GENOMES 4
About the Author

I became fascinated with the natural world when I was very young. I began my research career studying the effects of metal pollution on microorganisms and the tolerance that some plants display to high concentrations of toxic metals. I then became excited by DNA and worked on mitochondrial genes in fungi in order to learn the new (in those days) techniques for gene cloning and DNA sequencing. I contributed to the discovery of mitochondrial introns and to work that described the base-paired structure of these introns. I then became interested in ancient DNA and was one of the first people to carry out DNA extractions with bones and preserved plant remains. This work has required close collaboration with archaeologists, and has led to my current interests in paleogenomics, the origins of agriculture, and the evolution of domesticated plants.

I obtained my PhD from University College London in 1977 and then worked in New York, Oxford, Colchester, and Manchester before beginning in 1984 as a Lecturer in Biotechnology at the University of Manchester Institute of Science and Technology (UMIST). I was appointed Professor of Biomolecular Archaeology in 2000 and was Head of Biomolecular Sciences at UMIST from 2002–2004. I was then Associate Dean in the Faculty of Life Sciences of the University of Manchester until 2006, before taking a break from administration in order to have more time to do research.

My other undergraduate textbooks include Introduction to Genetics, A Molecular Approach (Garland Science).
PREFACE

There have been remarkable advances in our knowledge of genomes since the previous edition of this book was published ten years ago. Back in 2007, next-generation sequencing was in its infancy and high-throughput methods for transcriptomics and proteomics were only beginning to be exploited. The application of these methods over the last ten years has resulted in an exponential increase in the number of species for which genome sequences and annotations are now available, and has enabled multiple versions of the genome of a single species to be examined. The profusion of new sequences has had a particularly dramatic impact on bacterial genomics, with introduction of the pan-genome concept and the discovery of extensive lateral transfer of genes between species. Our knowledge of eukaryotic genomes has undergone equally dramatic change, with the discovery of new types of noncoding RNA, including the vast numbers of long RNAs that are transcribed from the supposedly intergenic regions of many genomes.

*Genomes 4* retains the overall structure of the previous editions, with the book divided in four parts, on genome sequencing and annotation, genome anatomies, genome expression, and genome replication and evolution. With some small changes, the order of chapters remains unchanged. However, the text throughout has been completely updated and, in many chapters, substantially revised. In particular, the development of transcriptomics and proteomics has reached the point where in *Genomes 4* it is possible to describe the processes of transcription and translation from a genomewide perspective, rather than simply through an examination of the expression of individual genes. This was my aim when I wrote the first edition of *Genomes* way back in 1999, but the information available at that time meant that these core chapters were fairly orthodox treatments of gene rather than genome expression. We are still some way from being able to describe the entire expression of a genome as a single integrated process, but we are getting there and I hope that in *Genomes 4* I have been able to convey to the reader at least some aspects of the joined-up nature of genome expression.

*Genomes 4* has been a long time in the making and I would like to thank Liz Owen of Garland Science for her continued enthusiasm for the book and her gentle reminders about approaching deadlines. I also wish to thank David Borrowdale and Georgina Lucas at Garland for managing the production of the book, and Matthew McClements for his splendid artwork. As with the previous editions, *Genomes 4* would not have been finished without the support of my wife, Keri. The acknowledgment in the first edition that “if you find this book useful then you should thank Keri, not me, because she is the one who ensured that it was written” is equally true for the fourth edition.
A NOTE TO THE READER

I have tried to make the fourth edition of *Genomes* as user friendly as possible. The book therefore includes a number of devices intended to help the reader and to make the book an effective teaching and learning aid.

Organization of the Book

*Genomes 4* is divided into four parts:

**Part I – Studying Genomes** begins with an orientation chapter that introduces the reader to genomes, transcriptomes, and proteomes, and then in Chapter 2 moves on to the methods, centered on PCR and cloning, that were used in the pre-genome era to examine individual genes. The techniques that are used for constructing genetic and physical maps, which are still important in many genome projects, are then described in Chapter 3, followed in Chapter 4 by the methodology for obtaining DNA sequences and assembling reads into draft and finished genomes sequences. Two chapters are then devoted to analysis of genome sequences: Chapter 5 on the annotation of a genome by identification of genes and other features, and Chapter 6 on functional analysis of the genes that are discovered.

**Part II – Genome Anatomies** surveys the anatomies of the various types of genome that are found on our planet. Chapter 7 covers eukaryotic nuclear genomes, with emphasis on the human genome, partly because of the importance of the human genome in so many areas of research, but also because our genome is the best studied of all those for which sequences are available. Chapter 8 deals with the genomes of prokaryotes and of eukaryotic organelles, the latter included here because of their prokaryotic origins, and Chapter 9 describes viral genomes and mobile genetic elements, these being grouped together because some types of mobile element are related to viral genomes.

**Part III – How Genomes are Expressed** describes how the biological information contained in a genome is utilized by the cell within which that genome resides. Chapter 10 addresses the important issue of how the packaging of DNA into chromatin affects expression of different parts of the genome, and Chapter 11 then describes the central role that DNA-binding proteins play in expressing those parts of the genome that are active at a particular time. Chapter 12 moves on to the transcriptome, describing how transcriptomes are studied, their compositions, and how a cell’s transcriptome is synthesized and maintained. Chapter 13 gives an equivalent description of proteomics and the proteome, and Chapter 14 concludes this part of the book by exploring how the genome acts within the context of the cell and organism, responding to extracellular signals and driving the biochemical changes that underlie differentiation and development.

**Part IV – How Genomes Replicate and Evolve** links DNA replication, mutation, and recombination with the gradual evolution of genomes over time. In Chapters 15–17 the molecular processes responsible for replication, mutation, repair, and recombination are described, and in Chapter 18 the ways in which these processes are thought to have shaped the structures and genetic contents of genomes over evolutionary time are considered. Chapter 18 then ends with a small number of case studies to illustrate how molecular phylogenomics and population genomics are being used in research and biotechnology.
LEARNING AIDS

Each chapter has a set of Short Answer Questions and In-Depth Problems, as well as an annotated Further Reading list. At the end of the book there is an extensive Glossary.

**Short answer questions** require 50- to 500-word answers. The questions cover the entire content of each chapter in a fairly straightforward manner, and most can be marked simply by checking each answer against the relevant part of the text. A student can use the short answer questions to work systematically through a chapter, or can select individual ones in order to evaluate their ability to answer questions on specific topics. The short answer questions could also be used in closed-book tests.

**In-depth problems** require a more detailed answer. They vary in nature and in difficulty, the simplest requiring little more than a literature survey, the intention of these particular problems being that the student advances his or her learning a few stages from where *Genomes 4* leaves off. Other problems require that the student evaluates a statement or a hypothesis, based on their understanding of the material in the book, possibly supplemented by reading around the subject. These problems will, hopefully, engender a certain amount of thought and critical awareness. A few problems are difficult, in some cases to the extent that there is no solid answer to the question posed. These are designed to stimulate debate and speculation, which stretches the knowledge of each student and forces them to think carefully about their statements. The in-depth problems can be tackled by students working individually, or alternatively can form the starting point for a group discussion.

**Further Reading** lists at the end of each chapter include those research papers, reviews, and books that I look on as the most useful sources of additional material. My intention throughout *Genomes 4* has been that students should be able to use the reading lists to obtain further information when writing extended essays or dissertations on particular topics. Research papers are therefore included, but only if their content is likely to be understandable to the average reader of the book. Emphasis is also placed on accessible reviews, one strength of these general articles being the context and relevance that they provide to a piece of work. The reading lists are divided into sections reflecting the organization of information in the chapter, and in some cases I have appended a few words summarizing the particular value of each item to help the reader decide which ones he or she wishes to seek out. In some cases, Further Reading also includes URLs for databases and other online resources relevant to the material covered in a chapter.

The **Glossary** defines every term that is highlighted in bold in the text, along with a number of additional terms that the reader might come across when referring to books or articles in the reading lists. The glossary therefore provides a quick and convenient means by which the reader can remind themselves of the technical terms relevant to the study of genomes, and also acts as a revision aid to make sure those definitions are clearly understood during the minutes of uncertainty that many students experience immediately before an exam.
INSTRUCTOR RESOURCES

The images from the book are available through www.garlandscience.com in two convenient formats: PowerPoint® and JPEG. They have been optimized for display on a computer. Figures are searchable by figure number, by figure name, or by keywords used in the figure legend from the book. Help on answering the In-Depth Problems, found at the end of each chapter, is also available.

ACKNOWLEDGMENTS

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## CONTENTS IN BRIEF

<table>
<thead>
<tr>
<th>CHAPTER</th>
<th>TITLE</th>
<th>PAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>GENOMES, TRANSCRIPTOMES, AND PROTEOMES</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>STUDYING DNA</td>
<td>27</td>
</tr>
<tr>
<td>3</td>
<td>MAPPING GENOMES</td>
<td>55</td>
</tr>
<tr>
<td>4</td>
<td>SEQUENCING GENOMES</td>
<td>87</td>
</tr>
<tr>
<td>5</td>
<td>GENOME ANNOTATION</td>
<td>119</td>
</tr>
<tr>
<td>6</td>
<td>IDENTIFYING GENE FUNCTIONS</td>
<td>135</td>
</tr>
<tr>
<td>7</td>
<td>EUKARYOTIC NUCLEAR GENOMES</td>
<td>155</td>
</tr>
<tr>
<td>8</td>
<td>GENOMES OF PROKARYOTES AND EUKARYOTIC ORGANELLES</td>
<td>181</td>
</tr>
<tr>
<td>9</td>
<td>VIRAL GENOMES AND MOBILE GENETIC ELEMENTS</td>
<td>203</td>
</tr>
<tr>
<td>10</td>
<td>ACCESSING THE GENOME</td>
<td>219</td>
</tr>
<tr>
<td>11</td>
<td>THE ROLE OF DNA-BINDING PROTEINS IN GENOME EXPRESSION</td>
<td>241</td>
</tr>
<tr>
<td>12</td>
<td>TRANSCRIPTOMES</td>
<td>257</td>
</tr>
<tr>
<td>13</td>
<td>PROTEOMES</td>
<td>293</td>
</tr>
<tr>
<td>14</td>
<td>GENOME EXPRESSION IN THE CONTEXT OF CELL AND ORGANISM</td>
<td>329</td>
</tr>
<tr>
<td>15</td>
<td>GENOME REPLICATION</td>
<td>357</td>
</tr>
<tr>
<td>16</td>
<td>MUTATIONS AND DNA REPAIR</td>
<td>389</td>
</tr>
<tr>
<td>17</td>
<td>RECOMBINATION AND TRANSPOSITION</td>
<td>411</td>
</tr>
<tr>
<td>18</td>
<td>HOW GENOMES EVOLVE</td>
<td>429</td>
</tr>
<tr>
<td></td>
<td>GLOSSARY</td>
<td>463</td>
</tr>
<tr>
<td></td>
<td>INDEX</td>
<td>491</td>
</tr>
</tbody>
</table>
CHAPTER 1
GENOMES, TRANSCRIPTOMES, AND PROTEOMES

1.1 DNA
Genes are made of DNA
DNA is a polymer of nucleotides
The double helix is stabilized by base pairing and base stacking
The double helix has structural flexibility

1.2 RNA AND THE TRANSCRIPTOME
RNA is a second type of polynucleotide
The RNA content of the cell
Many RNAs are synthesized as precursor molecules
There are different definitions of the transcriptome

1.3 PROTEINS AND THE PROTEOME
There are four hierarchical levels of protein structure
Amino acid diversity underlies protein diversity
The link between the transcriptome and the proteome
The genetic code is not universal
The link between the proteome and the biochemistry of the cell

CHAPTER 2
STUDYING DNA

2.1 ENZYMES FOR DNA MANIPULATION
The mode of action of a template-dependent DNA polymerase
The types of DNA polymerase used in research
Restriction endonucleases enable DNA molecules to be cut at defined positions
Gel electrophoresis is used to examine the results of a restriction digest
Interesting DNA fragments can be identified by Southern hybridization
Ligases join DNA fragments together
End-modification enzymes

2.2 THE POLYMERASE CHAIN REACTION
Carrying out a PCR
The rate of product formation can be followed during a PCR
PCR has many and diverse applications

2.3 DNA CLONING
Why is gene cloning important?
The simplest cloning vectors are based on E. coli plasmids
Bacteriophages can also be used as cloning vectors
Vectors for longer pieces of DNA
DNA can be cloned in organisms other than E. coli

CHAPTER 3
MAPPING GENOMES

3.1 WHY A GENOME MAP IS IMPORTANT
Genome maps are needed in order to sequence the more complex genomes
Genome maps are not just sequencing aids

3.2 MARKERS FOR GENETIC MAPPING
Genes were the first markers to be used
RFLPs and SSLPs are examples of DNA markers
Single-nucleotide polymorphisms are the most useful type of DNA marker

3.3 THE BASIS TO GENETIC MAPPING
The principles of inheritance and the discovery of linkage
Partial linkage is explained by the behavior of chromosomes during meiosis
From partial linkage to genetic mapping
3.4 LINKAGE ANALYSIS WITH DIFFERENT TYPES OF ORGANISMS
Linkage analysis when planned breeding experiments are possible 69
Gene mapping by human pedigree analysis 69
Genetic mapping in bacteria 71
The limitations of linkage analysis 73

3.5 PHYSICAL MAPPING BY DIRECT EXAMINATION OF DNA MOLECULES
Conventional restriction mapping is applicable only to small DNA molecules 75
Optical mapping can locate restriction sites in longer DNA molecules 75
Optical mapping can be used to map other features in a DNA molecule 79

3.6 PHYSICAL MAPPING BY ASSIGNING MARKERS TO DNA FRAGMENTS
Any unique sequence can be used as an STS 81
DNA fragments for STS mapping can be obtained as radiation hybrids 82
A clone library can be used as the mapping reagent 83

SUMMARY 84
SHORT ANSWER QUESTIONS 85
IN-DEPTH PROBLEMS 85
FURTHER READING 86

CHAPTER 4
SEQUENCING GENOMES 87

4.1 CHAIN-TERMINATION SEQUENCING
Chain-termination sequencing in outline 87
Not all DNA polymerases can be used for sequencing 89
Chain-termination sequencing with Taq polymerase 90
Strengths and limitations of chain-termination sequencing 91

4.2 NEXT-GENERATION SEQUENCING
Preparation of a sequencing library is the common feature of next-generation methods 92
Various next-generation sequencing methods have been devised 93
Third- and fourth-generation methods enable sequencing in real time 97

4.3 HOW TO SEQUENCE A GENOME
The potential of the shotgun method was proven by the Haemophilus influenzae sequence 98
Many prokaryotic genomes have been sequenced by the shotgun method 99
Shotgun sequencing of eukaryotic genomes requires sophisticated assembly programs 100
More complex genomes can be sequenced by a hierarchical shotgun approach 104
What is a genome sequence and do we always need one? 107

4.4 A SURVEY OF EUKARYOTIC GENOME SEQUENCING PROJECTS
The Human Genome Project: genome sequencing in the heroic age 109
The Neanderthal genome: assembly of an extinct genome by use of the human sequence as a reference 110
The giant panda genome: shotgun sequencing based entirely on next-generation data 111
The barley genome: the concept of gene space 113

SUMMARY 115
SHORT ANSWER QUESTIONS 115
IN-DEPTH PROBLEMS 116
FURTHER READING 117

CHAPTER 5
GENOME ANNOTATION 119

5.1 GENOME ANNOTATION BY COMPUTER ANALYSIS OF THE DNA SEQUENCE
The coding regions of genes are open reading frames 119
Simple ORF scans are less effective with genomes of higher eukaryotes 120
Locating genes for noncoding RNA 122
Homology searches and comparative genomics give an extra dimension to gene prediction 123

5.2 GENOME ANNOTATION BY ANALYSIS OF GENE TRANSCRIPTS
Hybridization tests can determine if a fragment contains transcribed sequences 124
Methods are available for precise mapping of the ends of transcripts 125
Exon–intron boundaries can also be located with precision 126

5.3 ANNOTATION BY GENOMEWIDE RNA MAPPING
Tiling arrays enable transcripts to be mapped onto chromosomes or entire genomes 127
Transcript sequences can be directly mapped onto a genome 128

5.4 GENOME BROWSERS 131
6.1 COMPUTER ANALYSIS OF GENE FUNCTION
Homology reflects evolutionary relationships
Homology analysis can provide information on the function of a gene
Identification of protein domains can help to assign function to an unknown gene
Annotation of gene function requires a common terminology

6.2 ASSIGNING FUNCTION BY GENE INACTIVATION AND OVEREXPRESSION
Functional analysis by gene inactivation
Individual genes can be inactivated by homologous recombination
Gene inactivation without homologous recombination
Gene overexpression can also be used to assess function
The phenotypic effect of gene inactivation or overexpression may be difficult to discern

6.3 UNDERSTANDING GENE FUNCTION BY STUDIES OF EXPRESSION PATTERN AND PROTEIN PRODUCT
Reporter genes and immunocytochemistry can be used to locate where and when genes are expressed
Directed mutagenesis can be used to probe gene function in detail

6.4 USING CONVENTIONAL GENETIC ANALYSIS TO IDENTIFY GENE FUNCTION
Identification of human genes responsible for inherited diseases
Genomewide association studies can also identify genes for diseases and other traits

CHAPTER 7
EUKARYOTIC NUCLEAR GENOMES 155
7.1 NUCLEAR GENOMES ARE CONTAINED IN CHROMOSOMES 155
Chromosomes are much shorter than the DNA molecules they contain
Special features of metaphase chromosomes
DNA–protein interactions in centromeres and telomeres

7.2 HOW ARE THE GENES ARRANGED IN A NUCLEAR GENOME? 161
Genes are not evenly distributed within a genome
A segment of the human genome
The yeast genome is very compact
Gene organization in other eukaryotes

7.3 HOW MANY GENES ARE THERE AND WHAT ARE THEIR FUNCTIONS? 167
Gene numbers can be misleading
Gene catalogs reveal the distinctive features of different organisms
Families of genes
Pseudogenes and other evolutionary relics

7.4 THE REPETITIVE DNA CONTENT OF EUKARYOTIC NUCLEAR GENOMES 176
Tandemly repeated DNA is found at centromeres and elsewhere in eukaryotic chromosomes
Minisatellites and microsatellites
Interspersed repeats

SUMMARY 178
SHORT ANSWER QUESTIONS 178
IN-DEPTH PROBLEMS 179
FURTHER READING 179

CHAPTER 8
GENOMES OF PROKARYOTES AND EUKARYOTIC ORGANELLES 181
8.1 PHYSICAL FEATURES OF PROKARYOTIC GENOMES 181
The traditional view of the prokaryotic chromosome
Some bacteria have linear or multipartite genomes

8.2 GENETIC FEATURES OF PROKARYOTIC GENOMES 186
Gene organization in the E. coli K12 genome
Operons are characteristic features of prokaryotic genomes. Prokaryotic genome sizes and numbers of genes vary according to biological complexity. Genome sizes and numbers of genes vary within individual species. Distinctions between prokaryotic species are further blurred by lateral gene transfer. Metagenomes describe the members of a community. 

8.3 EUKARYOTIC ORGANELLAR GENOMES
The endosymbiont theory explains the origin of organellar genomes. Most organellar genomes are circular. The gene catalogs of organellar genomes are

SUMMARY
SHORT ANSWER QUESTIONS
IN-DEPTH PROBLEMS
FURTHER READING

CHAPTER 9
VIRAL GENOMES AND MOBILE GENETIC ELEMENTS

9.1 THE GENOMES OF BACTERIOPHAGES AND EUKARYOTIC VIRUSES
Bacteriophage genomes have diverse structures and organizations. Replication strategies for bacteriophage genomes. Structures and replication strategies for eukaryotic viral genomes. Some retroviruses cause cancer. Genomes at the edge of life.

9.2 MOBILE GENETIC ELEMENTS
RNA transposons with long terminal repeats are related to viral retroelements. Some RNA transposons lack long terminal repeats. DNA transposons are common in prokaryotic genomes. DNA transposons are less common in eukaryotic genomes.

SUMMARY
SHORT ANSWER QUESTIONS
IN-DEPTH PROBLEMS
FURTHER READING

CHAPTER 10
ACCESSING THE GENOME

10.1 INSIDE THE NUCLEUS
The nucleus has an ordered internal structure. The DNA content of a nondividing nucleus displays different degrees of packaging. The nuclear matrix is thought to provide attachment points for chromosomal DNA. Each chromosome has its own territory within the nucleus. Each chromosome comprises a series of topologically associated domains. Insulators mark the boundaries of topologically associated domains.

10.2 NUCLEOSOME MODIFICATIONS AND GENOME EXPRESSION
Acetylation of histones influences many nuclear activities including genome expression. Histone deacetylation represses active regions of the genome. Acetylation is not the only type of histone modification. Nucleosome repositioning also influences gene expression.

10.3 DNA MODIFICATION AND GENOME EXPRESSION
Genome silencing by DNA methylation. Methylation is involved in genomic imprinting and X inactivation.

