267 hobo element, 485 Holley, Robert W., 297, 309, 398 Holliday, Robin, 354 Holliday model, 354–357 Holoenzyme, 240-241, 263, 264 Homeodomain, 540, 566 Homeotic genes, 565-566 Homeotic mutations, 566 Hominin, 676-677 Homoalleles, 346 Homo erectus, 676, 677 Homo ergaster, 676, 677 Homogametic sex, 102-103 Homo habilis, 676 Homologous, 423-424

Homologous chromosomes, 27, 28, 30, 31, 128 Homologous DNA sequences, 664 Homologues: defined, 27, 423 of invertebrate genes, 571-572 of viral oncogenes, 586-587 Homo neanderthalensis, 397-398 Homo rudolfensis, 676 Homo sapiens, see Human beings Homozygosity, 650 Homozygous genes, 42 Honeybees, 103 Hormones, 534-537 Hormone receptors, 535 Hormone response elements (HREs), 535, 537 HSEs (heat-shock response elements), 535 HSV, see Herpes simplex virus Hubby, J. L., 661 Huberman, Joel, 245 Hughes, Walter, 223 HUGO (Human Genome Organization), 409 Human beings (Homo sapiens): aging in, 249, 352 and apes, 676 behavioral traits of, 628-630 chromosomes of, 22, 27, 90-91 DMD gene of, 276 DNA base composition in, 200 and DNA repair defects, 351-354 DNA sequence variation, 216 gene therapy for, 450-455 globin mutation in, 338-339 identifying defective genes, 440-448 immune cell differentiation in, 577-578 inactivation of X-linked genes in, 105 linkage analysis in, 150-153 metabolic blocks in, 339-340 as model organism, 36 mtDNA of, 428 mutation of, 339-340 and Neanderthals, 398 number of chromosomes in, 111, 113 protein domains in, 431

recombinant DNA technology in, 440-448 sex-linked genes in, 98-100 telomere length and aging in, 249 transposable elements in, 494-496 trisomy in, 120-122 variation among, 660 Human diseases. See also specific diseases, e.g.: Tay-Sachs disease DNA repair defects and inherited, 351-354 molecular diagnosis of, 448-450 recombinant DNA technology for detection of, 440-448 and trinucleotide repeats, 443-444 Human evolution, 676–679 DNA sequence variation and human origins, 677-679 in fossil record, 676-677 and great apes, 676 Human genetics, Mendel's principles applied to, 52-57 Human genome: chromosomes in, 7 mapping, 409-410 sequencing, 386, 410-414 variations in, 412 Human Genome Organization (HUGO), 409 Human Genome Project, 1, 4–5, 56, 398, 409-415 HapMap Project, 414-415 mapping human genome, 409-410 sequencing human genome, 410-414 Human β-globin, 8, 9, 274–275, 314, 323, 546 Human growth hormone (hGH), 14-15, 461-462 Human immunodeficiency virus (HIV), 488-492 DNA integration into host cell, 490, 492 genome replication, 490, 491 life cycle of, 489–490 Human insulin, 14 Human karyotype, cytological analysis of, 113-114

Human origins, DNA sequence variation and, 677–679

Human Proteome Organization (HUPO), 413 hunchback gene, 563, 564 Huntington's disease (HD): behavior phenotype for, 628 DNA testing for, 450 as inherited condition, 53 recombinant DNA technology with, 440-442, 445 and trinucleotide repeats, 444 HUPO (Human Proteome Organization), 413 Hutchinson-Gilford syndrome, 249 Hybrid dysgenesis, 485-487 Hybridization: fluorescent in situ, 405, 409 molecular, 379-380 northern blot, 382, 384 Southern blot, 381–383 Hydrogen bonding, 201 Hydrogen bonds, 288-289 Hydrophilic (term), 19 Hydrophobic (term), 19 Hydrophobic bonds, 201, 203 Hydrophobic interactions, 288, 289 Hydroxylamine, 326, 328 Hydroxylating agent, 326, 328 Hylobates concolor, 434 Hyperactivation: of X chromosomes, 552 of X-linked genes, 104 Hyperploid, 119 Hypomorphic alleles, 65 Hypoploid, 119 Hypothesis, testing, 48–52

Identical twins, *see* Monozygotic twins Identification, DNA profiling for, 459 Identity by descent, 79 Ideogram, 114 IFs (initiation factors), 299–301 Illumina sequencing, 391 Imaginal discs, 560 Imago, 560 Immune cells, differentiation of, 575–578 Immune system, 280 Immunoglobins, 575 Imprinting, 48–550 Inactivation: of whole chromosomes, 550-553 of X-linked genes, 104-105 Inborn Errors of Metabolism (Archibald Garrod), 14 Inbred line, 77 Inbreeding, 77-82 compound, 81 effects of, 77-78 genetic analysis of, 78-82 and Hardy-Weinberg frequencies, 639 and measuring genetic relationships, 82 Inbreeding coefficient, 79-82 Inbreeding depression, 77-78 Incomplete dominance, 63 Incomplete P elements, 87 Incomplete penetrance, 71, 72 Independent assortment (principle of inheritance), 44-46, 95-97 Induced mutation, 324-326, 331 by chemicals, 326-328 by radiation, 328-331 spontaneous vs., 333 by transposable genetic elements, 331 Inducers, 507, 534 Inducible enzymes, 506 Inducible genes, 506 Inducible operon, 513-514 Induction: and cell type specification, 570 and gene expression, 506 lactose operon in E. coli, 513-514 and operons, 510 Induction of transcriptional activity, 534-537 and signal molecules, 535-537 and temperature, 535 Industrial applications, of eukaryotic protein production, 462-463 Inheritance, Mendel's principles of, see Mendelian principles Inheritance of complex traits, 607-630 artificial selection, 618-620 broad-sense heritability, 615-616 cardiovascular disease, 607 correlations between relatives, 624-628

genetic/environmental factors influencing, 608 human behavioral traits, 628-630 intelligence, 628-629 multiple factor hypothesis, 614 narrow-sense heritability, 616-618 partitioning of phenotypic variance, 614-615 personality, 629-630 predicting phenotypes, 617-618 quantifying traits, 608 quantitative trait loci, 620-624 statistics of quantitative genetics, 611-613 threshold traits, 610 Inherited human diseases, 351-354 Initiation: of polypeptide chain translation, 298-302 of RNA chains, 263-264, 270-271 Initiation codon, 299 Initiation complex, DNA replication, 234 Initiation factors (IFs), 299-301 Inman, Ross, 228 Inosine, 313 Insensitivity, to growth-inhibitor signals, 602 Insertional mutagenesis, 469 Insertion mutations, 572-573 Insertion sequences (IS elements), 181, 479-481 in situ colony hybridization, 379-380 Insulin, 14, 461, 535, 668 Integrated state, 176 Inteins, 289 Intelligence, 617-618, 628-629 Intelligence quotient (IQ), 617-618, 628-629 Intercellular communication, 533 Intercross, 46 Interference, 145 International HapMap Project, 415 International Human Genome Sequencing Consortium, 386, 411 Interphase, 24, 25 Interrupted genes, 274-276

Introns, 257, 274, 276 biological significance of, 276 in eukaryotic DNA, 431 excision by RNA splicing, 277-280 RNA splicing, 533 Inversion(s): of chromosome structure, 126-127 suppression of recombination by, 153–154 Invertebrate genes, vertebrate homologues of, 571-572 Inverted terminal repeats, 479-480 in vitro culture: of DNA polymerases/ synthesis, 231 RNA interference with, 471, 472 Ionic bonds, 288 Ionizing radiation, 328-331 IQ (intelligence quotient), 617-618, 628-629 Irons, Ernest, 285 IS elements (insertion sequences), 181, 479-481 Isoalleles, 337 Isochromosome, 128 Isoforms, 423 Isoleucine, 310

J

Jacob, François, 177, 178, 181, 509, 511, 512, 518, 534 Jellyfish, 419 Jimson weed, 119 Johannsen, Wilhelm, 42, 608, 614 Jumping genes, 187 Jürgens, Gerd, 562