SUMMARY
SHORT ANSWER QUESTIONS
IN-DEPTH PROBLEMS
FURTHER READING

CHAPTER 11
THE ROLE OF DNA-BINDING PROTEINS IN GENOME EXPRESSION

11.1 METHODS FOR STUDYING DNA-BINDING PROTEINS AND THEIR ATTACHMENT SITES
X-ray crystallography provides structural data for any protein that can be crystallized. NMR spectroscopy is used to study the structures of small proteins. Gel retardation identifies DNA fragments that bind to proteins.
Protection assays pinpoint binding sites with greater accuracy
Modification interference identifies nucleotides central to protein binding
Genomewide scans for protein attachment sites

11.2 THE SPECIAL FEATURES OF DNA-BINDING PROTEINS
The helix–turn–helix motif is present in prokaryotic and eukaryotic proteins
Zinc fingers are common in eukaryotic proteins
Other nucleic acid-binding motifs

11.3 INTERACTION BETWEEN DNA AND ITS BINDING PROTEINS
Direct readout of the nucleotide sequence
The nucleotide sequence has a number of indirect effects on helix structure
Contacts between DNA and proteins

SUMMARY
SHORT ANSWER QUESTIONS
IN-DEPTH PROBLEMS
FURTHER READING

CHAPTER 12
TRANSCRIPTOMES

12.1 COMPONENTS OF THE TRANSCRIPTOME
The mRNA fraction of a transcriptome is small but complex
Short noncoding RNAs have diverse functions
Long noncoding RNAs are enigmatic transcripts
Microarray analysis and RNA sequencing are used to study the contents of transcriptomes

12.2 SYNTHESIS OF THE COMPONENTS OF THE TRANSCRIPTOME
RNA polymerases are molecular machines for making RNA
Transcription start points are indicated by promoter sequences
Synthesis of bacterial RNA is regulated by repressor and activator proteins
Synthesis of bacterial RNA is also regulated by control over transcription termination
Synthesis of eukaryotic RNA is regulated primarily by activator proteins

12.3 DEGRADATION OF THE COMPONENTS OF THE TRANSCRIPTOME
Several processes are known for nonspecific RNA turnover

RNA silencing was first identified as a means of destroying invading viral RNA
MicroRNAs regulate genome expression by causing specific target mRNAs to be degraded

12.4 INFLUENCE OF RNA PROCESSING ON THE COMPOSITION OF A TRANSCRIPTOME
The splicing pathway for eukaryotic pre-mRNA introns
The splicing process must have a high degree of precision
Enhancer and silencer elements specify alternative splicing pathways

12.5 TRANSCRIPTOMES IN RESEARCH
Transcriptome analysis as an aid to genome annotation
Cancer transcriptomes
Transcriptomes and the responses of plants to stress

SUMMARY
SHORT ANSWER QUESTIONS
IN-DEPTH PROBLEMS
FURTHER READING

CHAPTER 13
PROTEOMES

13.1 STUDYING THE COMPOSITION OF A PROTEOME
The separation stage of a protein profiling project
The identification stage of a protein profiling project
Comparing the compositions of two proteomes
Analytical protein arrays offer an alternative approach to protein profiling

13.2 IDENTIFYING PROTEINS THAT INTERACT WITH ONE ANOTHER
Identifying pairs of interacting proteins
Identifying the components of multiprotein complexes
Identifying proteins with functional interactions
Protein interaction maps display the interactions within a proteome

13.3 SYNTHESIS AND DEGRADATION OF THE COMPONENTS OF THE PROTEOME
Ribosomes are molecular machines for making proteins
Yeast mating types are determined by gene conversion events
Genome rearrangements are responsible for immunoglobulin and T-cell receptor diversity

14.3 CHANGES IN GENOME ACTIVITY UNDERLYING DEVELOPMENT
Bacteriophage λ: a genetic switch enables a choice to be made between alternative developmental pathways
* Bacillus* sporation: coordination of activities in two distinct cell types
* Caenorhabditis elegans*: the genetic basis of positional information and the determination of cell fate
* Fruit flies*: conversion of positional information into a segmented body plan
Homeotic selector genes are universal features of higher eukaryotic development
Homeotic genes also underlie plant development

SUMMARY

SHORT ANSWER QUESTIONS
IN-DEPTH PROBLEMS
FURTHER READING

CHAPTER 14

GENOME EXPRESSION IN THE CONTEXT OF CELL AND ORGANISM

14.1 THE RESPONSE OF THE GENOME TO EXTERNAL SIGNALS
Signal transmission by import of the extracellular signaling compound
Receptor proteins transmit signals across cell membranes
Some signal transduction pathways have few steps between receptor and genome
Some signal transduction pathways have many steps between receptor and genome
Some signal transduction pathways operate via second messengers

14.2 CHANGES IN GENOME ACTIVITY RESULTING IN CELLULAR DIFFERENTIATION
Some differentiation processes involve changes to chromatin structure

During stress, bacteria inactivate their ribosomes in order to downsize the proteome
Initiation factors mediate large-scale remodeling of eukaryotic proteomes
The translation of individual mRNAs can also be regulated
Degradation of the components of the proteome

13.4 INFLUENCE OF PROTEIN PROCESSING ON THE COMPOSITION OF THE PROTEOME
The amino acid sequence contains instructions for protein folding
Some proteins are activated by proteolytic cleavage
Important changes in protein activity can be brought about by chemical modification

13.5 BEYOND THE PROTEOME
The metabolome is the complete set of metabolites present in a cell
Systems biology provides an integrated description of cellular activity

SUMMARY
SHORT ANSWER QUESTIONS
IN-DEPTH PROBLEMS
FURTHER READING

CHAPTER 15

GENOME REPLICATION

15.1 THE TOPOLOGY OF GENOME REPLICATION
The double-helical structure complicates the replication process
The Meselson–Stahl experiment proved that replication is semiconservative
DNA topoisomerases provide a solution to the topological problem
Variations on the semiconservative theme

15.2 THE INITIATION PHASE OF GENOME REPLICATION
Initiation at the E. coli origin of replication
Origins of replication have been clearly defined in yeast
Origins in higher eukaryotes have been less easy to identify

15.3 EVENTS AT THE REPLICATION FORK
DNA polymerases are molecular machines for making (and degrading) DNA
DNA polymerases have limitations that complicate genome replication
Okazaki fragments must be joined together to complete lagging-strand replication

**15.4 TERMINATION OF GENOME REPLICATION**
Replication of the *E. coli* genome terminates within a defined region
Little is known about termination of replication in eukaryotes
Telomerase completes replication of chromosomal DNA molecules, at least in some cells
Telomere length is implicated in cell senescence and cancer
*Drosophila* has a unique solution to the end-shortening problem

**15.5 REGULATION OF EUKARYOTIC GENOME REPLICATION**
Genome replication must be synchronized with the cell cycle
Origin licensing is the prerequisite for passing the G1–S checkpoint
Replication origins do not all fire at the same time
The cell has various options if the genome is damaged

**SUMMARY**

**SHORT ANSWER QUESTIONS**

**IN-DEPTH PROBLEMS**

**FURTHER READING**

---

Defects in DNA repair underlie human diseases, including cancers

**CHAPTER 16 MUTATIONS AND DNA Repair**

**16.1 THE CAUSES OF MUTATIONS**
Errors in replication are a source of point mutations
Replication errors can also lead to insertion and deletion mutations
Mutations are also caused by chemical and physical mutagens

**16.2 REPAIR OF MUTATIONS AND OTHER TYPES OF DNA DAMAGE**
Direct repair systems fill in nicks and correct some types of nucleotide modification
Base excision repairs many types of damaged nucleotide
Nucleotide excision repair is used to correct more extensive types of damage
Mismatch repair corrects replication errors
Single- and double-strand breaks can be repaired
If necessary, DNA damage can be bypassed during genome replication

---

**CHAPTER 17 RECOMBINATION AND TRANPOSITION**

**17.1 HOMOLOGOUS RECOMBINATION**
The Holliday and Meselson–Radding models for homologous recombination
The double-strand break model for homologous recombination
RecBCD is the most important pathway for homologous recombination in bacteria
*E. coli* can also carry out homologous recombination by the RecFOR pathway
Homologous recombination pathways in eukaryotes
The primary role of homologous recombination is thought to be DNA repair

**17.2 SITE-SPECIFIC RECOMBINATION**
Bacteriophage λ uses site-specific recombination during the lysogenic infection cycle
Site-specific recombination is an aid in construction of genetically modified plants

**17.3 TRANSPOSITION**
Replicative and conservative transposition of DNA transposons
Retroelements transpose replicatively via an RNA intermediate

**SUMMARY**

**SHORT ANSWER QUESTIONS**

**IN-DEPTH PROBLEMS**

**FURTHER READING**

---

**CHAPTER 18 HOW GENOMES EVOLVE**

**18.1 GENOMES: THE FIRST 10 BILLION YEARS**
The first biochemical systems were centered on RNA
The first DNA genomes
How unique is life?
18.2 EVOLUTION OF INCREASINGLY COMPLEX GENOMES
Genome sequences provide extensive evidence of past gene duplications 434
A variety of processes could result in gene duplication 434
Whole-genome duplication is also possible 438
Smaller duplications can also be identified in the human genome and other genomes 442
Both prokaryotes and eukaryotes acquire genes from other species 444
Genome evolution also involves rearrangement of existing genes 445
There are competing hypotheses for the origins of introns 448
The evolution of the epigenome 449

18.3 GENOMES: THE LAST 6 MILLION YEARS
The human genome is very similar to that of the chimpanzee 450
Paleogenomics is helping us understand the recent evolution of the human genome 451

18.4 GENOMES TODAY: DIVERSITY IN POPULATIONS
The origins of HIV/AIDS 453
The first migrations of humans out of Africa 454
The diversity of plant genomes is an aid in crop breeding 455

SUMMARY 458
SHORT ANSWER QUESTIONS 459
IN-DEPTH PROBLEMS 460
FURTHER READING 460

GLOSSARY 463
INDEX 491
Life as we know it is specified by the genomes of the myriad organisms with which we share the planet. Every organism possesses a genome that contains the biological information needed to construct and maintain a living example of that organism. Most genomes, including the human genome and those of all other cellular life forms, are made of DNA (deoxyribonucleic acid), but a few viruses have RNA (ribonucleic acid) genomes. DNA and RNA are polymeric molecules made up of chains of monomeric subunits called nucleotides. Each molecule of DNA comprises two polymers wound around one another to form the famous double helix, in which the two strands are held together by chemical bonds that link adjacent nucleotides into structures called base pairs.

The human genome, which is typical of the genomes of all multicellular animals, consists of two distinct parts (Figure 1.1):

- The nuclear genome comprises approximately 3,235,000,000 base pairs of DNA, divided into 24 linear molecules, the shortest 48,000,000 base pairs in length and the longest 250,000,000 base pairs, each contained in a different chromosome. These 24 chromosomes consist of 22 autosomes and the two sex chromosomes, X and Y. Altogether, some 45,500 genes are present in the human nuclear genome.

- The mitochondrial genome is a circular DNA molecule of 16,569 base pairs, up to 10 copies of which are present in each of the energy-generating organelles called mitochondria. The human mitochondrial genome contains just 37 genes.

Each of the approximately $10^{13}$ cells in the adult human body has its own copy or copies of the nuclear genome, the only exceptions being those few cell types, such as red blood cells, that lack a nucleus in their fully differentiated state. The vast majority of cells are diploid and so have two copies of each autosome, plus two sex chromosomes, XX for females or XY for males—46 chromosomes in all. These are called somatic cells, in contrast to sex cells, or gametes, which are haploid and have just 23 chromosomes, one of each autosome and one sex chromosome. Each cell also has multiple copies of the mitochondrial genome: 2000–7000 copies in somatic cells, such as those in the liver and heart tissue, and over 100,000 copies in each female oocyte.
The genome is a store of biological information, but on its own it is unable to release that information to the cell. Utilization of the biological information contained in the genome requires the coordinated activity of enzymes and other proteins, which participate in a complex series of biochemical reactions referred to as genome expression (Figure 1.2). The initial product of genome expression is the transcriptome, a collection of RNA molecules derived from those genes that are active in the cell at a particular time. The transcriptome is maintained by the process called transcription, in which individual genes are copied into RNA molecules. The second product of genome expression is the proteome, the cell’s repertoire of proteins, which specifies the nature of the biochemical reactions that the cell is able to carry out. The proteins that make up the proteome are synthesized by translation of some of the individual RNA molecules present in the transcriptome.

This book is about genomes and genome expression. It explains how genomes are studied (Part I), how they are organized (Part II), how they function (Part III), and how they replicate and evolve (Part IV). It was not possible to write this book until quite recently. Since the 1950s, molecular biologists have studied individual genes or small groups of genes, and from these studies they have built up a wealth of knowledge about how genes work. But only during the last few years have techniques been available that make it possible to examine entire genomes. Individual genes are still intensively studied, but information about individual genes is now interpreted within the context of the genome as a whole. This new, broader emphasis applies not just to genomes but to all of biochemistry and cell biology. No longer is it sufficient simply to understand individual biochemical pathways or subcellular processes. The challenge now is provided by systems biology, which attempts to link together these pathways and processes into networks that describe the overall functioning of living cells and living organisms.

This book will lead you through our knowledge of genomes and show you how this exciting area of research is underpinning our developing understanding of biological systems. First, however, we must pay attention to the basic principles of molecular biology by reviewing the key features of the three types of biological molecule involved in genomes and genome expression: DNA, RNA, and protein.

### 1.1 DNA

DNA was discovered in 1869 by Friedrich Miescher, a Swiss biochemist working in Tübingen, Germany. The first extracts that Miescher made from human white blood cells were crude mixtures of DNA and chromosomal proteins, but the following year he moved to Basel, Switzerland (where the research institute
named after him is now located), and prepared a pure sample of nucleic acid from salmon sperm. Miescher’s chemical tests showed that DNA is acidic and rich in phosphorus and also suggested that the individual molecules are very large, although it was not until the 1930s, when biophysical techniques were applied to DNA, that the huge lengths of the polymeric chains were fully appreciated.

**Genes are made of DNA**

The fact that genes are made of DNA is so well known today that it can be difficult to appreciate that for the first 75 years after its discovery the true role of DNA was unsuspected. As early as 1903, W. S. Sutton had realized that the inheritance patterns of genes parallel the behavior of chromosomes during cell division, an observation that led to the **chromosome theory**, the proposal that genes are located in chromosomes. Examination of cells by **cytochemistry**, which makes use of stains that bind specifically to just one type of biochemical, showed that chromosomes are made of DNA and protein, in roughly equal amounts. Biologists at that time recognized that billions of different genes must exist and the genetic material must therefore be able to take many different forms. But this requirement appeared not to be satisfied by DNA, because in the early part of the twentieth century it was thought that all DNA molecules were the same. On the other hand, it was known, correctly, that proteins are highly variable, polymeric molecules, each one made up of a different combination of 20 chemically distinct amino acid monomers (**Section 1.3**). Genes simply had to be made of protein, not DNA.

The errors in understanding DNA structure lingered on, but by the late 1930s it had become accepted that DNA, like protein, has immense variability. The notion that protein was the genetic material initially remained strong but was eventually overturned by the results of two important experiments:

- **Oswald Avery**, Colin MacLeod, and Maclyn McCarty showed that DNA is the active component of the **transforming principle**, a bacterial cell extract that, when mixed with a harmless strain of *Streptococcus pneumoniae*, converts these bacteria into a virulent form capable of causing pneumonia when injected into mice (**Figure 1.3A**). In 1944, when the results of this experiment were published, only a few microbiologists appreciated that transformation involves transfer of genes from the cell extract into the living bacteria. However, once this point had been accepted, the true meaning of the Avery experiment became clear: bacterial genes must be made of DNA.

- **Alfred Hershey** and Martha Chase used **radiolabeling** to show that when a bacterial culture is infected with **bacteriophages** (also called phages, a type of virus), DNA is the major component of the bacteriophages that enters the cells (**Figure 1.3B**). This was a vital observation because it was known that, during the infection cycle, the genes of the infecting bacteriophages are used to direct synthesis of new bacteriophages, and this synthesis occurs within the bacteria. If only the DNA of the infecting bacteriophages enters the cells, then it follows that the genes of these bacteriophages must be made of DNA.

Although from our perspective these two experiments provide the key results that tell us that genes are made of DNA, biologists at the time were not so easily convinced. Both experiments have limitations that leave room for skeptics to argue that protein could still be the genetic material. For example, there were worries about the specificity of the **deoxyribonuclease** enzyme that Avery and colleagues used to inactivate the transforming principle. This result, a central part of the evidence for the transforming principle being DNA, would be invalid if, as seemed possible, the enzyme contained trace amounts of a contaminating protease and hence was also able to degrade protein. Neither is the bacteriophage experiment conclusive, as Hershey and Chase stressed when they published their results: “Our experiments show clearly that a physical separation of phage T2 into genetic and nongenetic parts is possible ... The chemical identification of the
In retrospect, these two experiments are important not because of what they tell us but because they alerted biologists to the fact that DNA might be the genetic material and was therefore worth studying. This is what influenced Watson and Crick to work on DNA, and as we will see, it was their discovery of the double-helix structure, which solved the puzzling question of how genes can replicate, that really convinced the scientific world that genes are made of DNA.

**DNA is a polymer of nucleotides**

The names of James Watson and Francis Crick are so closely linked with DNA that it is easy to forget that when they began their collaboration in October 1951, the detailed structure of the DNA polymer was already known. Their contribution was
not to determine the structure of DNA per se but to show that in living cells two DNA chains are intertwined to form the double helix. First, therefore, we should examine what Watson and Crick knew before they began their work.

DNA is a linear, unbranched polymer in which the monomeric subunits are four chemically distinct nucleotides that can be linked together in any order in chains that are hundreds, thousands, or even millions of units in length. Each nucleotide in a DNA polymer is made up of three components (Figure 1.4):

- **2′-Deoxyribose**, which is a pentose, a type of sugar composed of five carbon atoms. These five carbons are numbered 1′ (spoken as one-prime), 2′, and so on. The name 2′-deoxyribose indicates that this particular sugar is a derivative of ribose, in which the hydroxyl (–OH) group attached to the 2′-carbon of ribose has been replaced by a hydrogen (–H) group.

- A **nitrogenous base**, one of cytosine or thymine (single-ring pyrimidines) or adenine or guanine (double-ring purines). The base is attached to the 1′-carbon of the sugar by a β-N-glycosidic bond attached to nitrogen number one of the pyrimidine or number nine of the purine.

- A **phosphate group**, comprising one, two, or three linked phosphate units attached to the 5′-carbon of the sugar. The phosphates are designated α, β, and γ, with the α-phosphate being the one directly attached to the sugar.

A molecule made up of just the sugar and base is called a **nucleoside**; addition of the phosphates converts this to a nucleotide. Although cells contain nucleotides with one, two, or three phosphate groups, only the nucleoside triphosphates act as substrates for DNA synthesis. The full chemical names of the four nucleotides that polymerize to make DNA are

- 2′-deoxyadenosine 5′-triphosphate
- 2′-deoxycytidine 5′-triphosphate
- 2′-deoxyguanosine 5′-triphosphate
- 2′-deoxythymidine 5′-triphosphate

The abbreviations of these four nucleotides are dATP, dCTP, dGTP, and dTTP, respectively, or when referring to a DNA sequence, A, C, G, and T, respectively.

In a polynucleotide, individual nucleotides are linked together by **phosphodiester bonds** between their 5′- and 3′-carbons (Figure 1.5). From the structure
of this linkage, we can see that the polymerization reaction (Figure 1.6) involves removal of the two outer phosphates (the β- and γ-phosphates) from one nucleotide and replacement of the hydroxyl group attached to the 3'-carbon of the second nucleotide. Note that the two ends of the polynucleotide are chemically distinct, one having an unreacted triphosphate group attached to the 5’-carbon (the 5’-P terminus) and the other having an unreacted hydroxyl attached to the 3’-carbon (the 3'-OH terminus). This means that the polynucleotide has a chemical direction, expressed as either 5’→3’ (down in Figure 1.5) or 3’→5’ (up in Figure 1.5). An important consequence of the polarity of the phosphodiester bond is that the chemical reaction needed to extend a DNA polymer in the 5’→3’ direction is different from that needed to make a 3’→5’ extension. The DNA polymerase enzymes present in living organisms are only able to carry out 5’→3’ synthesis, which adds significant complications to the process by which double-stranded DNA is replicated (Section 15.3).

In the years before 1950, various lines of evidence had shown that cellular DNA molecules are composed of two or more polynucleotides assembled together in some way. The possibility that unraveling the nature of this assembly might provide insights into how genes work prompted Watson and Crick, among others, to try to solve the structure. According to Watson in his book The Double Helix, their work was a desperate race against the famous American biochemist Linus Pauling, who initially proposed an incorrect triple-helix model, giving Watson and Crick the time they needed to complete the double-helix structure. It is now difficult to separate fact from fiction, especially regarding the part played by Rosalind Franklin, whose X-ray diffraction studies provided the bulk of the experimental data in support of the double helix and who was herself very close to solving the structure. The one thing that is clear is that the double helix, discovered by Watson and Crick on Saturday, March 7, 1953, was the single most important breakthrough in biology during the twentieth century.

The discovery of the double helix can be looked on as one of the first multidisciplinary biological research projects. Watson and Crick used four quite different types of information to deduce the double-helix structure:

- Biophysical data of various kinds were used to infer some of the key features of the structure. The water content of DNA fibers was particularly important because it enabled the density of the DNA in a fiber to be estimated. The number of strands in the helix and the spacing between the nucleotides had to be compatible with the fiber density. Pauling’s triple-helix model was based on an incorrect density measurement that suggested that the DNA molecule was more closely packed than is actually the case.
1.1 DNA

Chapter 1: Genomes, Transcriptomes, and Proteomes

- X-ray diffraction patterns (Section 11.1), most of which were produced by Rosalind Franklin, revealed the detailed helical structure (Figure 1.7).

- The base ratios, which had been discovered by Erwin Chargaff of Columbia University in New York, enabled the pairing between the polynucleotides in the helix to be deduced. Chargaff had carried out a lengthy series of chromatographic studies of DNA samples from various sources and showed

Figure 1.6 The polymerization reaction that results in synthesis of a DNA polynucleotide. Synthesis occurs in the 5' → 3' direction, with the new nucleotide being added to the 3'-carbon at the end of the existing polynucleotide. The β- and γ-phosphates of the nucleotide are removed as a pyrophosphate molecule.

Figure 1.7 Franklin’s photo 51 showing the X-ray diffraction pattern obtained with a fiber of DNA. The cross shape indicates that DNA has a helical structure, and the extent of the shadowing within the diamond spaces above, below, and to either side of the cross show that the sugar–phosphate backbone is on the outside of the helix (see Figure 1.9). The positions of the various smears that make up the arms of the cross enable dimensions such as the diameter, rise per base pair, and pitch (see Table 1.1) of the molecule to be calculated. The missing smears (the gap in each arm of the cross, marked by the arrows) indicate the relative positioning of the two polynucleotides. These missing smears enabled Watson and Crick to recognize that there are two grooves of different depths on the outer surface of the helix (see Figure 1.9). (From Franklin R & Gosling RG [1953] Nature 171:740–741. With permission from Macmillan Publishers Ltd.)
1. **DNA**

**Chapter 1: Genomes, Transcriptomes, and Proteomes**

... that although the values are different in different organisms, the amount of adenine is always the same as the amount of thymine and the amount of guanine is the same as that of cytosine (Figure 1.8). These base ratios led to the base-pairing rules, which were the key to the discovery of the double-helix structure.

- The construction of scale models of possible DNA structures, which was the only major technique that Watson and Crick performed themselves, enabled the relative positioning of the various atoms to be checked, to ensure that pairs that formed bonds were not too far apart and that other atoms were not so close together as to interfere with one another.

**The double helix is stabilized by base pairing and base stacking**

The double helix is right-handed, which means that if it were a spiral staircase and you were climbing upward, then the rail on the outside of the staircase would be on your right-hand side. The two strands run in opposite directions (Figure 1.9A). The helix is stabilized by two types of chemical interaction:

- **Base pairing** between the two strands involves the formation of hydrogen bonds between an adenine on one strand and a thymine on the other strand, or between a cytosine and a guanine (Figure 1.9B). Hydrogen bonds are weak electrostatic interactions between an electronegative atom (such as oxygen or nitrogen) and a hydrogen atom attached to a second electronegative atom. Hydrogen bonds are longer than covalent bonds and are much weaker; typical bond energies are 8-29 kJ mol⁻¹ at 25°C, compared with up to 348 kJ mol⁻¹ for a single covalent bond between a pair of carbon atoms. As well as their role in the DNA double helix, hydrogen bonds stabilize protein secondary structures. The two base-pair combinations—A base-paired with T and G base-paired with C—explain the base ratios discovered by Chargaff. These are the only pairs that are permissible, partly because of the geometries of the nucleotide bases and the relative positions of the atoms that are able to participate in hydrogen bonds, and partly because the pair must be between a purine and a pyrimidine: a purine-purine pair would be too big to fit within the helix, and a pyrimidine-pyrimidine pair would be too small.

- **Base stacking** involves attractive forces between adjacent base pairs and adds stability to the double helix once the strands have been brought together by base pairing. Base stacking is sometimes called π-π interactions, because it is thought to involve the π electrons associated with the double bonds of the purine and pyrimidine structures. However, this hypothesis is now being questioned, and the possibility that base stacking involves a type of electrostatic interaction is being explored.

Both base pairing and base stacking are important in holding the two polynucleotides together, but base pairing has added significance because of its biological implications. The limitation that A can only base-pair with T and G can only base-pair with C means that DNA replication can result in perfect copies of a parent molecule through the simple expedient of using the sequences of the preexisting strands to dictate the sequences of the new strands. This is template-dependent DNA synthesis, the system used by all cellular DNA polymerases.
The double helix has structural flexibility  

The double helix described by Watson and Crick, and shown in Figure 1.9A, is called the B-form of DNA or B-DNA. Its characteristic features lie in its dimensions: a helical diameter of 2.37 nm, a rise of 0.34 nm per base pair, and a pitch (the distance taken up by a complete turn of the helix) of 3.4 nm, corresponding to 10 base pairs (bp) per turn. The DNA in living cells is thought to be predominantly in this B-form, but it is now clear that genomic DNA molecules are not entirely uniform in structure. This is mainly because each nucleotide in the helix has the flexibility to take up a slightly different molecular shape. To adopt these different conformations, the relative positions of the atoms in the nucleotide must change slightly. There are a number of possibilities but the most important conformational changes are as follows:

- Rotation around the β-N-glycosidic bond changes the orientation of the base relative to the sugar: the two possibilities are called the anti- and syn-conformations (Figure 1.10A). Base rotation influences the positioning of the two polynucleotides.
Figure 1.10 Changes in nucleotide configuration that can affect the conformation of the double helix. (A) Structures of anti- and syn-deoxyadenosine. The two structures differ in the orientation of the base relative to the sugar component of the nucleoside; rotation around the β-N-glycosidic bond converts one form into the other. The three other nucleosides also have anti- and syn-conformations. (B) Sugar pucker, illustrating the positioning of the sugar carbons in the C2'-endo- and C3'-endo-configurations.

- **Sugar pucker** refers to the three-dimensional shape of the sugar. The ribose component of the nucleotide does not have a planar structure: when it is viewed from the side, one or two of the carbon atoms are either above or below the plane of the sugar (Figure 1.10B). In the C2'-endo-configuration, the 2'-carbon is above the plane and the 3'-carbon is slightly below, and in the C3'-endo-configuration, the 3'-carbon is above the plane and the 2'-carbon is below. Because the 3'-carbon participates in the phosphodiester bond with the adjacent nucleotide, the two pucker configurations have different effects on the conformation of the sugar–phosphate backbone.

Conformation changes resulting from rotation around the β-N-glycosidic bond and sugar pucker can give rise to major changes in the overall structure of the helix. It has been recognized since the 1950s that changes in the dimensions of the double helix occur when fibers containing DNA molecules are exposed to different relative humidities. For example, the modified version of the double helix called A-DNA has a diameter of 2.55 nm, a rise of 0.23 nm per base pair, and a pitch of 2.5 nm, corresponding to 11 base pairs per turn (Table 1.1). Like the B-form, A-DNA is a right-handed helix and the bases are in the anti-conformation relative to the sugar. The main difference lies with the sugar pucker: the sugars in the B-form are in the C2'-endo-configuration, and those in A-DNA are in the C3'-endo-configuration. Other right-handed variations of the double helix include B', C', C'', D', E', and T-DNAs.

A more drastic reorganization is also possible, leading to the left-handed Z-DNA, in which the sugar–phosphate backbone adopts an irregular zigzag conformation. Z-DNA is a more tightly wound version of the double helix with 12 bp per turn and a diameter of only 1.84 nm (Table 1.1). It is known to occur in regions of a double helix that contain repeats of the motif GC (that is, the sequence of each strand is ...GCGGGGCGG...). In these regions, each G nucleotide has the syn- and C3'-endo-conformations and each C has the anti- and C2'-endo-conformations.

### TABLE 1.1 FEATURES OF DIFFERENT CONFORMATIONS OF THE DNA DOUBLE HELIX

<table>
<thead>
<tr>
<th>Feature</th>
<th>A-DNA</th>
<th>B-DNA</th>
<th>Z-DNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type of helix</td>
<td>Right-handed</td>
<td>Right-handed</td>
<td>Left-handed</td>
</tr>
<tr>
<td>Helical diameter (nm)</td>
<td>2.55</td>
<td>2.37</td>
<td>1.84</td>
</tr>
<tr>
<td>Distance between base pairs (nm)</td>
<td>0.23</td>
<td>0.34</td>
<td>0.38</td>
</tr>
<tr>
<td>Distance per complete turn (nm)</td>
<td>2.5</td>
<td>3.4</td>
<td>4.6</td>
</tr>
<tr>
<td>Number of base pairs per turn</td>
<td>11</td>
<td>10</td>
<td>12</td>
</tr>
<tr>
<td>Base orientation</td>
<td>anti</td>
<td>anti</td>
<td>mixture</td>
</tr>
<tr>
<td>Sugar pucker</td>
<td>C3'-endo</td>
<td>C2'-endo</td>
<td>mixture</td>
</tr>
</tbody>
</table>
The bare dimensions of the various forms of the double helix do not reveal what are probably the most significant differences between them. These relate not to diameter and pitch but to the extent to which internal regions of the helix are accessible from the surface of the structure. As shown in Figure 1.9A, the B-form of DNA does not have an entirely smooth surface: instead, two grooves spiral along the length of the helix. One of these grooves is relatively wide and deep and is called the major groove; the other is narrow and less deep and is called the minor groove. A-DNA also has two grooves (Figure 1.11), but with this conformation the major groove is even deeper and the minor groove is shallower compared with B-DNA. Z-DNA is different again, with the major groove virtually nonexistent but the minor groove very narrow and deep. In each form of DNA, part of the internal surface of at least one of the grooves is formed by chemical groups attached to the nucleotide bases. In Chapter 11 we will see that expression of the biological information contained within a genome is mediated by DNA-binding proteins that attach to the double helix and regulate the activity of the genes contained within it. To carry out its function, each DNA-binding protein must attach at a specific position near the gene whose activity it will influence. This can be achieved, with at least some degree of accuracy, by the protein reaching down into a groove, within which the DNA sequence can be read without the helix being opened up by breaking the base pairs. A corollary of this is that a DNA-binding protein whose structure enables it to recognize a specific nucleotide sequence within B-DNA, for example, might not be able to recognize that sequence if the DNA has taken up a different conformation. As we will see in Section 11.3, conformational variations along the length of a DNA molecule, together with other structural polymorphisms caused by the nucleotide sequence, could be important in determining the specificity of interactions between the genome and its DNA-binding proteins.

### 1.2 RNA AND THE TRANSCRIPTOME

The initial product of genome expression is the transcriptome (see Figure 1.2), the collection of RNA molecules derived from those genes that are active in the cell at a particular time. The RNA molecules of the transcriptome are synthesized by the process called transcription. In this section we will examine the structure of RNA and then look more closely at the various types of RNA molecules that are present in living cells.

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**Figure 1.11** A-, B-, and Z-forms of the double helix. The major and minor grooves on each molecule are indicated by M and m, respectively. (Courtesy of Richard Wheeler under GFDL 1.2.)
RNA is a second type of polynucleotide

RNA is a polynucleotide similar to DNA but with two important chemical differences (Figure 1.12). First, the sugar in an RNA nucleotide is ribose, and second, RNA contains uracil instead of thymine. The four nucleotide substrates for synthesis of RNA are therefore:

- adenosine 5’-triphosphate
- cytidine 5’-triphosphate
- guanosine 5’-triphosphate
- uridine 5’-triphosphate

These nucleotides are abbreviated as ATP, CTP, GTP, and UTP or as A, C, G, and U, respectively.

As with DNA, RNA polynucleotides contain 3’–5’ phosphodiester bonds, but these phosphodiester bonds are less stable than those in a DNA polynucleotide because of the indirect effect of the hydroxyl group at the 2’-position of the sugar. RNA molecules are rarely more than a few thousand nucleotides in length, and although many form intramolecular base pairs (for example, see Figure 5.6A), most are single- rather than double-stranded.

The enzymes responsible for transcription of DNA into RNA are called DNA-dependent RNA polymerases. The name indicates that the enzymatic reaction they catalyze results in polymerization of RNA from ribonucleotides and occurs in a DNA-dependent manner, meaning that the sequence of nucleotides in a DNA template dictates the sequence of nucleotides in the RNA that is made (Figure 1.13). It is permissible to shorten the enzyme name to RNA polymerase, as the context in which the name is used means that there is rarely confusion with the RNA-dependent RNA polymerases that are involved in replication and expression of some viral genomes. The chemical basis of template-dependent RNA synthesis is equivalent to that shown for the synthesis of DNA in Figure 1.6. Ribonucleotides are added one after another to the growing 3’-end of the RNA transcript, with the identity of each nucleotide being specified by the base-pairing rules: A base-pairs with T or U, and G base-pairs with C. During each nucleotide addition, the β- and γ-phosphates are removed from the incoming nucleotide, and the hydroxyl group is removed from the 3’-carbon of the nucleotide at the end of the chain, precisely the same as for DNA polymerization.

The RNA content of the cell

A typical bacterium contains 0.05–0.10 pg of RNA, making up about 6% of its total weight. A mammalian cell, being much larger, contains more RNA, 20–30 pg in all, but this represents only 1% of the cell as a whole.

The best way to understand the RNA content of a cell is to divide it into categories and subcategories depending on function. There are several ways of doing this; the most informative scheme is the one shown in Figure 1.14. The primary division is between coding RNA and noncoding RNA. Coding RNA is made up of just one class of molecule, the messenger RNAs (mRNAs), which are transcripts of protein-coding genes and hence are translated into protein in the second stage of genome expression. Messenger RNAs rarely make up more than 4% of the total RNA and are short-lived, being degraded soon after synthesis. Bacterial mRNAs have half-lives of no more than a few minutes, and in euukaryotes most mRNAs are degraded within a few hours after synthesis. This rapid turnover means that the mRNA composition of the cell is not fixed and can quickly be restructured by changing the rate of synthesis of individual mRNAs.

The second type of RNA is referred to as noncoding, as these molecules are not translated into protein. An alternative name is functional RNA, which emphasizes that the noncoding RNAs still have essential roles within the cell. There are several diverse types of noncoding RNA, the two most important being as follows:
• **Ribosomal RNAs (rRNAs)** are present in all organisms and are usually the most abundant RNAs in the cell, making up over 80% of the total RNA in actively dividing bacteria. These molecules are components of **ribosomes**, the structures within which protein synthesis takes place (Section 13.3).

• **Transfer RNAs (tRNAs)** are small molecules that are also involved in protein synthesis and, like rRNA, are found in all organisms. The function of tRNAs is to carry amino acids to the ribosome and ensure that the amino acids are linked together in the order specified by the nucleotide sequence of the mRNA that is being translated (Section 13.3).

These are the two most important categories of noncoding RNA, but there are several other types with specialist roles in eukaryotic or bacterial cells. In eukaryotes, these RNAs are usually divided into two groups, the **short noncoding RNAs (sncRNAs)**, comprising RNAs less than 200 nucleotides in length, and the **long noncoding RNAs (lncRNAs)**, made up of molecules longer than 200 nucleotides. We will examine the roles of these various types of noncoding RNA in Chapter 12.

**Many RNAs are synthesized as precursor molecules**

As well as the mature RNAs described above, cells also contain precursor molecules. Many RNAs, especially in eukaryotes, are initially synthesized as precursor or **pre-RNA**, which has to be processed in order to release the functional molecules.

The most important of these processing events is **splicing**. Some eukaryotic genes contain internal segments that are copied during transcription but then excised from the pre-RNA (Figure 1.15). These excised segments are called **introns**, in contrast to the **exons**, which are spliced together to form the mature RNA. Introns are present in some rRNA and tRNA genes but are particularly common in protein-coding genes. Splicing of **pre-mRNA** is therefore a major part of the process that results in synthesis of the protein-coding component of the

---

**Figure 1.14** The RNA content of a cell. This scheme shows the types of RNA present in all organisms and those categories found only in eukaryotic cells. Precursor RNAs are included.

**Figure 1.15** Splicing of a eukaryotic pre-mRNA. The introns are cut out of the pre-mRNA and the exons are rejoined to give the functional mRNA.
transcriptome (Section 12.4). Splicing occurs in the nucleus, where the unspliced pre-mRNA forms the nuclear RNA fraction called heterogeneous nuclear RNA (hnRNA).

Splicing is not the only type of cutting event that occurs during processing of pre-RNA. Many rRNAs and tRNAs are initially synthesized as precursors that contain copies of more than one molecule. The pre-rRNAs and pre-tRNAs must therefore be cut into pieces to produce the mature RNAs (Figure 1.16). This type of processing occurs in both prokaryotes and eukaryotes.

Other processing events result in changes occurring at the ends of RNA molecules. These end modifications occur during the synthesis of eukaryotic mRNAs, most of which have a structure called a cap attached at the 5′-end and a poly(A) tail attached to the 3′-end. The cap structure comprises the modified nucleotide 7-methylguanosine linked to the first nucleotide in the pre-mRNA by a 5′–5′ triple-phosphate bond (Figure 1.17A). The first and second nucleotides in the pre-mRNA might also be modified by addition of methyl groups. The cap structure is needed to help initiate translation of the mRNA into a protein. The poly(A) tail is a series of up to 250 adenine nucleotides that are present at the 3′-end of the mRNA. The pre-mRNA is cut at a position close to its 3′-end, and the adenines are added to this new end by a template-independent RNA polymerase called poly(A) polymerase (Figure 1.17B). The function of this polyadenylation process is not fully understood, but we do know that if the poly(A) tail is absent or shorter than usual, then the mRNA is degraded.
The final type of processing event is chemical modification. Some bases within the rRNAs and tRNAs of all organisms are modified by methylation, deamination (removal of a -NH₂ group), and/or thio-substitution (replacement of oxygen with sulfur), and some bases undergo internal rearrangements that change the positions of particular groups and/or convert double bonds to single bonds (Figure 1.18). The reasons for many of these modifications are unknown, but functions have been assigned for specific cases. In tRNA, some of the modified nucleotides are recognized by the enzymes that attach an amino acid to the 3′-end of the molecule. This reaction is central to the role that tRNA plays during protein synthesis. The correct amino acid has to be attached to the correct tRNA, and the modifications within the tRNA are thought to provide some of the specificity that ensures that this happens.

A few eukaryotic mRNAs also undergo chemical modification. This process, called RNA editing, is uncommon but important because it can change the biological information in an mRNA in such a way that the protein coded by the mRNA has an altered structure. A notable example of RNA editing occurs with the human mRNA for apolipoprotein B. There are two versions of this protein: apolipoprotein B48, which is synthesized by intestinal cells, and apolipoprotein B100, which is about twice the size of B48 and is made in the liver. Both proteins are involved in the transport of lipids around the body, but their exact roles are different. The B48 protein forms part of the transport structure called a chylomicron, and B100 combines with other proteins to form a complex called a very low density lipoprotein. Both types of apolipoprotein B are specified by the same gene, but in intestinal cells a cytosine at position 6666 in the 14,000-nucleotide mRNA is edited by deamination. This change is sufficient to convert the mRNA for apolipoprotein B100 into an mRNA specifying apolipoprotein B48 (Figure 1.19).

There are different definitions of the transcriptome

Although most biologists now define the transcriptome as the total RNA content of a cell, the term, when first introduced in 1997, was initially used to describe just the mRNA component. The mRNA makes up less than 4% of the total cell RNA but is often looked on as the most significant component because it comprises the coding RNAs that are used in the next stage of genome expression. Even in the simplest organisms, such as bacteria and yeast, many different protein-coding genes are active at any one time. The mRNA content of a cell is therefore complex, containing copies of hundreds, if not thousands, of different genes. By specifying the set of proteins that the cell is able to make, the mRNA content determines the biochemical features of a cell. Many of the early studies of transcriptomes aimed to identify all, or as many as possible, of the mRNAs in a cell, in order to understand the overall pattern of gene expression and how that pattern changes when, for example, a cell becomes cancerous. These types of study are still important today (Section 12.5), and because they focus on mRNA, there remains a tendency to look on the transcriptome as just referring to the mRNA content of the cell.

The broader definition of the transcriptome to include all the RNA in a cell reflects our growing awareness of the important roles that noncoding RNAs play in specifying the biochemical properties of a cell. In particular, the small non-coding RNAs called microRNAs (miRNAs) regulate gene expression in eukaryotic cells by degrading...
those mRNAs whose products are no longer needed (Section 12.3). Human cells are able to make about 1000 miRNAs, each one specific for a single mRNA or small group of mRNAs. Understanding which miRNAs are synthesized in a particular cell, and how the pattern of miRNA synthesis changes in diseased cells, is an essential complement to the equivalent studies of mRNAs. It is therefore sensible to extend the term transcriptome to include all of the RNA in a cell, because a focus just on mRNA misses the vital role that other parts of the transcriptome play in mediating expression of the biological information contained in the genome.

1.3 PROTEINS AND THE PROTEOME

The second product of genome expression is the proteome (see Figure 1.2), the cell’s repertoire of proteins, which specifies the nature of the biochemical reactions the cell is able to carry out. These proteins are synthesized by translation of the mRNA component of the transcriptome.

There are four hierarchical levels of protein structure

A protein, like a DNA molecule, is a linear, unbranched polymer. In proteins, the monomeric subunits are called amino acids (Figure 1.20) and the resulting polymers, or polypeptides, are rarely more than 2000 units in length.

Proteins are traditionally looked upon as having four distinct levels of structure. These levels are hierarchical: meaning the protein is built up stage-by-stage, with each level of structure depending on the one below it.