Κ

Kappa polypeptide, 576, 577 Karyotype, human, 113–114 Kaufman, Thomas, 566 Kennedy's disease, 444 Khorana, Ghobind, 309 Kidd, Jeffrey M., 412 Kidwell, James, 485 Kidwell, Margaret, 485 Kimura, Motoo, 645 Kinases, 547 Kinetochores, 25, 26, 30 King, Jonathan, 341, 342 Klebsiella pneumonia, 186 Klinefelter, H. F., 121 Klinefelter syndrome, 121, 122 Knockout Mouse Project, 469 Knockout mutations, 467–469, 572–573 Knudson, Alfred, 591 Knudson's two-hit hypothesis, 590–593 Kornberg, Arthur, 231 Kozak, Marilyn, 301 Kozak's rules, 301 Kreitman, Martin, 661

L

L1 element, 494, 495 Lactose metabolism, 534 Lactose operon, in E. coli, 511-519 catabolite repression, 514-515, 517 constitutive mutations in, 513 induction, 513-514 protein-DNA interactions controlling, 518–519 Lagging strand, 232, 233 Lahn, Bruce, 154 Lamarck, Jean, 334 Lamarckism, 334 Landsteiner, Karl, 63-64 LANL (Los Alamos National Laboratory), 400 Larva, 560 Lateral elements, 30 Lathyrus odoratus, 73–74 Lawler, S. D., 151, 152 LCR (locus control region), 546 Leading strand, 232 Leber's congenital amaurosis type II, 439 Leder, Philip, 275, 309 Lederberg, Esther, 334, 335 Lederberg, Joshua, 175, 182, 183, 334, 335 Leptonema, 28, 30 Lethal mutations, 67, 325 Leucine zipper motif, 540 Levan sucrase, 374 Levis, Robert, 493 Lewis, Edward B., 342, 343, 566 Lewontin, R. C., 661 Liability variable, 610 Li-Fraumeni syndrome, 592

Ligand, 70

Ligases, splicing, 278 Light chains, 576-577 Light-dependent DNA repair, 348 Lilium longiflorum, 28–29 Lim, Johng, 486, 498 LINEs (long interspersed nuclear elements), 494, 496 Linkage, recombination and, 136-138 Linkage analysis, in humans, 150-153 Linkage phase, 138 Linkers, 209 Lipids, 19 Lippman, Zachary, 621-623 Locus, 620 Locus control region (LCR), 546 Locusts, 504 Long interspersed nuclear elements (LINEs), 494, 496 Long-lived mRNAs, 533-534 Long terminal repeats (LTRs), 490, 492 Loops, 206 Los Alamos National Laboratory (LANL), 400 Loss-of-function alleles, 68-69 LTRs (long terminal repeats), 490, 492 LTR retrotransposons, 492 Lwoff, André, 169 Lycopersicon, 620–623 Lyon, Mary, 104 Lysenko, Trofim, 334 Lysine, 207 Lysine riboswitch, 524-525 Lysogenic phage, 168, 169 Lysosomes, 20, 21 Lytic phage, 166, 168–169 М McCarty, Maclyn, 174, 194, 195 McClintock, Barbara, 139, 213, 483-485

McClung, C. E., 91

Maize:

139

81-82

in, 477

MacLeod, Colin, 174, 194, 195

Ac and Ds elements in, 483-485

homologous chromosomes in,

transposable genetic elements

genome sequencing of, 432

inbred varieties of, 78-79,

Male gametophyte, 33, 34 Male meiosis, 33, 35, 36 Male-specific-lethal (MSL) proteins, 552 Mammals: genome evolution in, 432-434 inactivation of X chromosomes in, 551-552 inactivation of X-linked genes in, 104–105 stem cells in, 573-574 Mammuthus prinigenius, 429 Map distances, 166 Mapping genes, see Gene mapping Marchantia polymorpha, 428, 429 Marfan syndrome, 53 Mason, James, 493 Mastectomy, 600 Maternal-effect genes, 561 Maternal-effect mutations, 561 Maternal gene activity, 561-565 in Drosophila, 562-565 maternal-effect genes, 561 Mating: consanguineous, 77, 79-81, 638 nonrandom, 638-639 in S. cerevisiae, 33 Maxam, Allan, 388 Maxam and Gilbert procedure, 388 MDR (multi-drug-resistant), 163-164, 186-187 Mean class, 612 Measure of genetic distance, 141 Measuring genetic relationships, 82 MECP2 protein, 270 Mediator complex, 538 Medicine: and antibiotic-resistant bacteria, 187 and drug-resistant bacteria, 186 genetics in, 14-15 Megagametogenesis, 34 Megaspores, 33, 34 Megasporogenesis, 34 Meiosis, 27–32 first stage, 27-28, 30-31, 34, 35 independent assortment during, 95-97 in Lilium longiflorum, 28-29 second stage, 29, 31-32, 34, 35 segregation of alleles during, 93-94

Malaria, 10, 398, 649

Meiosis I, 27-28, 30-31, 34, 35 Meiosis II, 29, 31-32, 34, 35 Melanocytes, 104 Melanoma, 592 Mello, Craig, 471 Membranes, cell, 19, 21 Membrane-bound receptor proteins, 535 Mendel, Gregor, 2-3, 40, 41, 67, 398 1866 paper of, 3, 40, 41 and evolutionary genetics, 658 experiments of, 41-46 Mendelian principles, 2–3, 40-57, 62-82 allelic series, 65 chi-square test, 50-52 dominant vs. recessive genes, 68 - 70effects of mutation, 66-67 environmental influence, 70-71 epistasis, 72-75 expressivity, 71-72 forked-line method, 46-47 gene interactions, 72 genetic counseling, 54-57 in human genetics, 52-57 inbreeding, 77-82 incomplete dominance and codominance, 63-64 multiple alleles, 64-65 pedigrees, 53, 56 penetrance and expressivity, 71-72 pleiotropy, 75-76 polypeptides, 67-68 probability method, 47-48 Punnett square method, 46 segregation in human families, 54 testing mutations for allelism, 65-66 "Mendelism-the Principles of Dominance, Segregation, and Independent Assortment" (William Bateson), 62 Mendel's Principles of Heredity (William Bateson), 62 Meselson, Matthew, 221–224 Messenger RNA (mRNA), 7 altering information content of, 273-274 cytoplasmic control of stability of, 533–534 in gene translation, 257, 260

as intermediary in gene expression, 259, 261 and intron splicing, 533 long-/short-lived, 533-534 in RNA synthesis, 261-262 Messing, Joachim, 483 Metabolic block, 337, 339-340 Metabolic pathways, mutation in human, 339-340 Metabolite, 170-171 Metafemales, 94 Metaphase, 25, 26, 28, 29, 31 Metaphase I, 28, 31 Metaphase II, 29, 31 Metaphase plate, 26 Metastasis, 582 Methionine (Met), 8 Methylation, 269, 368-370, 547-548 Methyl guanosine caps, 272 Microarrays, 400, 416-419 Micrococcus lysodeikticus, 200 Microfilaments, 21 Microprojectile bombardment, 464 Micropyle, 559 MicroRNAs (miRNAs), 259, 260, 471, 534, 541-544 Microsatellites, 405 Microspores, 33 Microtubules, 21, 24-26, 30 Microtubule organizing centers (MTOCs), 24, 25 Midparent value, 617 Miescher, Johann Friedrich, 192 Migration, 640 Milk, 464 Miller, Oscar, 266, 303 Minimal medium, 289 Minisatellites, 405 Minnesota Study of Twins Reared Apart, 629 mir genes, 542, 543 miRNAs, see MicroRNAs Mismatch DNA repair, 349-350 Missense mutations, 313 Mitochondria: in cell division, 22, 23 in cells, 20, 21 differences in, 312 function of, 532 genomes of, 427-429 and RNA editing, 273-274 Mitochondrial DNA (mtDNA), 427-428,665