- The primary structure of the protein is formed by joining amino acids into a polypeptide. The amino acids are linked by peptide bonds that are formed by a condensation reaction between the carboxyl group of one amino acid and the amino group of a second amino acid (Figure 1.21). Note that, as with a polynucleotide, the two ends of the polypeptide are chemically distinct: one has a free amino group and is called the amino-, NH2-, or N-terminus; the other has a free carboxyl group and is called the carboxyl-, COOH-, or C-terminus. The direction of the polypeptide can therefore be expressed as either N → C (left to right in Figure 1.21) or C → N (right to left in Figure 1.21).

- The secondary structure refers to the different conformations that can be taken up by the polypeptide. The two main types of secondary structure are the α-helix and β-sheet (Figure 1.22). These are stabilized mainly by hydrogen bonds that form between different amino acids in the polypeptide. Most polypeptides are long enough to be folded into a series of secondary structures, one after another, along the molecule.

- The tertiary structure results from folding the secondary structural components of the polypeptide into a three-dimensional configuration (Figure 1.23). The tertiary structure is stabilized by various chemical
forces, notably hydrogen bonding between individual amino acids, electrostatic interactions between the R groups of charged amino acids, and hydrophobic effects, which dictate that amino acids with nonpolar (water-hating) side groups must be shielded from water by embedding them within the internal regions of the protein. There may also be covalent linkages called disulfide bridges between cysteine amino acid residues at various places in the polypeptide.

- The quaternary structure involves the association of two or more polypeptides, each folded into its tertiary structure, into a multisubunit protein. Not all proteins form quaternary structures, but it is a feature of many proteins with complex functions, including several involved in genome expression. Some quaternary structures are held together by disulfide bridges between the different polypeptides, resulting in stable multisubunit proteins that cannot easily be broken down to the component parts. Other quaternary structures comprise looser associations of subunits stabilized by hydrogen bonding and hydrophobic effects, which means that these proteins can revert to their component polypeptides or change their subunit composition, according to the functional requirements of the cell.

**Amino acid diversity underlies protein diversity**

Proteins are functionally diverse because the amino acids from which proteins are made are themselves chemically diverse. Different sequences of amino acids therefore result in different combinations of chemical reactivities, and these combinations dictate not only the overall structure of the resulting protein but also the positioning on the surface of the structure of reactive groups that determine the chemical properties of the protein.

Amino acid diversity derives from the R group because this part is different in each amino acid and varies greatly in structure. Proteins are made up from a set of 20 amino acids (Figure 1.24; Table 1.2). Some of these have R groups that are small, relatively simple structures, such as a single hydrogen atom (in the amino acid called glycine) or a methyl group (alanine). Other R groups are large, complex aromatic side chains (phenylalanine, tryptophan, and tyrosine). Most amino acids are uncharged at pH 7.4 (the physiological pH of most cells and tissues), but two are negatively charged (aspartic acid and glutamic acid) and
Figure 1.24 Structures of the amino acid R groups. These 20 amino acids are the ones that are conventionally looked on as being specified by the genetic code. Note that the entire structure of proline is shown, not just its R group. This is because proline has an unusual structure in which the R group forms a bond not just to the α-carbon but also with the amino group attached to this carbon.

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Abbreviation</th>
<th>Three-letter</th>
<th>One-letter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine</td>
<td>Ala</td>
<td>A</td>
<td></td>
</tr>
<tr>
<td>Arginine</td>
<td>Arg</td>
<td>R</td>
<td></td>
</tr>
<tr>
<td>Asparagine</td>
<td>Asn</td>
<td>N</td>
<td></td>
</tr>
<tr>
<td>Aspartic acid</td>
<td>Asp</td>
<td>D</td>
<td></td>
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<tr>
<td>Cysteine</td>
<td>Cys</td>
<td>C</td>
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<tr>
<td>Glutamic acid</td>
<td>Glu</td>
<td>E</td>
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<tr>
<td>Glutamine</td>
<td>Gln</td>
<td>Q</td>
<td></td>
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<tr>
<td>Glycine</td>
<td>Gly</td>
<td>G</td>
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</tr>
<tr>
<td>Histidine</td>
<td>His</td>
<td>H</td>
<td></td>
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<tr>
<td>Isoleucine</td>
<td>Ile</td>
<td>I</td>
<td></td>
</tr>
<tr>
<td>Leucine</td>
<td>Leu</td>
<td>L</td>
<td></td>
</tr>
<tr>
<td>Lysine</td>
<td>Lys</td>
<td>K</td>
<td></td>
</tr>
<tr>
<td>Methionine</td>
<td>Met</td>
<td>M</td>
<td></td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>Phe</td>
<td>F</td>
<td></td>
</tr>
<tr>
<td>Proline</td>
<td>Pro</td>
<td>P</td>
<td></td>
</tr>
<tr>
<td>Serine</td>
<td>Ser</td>
<td>S</td>
<td></td>
</tr>
<tr>
<td>Threonine</td>
<td>Thr</td>
<td>T</td>
<td></td>
</tr>
<tr>
<td>Tryptophan</td>
<td>Trp</td>
<td>W</td>
<td></td>
</tr>
<tr>
<td>Tyrosine</td>
<td>Tyr</td>
<td>Y</td>
<td></td>
</tr>
<tr>
<td>Valine</td>
<td>Val</td>
<td>V</td>
<td></td>
</tr>
</tbody>
</table>
three are positively charged (arginine, histidine, and lysine). Some amino acids are polar (e.g. serine and threonine), while others are nonpolar (e.g. alanine, leucine, and valine).

The 20 amino acids shown in Figure 1.24 are conventionally looked upon as being specified by the genetic code. Therefore, they are the amino acids that are linked together when mRNA molecules are translated into proteins. However, these 20 amino acids do not, on their own, represent the limit of the chemical diversity of proteins. The diversity is even greater because of two factors:

- At least two additional amino acids—selenocysteine and pyrrolysine (Figure 1.25)—can be inserted into a polypeptide chain during protein synthesis. Their insertion is directed by a modified reading of the genetic code.

- During protein processing, some amino acids are modified by the addition of new chemical groups, for example, by acetylation or phosphorylation, or by attachment of large side chains made up of sugar units (Section 13.4).

Proteins therefore have an immense amount of chemical variability, some of this directly specified by the genome and the remainder arising by protein processing.

**The link between the transcriptome and the proteome**

The proteome comprises all the proteins present in a cell at a particular time. A typical mammalian cell, for example, a liver hepatocyte, is thought to contain 10,000–20,000 different proteins, about $8 \times 10^9$ individual molecules in all, representing approximately 0.5 ng of protein or 18–20% of the total cell weight. The copy numbers of individual proteins vary enormously, from less than 20,000 molecules per cell for the rarest types to 100 million copies for the commonest ones. Any protein that is present at a copy number of greater than 50,000 per cell is considered to be relatively abundant, and in the average mammalian cell some 2000 proteins fall into this category. When the proteomes of different types of mammalian cells are examined, very few differences are seen among these abundant proteins, suggesting that most of them are housekeeping proteins that perform general biochemical activities that occur in all cells. The proteins that provide the cell with its specialized function are often quite rare, although there are exceptions, such as the vast amounts of hemoglobin that are present only in red blood cells.

The proteome is synthesized by translation of the mRNA component of the transcriptome. In the early 1950s, shortly after the double-helix structure of DNA had been discovered, several molecular biologists attempted to devise ways in which amino acids could attach directly to mRNAs in an ordered fashion, but in all of these schemes at least some of the bonds had to be shorter or longer than was possible according to the laws of physical chemistry, and each idea was quietly dropped. Eventually, in 1957, Francis Crick cut a way through the confusion by predicting the existence of an adaptor molecule that would form a bridge between the mRNA and the polypeptide being synthesized. Soon afterward it was realized that the tRNAs are these adaptor molecules. Once this fact had been established, attention turned to the ribosomes, the structures within which proteins are synthesized. Gradually, a detailed understanding of the mechanism by which mRNAs are translated into polypeptides was built up (Section 13.3).

The other aspect of protein synthesis that interested molecular biologists in the 1950s was the informational problem. This refers to the second important component of the link between the transcriptome and proteome: the genetic code, which specifies how the nucleotide sequence of an mRNA is translated into the amino acid sequence of a protein. It was recognized in the 1950s that a triplet genetic code—one in which each code word, or codon, comprises three nucleotides—is required to account for all 20 amino acids found in proteins. A two-letter code would have only $4^2 = 16$ codons, which is not enough to specify all 20 amino acids, whereas a three-letter code would give $4^3 = 64$ codons. The genetic
code was worked out in the 1960s, partly by analysis of polypeptides arising from translation of artificial mRNAs of known or predictable sequence in cell-free protein-synthesizing systems, and partly by determining which amino acids associated with which RNA sequences in an assay based on purified ribosomes. When this work was completed, it was realized that the 64 codons fall into groups, where the members of each group code for the same amino acid (Figure 1.26). Only tryptophan and methionine have just a single codon each: all other amino acids are coded by two, three, four, or six codons. This feature of the code is called degeneracy. The code also has four punctuation codons, which indicate the points within an mRNA where translation of the nucleotide sequence should start and finish (Figure 1.27). The initiation codon is usually 5'-AUG-3', which also specifies methionine (so most newly synthesized polypeptides start with methionine), although other codons such as 5'-GUG-3' and 5'-UUG-3' are also used, especially in bacteria. The three termination codons are 5'-UAG-3', 5'-UAA-3', and 5'-UGA-3'.

**The genetic code is not universal**

It was originally thought that the genetic code must be the same in all organisms. The argument was that, once established, it would be impossible for the code to change because giving a new meaning to any single codon would result in widespread disruption of the amino acid sequences of proteins. This reasoning seems sound, so it is surprising that, in reality, the genetic code is not universal. The code shown in Figure 1.26 holds for the vast majority of genes in the vast majority of organisms, but deviations are widespread. In particular, mitochondrial genomes often use a nonstandard code (Table 1.3A). This was first discovered in 1979 by Frederick Sanger’s group in Cambridge, UK, who found that several human mitochondrial mRNAs contain the sequence UGA, which normally codes for termination, at internal positions where protein synthesis was not expected to stop. Comparisons with the amino acid sequences of the proteins coded by these mRNAs showed that 5'-UGA-3' is a tryptophan codon in human mitochondria and that this is just one of four code deviations in this particular genetic system. Mitochondrial genes in other organisms also display code deviations, although at least one of these—the use of 5'-CGG-3' as a tryptophan codon in plant mitochondria—is probably corrected by RNA editing before translation occurs.

**Figure 1.26** The genetic code. The codons are read in the 5' → 3' direction in an mRNA. Amino acids are designated by the standard three-letter abbreviations (see Table 1.2).

**Figure 1.27** Positions of the punctuation codons in an mRNA.
Nonstandard codes are also known for the nuclear genomes of lower eukaryotes. Often a modification is restricted to just a small group of organisms, and frequently it involves reassignment of the termination codons (Table 1.3B). Modifications are less common among prokaryotes, but examples are known in *Mycoplasma* and *Micrococcus*. A more important type of code variation is **context-dependent codon reassignment**, which occurs when the protein to be synthesized contains either selenocysteine or pyrrolysine. Proteins containing pyrrolysine are rare and are probably only present in the group of prokaryotes called the *archaea* (Chapter 8), but selenoproteins are widespread in many organisms, one example being the enzyme glutathione peroxidase, which helps protect the cells of humans and other mammals against oxidative damage. Selenocysteine is coded by 5′-UGA-3′ and pyrrolysine by 5′-UAG-3′. These codons therefore have a dual meaning because they are still used as termination codons in the organisms concerned (Table 1.3C). A 5′-UGA-3′ codon that specifies selenocysteine is distinguished from true termination codons by the presence of a **stem–loop** structure in the mRNA, where the loop is formed by bending the mRNA back on itself, with a short stretch of base pairing making up the stem that holds the conformation together (Figure 1.28). The stem–loop is positioned just **downstream** of the selenocysteine codon in prokaryotes, and in the 3′-**untranslated region** (the part of the mRNA after the termination codon) in eukaryotes. Recognition of the selenocysteine codon requires interaction between the stem–loop structure and a special protein that is involved in translation of these mRNAs. A similar system probably operates for recognition of a pyrrolysine codon.

<table>
<thead>
<tr>
<th>Organism</th>
<th>Codon</th>
<th>Should code for</th>
<th>Actually codes for</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>(A) Mitochondrial genomes</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mammals</td>
<td>UGA</td>
<td>Stop</td>
<td>Trp</td>
</tr>
<tr>
<td></td>
<td>AGA, AGG</td>
<td>Arg</td>
<td>Stop</td>
</tr>
<tr>
<td></td>
<td>AUA</td>
<td>Ile</td>
<td>Met</td>
</tr>
<tr>
<td><em>Drosophila</em></td>
<td>UGA</td>
<td>Stop</td>
<td>Trp</td>
</tr>
<tr>
<td></td>
<td>AGA</td>
<td>Arg</td>
<td>Ser</td>
</tr>
<tr>
<td></td>
<td>AUA</td>
<td>Ile</td>
<td>Met</td>
</tr>
<tr>
<td><em>Saccharomyces cerevisiae</em></td>
<td>UGA</td>
<td>Stop</td>
<td>Trp</td>
</tr>
<tr>
<td></td>
<td>CUN</td>
<td>Leu</td>
<td>Thr</td>
</tr>
<tr>
<td></td>
<td>AUA</td>
<td>Ile</td>
<td>Met</td>
</tr>
<tr>
<td>Fungi</td>
<td>UGA</td>
<td>Stop</td>
<td>Trp</td>
</tr>
<tr>
<td>Maize</td>
<td>CGG</td>
<td>Arg</td>
<td>Trp</td>
</tr>
<tr>
<td><strong>(B) Nuclear and prokaryotic genomes</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Several protozoa</td>
<td>UAA, UAG</td>
<td>Stop</td>
<td>Gln</td>
</tr>
<tr>
<td><em>Candida cylindracea</em></td>
<td>CUG</td>
<td>Leu</td>
<td>Ser</td>
</tr>
<tr>
<td><em>Micrococcus sp.</em></td>
<td>AGA</td>
<td>Arg</td>
<td>Stop</td>
</tr>
<tr>
<td></td>
<td>AUA</td>
<td>Ile</td>
<td>Stop</td>
</tr>
<tr>
<td><em>Euplotes sp.</em></td>
<td>UGA</td>
<td>Stop</td>
<td>Cys</td>
</tr>
<tr>
<td><em>Mycoplasma sp.</em></td>
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<td>Stop</td>
<td>Trp</td>
</tr>
<tr>
<td></td>
<td>CGG</td>
<td>Arg</td>
<td>Stop</td>
</tr>
<tr>
<td><strong>(C) Context-dependent codon reassignments</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>UGA</td>
<td>Stop</td>
<td>Selenocysteine</td>
</tr>
<tr>
<td>Archaea</td>
<td>UAG</td>
<td>Stop</td>
<td>Pyrrolysine</td>
</tr>
</tbody>
</table>

Abbreviation: N, any nucleotide.
The link between the proteome and the biochemistry of the cell

The biological information encoded by the genome finds its final expression in a protein whose biological properties are determined by its folded structure and by the spatial arrangement of chemical groups on its surface. By specifying proteins of different types, the genome is able to construct and maintain a proteome whose overall biological properties form the underlying basis of life. The proteome can play this role because of the huge diversity of protein structures that can be formed, which enables proteins to carry out a variety of biological functions. These functions include the following:

- **Biochemical catalysis** is the role of the special type of proteins called enzymes. The central metabolic pathways, which provide the cell with energy, are catalyzed by enzymes, as are the biosynthetic processes that result in the construction of nucleic acids, proteins, carbohydrates, and lipids. Biochemical catalysis also drives genome expression through the activities of enzymes such as RNA polymerase.

- **Structure**, which at the cellular level is determined by the proteins that make up the cytoskeleton, is also the primary function of some extracellular proteins. An example is collagen, which is an important component of bones and tendons.

- **Movement** is conferred by contractile proteins, of which actin and myosin in cytoskeletal fibers are the best-known examples.

- **Transport** of materials around the body is an important protein activity. For example, hemoglobin transports oxygen in the bloodstream, and serum albumin transports fatty acids.

- **Regulation** of cellular processes is mediated by proteins such as **transcription factors** that bind to the genome and influence the expression levels of individual genes and groups of genes (Section 12.2). The activities of groups of cells are regulated and coordinated by extracellular hormones and cytokines, many of which are proteins (e.g., insulin, the hormone that controls blood sugar levels, and interleukins, a group of cytokines that regulate cell division and differentiation).

- **Protection** of the body and of individual cells is the function of a range of proteins, including antibodies and those proteins involved in the blood-clotting response.

- **Storage** functions are performed by proteins such as ferritin, which acts as an iron store in the liver, and gliadins, which store amino acids in dormant wheat seeds.

This multiplicity of protein function provides the proteome with its ability to convert the blueprint contained in the genome into the essential features of life.
Chapter 1: Genomes, Transcriptomes, and Proteomes

SUMMARY

- The genome is the store of biological information possessed by every organism on the planet.
- The vast majority of genomes are made of DNA, the few exceptions being those viruses that have RNA genomes.
- Genome expression is the process by which the information contained in the genome is released to the cell.
- The first product of genome expression is the transcriptome, the collection of RNAs derived from those genes that are active at a particular time.
- The second product is the proteome, the cell’s repertoire of proteins that specify the nature of the biochemical reactions that the cell is able to carry out.
- Experimental evidence showing that genes are made of DNA was first obtained in 1945–1952, but it was the discovery of the double-helix structure by Watson and Crick in 1953 that convinced biologists that DNA is indeed the genetic material.
- A DNA polynucleotide is an unbranched polymer made up of multiple copies of four chemically different nucleotides.
- In the double helix, two polynucleotides are wound around one another, with the nucleotide bases on the inside of the molecule.
- The polynucleotides are linked by hydrogen bonding between the bases, with A always base-paired to T and G always base-paired to C.
- RNA is also a polynucleotide but the individual nucleotides have different structures compared with those found in DNA, and RNA is usually single-stranded.
- A cell contains various types of RNA, including mRNAs, which are transcripts of protein-coding genes, and several types of noncoding RNA.
- Many RNAs are initially synthesized as precursor molecules, which are processed by cutting and joining reactions and by chemical modification to give the mature forms.
- Proteins are also unbranched polymers, but in proteins the units are amino acids linked by peptide bonds.
- The amino acid sequence is the primary structure of a protein, the higher levels of structure—secondary, tertiary, and quaternary—being formed by folding of the primary structure into three-dimensional conformations and by association of individual polypeptides into multiprotein structures.
- Proteins are functionally diverse because individual amino acids have different chemical properties that, when combined in different ways, result in proteins with a range of chemical features.
- Proteins are synthesized by translation of mRNAs, with the rules of the genetic code specifying which triplet of nucleotides codes for which amino acid.
- The genetic code is not universal: variations occur in mitochondria and in lower eukaryotes, and some codons can have two different meanings in a single gene.
SHORT ANSWER QUESTIONS

1. Provide a timeline for the discovery of DNA, the discovery that DNA is the genetic material, the discovery of the structure of DNA, and the characterization of the first genome.

2. Which two types of chemical interaction stabilize the double helix?

3. Why does the specific base pairing between A and T, and G and C, provide a basis for the fidelity of DNA replication?

4. What are the two important chemical differences between RNA and DNA?

5. Why is noncoding RNA also called functional RNA?

6. Outline the various ways in which RNA molecules are processed.

7. Do cells ever lack a transcriptome? Explain the significance of your answer.

8. How do hydrogen bonds, electrostatic interactions, and hydrophobic forces play important roles in the secondary, tertiary, and quaternary structures of proteins?

9. How can proteins have so many diverse structures and functions when they are all synthesized from just 20 amino acids?

10. In addition to the 20 amino acids, proteins have additional chemical diversity because of two factors. What are these two factors, and what is their importance?

11. How can the codon 5′–UGA–3′ function as both a stop codon and as a codon for the modified amino acid selenocysteine?

12. How does the genome direct the biological activity of a cell?

IN-DEPTH PROBLEMS

1. The text (page 6) states that Watson and Crick discovered the double-helix structure of DNA on Saturday, March 7, 1953. Justify this statement.

2. Discuss why the double helix gained immediate universal acceptance as the correct structure for DNA.

3. What experiments led to elucidation of the genetic code in the 1960s?

4. Discuss the reasons why polypeptides can take up a large variety of structures whereas polynucleotides cannot.

5. The transcriptome and proteome are looked on as, respectively, an intermediate and the end-product of genome expression. Evaluate the strengths and limitations of these terms for our understanding of genome expression.
FURTHER READING

Books and articles on the discovery of the double helix and other important landmarks in the study of DNA


Research papers and reviews describing important aspects of DNA, RNA, or proteins


Virtually everything we know about genomes and genome expression has been discovered by scientific research: theoretical studies have played very little role in this or any other area of molecular and cell biology. It is possible to learn facts about genomes without knowing very much about how those facts were obtained, but in order to gain a real understanding of the subject we must examine in detail the techniques and scientific approaches that have been used to study genomes. The next five chapters cover these research methods. First, in this chapter, we examine the techniques, centered on the polymerase chain reaction and DNA cloning, that are used to study DNA molecules. These techniques are very effective with short segments of DNA, including individual genes, enabling a wealth of information to be obtained at this level. Chapter 3 then covers the methods that are used to construct maps of genomes, and Chapter 4 describes the methods used to sequence DNA molecules and to assemble the short sequences generated by these methods into the immensely long sequences that make up individual chromosomes and entire genomes. Finally, in Chapters 5 and 6, we will look at the various approaches that are used to locate the positions of genes in a genome sequence and to identify the functions of those genes. As you read through these chapters, you will begin to appreciate that understanding the structure and function of an individual genome is a major undertaking and that research is currently in the middle of an exciting discovery phase, with new techniques and new approaches revealing novel and unexpected aspects of genomes almost every week.

The toolkit of techniques used by molecular biologists to study DNA molecules was assembled during the 1970s and 1980s. Before then, the only way in which individual genes could be studied was by classical genetics, using techniques that originated with Mendel in the middle part of the nineteenth century. The development of more direct methods for studying DNA was stimulated by breakthroughs in biochemical research that, in the early 1970s, provided molecular biologists with enzymes that could be used to manipulate DNA molecules in the test tube. These enzymes occur naturally in living cells and are involved in processes such as DNA replication, repair, and recombination, which we will discuss in Chapters 15-17. In order to determine the functions of these enzymes, many of them were purified and the reactions that they catalyze were studied. Molecular biologists then adopted the pure enzymes as tools for manipulating DNA molecules in predetermined ways, using them to make copies of DNA molecules, to cut DNA molecules into shorter fragments, and to join them together again in combinations that do not exist in nature (Figure 2.1). These manipulations form the basis of

Figure 2.1 Examples of manipulations that can be carried out with DNA molecules.
recombinant DNA technology, in which new or recombinant DNA molecules are constructed from pieces of naturally occurring chromosomes and plasmids.

Recombinant DNA methodology led to development of the polymerase chain reaction (PCR). PCR is a deceptively simple technique—all that it achieves is the repeated copying of a short segment of a DNA molecule (Figure 2.2)—but it has become immensely important in many areas of biological research, not least the study of genomes. PCR is covered in detail in Section 2.2. Recombinant DNA techniques also underlie DNA cloning, or gene cloning, in which a DNA fragment is inserted into a plasmid or virus chromosome and then replicated in a bacterial or eukaryotic host (Figure 2.3). We will examine exactly how gene cloning is performed, and the reasons why this technique is important in genome research, in Section 2.3.

2.1 ENZYMES FOR DNA MANIPULATION

Recombinant DNA technology was one of the main factors that contributed to the rapid advance in knowledge concerning gene expression that occurred during the 1970s and 1980s. The basis of recombinant DNA technology is the ability to manipulate DNA molecules in the test tube. This, in turn, depends on the availability of purified enzymes whose activities are known and can be controlled, and which can therefore be used to make specified changes to the DNA molecules that are being manipulated. The enzymes available to the molecular biologist fall into four broad categories:

- **DNA polymerases**, which are enzymes that synthesize new polynucleotides complementary to an existing DNA or RNA template (Figure 2.4A)

- **Nucleases**, which degrade DNA molecules by breaking the phosphodiester bonds that link one nucleotide to the next (Figure 2.4B)

- **Ligases**, which join DNA molecules together by synthesizing phosphodiester bonds between nucleotides at the ends of two different molecules or at the two ends of a single molecule (Figure 2.4C)

- **End-modification enzymes**, which make changes to the ends of DNA molecules (Figure 2.4D)

We begin our study of recombinant DNA techniques by examining how each of these types of enzyme is used to make specified changes to DNA molecules.

**The mode of action of a template-dependent DNA polymerase**

Many of the techniques used to study DNA depend on the synthesis of DNA copies of all or part of existing DNA or RNA molecules. This is an essential requirement for PCR (Section 2.2), DNA sequencing (Sections 4.1 and 4.2), and many other procedures that are central to molecular biology research. An enzyme that synthesizes DNA is called a DNA polymerase, and one that copies an existing DNA or RNA molecule is called a template-dependent DNA polymerase. A template-dependent DNA polymerase makes a new DNA polynucleotide whose sequence is dictated, via the base-pairing rules, by the sequence of nucleotides in
the DNA or RNA molecule that is being copied (Figure 2.5). The new polynucleotide is always synthesized in the 5′ → 3′ direction: DNA polymerases that make DNA in the other direction are unknown in nature.

An important feature of template-dependent DNA synthesis is that a DNA polymerase is unable to use an entirely single-stranded molecule as the template. In order to initiate DNA synthesis, there must be a short, double-stranded region to act as a primer for DNA polymerase action. This is accomplished by an RNA primer synthesized by primase, which is then removed by RNase H.

Figure 2.5 Activity of a DNA-dependent DNA polymerase. New nucleotides are added onto the 3′-end of the growing polynucleotide, with the sequence of this new polynucleotide being determined by the sequence of the template DNA. Compare with the process of transcription (DNA-dependent RNA synthesis) shown in Figure 1.13.
Provide a 3’-end onto which the enzyme will add new nucleotides (Figure 2.6A). The way in which this requirement is met in living cells when the genome is replicated is described in Chapter 15. In the test tube, a DNA-copying reaction is initiated by attaching to the template a short, synthetic oligonucleotide, usually about 20 nucleotides in length, which acts as a primer for DNA synthesis. At first glance, the need for a primer might appear to be an undesired complication in the use of DNA polymerases in recombinant DNA technology, but nothing could be further from the truth. Because annealing of the primer to the template depends on complementary base pairing, the position within the template molecule at which DNA copying is initiated can be specified by synthesizing a primer with the appropriate nucleotide sequence (Figure 2.6B). A short, specific segment of a much longer template molecule can therefore be copied, which is much more valuable than the random copying that would occur if DNA synthesis did not need to be primed. You will fully appreciate the importance of priming when we deal with PCR in Section 2.2.

A second general feature of template-dependent DNA polymerases is that many of these enzymes are multifunctional and are able to degrade DNA molecules as well as synthesize them. This is a reflection of the way in which DNA polymerases act in the cell during genome replication (Section 15.3). As well as their 5’ → 3’ DNA synthesis capability, DNA polymerases can also have one or both of the following exonuclease activities (Figure 2.7):

- A 3’ → 5’ exonuclease activity enables the enzyme to remove nucleotides from the 3’-end of the strand that it has just synthesized. This is called the proofreading activity because it allows the polymerase to correct errors by removing a nucleotide that has been inserted incorrectly.

- A 5’ → 3’ exonuclease activity is less common but is possessed by some DNA polymerases whose natural function in genome replication requires that they must be able to remove at least part of a polynucleotide that is already attached to the template strand that the polymerase is copying.

The types of DNA polymerase used in research

Several of the template-dependent DNA polymerases that are used in molecular biology research (Table 2.1) are versions of the Escherichia coli DNA polymerase I enzyme, which plays a central role in replication of this bacterium’s genome (Section 15.3). This enzyme, sometimes called the Kornberg polymerase, after its discoverer Arthur Kornberg, has both the 3’ → 5’ and 5’ → 3’ exonuclease activities, which limits its usefulness in DNA manipulation. Its main application is in the synthesis of DNA molecules containing radioactive or fluorescent nucleotides: the process called DNA labeling.
2.1 ENZYMES FOR DNA MANIPULATION

### TABLE 2.1 FEATURES OF TEMPLATE-DEPENDENT DNA POLYMERASES USED IN MOLECULAR BIOLOGY RESEARCH

<table>
<thead>
<tr>
<th>Polymerase</th>
<th>Description</th>
<th>Main uses</th>
<th>Cross reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA polymerase I</td>
<td>Unmodified <em>E. coli</em> enzyme</td>
<td>DNA labeling</td>
<td>Section 2.1</td>
</tr>
<tr>
<td>Klenow polymerase</td>
<td>Modified version of <em>E. coli</em> DNA polymerase I</td>
<td>DNA labeling, chain termination DNA sequencing</td>
<td>Sections 2.1 and 4.1</td>
</tr>
<tr>
<td><em>Taq</em> polymerase</td>
<td><em>Thermus aquaticus</em> DNA polymerase I</td>
<td>PCR</td>
<td>Section 2.2</td>
</tr>
<tr>
<td>Reverse transcriptase</td>
<td>RNA-dependent DNA polymerase, obtained from various retroviruses</td>
<td>cDNA synthesis</td>
<td>Sections 3.6 and 5.3</td>
</tr>
</tbody>
</table>

Of the two exonuclease activities, the 5’ → 3’ version causes most problems when a DNA polymerase is used to manipulate molecules in the test tube. This is because an enzyme with this activity is able to remove nucleotides from the 5’-ends of polynucleotides that have just been synthesized (Figure 2.8). It is unlikely that the polynucleotides will be completely degraded, because the polymerase function is usually much more active than the exonuclease function, but some techniques will not work if the 5’-ends of the new polynucleotides are shortened in any way. In particular, some of the older DNA sequencing methods are based on synthesis of new polynucleotides, all of which share exactly the same 5’-end, marked by the primer used to initiate the sequencing reactions. If any nibbling of the 5’-ends occurs, then it is impossible to determine the correct DNA sequence. Because of this problem, when DNA sequencing was first introduced in the late 1970s, a modified version of the Kornberg enzyme called the Klenow polymerase was used. The Klenow polymerase was initially prepared by cutting the natural *E. coli* DNA polymerase I enzyme into two segments by use of a protease. One of these segments retained the polymerase and 3’ → 5’ exonuclease activities but lacked the 5’ → 3’ exonuclease function of the untreated enzyme. Nowadays, the enzyme is almost always prepared from *E. coli* cells whose polymerase gene has been engineered so that the resulting enzyme has the desired properties.

The *E. coli* DNA polymerase I enzyme has an optimum reaction temperature of 37°C, which is the usual temperature of the natural environment of the bacterium, inside the lower intestines of mammals such as humans. Test-tube reactions with either the Kornberg or Klenow polymerases are therefore incubated at 37°C and terminated by raising the temperature to 75°C or above, which causes the protein to unfold, or denature, destroying its enzymatic activity. This regimen is perfectly adequate for most molecular biology techniques, but for reasons that will become clear in Section 2.2, PCR requires a thermostable DNA polymerase, one that is able to function at temperatures much higher than 37°C. Suitable enzymes can be obtained from bacteria such as *Thermus aquaticus*, which live in hot springs at temperatures up to 95°C, and whose DNA polymerase I enzyme has an optimum working temperature of 75–80°C. The biochemical basis of protein thermostability is not fully understood but probably centers on structural features that reduce the amount of protein unfolding that occurs at elevated temperatures.

One additional type of DNA polymerase is important in molecular biology research. This is reverse transcriptase, which is an RNA-dependent DNA polymerase and so makes DNA copies of RNA rather than DNA templates. Reverse transcriptases are involved in the replication cycles of retroviruses (Section 9.1), including the human immunodeficiency viruses, which have RNA genomes that are copied into DNA after infection of the host. In the test tube, a reverse transcriptase can be used to make DNA copies of mRNA molecules. These copies are called complementary DNAs (cDNAs). Their synthesis is important in some types of gene cloning and in techniques used to map the regions of a genome that specify particular mRNAs (Section 5.3).
Restriction endonucleases enable DNA molecules to be cut at defined positions

A variety of nucleases have found applications in recombinant DNA technology (Table 2.2). Some nucleases have a broad range of activities but most are either exonucleases, removing nucleotides from the ends of DNA and/or RNA molecules, or endonucleases, making cuts at internal phosphodiester bonds. Some nucleases are specific for DNA and some for RNA, some work only on double-stranded DNA and others only on single-stranded DNA, and some are not fussy what they work on. We will encounter various examples of nucleases in later chapters when we deal with the techniques in which they are used. Only one type of nuclease will be considered in detail here: the restriction endonucleases, which play a central role in all aspects of recombinant DNA technology.

A restriction endonuclease is an enzyme that binds to a DNA molecule at a specific sequence and makes a double-stranded cut at or near that sequence. Because of the sequence specificity, the positions of cuts within a DNA molecule can be predicted if the DNA sequence is known, enabling defined segments to be excised from a larger molecule. This ability underlies gene cloning and all other aspects of recombinant DNA technology in which DNA fragments of known sequence are required.

There are three main types of restriction endonuclease. With types I and III, there is no strict control over the position of the cut relative to the specific sequence in the DNA molecule that is recognized by the enzyme. These enzymes are therefore less useful because the sequences of the resulting fragments are not precisely known. Type II enzymes do not suffer from this disadvantage because the cut is always at the same place, either within the recognition sequence or very close to it (Figure 2.9). For example, the type II enzyme called EcoRI (isolated from Escherichia coli) cuts DNA only at the hexanucleotide 5’-GAATTC-3’. Digestion of DNA with a type II enzyme therefore gives a reproducible set of fragments whose sequences are predictable if the sequence of the target DNA molecule is known. Almost 4000 type II enzymes have been isolated, and more than 600 are available for use in the laboratory. Many enzymes have hexanucleotide target sites, but others recognize shorter or longer sequences (Table 2.3). There are also examples of enzymes with degenerate recognition sequences, meaning that they cut DNA at any of a family of related sites. HinfI (from Haemophilus influenzae), for example, recognizes 5’-GANTC-3’, where N is any nucleotide, and so it cuts at 5’-GAATC-3’, 5’-GATTC-3’, 5’-GAGTC-3’, and 5’-GACTC-3’. Most enzymes cut within the recognition sequence, but a few, such as BsrBI, cut at a specified position outside of this sequence.

Restriction enzymes cut DNA in two different ways. Many make a simple double-stranded cut, giving a blunt or flush end, but others cut the two DNA strands at different positions, usually two or four nucleotides apart, so that the resulting DNA fragments have short, single-stranded overhangs at each end. These are called sticky or cohesive ends because base pairing between them can stick the DNA molecule back together again (Figure 2.10A). Some sticky-end cutters
Studying DNA

Recognition sequence

Type of ends

End sequences

Table 2.3 Some Examples of Restriction Endonucleases

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Recognition sequence</th>
<th>Type of ends</th>
<th>End sequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alu</td>
<td>5'-AGCT-3' 3'-TCGA-5'</td>
<td>Blunt</td>
<td>5'-AG CT-3' 3'-TC GA-5'</td>
</tr>
<tr>
<td>Sau3AI</td>
<td>5'-GATC-3' 3'-CTAG-5'</td>
<td>Sticky, 5'-overhang</td>
<td>5'-GATC-3' 3'-CTAG G-5'</td>
</tr>
<tr>
<td>HinfI</td>
<td>5'-GANTC-3' 3'-CTNAG-5'</td>
<td>Sticky, 5'-overhang</td>
<td>5'-G ANTC-3' 3'-CTNA G-5'</td>
</tr>
<tr>
<td>BamHI</td>
<td>5'-GGATCC-3' 3'-CCTAGG-5'</td>
<td>Sticky, 5'-overhang</td>
<td>5'-GGATCC-3' 3'-CCTAG G-5'</td>
</tr>
<tr>
<td>BsrBI</td>
<td>5'-CCGCTC-3' 3'-GCGAGG-5'</td>
<td>Sticky, 5'-overhang</td>
<td>5'-CCGCTC-3' 3'-GCGAG G-5'</td>
</tr>
<tr>
<td>EcoRI</td>
<td>5'-GAATTC-3' 3'-CTTAAG-5'</td>
<td>Sticky, 5'-overhang</td>
<td>5'-GAATTC-3' 3'-CTTAAG G-5'</td>
</tr>
<tr>
<td>PstI</td>
<td>5'-CTGCA-3' 3'-GACGTC-5'</td>
<td>Sticky, 3'-overhang</td>
<td>5'-CTGCA-3' 3'-GACGTC G-5'</td>
</tr>
<tr>
<td>NotI</td>
<td>5'-GGCGGCCG-3' 3'-CCCGGCGG-5'</td>
<td>Sticky, 3'-overhang</td>
<td>5'-GGCGGCCG-3' 3'-CCCGGCGG G-5'</td>
</tr>
<tr>
<td>BglII</td>
<td>5'-GCNNNNNNGC-3' 3'-CGGNINNCCG-5'</td>
<td>Sticky, 3'-overhang</td>
<td>5'-GCNNNN NNGC-3' 3'-CGGN NNNCCG G-5'</td>
</tr>
</tbody>
</table>

Abbreviation: N, any nucleotide.

Note that most, but not all, recognition sequences have inverted symmetry: when read in the 5'→3' direction, the sequence is the same on both strands.

Figure 2.10 Results of digestion of DNA with different restriction endonucleases. (A) Blunt ends and sticky ends. (B) Different types of sticky ends: the 5'-overhangs produced by BamHI and the 3'-overhangs produced by PstI. (C) The same sticky ends produced by two different restriction endonucleases: a 5'-overhang, 5'-GATC-3', is produced by both BamHI (which recognizes the sequence 5'-GGATCC-3') and Sau3AI (which recognizes the sequence 5'-GATC-3').
give 5'-overhangs (for example, *Sau*3AI, *Hinf*I), whereas others leave 3'-overhangs (for example, *Pst*I) (Figure 2.10B). One feature that is particularly important in recombinant DNA technology is that some pairs of restriction enzymes have different recognition sequences but give the same sticky ends: for example, *Sau*3AI and *Bam*HI both give a 5'-GATC-3' sticky end, even though *Sau*3AI has a four-base-pair recognition sequence and *Bam*HI recognizes a six-base-pair sequence (Figure 2.10C).

**Gel electrophoresis is used to examine the results of a restriction digest**

Treatment with a restriction endonuclease results in a larger DNA molecule being cut into smaller fragments. How do we measure the sizes of these fragments? The answer is by **gel electrophoresis**. This is the standard method for separating DNA molecules of different lengths. It has many applications in size analysis of DNA fragments and can also be used to separate RNA molecules.

**Electrophoresis** is the movement of charged molecules in an electric field: negatively charged molecules migrate toward the positive electrode, and positively charged molecules migrate toward the negative electrode. The technique was originally carried out in aqueous solution, in which the predominant factors influencing migration rate are the shape of a molecule and its electric charge. This is not particularly useful for DNA separations because most DNA molecules are the same shape (linear), and although the charge of a DNA molecule is dependent on its length, the differences in charge are not sufficient to result in effective separation (Figure 2.11A). The situation is different when electrophoresis is carried out in a gel, because now shape and charge are less important and molecular length is the critical determinant of migration rate. This is because the gel is a network of pores through which the DNA molecules have to travel to reach the positive electrode. Shorter molecules are less impeded by the pores than are longer molecules and so move through the gel more quickly. Molecules of different lengths therefore form bands in the gel (Figure 2.11B).

Two types of gel are used in molecular biology: **agarose gels**, as described here, and **polyacrylamide gels**, which are mainly used in DNA sequencing (Section 4.1). Agarose is a polysaccharide that forms gels with pores ranging from 100 to 300 nm in diameter; the pore size depends on the concentration of agarose in the gel. Gel concentration therefore determines the range of DNA fragments that can be separated. The separation range is also affected by the **electroendosmosis** (EEO) value of the agarose, which is a measure of the amount of bound sulfate and pyruvate anions. The greater the EEO, the slower the migration rate for a negatively charged molecule such as DNA.

An agarose gel is prepared by mixing the appropriate amount of agarose powder in a buffer solution, heating the mixture to dissolve the agarose, and then...
pouring the molten gel onto a Perspex plate with tape around the sides to prevent spillage. A comb is placed in the gel to form wells for the samples. The gel is allowed to set, and electrophoresis then carried out with the gel submerged under buffer. In order to follow the progress of the electrophoresis, one or two dyes of known migration rates are added to the DNA samples before loading. The bands of DNA can be visualized by soaking the gel in ethidium bromide solution. This compound intercalates between DNA base pairs and fluoresces when activated with ultraviolet radiation (Figure 2.12). Unfortunately, the procedure is hazardous because ethidium bromide is a powerful mutagen. Nonmutagenic dyes that stain DNA green, red, or blue are therefore now used in many laboratories. The most sensitive dyes are able to detect bands that contain less than 1 ng of DNA, compared to a minimum of 10 ng of DNA when ethidium bromide is used.

Depending on the concentration of agarose in the gel, fragments between 100 bp and 50 kilobase pairs (kb) in length can be separated into sharp bands after electrophoresis (Figure 2.13). For example, a 0.5 cm thick slab of 0.5% agarose, which has relatively large pores, would be used for molecules in the size range 1–30 kb, allowing, for example, molecules of 10 and 12 kb to be clearly distinguished. Alternatively, a 0.3% gel can be used for longer molecules up to 50 kb, and a 5% gel can be used for shorter molecules 100–500 bp in length.

**Interesting DNA fragments can be identified by Southern hybridization**

If the DNA that is cut with a restriction endonuclease is a relatively short molecule, and twenty or fewer fragments are produced after restriction, then usually it is possible to select an agarose concentration that results in each fragment being visible as a separate band in the gel. If the starting DNA is long and gives rise to many fragments after digestion, then regardless of the agarose concentration used, the gel may simply show a smear of DNA, because there are fragments of every possible length that all merge together. This is the usual result when genomic DNA is cut with a restriction enzyme.