Mitochondrial RNA, 428 Mitochondrion, 21 Mitosis, 24-26 in cell cycle, 23 cytological analysis of chromosomes in, 111-113 meiosis vs., 27 process of, 24-26 Mizutani, Satoshi, 488 Modal class, 612 Model organisms, 32-36 Arabidopsis thaliana, 32-34 Caenorhabditis elegans, 32, 559 Danio rerio, 32 defined, 32 Drosophila, 559-560 Drosophila melanogaster, 32 Mus musculus, 32, 34-36 Saccharomyces cerevisiae, 32–33 Molecular analysis, 380-391 for cystic fibrosis, 383 northern blot hybridization RNA analysis, 382, 384 nucleotide sequence mapping, 387-391 restriction endonuclease mapping, 386-387 reverse transcriptase-PCR RNA analysis, 384, 385 Southern blot hybridization DNA analysis, 381–383 western blot protein analysis, 384-385 Molecular biology, central dogma of, 9, 257 Molecular clock, 667 Molecular control, of transcription, 537-541 DNA sequences involved in, 537-538 proteins involved in, 538-540 Molecular diagnosis, of human diseases, 448-450 Molecular evolution, 662-672 molecular clock, 667 molecular phylogenies, 664 Neutral Theory of, 669-670 and phenotypic evolution, 670-672 rates of, 664-667 variation in DNA sequences, 668-669 variation in protein sequences, 667-668 Molecular genetics, 11-13, 439-472

cystic fibrosis, 445-448 DNA profiling, 455-461 eukaryotic protein production in bacteria, 461-463 gene therapy, 450-455 human growth hormone, 461-462 Huntington's disease, 440-442, 445 industrial applications, 462-463 molecular diagnosis of human diseases, 448-450 mouse knockout mutations, 467-469 recombinant DNA technology, 440-448 reverse genetics, 467-473 RNA interference, 471, 472 T-DNA and transposon insertions, 469-471 transgenic animals and plants, 463-466 Molecular hybridization, of DNA libraries, 379-380 Molecular phylogenies, 664 Molecules, as documents of evolutionary history, 663 Monod, Jacques, 509, 511, 512, 518, 534 Monogenic mRNAs, 267 Monohybrid cross, 42-44 Monoploid, 205 Monosomy, 121–124 Monozygotic (MZ) twins, 220, 549, 610, 623, 625, 627, 629,630 Moore, Graham, 431 Morgan, Lillian, 128, 129 Morgan, Thomas Hunt, 89, 92-94, 135 Morphogens, 565 Morphogenesis, 341-342 Mosaicism, 122 Mother cell, 23, 27 Motility, cell, 20 Mouse: eve development, 569 genome sequencing of, 431 and human chromosomes, 433 inactivation of X-linked genes in, 104 knockout mutations in, 467-469 as model organism, 32, 34-36 mutations in the, 572-573

Mouse: (Continued) pneumococcal experiments with, 173, 174 transgenic, 463-464 yellow-lethal mutation in, 67 Mouse ear cress, see Arabidopsis thaliana M phase, cell cycle, 23 mRNA, see Messenger RNA mRNA degradation, 266 mRNA templates, 298-305 MSL proteins (male-specificlethal proteins), 552 mtDNA (mitochondrial DNA), 427-428,665 MTOCs (microtubule organizing centers), 24, 25 Muller, Hermann J., 212, 324–325 Müller-Hill, Benno, 512 Multi-drug-resistant (MDR), 163-164, 186-187 Multigenic mRNAs, 267 Multimers, 19 Multinemes, 208 Multiple alleles, 64-65 Multiple Factor Hypothesis, 614 Multiple replicons per chromosome, 245-246 Multiregional Hypothesis, 677 Muntjac (Asian deer), 90, 112, 433 Murein, 19 Muscular dystrophy, 53, 276 Mus musculus, see Mouse Mustard gas, 326 μ symbol, 614 Mutagens, 333, 496, 497 Mutagenicity: Ames test for, 346–348 of X rays, 324-325 Mutants, 321 Mutant genes: and alleles, 64 in bacteria, 170-171 blocked in ability to utilize specific energy sources, 170 drug-/antibiotic-resistance in, 171, 333-335 and synthesis of essential metabolite, 170-171 Mutation(s), 320-358 Ames test, 346-348 of cellular oncogenes, 587-589 as changing genetic information, 9-10

chemical-induced, 326-328 complementation test for, 342-346 conditional lethal, 340-342 defined, 9 and DNA recombination mechanisms, 354-358 and DNA repair mechanisms, 348-351 dominant vs. recessive, 68-70 and evolution, 669-670 eye color, 92-95 and eye development, 571 gap-gene, 567 as gene function, 193 homeotic, 566 in human globin genes, 338-339 in human metabolic pathways, 339-340 induced, 324-326, 331 inherited human diseases, 351-354 knockout, 467-469 in lactose operon, 513 maternal-effect, 561 missense, 313 molecular basis of, 321-332 nonsense, 313 pair-rule, 567 phenotypic effects of, 337-342 radiation-induced, 328-331 as random/nonadaptive process, 333-335 recessive, 337-338 reverse, 336 segment-polarity, 567 sickle cells, 9-10 somatic vs. germinal, 332-333 spontaneous vs. induced, 333 suppressor, 307, 313-314 Tay-Sachs disease, 340 testing for allelism, 65-66 tfm, 102 transposon-induced, 331 trinucleotide repeats, 331-332 variability through, 321 variation as effect of, 66-67 in viruses, 166 xeroderma pigmentosum, 320 Mutation-drift balance, 650-651 Mutation-selection balance, 649-650 Mycobacterium tuberculosis, 163-164, 186, 424

Mycoplasma genitalium, 7, 425 Mycoplasm mycoides, 425–426 Myoglobin, 671 Myotonic dystrophy, 442 MZ twins, *see* Monozygotic twins

Ν

n (chromosome number), 90 Nail-patella syndrome, 151–152 Nalidixic acid, 238 nanos gene, 563, 564 Narrow-sense heritability, 616-618, 626 NASC (Nottingham Arabidopsis Stock Centre), 470 Nathans, Daniel, 368 National Center for Biotechnology Information (NCBI), 5, 216,400 National Institutes of Health (NIH), 52, 400 National Library of Medicine (NLM), 400 National Science Foundation (NSF), 400 Natural selection, 641-645 Darwin's theory of, 657-658 at gene level, 642-644 Navel orange, 332 NCBI, see National Center for Biotechnology Information NDM-1 gene, 186 Neanderthals, 397-398, 677 Negative control mechanisms, 507-509 Negative supercoils, 204-205, 238 Neisseria gonorrhoeae, 174 Neomycin, 467-469 Neomycin phosphotransferase type II (NPTII), 466 Neurofibromatosis, 53, 592 Neurospora crassa, 289-291 Neutral mutations, 337 Neutral Theory of molecular evolution, 669-670 Nicholas, Czar, 98, 99 Nicked domains, 206 Nicotiana tabacum, 429 Night blindness, 53 NIH (National Institutes of Health), 52, 400 Nijmegan breakage syndrome, 352, 353 Nilsson-Ehle, Herman, 608-609, 614

Nirenberg, Marshall, 309 Nitrates, 347 Nitrous acid, 326-328 NLM (National Library of Medicine), 400 Nomenclature, 64, 67, 68, 170 - 171Nomura, Masayasu, 295 Noncomplementation, 345 Nonconjugative plasmids, 181 Nondisjunction, 93-95, 124 Nonheritable gene therapy, 451 Nonhistone chromosomal proteins, 207, 208, 211 Nonionizing radiation, 328-330 Nonpolypoid colorectal cancer, 55 Nonrandom mating, 638–639 Nonsense mutations, 313 Nonsynonymous substitution, 669 Nontemplate strand, 261 Nopaline, 465 Normal cellular oncogenes, 587 Normal distribution, 613 North American Conditional Mouse Mutagenesis Project, 469 Northern blot hybridization, 382, 384 NotI, 368, 380 Nottingham Arabidopsis Stock Centre (NASC), 470 NPTII (neomycin phosphotransferase type II), 466 NSF (National Science Foundation), 400 Nuclear envelope, 21 Nuclear membrane, 25 Nuclear pore, 21 Nucleic acids, 3, 19, 198 Nuclein, 192 Nucleolar organizer regions, 296 Nucleolus, 21, 25, 295-296 Nucleosomes, 207, 209-210, 247-248, 269 Nucleosome core, 209 Nucleotides: components of, 3, 198-199 recombination between adjacent pairs of, 389-391 structure of, 3 three nucleotides per codon, 306-309 types of, 4 Nucleotide excision repair, 348

Nucleotide repeats, 331–332 Nucleotide sequences: mapping of, 387–391 variation in, 661–662 Nucleus, 20, 21, 532–533 Null alleles, 65, 337 Nullo-X egg, 94 Nüsslein-Volhard, Christiane, 562, 566

0

Ochoa, Severo, 309 Octamer box, 270, 271 Octaploid, 115 Octopine, 465 Octoploid, 90 OD (oligomerization domain), 595, 596 Okazaki, Reiji, 232, 233 Okazaki, Tuneko, 232, 233 Okazaki fragment, 232–235 Olbrycht, T. M., 142-145 Oligomerization domain (OD), 595, 596 Oncogenes, 585-590 mutant cellular, 587-589 proto-, 586-587 tumor-inducing retroviruses and viral, 585-586 1000 Genomes Project, 216 One gene-one enzyme, 289-291 One gene-one polypeptide, 289-292 Oogenesis, 35 Oogonia, 35 Oophorectomy, 600 Open reading frames (ORFs), 421-422 Operators, 511, 513, 515 Operons, 509-523 histidine operon in Salmonella typhimurium, 522 lactose operon in E. coli, 511-519 mechanism of, 509-511 tryptophan operon in E. coli, 519-523 Operon model, 509-511 Opines, 465 Orange, 332 Orangutans, 380 Order, in genetic code, 311–312 ORFs (open reading frames), 421-422 Organelles, 19, 21, 25, 532-533

Organ formation, 567–569, 584 Origin, of replication, 224–227 *The Origin of Species* (Charles Darwin), 645, 656–658, 663 *OriT* (origin of transfer) site, 176 Orthologues, 424 Outer membrane, bacterial cell, 21 Out-of-Africa Hypothesis, 677 Ovalbumin, 274, 546 Ovarian cancer, 599 Ovary, 33–35 Overdominance, 648

Ρ

P1 artificial chromosomes (PACs), 373, 374 p21 protein, 595 p53 (tumor suppressor protein), 595-597,602 Pääbo, Svante, 397 PACs (P1 artificial chromosomes), 373, 374 Pachynema, 28, 30 Page, David, 154 Pair-rule genes, 567 Palindromes, 369 Panmictic population, 640 Panmixis, 640 Pan troglodytes, 368 pAPC (tumor suppressor protein), 596-598 Paracentric inversions, 126, 127 Paralogues, 424 Parasegments, 567 Parasexual processes, 172-185 conjugation, 175-179 and drug-resistant bacteria, 186 transduction, 182-185 transformation, 173-175 types of, 172 Pardue, Mary Lou, 493 Parental strains, 43 Partial diploids, 183 Partial dominance, 63 Patau syndrome, 122 Paternity tests, 459 Pathways: anabolic, 507 to cancer, 600-603 catabolic, 506 metabolic, 339-340 RNAi, 541-542 Pattern baldness, 71

pBRCA1 (tumor suppressor protein), 599 pBRCA2 (tumor suppressor protein), 599 PCNA (proliferating cell nuclear antigen), 246-247 PCR, see Polymerase chain reaction Peas: Mendel's experiments on, 41-46 sweet, 73-74, 136-138 Pedigrees, 53 P elements, 485-487, 497-498 Penetrance, 71, 72 Penicillin, 163, 187 Peppered moth, 643, 644, 660 Peptides, 286, 287 Peptide bonds, 286, 287 Peptide hormones, 535–537 Peptidyl (P) site, 297, 298 Peptidyl transferase, 302, 303 Pericentric inversions, 126-127 Pericentriolar material, 25 Perlegen Sciences, Inc., 414 Permissive condition, 341 Peroxisomes, 20 Personality, 629-630 Pesticides, 333-334 Petals, of flower, 33 Phage, see Bacteriophage Phagemid vectors, 372, 373 Phagocytes, 584 Phaseolus vulgaris, 608 Phenotype(s): and conditional lethal mutation, 340-342 correlating between relatives, 624-626 defined, 43 and human globin genes, 338-339 and human metabolic pathways, 339-340 mutation's effects on, 337-342 nomenclature for, 170-171 predicting, 617-618 and recessive mutation, 337-338 variation in, 659-660 Phenotypic evolution, 670–672 Phenotypic variance, partitioning of, 614-615 Phenylalanine, 71, 257, 297, 298, 339-340, 628

Phenylketonuria (PKU), 14, 53, 70-71, 75-76, 80-81, 339, 628, 637-638 Phenylthiocarbamide (PTC) tasting, 53 Pheochromocytoma, 581 Philadelphia chromosome, 589-590 pHMSH2 (tumor suppressor protein), 598-599 Phosphorylation, 269 Photreactivation, 348 Phylogenetic trees, 664 Phylogeny: defined, 10 of human populations, 679 and mitochondrial DNA, 665 Physical distance, 149-150 Physical gene mapping, 386-387, 403, 405-406 Pigs, 433, 464 Pilus, 21 piRNAs, 487 Pistil, 33, 34 Pisum sativum, 40, 41 Pitcairn Island, 634, 647 Piwi proteins, 487 pJCPAC-Mam1 shuttle vector, 374 PKU, see Phenylketonuria Plants: cell division in, 25, 26 cells of, 20, 21 cell walls of, 19 as eukarvotes, 20 transgenic, 464-466 Plant cells, 21, 26 Plant development, 22 Plaque hybridization, 379–380 Plasma membrane, 21 Plasmids: defined, 20 DNA in, 22 and genetic exchange in bacteria, 179-181 genetic information in, 169 - 170IS elements in, 480 Plasmid vectors, 372, 373 Plasmodium falciparum, 398 Plastids, 428 Pleiotropy, 75-76 Ploidy, 114 Pluripotent, 573 Pneumococci, 173-174

Point mutations, 321 Polar bodies, 35 Polar nuclei, 33, 34 Pole cells, 560 Pollen abortion, 129 Pollen grains, 33, 34 Pollen tube, 34 Pollination, 34 Polyacrylamide gel electrophoresis, 385, 386 Polyadenylation, 273 Polycloning site, 372 Polydactyly, 71 Polygenic trait, 614 Polvlinker, 372 Polymers, glucose, 19 Polymerases, DNA, see DNA polymerases Polymerase chain reaction (PCR): amplification of DNA sequences by, 374–376 in DNA profiling, 456-459 Polymorphism, 459, 648, 659-660 Polypeptide(s), 7-9 in cells, 19 colinear, 289-292 defined. 7 and genetic symbols, 68 Mendel's principles applied to, 67-68 one gene and one, 289–292 in proteins, 286-289 synthesis of, 298-305 Polypeptide chain(s): codons for initiation/termination of, 309-310 elongation of, 302-304 initiation of, 298-302 termination of, 304, 305 Polypeptide products, 291–292 **Polyploids:** chromosome pairing in, 117 fertile, 116-117 sterile, 115-116 Polyploidy, 115-119 defined, 114, 115 effects of, 115 and polyteny, 118-119 tissue-specific, 117-118 Poly(A) polymerase, 273 Poly(A) tails, 273, 294, 534 Polytene chromosome maps, 118

Polyteny, 118-119, 125 Pongo albelii, 380 Population genetics, 12, 634–651 allele frequencies, 635-641 balancing selection, 648-649 genetic counseling, 640-641 genetic equilibrium, 647-651 Hardy-Weinberg principle, 636-640 mutation-drift balance, 650-651 mutation-selection balance, 649-650 natural selection, 641-645 on Pitcairn Island, 634 random genetic drift, 645-647 Population size, random genetic drift and. 646 Population subdivision, 640 Positional cloning, 402, 407-409 chromosome jumps, 408-409 chromosome walks/jumps, 408-409 steps in, 407-408 Position effect, 343 Position-effect variegation, 545 Positive control mechanisms, 507-509 Posttranscriptional regulation by RNA interference, 541–544 Posttranslational regulatory mechanisms, 526 Postzygotic isolating mechanisms, 673 POT-1 protein, 213 PP (pyrophosphate), 261 pRB (tumor suppressor protein), 593-595 Pre-mRNAs, 257 Prepriming proteins, 234 Preproinsulin, 667-668 Prezygotic isolating mechanisms, 673 Primary structure, of polypeptides, 287, 288 Primary transcript, 257 Primer DNA, 232, 239, 240 Primosomes, 242 Private alleles, 600 Probability method, 47-48 Probe, 112–133 Profiling, DNA, 455-461 Proflavin, 306-307, 326-328 Progerias, 249