If the sequence of the starting DNA is known, then the sequences (and hence the sizes) of the fragments resulting from treatment with a particular restriction enzyme can be predicted. The band for a desired fragment (for example, one containing a gene) can then be identified and cut out of the gel, and the DNA can be purified. Even if its size is unknown, a fragment containing a gene or another segment of DNA of interest can be identified by a technique called **Southern hybridization**. The only requirement is that at least some of the sequence of the desired gene or DNA segment is known or can be predicted. The first step is to transfer the restriction fragments from the agarose gel to a nitrocellulose or nylon
membrane. This is done by placing the membrane on the gel and allowing buffer to soak through, taking the DNA from the gel to the membrane, where it becomes bound (Figure 2.14A). This process results in the DNA bands becoming immobilized in the same relative positions on the surface of the membrane.

The next step is to prepare a hybridization probe, which is a labeled DNA molecule whose sequence is complementary to the target DNA that we wish to detect. The label is often a radioactive marker. Nucleotides can be synthesized in which one of the phosphorus atoms is replaced with $^{32}\text{P}$ or $^{33}\text{P}$, one of the oxygen atoms in the phosphate group is replaced with $^{35}\text{S}$, or one or more of the hydrogen atoms is replaced with $^{3}\text{H}$. Radioactive nucleotides still act as substrates for DNA polymerases and so are incorporated into a DNA molecule by any strand-synthesis reaction catalyzed by a DNA polymerase. Alternatively, to avoid the health hazards and disposal issues associated with the use of radioactive chemicals, nucleotides carrying a fluorescent marker or one that emits chemiluminescence can be used.

The probe could be a synthetic oligonucleotide whose sequence matches part of an interesting gene. Because the probe and target DNAs are complementary, they will base-pair or hybridize, and the position of the hybridized probe on the membrane can then be identified by detecting the signal given out by the label attached to the probe. To carry out the hybridization, the membrane is placed in a glass bottle with the labeled probe and some buffer, and the bottle is gently rotated for several hours so that the probe has plenty of opportunity to hybridize to its target DNA. The membrane is then washed to remove any probe that has not become hybridized, and the signal from the label is detected. In the example shown in Figure 2.14B, the probe is radioactively labeled and the signal is detected by exposure of an X-ray-sensitive film (autoradiography). The band seen on the autoradiograph is the one that corresponds to the restriction fragment that hybridizes to the probe and therefore contains the gene that we are searching for. If a fluorescent marker is used, then the label is detected with a film sensitive to the emission spectrum of the fluorophore. Chemiluminescent markers can be detected in the same way, but these have the disadvantage that the signal is not generated directly by the label and instead must be developed by treatment of the labeled molecule with chemicals. A popular method involves labeling the DNA with the enzyme alkaline phosphatase, which is detected by applying dioxygen, which the enzyme dephosphorylates to produce the chemiluminescence. All three types of label—radioactive, fluorescent, and

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**Figure 2.14** Southern hybridization. (A) Transfer of DNA from the gel to the membrane. (B) The membrane is probed with a radioactively labeled DNA molecule. On the resulting autoradiograph, one hybridizing band is seen in lane 2 and two in lane 3.
Ligases join DNA fragments together

DNA fragments that have been generated by treatment with a restriction endonuclease can be joined back together again or attached to a new partner by a DNA ligase. The reaction requires energy, which is provided by adding either ATP or nicotinamide adenine dinucleotide (NAD) to the reaction mixture, depending on the type of ligase used.

The most widely used DNA ligase is obtained from *E. coli* cells infected with T4 bacteriophage. This enzyme is involved in replication of the phage DNA and is encoded by the T4 genome. The natural role of the enzyme is to synthesize phosphodiester bonds between unlinked nucleotides present in one polynucleotide of a double-stranded molecule (Figure 2.15A). In order to join together two restriction fragments, the ligase has to synthesize two phosphodiester bonds, one in each strand (Figure 2.15B). This is by no means beyond the capabilities of the enzyme, but the reaction can occur only if the ends to be joined come close enough to one another by chance: the ligase is not able to catch hold of them and bring them together. If the two molecules have complementary sticky ends, and the ends come together by random diffusion events in the ligation mixture, then transient base pairs might form between the two overhangs. These base pairs are not particularly stable, but they may persist for sufficient time for a ligase enzyme to attach to the junction and synthesize phosphodiester bonds to fuse the ends together (Figure 2.15C). If the molecules are blunt-ended, then they cannot base-pair to one another, not even temporarily, and ligation is a much less efficient process, even when the DNA concentration is high and ends are in relatively close proximity.

The greater efficiency of sticky-end ligation has stimulated the development of methods for converting blunt ends into sticky ends. In one method, short double-stranded molecules called linkers or adaptors are attached to the blunt ends. Linkers and adaptors work in slightly different ways, but both contain a recognition sequence for a restriction endonuclease and so produce a sticky end after treatment with the appropriate enzyme (Figure 2.16). Another way to create a sticky end is by homopolymer tailing, in which nucleotides are added one after the other to the 3’-terminus at a blunt end (Figure 2.17). The enzyme involved is called terminal deoxynucleotidyl transferase, which we will discuss in the next section. If the reaction mixture contains the DNA, enzyme, and only one of the four nucleotides, then the new stretch of single-stranded DNA that is made consists entirely of just that single nucleotide. It could, for example, be a poly(G) tail, which would enable the molecule to base-pair to other molecules that carry poly(C) tails, created in the same way but with dCTP, rather than dGTP, in the reaction mixture.
**End-modification enzymes**

Terminal deoxynucleotidyl transferase (see Figure 2.17), obtained from calf thymus tissue, is one example of an end-modification enzyme. It is, in fact, a **template-independent DNA polymerase**, because it is able to extend a DNA polynucleotide without base pairing of the incoming nucleotides to an existing strand of DNA or RNA. Its main role in recombinant DNA technology is in homopolymer tailing, as described above.

Two other end-modification enzymes are also frequently used. These are **alkaline phosphatase** and **T4 polynucleotide kinase**, which act in complementary ways. Alkaline phosphatase is obtained from various sources, including *E. coli*, calf thymus tissue, and Arctic shrimp. It removes phosphate groups from the 5'-ends of DNA molecules, which prevents these molecules from being ligated to one another. Two ends carrying 5'-phosphates can be ligated to one another, and an end lacking a phosphate group can be ligated to an end containing a phosphate, but a link cannot be formed between a pair of ends if neither carries a 5'-phosphate. Judicious use of alkaline phosphatase can therefore direct the action of a DNA ligase in a predetermined way so that only desired ligation products are obtained. T4 polynucleotide kinase, obtained from *E. coli* cells infected with T4 bacteriophage, performs the reverse reaction to alkaline phosphatase: it adds phosphates to 5'-ends. Like alkaline phosphatase, the enzyme is used during complicated ligation experiments, but its main application is in the **end-labeling** of DNA molecules.

**2.2 THE POLYMERASE CHAIN REACTION**

Although methods with similar outcomes were suggested as early as 1971, the invention of PCR is now credited to Kary Mullis, who describes how he experienced a eureka moment one evening in early 1983 as he drove along the Pacific Coast Highway in California. His brainwave was an exquisitely simple technique that results in repeated copying of a selected segment of a longer DNA molecule. The technique is so straightforward that it is sometimes difficult for students encountering it for the first time to appreciate why it has become so important in modern biology. We will first look at the technique itself and then explore some of its myriad applications.
Carrying out a PCR

PCR results in the repeated copying of a selected region of a DNA molecule (see Figure 2.2). The reaction is carried out by the purified, thermostable DNA polymerase of *T. aquaticus* (Section 2.1). The reason a thermostable enzyme is needed will become clear when we look in more detail at the events that occur during a PCR.

To carry out a PCR experiment, the target DNA is mixed with *Taq* DNA polymerase, a pair of oligonucleotide primers, and a supply of nucleotides. The amount of target DNA can be very small because PCR is extremely sensitive and will work with just a single starting molecule. The primers are needed to initiate the DNA synthesis reactions that will be carried out by the *Taq* polymerase (see Figure 2.6). They must attach to the target DNA at either side of the segment to be copied. This means that the sequences of these attachment sites must be known so that primers of the appropriate sequences can be synthesized.

The reaction is started by heating the mixture to 94°C. At this temperature the hydrogen bonds that hold together the two polynucleotides of the double helix are broken, so the target DNA becomes denatured into single-stranded molecules (Figure 2.18). The temperature is then reduced to 50–60°C, which results in some rejoining of the single strands of the target DNA but also allows the primers to attach to their annealing positions. DNA synthesis can now begin, so the temperature is raised to 72°C, the optimum for *Taq* polymerase. In this first stage of the PCR, a set of long products is synthesized from each strand of the target DNA. These polynucleotides have identical 5'-ends but random 3'-ends, the latter representing positions where DNA synthesis terminates by chance.

When the cycle of denaturation–annealing–synthesis is repeated, the long products act as templates for new DNA synthesis, giving rise, in the third cycle, to short products, the 5'- and 3'-ends of which are both set by the primer annealing positions (Figure 2.19). In subsequent cycles, the number of short products accumulates in an exponential manner.

**Figure 2.18** The first stage of PCR.

**Figure 2.19** Synthesis of short products during PCR. From the first-cycle products shown at the top of the diagram, the next cycle of denaturation–annealing–synthesis leads to four products, two of which are identical to the first-cycle products and two of which are made entirely of new DNA. During the third cycle, the latter give rise to short products that, in subsequent cycles, accumulate in an exponential fashion.
accumulates in an exponential fashion (doubling during each cycle) until one of the components of the reaction becomes depleted. This means that, after 30 cycles, there will be over 130 million short products derived from each starting molecule. In real terms, this equates to several micrograms of PCR product from a few nanograms or less of target DNA.

The rate of product formation can be followed during a PCR

Often a PCR is allowed to reach completion before the outcome is determined. After a preset number of cycles, usually 30–40, the reaction is halted and a sample is analyzed by agarose gel electrophoresis, which will reveal a single band if the PCR has worked as expected and has amplified a single segment of the target DNA (Figure 2.20). Alternatively, the sequence of the product can be determined, by techniques described in Section 4.1.

It is also possible to follow synthesis of the product as the PCR proceeds through its series of cycles. This is called real-time PCR, and it can be carried out in two different ways. In the simplest method, a dye that gives a fluorescent signal when it binds to double-stranded DNA is included in the PCR mixture. The gradual increase in the fluorescent signal given out by the mixture indicates the rate at which the product is being synthesized. The disadvantage of this approach is that it measures the total amount of double-stranded DNA in the PCR at any particular time, which may overestimate the actual amount of the product. This is because the primers sometimes anneal to themselves in various nonspecific ways, increasing the amount of double-stranded DNA that is present.

The second method for real-time PCR requires a short oligonucleotide called a reporter probe, which gives a fluorescent signal when it hybridizes to the PCR product. Because the probe hybridizes only to the PCR product, this method avoids the problems caused by primer–primer annealing. Several systems have been developed, one of which makes use of a pair of labels comprising a fluorescent dye and a compound that quenches the fluorescent signal when brought into close proximity with the dye. This quenching is brought about by a process called Förster resonance energy transfer (FRET). The dye is attached to one end of the reporter probe and the quenching compound is attached to the other end. Normally there is no fluorescence because the probe is designed in such a way that the two ends base-pair to one another, placing the quencher next to the dye and quenching the fluorescent signal (Figure 2.21). Hybridization between probe and PCR product disrupts this base pairing, moving the quencher away from the dye and enabling the fluorescent signal to be generated.
Both methods can be used as the basis for **quantitative PCR**, which enables the amount of target DNA present at the start of the PCR to be measured. The rate of product synthesis during the test PCR is compared with the progress of control PCRs with known amounts of starting DNA. The comparison is usually made by identifying the stage in the PCR at which the amount of fluorescent signal reaches a preset threshold (Figure 2.22). The more rapidly the threshold is reached, the greater the amount of target DNA in the starting mixture.

**PCR has many and diverse applications**

Why is PCR so important in modern research? First we will deal with its limitations. In order to synthesize primers that will anneal at the correct positions, the sequences of the boundary regions of the DNA to be amplified must be known. This means that PCR cannot be used to purify fragments of genes, or other parts of a genome, that have never been studied before. A second constraint is the length of DNA that can be copied. Regions of up to 5 kb can be amplified without too much difficulty, and longer amplifications—up to 40 kb—are possible by modifications of the standard technique. Fragments longer than about 40 kb are unattainable by PCR.

Now we will consider the strengths of PCR. Primary among these is the ease with which products representing a single segment of a genome can be obtained. PCR can therefore be used to screen human DNA samples for mutations associated with genetic diseases such as thalassemia and cystic fibrosis. It also forms the basis of **genetic profiling**, in which natural variations in genome sequences are typed in order to connect samples taken at crime scenes with suspects and to establish paternity in cases where parentage is disputed (Section 7.4).

A second important feature of PCR is its ability to work with minuscule amounts of starting DNA. This means that PCR can be used to obtain sequences from the trace amounts of DNA that are present in hairs, blood stains, and other forensic specimens and from bones and other remains preserved at archaeological sites. Our ability to use PCR to amplify DNA from preserved skeletons has led to genome sequences of extinct species such as Neanderthals (Section 4.4). In clinical diagnosis, PCR is able to detect the presence of viral DNA well before the virus has reached the levels needed to initiate a disease response. This is particularly important in the early identification of viral-induced cancers because it means that treatment programs can be initiated before the cancer becomes established.

The above are just a few of the applications of PCR. The technique is now a major component of the molecular biologist’s toolkit, and we will discover many more examples of its use as we progress through the chapters of this book.

### 2.3 DNA CLONING

DNA cloning was the first of the important new research tools developed during the early years of the recombinant DNA revolution. Cloning is a logical extension of the ability to manipulate DNA molecules with restriction endonucleases and ligases. First we will look at the reasons why DNA cloning is a central technique in genomics research, and then we will examine how the technique is carried out.

**Why is gene cloning important?**

Imagine that an animal gene has been obtained as a single restriction fragment after digestion of a larger molecule with the restriction enzyme **BamHI**, which leaves 5’-GATC-3’ sticky ends (Figure 2.23). Imagine also that a plasmid—a small circle of DNA capable of replicating inside a bacterium—has been purified from **E. coli** and treated with **BamHI**, which cuts the plasmid in a single position. The circular plasmid has therefore been converted into a linear molecule, again with 5’-GATC-3’ sticky ends. If we mix the two DNA molecules together and add DNA ligase, various recombinant ligation products will be obtained, one of which comprises the circularized plasmid with the animal gene inserted into the position...
originally taken by the BamHI restriction site. If the recombinant plasmid is now reintroduced into E. coli, and the inserted gene has not disrupted its replicative ability, then the plasmid plus inserted gene will be replicated and copies will be passed to the daughter bacteria after cell division. The plasmid therefore acts as a cloning vector, providing the replicative ability that enables the cloned gene to be propagated inside the host cell. More rounds of plasmid replication and cell division will result in a colony of recombinant E. coli bacteria, where each bacterium contains multiple copies of the animal gene. This series of events, as illustrated in Figure 2.23, constitutes the process called DNA cloning or gene cloning.

When DNA cloning was first invented in the early 1970s, it revolutionized molecular biology by making possible experiments that previously had been inconceivable. This is because cloning can provide a pure sample of an individual DNA fragment, separated from all the other fragments produced when one or more larger molecules are cut with a restriction enzyme. These larger molecules could be, for example, an entire genome. Each of the fragments resulting from treatment with the endonuclease becomes inserted into a different plasmid molecule to produce a family of recombinant plasmids (Figure 2.24). Usually only one recombinant molecule is transported into any single host cell, so that although the final set of clones may contain many different recombinant molecules, each individual clone contains multiple copies of just one. The end result is a clone library, whose inserted DNA fragments come from different parts of the starting DNA. If enough clones are obtained, then it is possible to have every part of a genome represented in the library.
Clone libraries are important for two reasons. First, it is often possible to identify from within the library the clone or clones that contain the DNA from a single gene, so that gene can be isolated and studied in detail. Second, a clone library is often the starting point for a genome sequencing project, because by sequencing the individual fragments contained in different clones, the genome sequence can gradually be built up (Section 4.3).

The simplest cloning vectors are based on E. coli plasmids

Plasmids replicate efficiently in their bacterial hosts because each plasmid possesses an origin of replication that is recognized by the DNA polymerases and other proteins that normally replicate the bacterium’s chromosome. The host cell’s replicative machinery will also propagate a plasmid cloning vector, plus any new genes that have been inserted into it, providing that the vector possesses an origin of replication. Cloning vectors based on bacterial plasmids are therefore simple to construct and relatively easy to use.

One of the most popular plasmid vectors is pUC8, a member of a series of vectors first introduced in the early 1980s. The pUC series is derived from an earlier cloning vector, pBR322, which was originally constructed by ligating together restriction fragments from three naturally occurring E. coli plasmids: R1, R6.5, and pMB1. pUC8 is a small plasmid, comprising just 2.7 kb. As well as its origin of replication, it carries two genes (Figure 2.25):

- A gene for ampicillin resistance. The presence of this gene means that a bacterium containing a pUC8 plasmid is able to synthesize an enzyme, called β-lactamase, that enables the cell to withstand the growth-inhibitory effect of the antibiotic ampicillin. This means that cells containing pUC8 plasmids can be distinguished from those lacking these plasmids by plating the bacteria onto agar medium containing ampicillin. Normal E. coli cells are sensitive to ampicillin and cannot grow when the antibiotic is present. Ampicillin resistance is therefore a selectable marker for pUC8.

- The lacZ gene, which codes for part of the enzyme β-galactosidase. This enzyme is involved in the breakdown of lactose to glucose plus galactose. It is normally coded by the gene lacZ, which resides on the E. coli chromosome. Some strains of E. coli have a modified lacZ gene, one that lacks the segment referred to as lacZ’, which codes for the α-peptide portion of β-galactosidase. These mutants can synthesize the enzyme only when they harbor a plasmid, such as pUC8, that carries the missing lacZ’ segment of the gene.

To carry out a cloning experiment with pUC8, the manipulations shown in Figure 2.23, which result in construction of a recombinant plasmid, are performed in the test tube with purified DNA. Pure pUC8 DNA can be obtained quite easily from extracts of bacterial cells, and after manipulation, the plasmids can be reintroduced into E. coli by transformation, the process by which naked DNA is taken up by a bacterial cell. This is the system studied by Avery and his colleagues in the experiments that showed that bacterial genes are made of DNA (Section 1.1). Transformation is not a particularly efficient process with many bacteria, including E. coli, but the rate of DNA uptake can be enhanced significantly by suspending the cells in calcium chloride before addition of the DNA and then briefly incubating the mixture at 42°C. Even after this enhancement, only a very small proportion of the cells take up a plasmid. This is why the ampicillin-resistance marker is so important: it allows the small number of transformants to be selected from the large background of nontransformed cells.

The map of pUC8 shown in Figure 2.25 indicates that the lacZ gene contains a cluster of unique restriction sites. Ligation of new DNA into any one of these sites results in insertional inactivation of the gene and hence loss of β-galactosidase activity. This is the key to distinguishing a recombinant plasmid—one that contains an inserted piece of DNA—from a nonrecombinant plasmid that has no new
DNA. Identifying recombinants is important because the manipulations illustrated in Figures 2.23 and 2.24 result in a variety of ligation products, including plasmids that have recircularized without insertion of new DNA. Screening for the presence or absence of β-galactosidase is, in fact, quite easy. Rather than assaying for lactose being split to glucose and galactose, the presence of functional β-galactosidase molecules in the cells is checked by a histochemical test with a compound called X-gal (5-bromo-4-chloro-3-indolyl β-d-galactopyranoside), which the enzyme converts into a blue product. If X-gal (plus an inducer of the enzyme such as isopropyl thiogalactoside, IPTG) is added to the agar, along with ampicillin, then nonrecombinant colonies, the cells of which synthesize β-galactosidase, will be colored blue, whereas recombinants with a disrupted lacZ gene, which are unable to make β-galactosidase, will be white (Figure 2.26). This system is called Lac selection.

Bacteriophages can also be used as cloning vectors

Bacteriophage (or phage) genomes can also be used as cloning vectors because they too possess origins of replication that enable them to be propagated inside bacteria, either by the bacterial enzymes or by DNA polymerases and other proteins specified by phage genes. E. coli bacteriophages were developed as cloning vectors back in the earliest days of the recombinant DNA revolution. The main reason for seeking a different type of vector was the inability of plasmids such as pUC8 to handle DNA fragments greater than about 10 kb in size. Larger inserts are likely to undergo rearrangements or interfere with the plasmid replication system in such a way that the recombinant DNA molecules become lost from the host cells. The first attempts to develop vectors able to handle larger fragments of DNA centered on the bacteriophage called lambda (λ).

To replicate, a bacteriophage must enter a bacterial cell and subvert the bacterial enzymes into expressing the information contained in the phage genes, so that the bacterium synthesizes new phages. Once replication is complete, the new phages leave the bacterium, usually causing its death as they do so, and move on to infect new cells (Figure 2.27A). This is called a lytic infection cycle because it results in lysis of the bacterium. As well as the lytic cycle, λ (unlike many other types of bacteriophage) can also follow a lysogenic infection cycle, during which the λ genome integrates into the bacterial chromosome, where it can remain quiescent for many generations, being replicated along with the host chromosome every time the cell divides (Figure 2.27B).