Progesterone, 535

Programmed cell death, 584, 602 Prokarvotes: cell division in, 23 cells of, 20, 21 chromosomes in, 20, 22 chromosome structure in, 205-206 DNA replication, 231-244 Prokaryote transcription and RNA processing, 263–267 concurrent transcription/ translation/mRNA degradation, 266 elongation of RNA chains, 263-265 initiation of RNA chains, 263, 264 RNA polymerases, 263-264 stages of, 263 termination of RNA chains, 263, 265-266 Prokaryotic gene expression, 504-527 constitutive/inducible/ repressive genes, 506-507 and d'Hérelle's work on dysentery, 504 lactose operon in E. coli, 511-519 operons, 509-511 pathway of, 505 positive/negative control of, 507-509 posttranslational regulatory mechanisms, 526 translational control of, 525-526 tryptophan operon in E. coli, 519-523 Prokaryotic genomes, 424–425 Prolactin, 535 Proliferating cell nuclear antigen (PCNA), 246-247 Promoters, 261, 263, 264, 270, 271 Proofreading, 238-242 Prophage, 168, 169 Prophase, 25, 28-31, 140 Prophase I, 28, 30-31 Prophase II, 29, 31 Prostate cancer, 582, 599, 602 Protamines, 207 Proteases, 194 Protein(s): cell division controlled by, 23 in cells, 19

and E. coli lactose operon, 518 eukaryotic production of, 461-463 evolution of protein sequences, 667-668 in molecular control of transcription, 538-540 polypeptide components of, 67 prepriming, 234 seleno, 315 tumor suppressor, 593-599 western blot analysis of, 384-385 Protein structure: complexity of, 286-289 polypeptides, 286-289 and sickle-cell anemia, 285 variation in, 661 Protein synthesis, 293–304 about, 293-294 GFP as reporter of, 419-420 ribosomes in, 294-296 steps in, 258, 260 transfer RNAs in, 296-302 translation, 298-305 Proteome, 7, 402 Proteomics, 8, 402 Protists, 20 Proto-oncogenes, 586-587 Protoplasts, 196-197 Prototrophs, 170, 341 Pseudoautosomal genes, 100 Pseudogenes, 668 PTC (phenylthiocarbamide) tasting, 53 PubMed, 400, 401 Punnett, R. C., 46, 72, 73, 136, 137 Punnett square method, 46, 47,636 Pupa, 560 Purifying selection, 649, 669 Purines, 199 Pyrimidines, 199 Pyrophosphate (PP), 261 Pyrosequencing, 391

C

QT (quantitative trait) loci, 620–624 Quantifying complex traits, 608 Quantitative traits: behavioral traits, 628–630 defined, 608

frequency distributions of, inversion, 126-127 genetic/environmental factors intelligence, 628-629 mean/modal class of, 612 multiple genes influencing, variance/standard deviation of, Quantitative trait analysis, 127-128 artificial selection, 618-620 broad-sense heritability,

615-616 multiple factor hypothesis, 614 narrow-sense heritability, 616-618 partitioning of phenotypic variance, 614-615 predicting phenotypes, 617-618 quantitative trait loci, 620-624 Quantitative trait (QT) loci, 620-624 Quaternary structure, of polypeptides, 288 Quinacrine stain, 112

611-612

608-610

612-613

613-624

influencing, 608

personality, 629-630

statistics of, 611-613

R

Rabbits, 64, 65 Radiation hybrid mapping, 410 Radiation-induced mutation, 328-331 Radical amino acids, 286 Ramibacterium ramosum, 200 Random genetic drift, 645-647, 670 Random insertion mutations, 572-573 Ras protein, 588, 589 RB gene product (pRB), 593-595 R-determinant, 481 RdRPs (RNA-dependent RNA polymerases), 544 Reading frame, 307-309 Rearrangement, of chromosome structure, 126-129 and cancer, 589-590 compound chromosome, 128-129 defined, 114

Robertsonian translocation, 129 translocation, 127-128 transposon-mediated, 498 Recessive genes, 42 dominant vs., 68-70 in pedigrees, 53 selection against, 643 Recessive lethals, 337 Recessive mutation, 337-338 Recipient cell, 171, 172 Reciprocal translocation, Recognition sequence, 264 Recombinant chromatids, 31 Recombinant-deficient mutants, 354 Recombinant DNA molecules, 12,368 amplification by PCR, 374-376 amplification in cloning vectors, 372-373 in vitro production of, 371 Recombinant DNA technology, 367 with cystic fibrosis, 445-448 with Huntington's disease, 440-442, 445 Recombination: between adjacent nucleotide pairs, 389-391 cleavage/rejoining, 354-356 and crossing over, 138-139 DNA mechanisms of, 354-358 and evolution, 153-155 gene conversion, 356–358 genetic control of, 155 genetic map distance and frequency of, 146-147 and linkage, 136-138 suppression by inversions, 153-154 Recombination frequency, 137-138 Recombination mapping, 141-146 Recombination signal sequences (RSSs), 577 Regulator genes, 507, 508 Regulator protein binding site (RPBS), 507-509 Regulatory binding site, 526 Reichman, Lee, 164

Rejoining, cleavage and, 354-356 Relationship(s). See also Inbreeding coefficient of, 82 correlations between, 624-628 pedigrees of, 53 Relative fitness, 642 Release factors (RFs), 304, 305 Renaturation, 215 Rennin, 462-463 Renwick, J. H., 151, 152 Repetitive DNA, 214-215 Replica plating, 334–335 Replication, of DNA and chromosomes, 220-250 autoradiography of, 227-228 bidirectional, 228-230 continuous/discontinuous synthesis, 232, 233 DNA helicase, 236-237 DNA ligase, 232–233 DNA topoisomerase, 237, 238 eukaryotic chromosome replication, 244-250 as gene function, 193 genetic information propagated by, 6 initiation of, 234-263 and methylation, 548 and monozygotic twins, 220 origins of, 224-227 primosomes, 242 in prokaryotes, 231–244 replisome, 242-243 RNA-primer initiation of DNA chains in, 234–236 rolling-circle, 243-244 semiconservative, 221-224 single-strand DNA-binding protein, 237 stages of, 221 unwinding DNA in, 236–238 in vivo, 221-230 Replication bubble, 234 Replication factor C (Rf-C), 246-247 Replication forks, 227 Replicative transposons, 478 Replicons, 245-246 Replisomes, 243 Repressed genes, 507 Repressible operon, 519–520 Repression, 507, 510, 519–520 Repressors, 507, 508

Repressor genes, 513, 515 Reproduction, 27, 35, 36 Reproductive cloning, 574-575 Reproductive isolation, 673 Repulsion linkage phase, 138 Resistance transfer factor (RTF), 481 Resolution site, 483 Response to selection, 619 Restriction endonucleases: discovery of, 368-371 mapping of, 386-387 recognition sequences/ cleavage sites of, 369 Restriction fragments, 368, 371 Restriction fragment-length polymorphism (RFLP): detection of, 403-405 HD gene linked to, 440-441 in tomatoes, 620-621 Restriction maps, 386–387 Restriction sites, 368 Restrictive condition, 341 Retinoblastoma, 584, 591-593 Retroposons (non-LTR retrotransposons), 493 Retrotransposons, 478-479, 488-493 retroposons, 493 retroviruses, 488-492 retroviruslike elements, 492-493 Retroviral oncogenes, 585-586 Retroviral vectors, 451-454 Retroviruses, 488-492, 585-586 Retroviruslike elements, 492-493 Rett syndrome, 270 Reverse genetics, 467-473 mouse knockout mutations, 467-469 RNA interference, 471, 472 T-DNA and transposon insertions, 469-471 Reverse mutation, 336 Reverse transcriptase, 488 Reverse transcriptase-PCR (RT-PCR), 384, 385 Reverse transcription, 9 Rf-C (replication factor C), 246-247 RFLP, see Restriction fragmentlength polymorphism RFs (release factors), 304, 305

Rho-dependent terminators, 265, 266 Rho-independent terminators, 265-266, 273 Ribonuclease (RNase), 194, 206 Ribonucleic acid, see RNA Ribonucleoside triphosphates, 261 Ribonucleotide monophosphate (RMP), 261 Ribonucleotide triphosphate (RTP), 261 Ribosomal RNAs (rRNAs), 258-260 Ribosomes, 20, 21 crystal structure of, 298, 299 in gene translation, 258-259 in protein synthesis, 294-296 tRNA binding sites on, 297-298 Riboswitches, 524-525 Rice, 432 Riggs, Arthur, 245 Riken BioResource Center, 470 RISC, see RNA-induced silencing complex R-loops, 275 RMP (ribonucleotide monophosphate), 261 RNA (ribonucleic acid), 197-199 alternate splicing of, 533 in cells, 19 chemical subunits of, 198-199 in chromosomes, 20 DNA vs., 3-4 genetic information carried by, 197 initiation of DNA chains with primers of, 234-236 structure of, 4 types of RNA molecules, 258-260 RNA analysis: by northern blot hybridization, 382, 384 by RT-PCR, 384, 385 **RNA** chains: elongation of, 263-265, 271-272 initiation of, 263, 264, 270-271 termination of, 263, 265-266, 272-273 RNA-dependent RNA polymerases (RdRPs), 544 RNA editing, 273-274

RNA-induced silencing complex (RISC), 260, 471, 472, 541-542 RNA interference (RNAi), 471, 472, 541-545 RNAi pathways, 541–542 RNA polymerases, 261, 263-264, 270-273 RNA polymerase I, 267-268, 271, 273 RNA polymerase II, 267-268, 270-273 RNA polymerase III, 267–268, 271, 273 RNA polymerase IV, 268 RNA polymerase V, 268 RNA primers, 234–236 RNase (ribonuclease), 194, 206 **RNA** splicing: alternate, 533 autocatalytic, 278-279 intron excision by, 277-280 introns before, 279-280 RNA synthesis, 261-262 RNA transcript, 7 Robertsonian translocation, 129 Roderick, Thomas, 402 Roentgen (r) units, 330 Rolling-circle replication, 176, 228, 243-244 Rooted tree, 664 Roslin Institute, 18, 574 Rothmund-Thomson syndrome, 352, 353 Rough endoplasmic reticulum, 21 Round worm, see Caenorhabditis elegans Rous, Peyton, 585 Rous sarcoma virus, 585 Royal hemophilia, 98, 99 RPBS (regulator protein binding site), 507-509 RPE64 gene, 439 R plasmids, 181, 187 rRNAs (ribosomal RNAs), 258-260 RSSs (recombination signal sequences), 577 RTF (resistance transfer factor), 481 RTP (ribonucleotide triphosphate), 261 RT-PCR (reverse transcriptase-PCR), 384, 385 Rubin, Gerald, 486, 497, 498, 570

S

Saccharomyces cerevisiae: alanine tRNA structure of, 297 centromeres in, 212 DNA base composition in, 200 GAL4 transcription factor from, 539 genome sequencing of, 430 as model organism, 32-33 phenylalanine of, 297, 298 RNA polymerase II in, 272 snRNAs in, 280 SWI/SNF complex in, 547 tRNA precursor splicing in, 278 Saedler, Heinz, 483 Salk Institute, 470 Salmonella typhimurium, 183, 522 Sample, 611 Sanger, Frederick, 388 Satellite bands, 214 Satellite DNAs, 214 Scaffold, 210, 212 Schnös, Maria, 228 Schüpbach, Trudi, 562 SCID (severe combined immunodeficiency disease), 452-455 Scott, Matthew, 566 Search tools, 400-401, 422-423 Sea urchin, 276 SECIS elements, 315 Secondary endosperm nucleus, 33, 34 Secondary structure, of polypeptides, 287, 288 Seed germination, 34 Seedling, 34 Segments, 565-567 Segmentation genes, 566–568 Segment-polarity genes, 567 Segregation (principle of inheritance), 44 chromosomal basis of, 95 experimental evidence of, 92-93 during gamete formation, 44 in human families, 54 Selectable marker genes, 372, 424, 466 Selection: artificial, 618-620 balancing, 648-649 natural, 641-645 purifying, 649, 669

Selection coefficient, 642 Selection differential, 618 Selective breeding, 13, 608 Selective media, 177 Selector genes, 566 Selenocysteine, 315 Selenoproteins, 315 Self-fertilization, 41 Self-splicing, 278, 279 Self-sufficiency, of cancer cells, 602 Semiconservative replication, 221-224 Semidominant, 63 Senescence, 249 Sense strands, 261 Sepals, 33 September 11 terrorist attack, 456 Sequence-specific protein-nucleic acid interactions, 518 Sequence-tagged sites (STSs), 403 Sequencing, DNA, see DNA sequencing Serine tRNAs, 313 Serratia marcescens, 170 7-methyl guanosine (7-MG), 272 Severe combined immunodeficiency disease (SCID), 452-455 SEV proteins, 571 Sex chromosomes: nondisjunction, 93-95, 124 research discoveries about, 89 - 91and sex determination. 101 - 103Sex determination: in Drosophila, 102 haplo-diplo system of, 102-103 in human beings, 101-102 Sex-determining region Y (SRY) gene, 101 Sexduction, 181–182 Sex-linked genes, 98–100 color blindness, 98-100 in fruit flies, 92-95 hemophilia, 98-100 on X and Y chromosomes, 100 on Y chromosome, 100 Sexual reproduction, 35, 36 Sheep, 18, 333 Shelterin, 213, 214 Sherman, Stephanie, 443 Sherman paradox, 443

Shigella, 187, 504 Shine-Dalgarno sequence, 301 Short fingers, 53 Short interfering RNAs (siRNAs), 541-544 Short interspersed nuclear elements (SINEs), 494, 496 Short-lived mRNAs, 533-534 Short tandem repeat (STR) mapping, 463 Short tandem repeats (STRs), 456-460 Shull, George, 74, 78 Shuttle vectors, 374 Sia, Richard, 173, 194 Sickle-cell disease, 9-10, 53, 285, 338-339, 448-450, 648 Side groups, 286 Sigma (σ) factor, 263–264 Σ symbol, 612 Signaling systems, 533 Signal molecules, 535-537 Signal sequence, 294 Signal transduction, 535 Silencers, 271 Silent polymorphisms, 661–662 Simian virus S40, 246 Simmons, Michael, 486 Simple tandem repeats, 331 SINEs (short interspersed nuclear elements), 494, 496 Single-nucleotide polymorphisms (SNPs), 414-415 Single-strand assimilation, 354 Single-strand DNA-binding (SSB) protein, 237 siRNAs (short interfering RNAs), 541-544 siRNAs (small interfering RNAs), 268, 534 Sister chromatids, 24-31, 139 Site-specific insertion, 183 Site-specific recombination, 176 Small interfering RNAs (siRNAs), 268, 534 Small nuclear RNA-protein complexes (snRNPs), 280 Small nuclear RNAs (snRNAs), 259, 260, 279, 280 Smith, Hamilton, 368 Smithies, Oliver, 455 Smooth endoplasmic reticulum, 21 snail gene, 562 SNPs (single-nucleotide polymorphisms), 414-415