The size of the λ genome is 48.5 kb, of which some 15 kb or so is optional in that it contains genes that are needed only for integration of the phage DNA into the E. coli chromosome (Figure 2.28A). These segments can therefore be deleted without impairing the ability of the phage to infect bacteria and direct synthesis of new λ particles by the lytic cycle. Two types of vector have been developed (Figure 2.28B):
• **Insertion vectors**, in which part or all of the optional DNA has been removed and a unique restriction site has been introduced at some position within the trimmed-down genome

• **Replacement vectors**, in which the optional DNA is contained within a stuffer fragment, flanked by a pair of restriction sites, that is replaced when the DNA to be cloned is ligated into the vector

The λ genome is linear, but the two natural ends of the molecule have 12-nucleotide single-stranded overhangs, called **cos sites**, which have complementary sequences and so can base-pair to one another. A λ cloning vector can therefore be obtained as a circular molecule that can be manipulated in the test tube in the same way as a plasmid and reintroduced into *E. coli* by **transfection**, the term used for uptake of naked phage DNA. Alternatively, a more efficient uptake system called **in vitro packaging** can be utilized. This procedure starts with a linear version of the cloning vector, which is cut into two segments, the left and right arms, each with a cos site at one end. Ligation is then carried out with carefully
measured quantities of each arm and the DNA to be cloned. The aim is to produce concatemers in which the different fragments are linked together in the order left arm-new DNA-right arm, as shown in Figure 2.29. The concatemers are then added to an \textit{in vitro} packaging mix, which contains all the proteins needed to make \(\lambda\) phage particles. These proteins form phage particles spontaneously and will place inside the particles any DNA fragment that is between 37 and 52 kb in length.

**Figure 2.28** Cloning vectors based on bacteriophage \(\lambda\). (A) In the \(\lambda\) genome, the genes are arranged into functional groups. For example, the region marked as ‘protein coat’ comprises genes coding for proteins that are either components of the phage capsid or are required for capsid assembly, and the cell lysis region comprises genes involved in lysis of the bacterium at the end of the lytic phase of the infection cycle. The regions of the genome that can be deleted without impairing the ability of the phage to follow the lytic cycle are indicated in green. (B) Differences between a \(\lambda\) insertion vector and a \(\lambda\) replacement vector.

**Figure 2.29** Cloning with a \(\lambda\) insertion vector. The linear form of the vector is shown at the top of the diagram. Treatment with the appropriate restriction endonuclease produces the left and right arms, both of which have one blunt end and one end with the 12-nucleotide overhang of the cos site. The DNA to be cloned is blunt-ended and so is inserted between the two arms during the ligation step. The arms also ligate to one another via their cos sites, forming a concatemer. Some parts of the concatemer are in the order left arm-insert DNA-right arm, and if this combination is 37–52 kb in length, it will be enclosed inside the capsid by the \textit{in vitro} packaging mix. In this example, parts of the concatemer made up of left arm ligated directly to right arm, without new DNA, are too short to be packaged.
length and is flanked by cos sites. The packaging mix therefore cuts left arm–new DNA-right arm combination 37–52 kb out of the concatemers and constructs λ phages around them. The phages are then mixed with E. coli cells, and the natural infection process transports the vector plus new DNA into the bacteria.

After infection, the cells are spread onto an agar plate. The objective is not to obtain individual colonies but to produce an even layer of bacteria across the entire surface of the agar. Bacteria that were infected with the packaged cloning vector die within about 20 minutes because the λ genes contained in the arms of the vector direct replication of the DNA and synthesis of new phages by the lytic cycle. Each of these new phages contains its own copy of the vector plus cloned DNA. Death and lysis of the bacterium releases these phages into the surrounding medium, where they infect new cells and begin another round of phage replication and lysis. The end result is a zone of clearing, called a plaque, which is visible on the lawn of bacteria that grows on the agar plate (Figure 2.30). With some λ vectors, all plaques are made up of recombinant phages because ligation of the two arms without insertion of new DNA results in a molecule too short to be packaged. With other vectors, it is necessary to distinguish recombinant plaques from nonrecombinant ones. Various methods are used, including the β-galactosidase system described for the plasmid vector pUC8 (see Figure 2.26), which is also applicable to those λ vectors that carry a fragment of the lacZ gene into which the DNA to be cloned is inserted.

Vectors for longer pieces of DNA

The λ phage particle can accommodate up to 52 kb of DNA, so if the genome has 15 kb removed, then up to 18 kb of new DNA can be cloned. This limit is higher than that for plasmid vectors but is still very small compared with the size of an intact bacterial or eukaryotic genome. The comparison is important if a clone library is to be used as the starting point for a project aimed at determining a genome sequence (Section 4.3). If a λ vector is used with human DNA, then over half a million clones are needed for there to be a 95% chance of any particular part of the genome being present in the library (Table 2.4). It is possible to prepare a library comprising half a million clones, especially if automated techniques are used, but such a large collection is far from ideal. It would be much better to reduce the number of clones by using a vector that is able to handle fragments of DNA longer than 18 kb. Many of the developments in cloning technology over the last 25 years have been aimed at finding ways of doing this.

### TABLE 2.4 SIZES OF HUMAN GENOMIC LIBRARIES PREPARED IN DIFFERENT TYPES OF CLONING VECTORS

<table>
<thead>
<tr>
<th>Type of vector</th>
<th>Insert size (kb)</th>
<th>( P = 95% )</th>
<th>( P = 99% )</th>
</tr>
</thead>
<tbody>
<tr>
<td>λ replacement</td>
<td>18</td>
<td>532,500</td>
<td>820,000</td>
</tr>
<tr>
<td>Cosmid, fosmid</td>
<td>40</td>
<td>240,000</td>
<td>370,000</td>
</tr>
<tr>
<td>P1</td>
<td>100</td>
<td>96,000</td>
<td>150,000</td>
</tr>
<tr>
<td>BAC, PAC</td>
<td>300</td>
<td>32,000</td>
<td>50,000</td>
</tr>
</tbody>
</table>

*Calculated from the equation

\[
N = \frac{\ln(1 - P)}{\ln(1 - \frac{a}{b})}
\]

where \( N \) is the number of clones required, \( P \) is the probability that any given segment of the genome is present in the library, \( a \) is the average size of the DNA fragments inserted into the vector, and \( b \) is the size of the genome.
One possibility is to use a cosmid, which is a special type of plasmid that carries a λ cos site (Figure 2.31). Concatemers of cosmid molecules, linked at their cos sites, act as substrates for in vitro packaging because the cos site is the only sequence that a DNA molecule needs in order to be recognized as a λ genome by the proteins that package DNA into λ phage particles. Particles containing cosmid DNA are as infective as real λ phages, but once inside the cell, the cosmid cannot direct synthesis of new phage particles and instead replicates as a plasmid. Recombinant DNA is therefore obtained from colonies rather than plaques. As with other types of λ vector, the upper limit for the length of cloned DNA is set by the space available within the λ phage particle. A cosmid can be 8 kb or less in size, so up to 44 kb of new DNA can be inserted before the packaging limit of the λ phage particle is reached. This reduces the size of the human genomic library to about a quarter of a million clones, which is an improvement compared with a λ library but still a massive number of clones to have to work with.

To reduce the size of clone libraries even further, other types of vector with even greater carrying capacities have been developed. The following are the most important of these vectors:

- **Bacterial artificial chromosomes**, or BACs, are based on the naturally occurring F plasmid of *E. coli*. Unlike the plasmids used to construct the early cloning vectors, the F plasmid is relatively large, and vectors based on it have a higher capacity for inserted DNA. BACs are designed so that recombinants can be identified by Lac selection (see Figure 2.26) and hence are easy to use. They can clone fragments of 300 kb and longer, and the inserts are very stable. BACs were used extensively in the Human Genome Project (Section 4.4), and they are currently the most popular vectors for cloning large pieces of DNA.

- **Bacteriophage P1 vectors** are very similar to λ vectors in that they are based on a deleted version of a natural phage genome, where the capacity of the cloning vector is determined by the size of the deletion and the space within the phage particle. P1 has the advantage over λ that it is able to squeeze 110 kb of DNA into its capsid structure, which results in P1 vectors having a higher capacity than those based on λ. Cosmid-type P1 vectors have been designed and used to clone DNA fragments ranging in size from 75 to 100 kb.

- **P1-derived artificial chromosomes**, or PACs, combine features of P1 vectors and BACs and have a capacity of up to 300 kb.

- **Fosmids** contain the F plasmid origin of replication and a λ cos site. They are similar to cosmids in the way they are used and in their capacity for inserted DNA, but they have a lower copy number in *E. coli*, which means that they are less prone to instability problems.

The sizes of human genome libraries prepared in these various types of vectors are given in Table 2.4.

**DNA can be cloned in organisms other than *E. coli***

Cloning is not merely a means of producing DNA for sequencing and other types of analysis. It is also the central part of techniques that are used to identify the function of an unknown gene and to study its mode of expression and the way in which its expression is regulated. In the wider research world, cloning is also used to carry out genetic engineering experiments aimed at modifying the biological characteristics of the host organism and to transfer genes for important animal proteins, such as pharmaceuticals, into a new host cell from which the proteins can be obtained in larger quantities than is possible by conventional purification from animal tissue. These various applications demand that genes must frequently be cloned in organisms other than *E. coli*. 
Cloning vectors based on plasmids or phages have been developed for most of the well-studied species of bacteria, such as *Bacillus*, *Streptomyces*, and *Pseudomonas*; these vectors are used in exactly the same way as the *E. coli* analogs. Plasmid vectors are also available for yeasts and fungi. Some of these carry the origin of replication from the 2μm plasmid, which is present in many strains of *Saccharomyces cerevisiae*, but other plasmid vectors for yeast and fungi have only an *E. coli* origin. An example is YIp5, an *S. cerevisiae* vector that is simply an *E. coli* plasmid that contains a copy of the yeast gene called URA3 (Figure 2.32A). The presence of the *E. coli* origin means that YIp5 is a shuttle vector that can be used with either *E. coli* or *S. cerevisiae* as the host. This is a useful feature because cloning in *S. cerevisiae* is a relatively inefficient process, and generating a large number of clones is difficult. If the experiment requires that the desired recombinant be identified from a mixture of clones, then it may not be possible to obtain enough recombinants to find the correct one. To avoid this problem, construction of recombinant DNA molecules and selection of the correct recombinant is carried out with *E. coli* as the host. When the correct clone has been identified, the recombinant YIp5 construct is purified and transferred into *S. cerevisiae*, usually by mixing the DNA with protoplasts, which are yeast cells whose walls have been removed by enzyme treatment. Without an origin of replication, the vector is unable to propagate independently inside yeast cells, but it can survive if it becomes integrated into one of the yeast chromosomes, which can occur by homologous recombination (Section 17.1) between the URA3 gene carried by the vector and the chromosomal copy of this gene (Figure 2.32B). YIp in fact stands for yeast integrative plasmid. Once integrated, the YIp, plus any DNA that has been inserted into it, replicates along with the host chromosomes.

Integration into chromosomal DNA is also a feature of many of the cloning systems used with animals and plants, and it forms the basis of the construction of knockout mice, which have been used to identify the functions of previously unknown genes discovered in the human genome (Section 6.2). The vectors are animal equivalents of YIps. A similar range of vectors has been developed for cloning genes in plants. Bacterial plasmids can be introduced into plant embryos by bombardment with DNA-coated microprojectiles, a process called biolistics. Integration of plasmid DNA into plant chromosomal DNA, followed by growth of the embryo, results in a plant that contains the cloned DNA in most or all of its cells. Some success has also been achieved with plant vectors based on the genomes of caulimoviruses and geminiviruses, but the most interesting types of plant cloning vector are those derived from the Ti plasmid, a large bacterial plasmid found in the soil microorganism *Agrobacterium tumefaciens*. Part of the Ti plasmid, the region called the T-DNA, becomes integrated into a plant chromosome when the bacterium infects a plant stem and causes crown gall disease. The T-DNA carries a number of genes that are expressed inside the plant cells and...
induce the various physiological changes that characterize the disease. Vectors such as pBIN19 (Figure 2.33) have been designed to make use of this natural genetic engineering system. The recombinant vector is introduced into A. tumefaciens cells, which are allowed to infect a cell suspension or plant callus culture, from which mature transformed plants can be regenerated (Figure 2.34).

**SUMMARY**

- The four main types of enzyme used in recombinant DNA technology are DNA polymerases, nucleases, ligases, and end-modification enzymes.
- DNA polymerases synthesize new DNA polynucleotides and are used in procedures such as DNA sequencing and PCR.
- The most important nucleases are the restriction endonucleases, which cut double-stranded DNA molecules at specific nucleotide sequences, and hence cut a molecule into a predicted set of fragments, the sizes of which can be determined by agarose gel electrophoresis.
- Ligases join molecules together and end-modification enzymes carry out a variety of reactions including several used to label DNA molecules.
- PCR results in the repeated copying of a selected region of a DNA molecule, but at least part of the DNA sequence of this region must be known.
- Starting with just a single target DNA molecule, over 130 million copies can be made during 30 cycles of a PCR.
- Real-time and quantitative methods enable the dynamics of product synthesis to be followed during a PCR.
DNA cloning is a means of obtaining a pure sample of an individual gene or other segment of a DNA molecule.

Many different types of cloning vector have been designed for use with *E. coli* as the host organism, the simplest being based on small plasmids that carry selectable markers such as the *lacZ*’ gene.

Bacteriophage *λ* has also been used as the basis for a series of *E. coli* cloning vectors, including the plasmid–phage hybrids called cosmids, which are used to clone fragments of DNA up to 44 kb in length.

Other types of vector, such as bacterial artificial chromosomes, can be used to clone even longer pieces of DNA up to 300 kb.

Organisms other than *E. coli* can also be used as the hosts for DNA cloning. Several types of vector have been designed for *Saccharomyces cerevisiae*, and specialized techniques are available for cloning DNA in animals and plants.

**SHORT ANSWER QUESTIONS**

1. Describe how a DNA fragment containing a single gene of interest would be obtained by (A) PCR or (B) gene cloning.

2. How can a researcher identify a single restriction enzyme fragment containing a gene of interest in a digest of genomic DNA that contains thousands of different restriction fragments?

3. Describe a useful and quick method for increasing the ligation efficiency of blunt-ended DNA molecules.

4. Why are the initial PCR products—produced in the first few cycles of the reaction—long and of varying sizes, and the final PCR products all of a shorter and uniform size?

5. How do the primers determine the specificity of a PCR?

6. Explain how the rate of product formation can be followed during a PCR.

7. Explain why bacterial plasmids are popular cloning vectors.

8. Distinguish between the ways in which antibiotic resistance and Lac selection are used in identification of recombinant cloning vectors.

9. List the features of bacteriophage *λ* that have led to this phage being used as a cloning vector.

10. Outline the key differences between a *λ* insertion vector and a *λ* replacement vector.

11. Why are vectors that can carry larger DNA inserts beneficial for the creation of clone libraries?

12. Describe the important features of the cloning systems used with (A) *Saccharomyces cerevisiae*, (B) animals, and (C) plants.

**IN-DEPTH PROBLEMS**

1. How might you determine the positions of the restriction sites in a DNA molecule, other than by working out the sequence of the molecule?

2. Calculate the numbers of short and long products that would be present after 20, 25, and 30 cycles of a PCR.
3. When DNA is cloned in pUC8, recombinant bacteria (those containing a circular pUC8 molecule that carries an inserted DNA fragment) are identified by plating onto an agar medium containing ampicillin and the lactose analog called X-gal. An older type of cloning vector, called pBR322, also had the gene for ampicillin resistance but did not carry the lacZ' gene. Instead, DNA was inserted into a gene for tetracycline resistance present in pBR322. Describe the procedure that would be needed to distinguish bacteria that had taken up a recombinant pBR322 plasmid from those that had taken up a plasmid that had circularized without insertion of new DNA.

4. What would be the features of an ideal cloning vector? To what extent are these requirements met by any of the existing cloning vectors?

5. Soon after the first gene cloning experiments were carried out in the early 1970s, a number of scientists argued that there should be a temporary moratorium on this type of research. What was the basis of these scientists’ fears and to what extent were these fears justified?

FURTHER READING

Textbooks and practical guides on the methods used to study DNA

Enzymes for DNA manipulation

PCR


DNA cloning in bacteria

High-capacity cloning vectors

Cloning in plants and animals


**Online resources**


REBASE. http://rebase.neb.com/rebase/rebase.html *A comprehensive list of all the known restriction endonucleases and their recognition sequences.*
In this chapter we will study the various ways in which genome maps are constructed. A genome map, like any other type of map, indicates the positions of interesting features and other important landmarks. In a genome map, these features and landmarks are genes and other distinctive DNA sequences. Although a variety of techniques can be used to map genes and other DNA landmarks, the convention is to look on genome mapping as comprising two complementary approaches:

- Genetic mapping (Sections 3.2–3.4), also called linkage analysis, is based on the use of genetic techniques, including planned breeding experiments or, in the case of humans, the examination of family histories (also called pedigrees).

- Physical mapping (Sections 3.5 and 3.6) uses molecular biology techniques to examine DNA molecules directly in order to identify the positions of sequence features, including genes.

Before exploring the various techniques involved in genetic and physical mapping, we must first understand why genome maps are important.

### 3.1 Why a Genome Map Is Important

The study of genomes is often looked on as a modern, edgy area of biological research, far divorced from the work of the old era geneticists such as Gregor Mendel. And yet many of the techniques used to construct genome maps are based directly on the discoveries of Mendel and the other early geneticists. We must therefore spend a few minutes understanding why genome mapping, despite being an old-fashioned type of biology, is still important in the fast-paced research of the genomic age.

**Genome maps are needed in order to sequence the more complex genomes**

During the early days of genome research, it was believed that possession of a detailed map would be an essential prerequisite for assembly of the correct sequence of a genome. This is because DNA sequencing has one major limitation: only with the most sophisticated and recently introduced technology is it possible to obtain a sequence of more than about 750 bp in a single experiment. This means that the sequence of a long DNA molecule has to be constructed from a series of shorter sequences. This is done by breaking the molecule into fragments, determining the sequence of each one, and using a computer to search for overlaps and build up the master sequence (Figure 3.1). This **shotgun method** is the standard approach for genome sequencing, but it suffers from two problems.

---

**Figure 3.1** The shotgun method for sequence assembly. The DNA molecule is broken into small fragments, each of which is sequenced. The master sequence is assembled by searching for overlaps between the sequences of individual fragments.
Chapter 3: Mapping Genomes

The first is that, especially with larger genomes, it might not be possible to obtain sufficient short sequences to produce a contiguous DNA sequence for the entire genome. Instead, the genome sequence might be made up of many short segments separated by gaps that represent parts of the genome that, by chance, are not covered by the sequences that have been obtained (Figure 3.2). If these segments are unconnected, then how can they be positioned correctly relative to one another in order to build up the genome sequence? The answer is to identify within those segments features that are located on the genome map. By anchoring the segments onto the map, the correct genome sequence can be obtained, even if that sequence still contains some gaps.

The second problem with the shotgun approach is that it can lead to errors if the genome contains repetitive DNA sequences. These are sequences, up to several kilobases in length, that are repeated at two or more places in a genome. When a genome containing repetitive DNA is broken into fragments, some of the resulting pieces will contain the same sequence motifs. It would be very easy to reassemble these sequences so that a portion of the DNA between the repeats is left out, or even to connect together two quite separate pieces of the same or different chromosomes (Figure 3.3A). Once again, a genome map enables errors of this type to be avoided. If the sequence features on either side of a repetitive region match the genome map, then the sequence in that region has been put together correctly. If the sequence and the map do not match, then a mistake has been made and the assembly must be revised (Figure 3.3B).

Over the years, sequencing technology has become more powerful, enabling ever-increasing numbers of short sequences to be generated from a single genome, which means that there is less likelihood that the final sequence will contain many gaps. At the same time, the computer algorithms used to assemble those sequences into contiguous segments have become more sophisticated. The latest algorithms are able to recognize when the assembly reaches a region of repetitive DNA and can take steps to ensure that the sequence around these regions is not put together incorrectly (Section 4.3). Many prokaryotic genomes (which are relatively small and have little repetitive DNA) have been sequenced without reference to a map, and an increasing

Figure 3.2 Using a genome map as an aid to sequence assembly. A genome has been broken into short DNA fragments, which have been sequenced by the shotgun method. When the sequences are assembled, a series of unconnected genome segments is obtained. The segments contain genes and other sequence features (A, B, C, etc.) whose positions in the genome have been mapped. The map can therefore be used to identify the positions of the segments in the genome sequence.

Figure 3.3 A possible error in sequence assembly caused by repetitive DNA. (A) The DNA molecule contains two copies of a repeat sequence. When the shotgun sequences are examined, two fragments appear to overlap, but one fragment contains the left-hand part of one repeat and the other fragment has the right-hand part of the second repeat. Failure to recognize this assembly error would lead to the segment of DNA between the two repeats being left out of the master sequence. If the two repeats were on different chromosomes, then the sequences of these chromosomes would mistakenly be linked together. (B) The error in sequence assembly is recognized because the relative positions of mapped features (A, B, C, etc.) in the assembled sequence do not correspond with the correct positions of these features in the genome map.
3.1 Why a Genome Map is Important

The number of eukaryotic genome projects are dispensing with them. But maps are not yet entirely redundant as aids to genome sequencing. One of the greatest challenges today is obtaining genome sequences for important crop plants. Many of these species have large genomes with a substantial repetitive DNA content. The sunflower, *Helianthus annuus*, which is a source of vegetable oil used as both food and biofuel, is an example. Its genome is only slightly larger than the human genome (3600 Mb for *H. annuus* compared with 3235 Mb for humans), but 80% of the sunflower genome is made up of repetitive DNA, compared with just 44% for the human genome. The barley genome also has about 80% repetitive DNA and is much larger, at 5100 Mb. An even greater challenge is presented by bread wheat, which is a hexaploid, meaning that it has three genomes, called A, B, and D. Each is about 5500 Mb (a massive 16,500 Mb in total) with a similar repetitive DNA content to barley. The genome projects for these and other important crops are still ongoing, and because of the complexities of their genomes, comprehensive maps are essential in order to assemble the sequences. This is a critical area of research: understanding all aspects of the biology of crops is essential for dealing with global hunger over the coming decades.