snRNPs (small nuclear RNAprotein complexes), 280 Society, genetics in, 15 Solenoid model, 210 Somatic-cell gene therapy, 451-453 Somatic cells, 22 Somatic mosaics, 122 Somatic mutation, 332-333 Somatotropin, 535 SOS response (DNA repair), 351 Southern, E. M., 382 Southern blot hybridization, 381-383 spätzle gene, 562 Specialized transduction, 183-185 Special transcription factors, 538 Speciation, 672-675 Species, 672-673 Spermatogenesis, 35 Spermatogonia, 35 Sperm cell, 3, 18, 22, 27, 34, 35 S phase, of cell cycle, 23 Spindle, 24-26, 30, 31 Spinobulbar muscular atrophy, 442,444 Spinocerebellar ataxia, 442 Spliceosomes, 257, 279, 280 Splicing: alternate, 533 autocatalytic, 278-279 precursor, 278 RNA, 277-280, 533 self-, 278, 279 Splicing endonuclease, 278 Splicing ligase, 278 Splicing reactions, 257 Split genes, 257 Spontaneous mutation, 333 Sporadic cancers, 591 Sporophytes, 33, 34 Sporulation, 33 Spradling, Allan, 497, 498 Squared deviated from the mean, 612 SSB (single-strand DNA-binding) protein, 237 Stahl, Franklin, 221-224, 356 Stamens, 33 Standard deviation, 613 Starlinger, Peter, 483 START checkpoint, 584

snRNAs, see Small nuclear RNAs

Statistics, of quantitative traits, 611-613 frequency distributions, 611-612 mean/modal class, 612 variance/standard deviation, 612-613 Stem cells: embryonic, 464 in mammals, 573-574 Stem cell therapy, 452–454, 558 Sterile mutations, 67 Sterile polyploids, 115–116 Sterility, 94 Steroid hormones, 535-537 Stevens, N. M., 91 Stigma, 33, 34 Stop codon, 8 Streptococcus pneumoniae, 173-174, 194-195 Streptomycin, 177, 186, 187, 334, 335 STR mapping, 463 STRs (short tandem repeats), 456-460 Structural allelism, 344, 346 Structural genomics, 402 STSs (sequence-tagged sites), 403 Sturtevant, Alfred H., 135, 136, 141 Style, of flower, 33, 34 Sumatran orangutans, 380 Supercoils, negative, 204-205, 238 Suppression, of recombination by inversions, 153–154 Suppressor mutations, 307, 313-314, 336 Suppressor-sensitive mutations, 341 Suppressor tRNAs, 313-314 Survival, unequal, 639 Sutton, W. S., 91 Sved, John, 485 Sweden, 187 Sweet pea, 73-74, 136-138 SWI/SNF complex, 547 Sympatric speciation, 674-675 Synapsis, 30, 31 Synaptonemal complex, 30 Syncytium, 559, 560 Synergid cells, 33, 34 Synonymous substitution, 669 Synteny, 433-434 Systemic lupus erythematosus, 280 Szostak, Jack, 248, 356

T.

T, see Thymine TACTAAC box, 277 TAD (transcription-activation domain), 595, 596 Tahiti, 634, 656 Tandem repeats, 331 Tanksley, Steven, 620-623 T antigen, 246 Taq polymerase, 375, 376 Targeted gene transfers, 455 Target site duplications, 480 TATA-binding protein (TBP), 271 TATA box, 270 Tatum, Edward, 67, 175, 289-291 Tautomeric shifts, 322-326 Tay, Warren, 340 Taylor, J. Herbert, 223 Tay-Sachs disease, 53, 340, 641 TB, see Tuberculosis **TBP** (TATA-binding protein), 271 T-cell acute lymphoblastic leukemia (T-cell ALL), 454 T-cell receptors, 575-576 TDF (testis-determining factor), 101 T-DNA, 465-466 T-DNA insertions, 469-471 Telomerase, 248-249 Telomeres, 212-214 Telomere length, 249 Telophase, 25, 26, 28, 29, 31 Telophase I, 28, 31 Telophase II, 29, 31 Temin, Howard, 488 Temperate bacteriophages, 165 Temperature: and induction of transcriptional activity, 535 and mutation, 70 Temperature-sensitive (ts) mutations, 166, 341 Template DNA, 234, 239 Template strand, 6, 261 -10 sequence, 264 Teosinte, 13 Terminalization, 31 Termination: of polypeptide chain translation, 304, 305 of RNA chains, 263, 265-266, 272-273

Termination signal, 265, 273 Tertiary structure, of polypeptides, 287-289 Testcross, 46, 137, 138 Testicular feminization, 102 Testing. See also DNA testing for aneuploidy, 123 of genetic hypotheses, 48-52 of mutations for allelism, 65-66 Testis-determining factor (TDF), 101 Testosterone, 71, 101-102, 535 Tetracycline, 187 Tetrad, 30, 31, 138 Tetrahymena thermophila, 278-279 Tetrameric core, 263 Tetraploid, 90, 115 TFIIX (transcription factor for polymerase II), 271 Tfm mutation, 102 Therapeutic cloning, 574 Thermus thermophilus, 298, 299 -35 sequence, 264 3' poly(A) tail, 273, 294 3' untranslated region (3' UTR), 534, 542 Three-point testcross, 142–146 Threshold traits, 610 Thymine (T), 4, 198–200, 202 Tilghman, Shirley, 275 Ti plasmid, 465-466 Tissue plasminogen activator (TPA), 672 Tissue-specific enhancers, 538 Tissue-specific polyploidy, 117-118 T-loops, 213-214 TMV (tobacco mosaic virus), 165, 197 Tn3 elements, 481-483 Tobacco, 429, 609, 610 Tobacco mosaic virus (TMV), 165, 197 toll gene, 562 Tomatoes, 620-623 Tomato plants, 52 Topoisomerase, 237, 238 Total phenotypic variance, 614 Totipotency, 465 TPA (tissue plasminogen activator), 672 Trafficking, 20 trans configuration, 343

Transcript, 7 Transcription-activation domain (TAD), 595, 596 Transcription and RNA processing: addition of 3' poly(A) tails, 273 addition of 5' methyl guanosine caps, 272 chain cleavage, 272-273 concurrent translation/mRNA degradation and, 266 controlled, 532-533 elongation of RNA chains, 263-265, 271-272 in eukarvotes, 267-274 factors in induction of, 534-537 in gene expression, 7 initiation of RNA chains, 263, 264, 270-271 interrupted genes, 274-276 intron excision, 277-280 molecular control of, 537-541 in prokaryotes, 263-267 reverse, 9 RNA editing, 273-274 RNA polymerases, 263–264, 267-270 RNA splicing, 277–280 termination of RNA chains, 263, 265-266, 272-273 and translation, 257-258 Transcription bubble, 262, 264, 266 Transcription factors, 261, 263, 268, 533, 538-540 Transcription factor for polymerase II (TFIIX), 271 Transcription-termination signals, 521-522 Transcription unit, 263 Transcriptome, 402 Transducing particles, 183 Transduction, 172, 182-185 Transfection, 197 Transfection test, 587–588 Transfer, of genetic information: direction of, 257 RNA classes for, 258, 260 transcription/translation, 257-258 Transfer RNAs (tRNAs), 258, 260 codon interactions with, 312-314 precursor splicing of, 278

in protein synthesis, 296-302 suppressor, 313-314 Transformation: in bacteria, 172-175 DNA as mediator of. 194-195 principle of, 194-195 with transposons, 496-498 Transformation vector, 498 Transgenes, 451, 452 Transgenic animals, 463-464 Transgenic organisms, 451 Transgenic plants, 464–466 trans heterozygote, 343-345 Transitions, 323 Translation, 293, 298-305 concurrent transcription/ mRNA degradation and, 266 elongation of polypeptide chain, 302-304 in gene expression, 7 gene expression controlled by, 525-526 initiation of polypeptide chain, 298-302 and sickle-cell anemia, 285-286 stages of, 98 termination of polypeptide chain, 304, 305 and transcription, 257-258 Translesion polymerases, 231 Translocation: of chromosome structure, 127-128 Robertsonian, 129 Transposase, 480 Transposition, 477 Transposons (transposable elements), 187, 213, 215, 477-499 about, 478-479 in bacteria, 479-483 composite, 481 cut-and-paste, 478 in eukarvotes, 483-488 genetic/evolutionary significance of, 496-499 genetic transformation with, 496-498 and genome organization, 498-499 in humans, 494-496 IS elements, 479-481 in maize, 477 mechanisms of, 478

as mutagens, 496, 497 replicative, 478 retro-, 478-479, 488-493 retroviruses and retrotransposons, 488-493 Tn3 elements, 481–483 Transposon-induced mutation, 331 Transposon insertions, 469–471 Transposon tagging, 496 trans test, 342-346 Transverse fibers, 30 Transversions, 323 "Tree of Life" program, 10-11 Tribolium castaneum, 619 Trichothiodystropy, 352, 353 Trinucleotide repeats: and human disease, 443-444 and Huntington's disease, 442, 445 mutation by, 331–332 Triplet code, 306-312 Triplet codon, 8 Triploid, 115 Triploid endosperm nucleus, 34 Triplo-X syndrome, 121, 122 Trisomy, 119-122, 125 Triticum aestivum, 110, 117 Triturus viridescens, 295 Trivalent, 116 tRNA;^{Met}, 301 tRNA^{Met}, 300, 301 tRNAs, see Transfer RNAs Troponin T gene, 533 True-breeding, 41 Trypanosomes, 274, 531 Tryptophan, 507, 511 Tryptophan operon, in E. coli, 519-523 Tryptophan synthetase, 291-292 Tschermak-Seysenegg, Eric von, 41 Tsetse, 531 ts (temperature-sensitive) mutations, 166, 341 Tsui, Lap-Chee, 383, 409, 446 Tuberculosis (TB): multi-drug-resistant, 163-164 resurgence of, 186 streptomycin treatment for, 186-187 Tubulins, 24, 539 Tumor-inducing retroviruses, 585-586

Tumor suppressor genes, 585, 593-599 cellular roles of proteins in, 593-599 inherited cancers and Knudson's two-hit hypothesis, 590-593 Tumor suppressor proteins, 593-599 p53, 595-597 pAPC, 596-598 pBRCA1 and pBRCA2, 599 pHMSH2, 598-599 pRB, 592-595 Turner syndrome, 122–123 Twins: correlating quantitative phenotypes between, 625-626 dizygotic, 610, 627, 629, 630 monozygotic, see Monozygotic twins trait inheritance in, 608, 623 twist gene, 562 Two-hit hypothesis (of cancer), 590-593 Two-point testcross, 141-142 Twort, Frederick W., 504 Tv1 element, 492-493 Tyrosine, 339-340 Tyrosinemia, 339 Tyrosinosis, 339

U

U (Uracil), 4, 198 ultrabithorax gene, 276 Ultraviolet (UV) light, 112, 169, 320, 329-331 Unequal survival, 639 Unidirectional gene transfer, 171 Uninemes, 208 U.S. Department of Energy (DOE), 400 Univalent, 116 Universal genetic code, nearly, 312 Unrooted tree, 664 Untranslated region (UTR), 534, 542 Unwinding DNA, 236-238 Upstream region, 263 Upstream sequences, 264 Uracil (U), 4, 198 Urinary tract infections, 186

UTR (untranslated region), 534, 542 UV light, see Ultraviolet light

V

Vacuoles, 21 Valine, 9, 311 Van der Waals interactions, 289 Variable expressivity, 71-72 Variable number tandem repeats (VNTRs), 405, 456 Variable region, 576 Variance, 612-615 Variant surface glycoproteins (VSGs), 531 Variation: among animals, 657-658 in chromosome structure, 660-661 as effect of mutation, 66-67 in evolution of DNA sequences, 668-669 in evolution of protein sequences, 667-668 and human origins, 677-679 in natural populations, 659-662 in nucleotide sequences, 661-662 in phenotypes, 659-660 in protein structure, 661 Vectors, 498 bacteriophage, 372 cloning, 367, 372-374 cosmid, 372-373 phagemid, 372, 373 plasmid, 372, 373 retroviral, 451-454 shuttle, 374 Venter, J. Craig, 5, 398, 410, 412, 425 v-erbB gene, 586 Versailles Genomic Resource Center (VGRC), 470 Vertebrate animals, α-globin variations among, 664-667 Vertebrate development, 571-578 immune cell differentiation, 575-578 mammalian stem cells,

573-574

mouse mutations, 572-573

reproductive cloning, 574-575

and vertebrate homologues of invertebrate genes, 571-572

Vesicle, 21 v-fms gene, 586 VGRC (Versailles Genomic Resource Center), 470 Vicia faba, 223, 224 Victoria, Queen, 98 Villi, 123 Viral oncogenes, 585–586 vir region, 465-466 Virulent bacteriophages, 165 Viruses, 164-169. See also Bacteriophages cancer-causing, 584 and cells, 19 chromosome structure in, 205-206 DNA base composition in, 200 genetic importance of, 164-165 mapping genes in bacteriophage, 166 retro-, 488-492 reverse transcription in, 9 RNA as carrier of genetic information in some, 197 Visible mutations, 66-67 VNTRs (variable number tandem repeats), 405, 456 VNTR fingerprints, 460 v-onc gene, 586 Von Hippel-Lindau disease, 592 VSGs (variant surface glycoproteins), 531 v-src gene, 585, 586 W Waksman, Selman, 186

Waldever, W., 90 Wallace, Alfred, 10 Wasps, 103 Water, 19 Watson, James, 3-4, 199-201, 221, 321, 322, 398, 409, 412 Weinberg, Robert, 587, 588, 602 Weinberg, Wilhelm, 636 Weintraub, Harold, 545 Weischaus, Eric, 562 Werner syndrome, 249, 352, 353 Wessler, Susan, 483 Western blot analysis, 384–385, 419 Wexler, Nancy, 440 Wheat, 110, 117, 608, 609 White blood cell cancers, 584 White-cheeked gibbon, 434

Widow's peak, 53 Wieschaus, Eric, 566 Wild-type alleles, 64, 68 Wilkins, Maurice, 200, 201 Wilm's tumor, 592 Wilson, Edmund Beecher, 89, 91 Wobble hypothesis, 312–313 Wollman, Elie, 177, 178, 181 Wood, William, 341, 342 Woods, Philip, 223 Woolly mammoth, 429 Wooly hair, 53 Wright, Seth, 333 Wright, Sewall, 79, 81, 645,658

Х

X-bearing eggs, 91 X chromosome(s), 90–93 attached-, 128-129 Caenorhabditis hypoactivation of, 553 Drosophila hyperactivation of, 552 genes on both Y and, 100 mammalian inactivation of, 551-552 mapping of, 409 nucleosomes in, 210 replication of, 246 XDR (extensively drug-resistant) bacteria, 163-164 Xenopus laevis, 276, 534 Xeroderma pigmentosum (XP), 231, 320, 351–353 X-gal, 373 X inactivation center (XIC), 551 XIST gene, 551 X-linked genes: activation/inactivation of, 550-551 and autosomal inheritance, 97 dosage compensation of, 104-105 experimental evidence of, 92-95 Hardy-Weinberg principle applied to, 638 X-linked SCID, 452–455 XO zygotes, 94, 101 XP, see Xeroderma pigmentosum X rays: effects of, 324, 325, 329, 330 measurement of, 330 mutagenicity of, 324-325

X-ray diffraction patterns, 201 XXX zygotes, 94 XXY zygotes, 94, 101

Y

YACs (yeast artificial chromosomes), 373, 374Yanofsky, Charles, 291Y-bearing sperm, 91Y chromosome(s), 90–92, 94

genes on human, 100 mapping of, 409 and sex determination, 101–102 Yeast, baker's, *see Saccharomyces cerevisiae* Yeast artificial chromosomes (YACs), 373, 374 YO zygotes, 94

genes on both X and, 100

Ζ

Zea mays, 200 Zebra fish, 32 zerknüllt gene, 562 Zigzag model, 210 Zinc finger motif, 540 Zinder, Norton, 182, 183 Zoo blots, 446 Zoo-FISH experiments, 432 Zuckerkandl, Emile, 663 Zygonema, 28, 30

Zygote, 3, 18, 34 Zygotic gene expression, 565–571 body segmentation, 565–567 cell type specification, 569–570 organ formation, 567–569

Index **767**