**Genome maps are not just sequencing aids**

Maps might have become less generally relevant as aids in the assembly of genome sequences, but their value in other aspects of genomics research is undiminished. It is important to recognize that completion of the nucleotide sequence of a genome is not an end in itself. Indeed, every genome is simply a series of As, Cs, Gs, and Ts, and working out the order of these letters does not tell us much, if anything, about the way in which a genome acts as a store of biological information or how that information is used to specify the characteristics of the species being studied. As we will see in Chapters 5 and 6, the first stage in understanding a genome sequence is to identify the genes that it contains and to assign functions to as many of these as possible. Many of the methods used to assign functions begin with a gene and ask what does this gene do, but the reverse process, in which we start with a function and ask what is the responsible gene, is equally important. As we will see in Section 6.4, a genome map is essential in order to answer this second question, because the approach used initially involves identifying the position of the gene being sought relative to other genes or sequence features whose locations on the map are already known. This process has been, and continues to be, the key to identification of genes responsible for human diseases such as cystic fibrosis and breast cancer. Similar methods are used to identify groups of genes, possibly spread around the genome, that do not directly cause a disease but confer differing degrees of susceptibility to that disease. One step further on are methods used to identify quantitative trait loci (QTLs), which are regions of a genome, each possibly containing several genes, that control variable traits such as meat productivity in farm animals and pest resistance in crop plants.

The information provided by a genome map on the locations of genes and QTLs controlling commercially important traits in crop plants is also utilized in breeding programs aimed at the development of new varieties with improved agricultural properties. These breeding programs typically generate thousands of seedlings, whose precise biological characteristics are unknown because of the randomness of the inheritance process. A seedling might combine the best features of the two parents and potentially be an important new crop variety, or it might combine the least useful properties of both parents and be of no commercial value. Many traits of interest to crop breeders are exhibited late in the life cycle of the plant—examples are seed or fruit yield—and can be assayed only by growing each seedling to maturity, which takes time and requires large amounts of growing space. We will see in Section 18.4 how the method called marker-assisted selection enables DNA screening to be used to identify those seedlings that possess a beneficial characteristic, so these can be retained and other, less interesting seedlings can be discarded. Marker-associated selection is possible only if a genome map is available. If a map is available, then it can be
carried out with success even if the complete genome sequence is unknown, as is the case for crops such as barley and wheat.

3.2 MARKERS FOR GENETIC MAPPING

As with any type of map, a genetic map must show the positions of distinctive features. In a geographic map, these markers are recognizable components of the landscape such as rivers, roads, and buildings. What markers can we use in a genetic landscape?

Genes were the first markers to be used

The first genetic maps, constructed in the early decades of the twentieth century for organisms such as the fruit fly, used genes as genetic markers. To be useful in genetic analysis, a gene must exist in at least two forms or alleles, each specifying a different phenotype; an example is tall or short stems in the pea plants originally studied by Gregor Mendel. To begin with, the only genes that could be studied were those specifying phenotypes that were distinguishable by visual examination. So, for example, the first fruit fly maps showed the positions of genes for body color, eye color, wing shape, and so forth, all of these phenotypes being visible simply by looking at the flies with a low-power microscope or the naked eye. This approach was fine in the early days, but geneticists soon realized that there were only a limited number of visual phenotypes whose inheritance could be studied, and in many cases their analysis was complicated because a single phenotype could be affected by more than one gene. For example, by 1922, over 50 genes had been mapped onto the four fruit fly chromosomes, but nine of these genes were for eye color. In later research, geneticists studying fruit flies had to learn to distinguish between fly eyes that were colored red, light red, vermilion, garnet, carnation, cinnabar, ruby, sepia, scarlet, pink, cardinal, claret, purple, or brown. To make gene maps more comprehensive, it was necessary to find characteristics that were more distinctive and less complex than visual ones.

The answer was to use biochemistry to distinguish phenotypes. This has been particularly important with two types of organisms: microbes and humans. Microbes, such as bacteria and yeast, have very few visual characteristics, so gene mapping with these organisms has to rely on biochemical phenotypes such as those listed in Table 3.1. For humans, it is possible to use visual characteristics, but since the 1920s, studies of human genetic variation have been based largely on biochemical phenotypes that can be scored by blood typing. These phenotypes include not only the standard blood groups, such as the ABO series, but also variants of blood serum proteins and of immunological proteins such as the human leukocyte antigens (the HLA system). A big advantage of these markers is that

<table>
<thead>
<tr>
<th>Marker</th>
<th>Phenotype</th>
<th>Method by which cells carrying the marker are identified</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADE2</td>
<td>Requires adenine</td>
<td>Grows only when adenine is present in the medium</td>
</tr>
<tr>
<td>CAN1</td>
<td>Resistant to canavanine</td>
<td>Grows in the presence of canavanine</td>
</tr>
<tr>
<td>CUP1</td>
<td>Resistant to copper</td>
<td>Grows in the presence of copper</td>
</tr>
<tr>
<td>CYH1</td>
<td>Resistant to cycloheximide</td>
<td>Grows in the presence of cycloheximide</td>
</tr>
<tr>
<td>LEU2</td>
<td>Requires leucine</td>
<td>Grows only when leucine is present in the medium</td>
</tr>
<tr>
<td>SUC2</td>
<td>Able to ferment sucrose</td>
<td>Grows if sucrose is the only carbohydrate in the medium</td>
</tr>
<tr>
<td>URA3</td>
<td>Requires uracil</td>
<td>Grows only when uracil is present in the medium</td>
</tr>
</tbody>
</table>
many of the relevant genes have multiple alleles. For example, the gene called \textit{HLA-DRB1} has over 1800 alleles and \textit{HLA-B} has 4200. This is relevant because of the way in which gene mapping is carried out with humans (Section 3.4). Rather than setting up planned breeding experiments, which is the procedure with experimental organisms such as fruit flies or mice, data on inheritance of human genes have to be gleaned by examining the phenotypes displayed by members of families in which the parents have come together for personal reasons rather than for the convenience of a geneticist. If all the members of a family have the same allele for the gene being studied, then no useful information can be obtained. For gene mapping purposes, it is therefore necessary to find families in which the parents, by chance, have different alleles. This is much more likely if the gene being studied has 1800 rather than 2 alleles.

**RFLPs and SSLPs are examples of DNA markers**

Genes are very useful markers but they are by no means ideal. One problem, especially with larger genomes such as those of vertebrates and flowering plants, is that a map based entirely on genes is not very detailed. This would be true even if every gene could be mapped, because in most eukaryotic genomes the genes are widely spaced out with large gaps between them. The problem is made worse by the fact that many genes do not exist in allelic forms that can be distinguished conveniently. Gene maps are therefore not very comprehensive. We need other types of markers.

Mapped features that are not genes are called DNA markers. As with gene markers, a DNA marker must have at least two alleles to be useful. Two examples of DNA markers are the sequences called \textit{restriction fragment length polymorphisms} (RFLPs) and \textit{simple sequence length polymorphisms} (SSLPs).

RFLPs were the first type of DNA marker to be studied. Recall that restriction enzymes cut DNA molecules at specific recognition sequences (Section 2.1). This sequence specificity means that treatment of a DNA molecule with a restriction enzyme should always produce the same set of fragments. This is not always the case with genomic DNA molecules because some restriction sites are polymorphic, existing as two alleles, with one allele displaying the correct sequence for the restriction site and therefore being cut when the DNA is treated with the enzyme, and the second allele having a sequence alteration so the restriction site is no longer recognized. The result of the sequence alteration is that the two adjacent restriction fragments remain linked together after treatment with the enzyme, leading to a length polymorphism (Figure 3.4). This is an RFLP, and its position on a genome map can be worked out by following the inheritance of its alleles, just as is done when genes are used as markers. There are thought to be about $10^5$ RFLPs in a mammalian genome.

With small DNA molecules, the two alleles of an RFLP can be distinguished simply by cutting with the appropriate restriction enzyme and identifying the sizes of the resulting fragments in an agarose gel. Typing an RFLP in genomic DNA is more difficult. An enzyme such as \textit{EcoRI}, with a six-nucleotide recognition sequence, should cut approximately once every $4^6 = 4096$ bp and so would give almost 800,000 fragments when used with human DNA. After separation by agarose gel electrophoresis, these 800,000 fragments produce a smear of DNA. Southern hybridization, using a probe that spans the polymorphic restriction site, would therefore have to be carried out in order to visualize the fragments relevant to the RFLP (Figure 3.5A). This is a lengthy process, and it is difficult to examine more than about 12 DNA samples in a single experiment. RFLP typing is an example of the many procedures that have been made easier since PCR was invented. By use of PCR, an RFLP can be typed in a sample of genomic DNA without the need to cut that DNA with the restriction enzyme. Instead, the primers for the PCR are designed so that they anneal on either side of the polymorphic site, and the RFLP is typed by treating the amplified fragment with the restriction enzyme (Figure 3.5B). Multiple PCRs can easily be set up in multiwell plates, so up to 96 DNA samples can now be typed in a single run.

![Figure 3.4 A restriction fragment length polymorphism (RFLP).](image)

The DNA molecule on the left has a polymorphic restriction site (marked with the asterisk) that is not present in the molecule on the right. The RFLP is revealed after treatment with the restriction enzyme because one of the molecules is cut into four fragments whereas the other is cut into three fragments.
SSLPs are quite different from RFLPs. SSLPs are arrays of repeat sequences that display length variations, with different alleles containing different numbers of repeat units (Figure 3.6A). Unlike RFLPs, SSLPs can be multiallelic, as each SSLP can have a number of different length variants. There are two types of SSLPs:

- **Minisatellites**, also known as **variable number of tandem repeats (VNTRs)**, in which the repeat unit is up to 25 bp in length
- **Microsatellites**, or **short tandem repeats (STRs)**, whose repeats are shorter, usually 13 bp or less

Microsatellites are more popular than minisatellites as DNA markers for two reasons. First, minisatellites are not spread evenly around the genome but tend to be found more frequently in the telomeric regions at the ends of chromosomes. In geographic terms, this is equivalent to trying to use a map of lighthouses to find one’s way around the middle of an island. Microsatellites are more conveniently spaced throughout the genome. Second, the fastest way to type a length polymorphism is by PCR, but PCR typing is much quicker and more accurate with sequences less than 300 bp in length. Most minisatellite alleles are longer than this because the repeat units are relatively large and there tend to be many of them in a single array, so PCR products several kilobases in length are needed to

![Figure 3.6 SSLPs and how they are typed.](image)

(A) Southern hybridization

(B) PCR

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type them. Microsatellites used as DNA markers typically consist of 10–30 copies of a repeat that is no longer than 6 bp in length, and so they are much more amenable to analysis by PCR. There are $2.86 \times 10^6$ microsatellites with repeat units of 2–6 bp in the human genome.

When examined by PCR, the allele present at an STR is revealed by the precise length of the PCR product (Figure 3.6B). The length variations can be visualized by agarose gel electrophoresis, but standard gel electrophoresis is a cumbersome procedure that is difficult to automate, which means that it is unsuitable for the high-throughput analyses that are demanded by modern genome research. Instead, STRs are usually typed by capillary electrophoresis in a polyacrylamide gel. Polyacrylamide gels have smaller pore sizes than agarose gels and allow greater precision in the separation of molecules of different lengths. Most capillary electrophoresis systems use fluorescence detection, so a fluorescent label is attached to one or both of the primers before the PCR is carried out. After PCR, the product is loaded into the capillary system and run past a fluorescence detector. A computer attached to the detector correlates the time of passage of the PCR product with equivalent data for a set of size markers and hence identifies the precise length of the product.

**Single-nucleotide polymorphisms are the most useful type of DNA marker**

RFLPs and SSLPs are useful in some types of genomic research, but most modern genetic mapping projects make use of a different type of DNA marker. These are called **single-nucleotide polymorphisms (SNPs)**. An SNP is a position in a genome where some individuals have one nucleotide (e.g., a G) and others have a different nucleotide (e.g., a C) (Figure 3.7). There are vast numbers of SNPs in every genome (approximately 10 million in the human genome), some of which also give rise to RFLPs, but many of which do not because the sequence in which they lie is not recognized by any restriction enzyme.

Any one of the four nucleotides could be present at any single position in a genome, so it might be imagined that each SNP should have four alleles. Theoretically this is possible but in practice most SNPs exist as just two variants. This is because each SNP originates when a **point mutation** (Chapter 16) occurs in a genome, converting one nucleotide into another. If the mutation occurs in the reproductive cells of an individual, then one or more of that individual’s offspring might inherit the mutation and, after many generations, the SNP may eventually become established in the population. But there are just two alleles: the original sequence and the mutated version. For a third allele to arise, a new mutation must occur at the same position in the genome in another individual, and this individual and his or her offspring must reproduce in such a way that the new allele becomes established. This scenario is not impossible but it is unlikely: consequently the vast majority of SNPs are biallelic. This disadvantage is more than outweighed by the huge number of SNPs present in each genome: in most eukaryotes, at least one for every 1000 bp of DNA. SNPs therefore enable very detailed genome maps to be constructed.

The frequency of SNPs in a genome means that these markers have assumed considerable importance in projects that utilize a genome map in order to identify genes or QTLs specifying particular characteristics (Section 6.4), as well as in crop breeding programs that use a map as an aid to marker-assisted selection (Section 18.4). These applications have driven the development of methods for rapid typing of individual SNPs as well as large sets of SNPs. Several of these typing methods are based on **oligonucleotide hybridization analysis**. An oligonucleotide is a short, single-stranded DNA molecule, usually less than 50 nucleotides long, that is synthesized in the test tube. If the conditions are just right, then an oligonucleotide will hybridize with another DNA molecule only if the oligonucleotide forms a completely base-paired structure with the second molecule. If there is a single mismatch—a single position within the oligonucleotide...
Figure 3.8 The basis of SNP typing by oligonucleotide hybridization analysis. Under highly stringent hybridization conditions, a stable hybrid occurs only when the oligonucleotide is able to form a completely base-paired structure with the target DNA. If there is a single mismatch, then the hybrid does not form. To achieve this level of stringency, the incubation temperature must be just below the melting temperature, or \( T_m \), of the oligonucleotide. At temperatures above \( T_m \), even the fully base-paired hybrid is unstable. At more than 5°C below \( T_m \), mismatched hybrids might be stable. \( T_m \) for the oligonucleotide shown in the figure would be about 58°C. \( T_m \) (in degrees Celsius) is calculated from the formula

\[
T_m = (4 \times \text{number of G and C nucleotides}) + (2 \times \text{number of A and T nucleotides}).
\]

This formula gives a rough indication of \( T_m \) for oligonucleotides of 15–30 nucleotides in length.

that does not form a base pair—then hybridization does not occur (Figure 3.8). Oligonucleotide hybridization can therefore discriminate between the two alleles of an SNP. Various SNP typing strategies based on oligonucleotide hybridization have been devised, including the following:

- **DNA chip** technology makes use of a wafer of glass or silicon, 2 cm\(^2\) or less in area, carrying many different oligonucleotides in a high-density array. The DNA to be tested is labeled with a fluorescent marker and pipetted onto the surface of the chip. Hybridization is detected by examining the chip with a fluorescence microscope. The positions at which the fluorescent signal is emitted indicate which oligonucleotides have hybridized with the test DNA (Figure 3.9). Hybridization requires a complete match between an oligonucleotide and its complementary sequence in the test DNA, and so indicates which of the two versions of a SNP is present in the test. A density of up to 300,000 oligonucleotides/cm\(^2\) is possible on the surface of the chip, so a chip of 2 cm\(^2\) can type 300,000 SNPs in a single experiment, if the chip carries oligonucleotides for both alleles of each SNP.

- **Solution hybridization** techniques are carried out in the wells of a microtiter tray, using a detection system that can discriminate between nonhybridized, single-stranded DNA and the double-stranded product that results when an oligonucleotide hybridizes to the test DNA. The most popular detection system makes use of dye quenching, which we encountered in Section 2.2 as the basis to the way in which a reporter probe is used to follow product formation during real-time PCR (see Figure 2.21). In SNP typing, the dye is attached to one end of the oligonucleotide and the quenching compound to the other end. Hybridization between oligonucleotide and the test DNA is indicated by generation of the fluorescent signal. When used in this context, the dye-quenching technique is sometimes called **molecular beacons**.

Other typing methods make use of an oligonucleotide whose mismatch with the SNP occurs at its extreme 5'- or 3'-end. Under the appropriate conditions, an oligonucleotide of this type will hybridize to the mismatched template DNA with a short, non-base-paired tail (Figure 3.10A). This feature is utilized in two different ways:

- The **oligonucleotide ligation assay** (OLA) makes use of two oligonucleotides that anneal adjacent to one another, with the 3'-end of one of these oligonucleotides positioned exactly at the SNP. This oligonucleotide will form a completely base-paired structure if one version of the SNP is present in the template DNA, and when this occurs, the oligonucleotide can be ligated to its partner (Figure 3.10B). If the DNA being examined contains the other allele of the SNP, then the 3'-nucleotide of the test oligonucleotide will not anneal to the template and no ligation occurs. The allele is therefore typed by determining if the ligation product is synthesized. If a
single SNP is being assayed, then formation of the ligation product can be identified by running the postreaction mixture in a capillary electrophoresis system, as described above for STR typing.

- In the amplification refractory mutation system, or ARMS test, the test oligonucleotide is one of a pair of PCR primers. If the 3'-nucleotide of the test primer anneals to the SNP, then it can be extended by Taq polymerase and the PCR can take place, but if it does not anneal because the alternative version of the SNP is present, then no PCR product is generated (Figure 3.10C).

### 3.3 THE BASIS TO GENETIC MAPPING

Now that we have assembled a set of markers with which to construct a genetic map, we can move on to look at the mapping techniques themselves. These techniques are all based on genetic linkage, which in turn derives from the seminal discoveries in genetics made in the mid-nineteenth century by Gregor Mendel.

**The principles of inheritance and the discovery of linkage**

Genetic mapping is based on the principles of inheritance as first described by Gregor Mendel in 1865. From the results of his breeding experiments with peas, Mendel concluded that each pea plant possesses two alleles for each gene but displays only one phenotype. This is easy to understand if the plant is pure-breeding, or homozygous, for a particular characteristic, as it then possesses two identical alleles and displays the appropriate phenotype (Figure 3.11A). However, Mendel

**Figure 3.11 Homozygosity and heterozygosity.** Mendel studied seven pairs of contrasting characteristics in his pea plants, one of which was violet or white flower color, as shown here. (A) Pure-breeding plants always give rise to flowers with the parental color. These plants are homozygotes, each possessing a pair of identical alleles, denoted here by $VV$ for violet flowers and $WW$ for white flowers. (B) When two pure-breeding plants are crossed, only one of the phenotypes is seen in the $F_1$ generation. Mendel deduced that the **genotype** of the $F_1$ plants was $VW$, so $V$ is the dominant allele and $W$ is the recessive allele.
Incomplete dominance

**Figure 3.12** Two types of allele interaction not encountered by Mendel. (A) Incomplete dominance of flower color in carnations. (B) Co-dominance of the \( M \) and \( N \) blood group alleles.

showed that if two pure-breeding plants with different phenotypes are crossed, then all the progeny (the \( F_1 \) generation) display the same phenotype. These \( F_1 \) plants must be **heterozygous**, meaning that they possess two different alleles, one for each phenotype: one allele inherited from the mother and one from the father. Mendel postulated that in this heterozygous condition one allele overrides the effects of the other allele; he therefore described the phenotype expressed in the \( F_1 \) plants as being **dominant** over the second, **recessive** phenotype (**Figure 3.11B**).

Mendel’s interpretation of the heterozygous condition is perfectly correct for the pairs of alleles that he studied, but we now appreciate that this simple dominant-recessive rule can be complicated by situations that he did not encounter. These include the following:

- **Incomplete dominance**, where the heterozygous form displays a phenotype intermediate between the two homozygous forms. Flower color in plants such as carnations (but not peas) is an example: when red carnations are crossed with white ones, the \( F_1 \) heterozygotes are neither red nor white but pink (**Figure 3.12A**).

- **Co-dominance**, where the heterozygous form displays both of the homozygous phenotypes. Human blood groups provide several examples of co-dominance. For example, the two homozygous forms of the MN series are \( M \) and \( N \), with these individuals synthesizing \( M \) or \( N \) blood glycoproteins, respectively. Heterozygotes, however, synthesize both glycoproteins and hence are designated MN (**Figure 3.12B**).

As well as discovering dominance and recessiveness, Mendel carried out additional experiments that enabled him to establish his two laws of genetics. The first law states that **alleles segregate randomly**. In other words, if the parent’s alleles are \( A \) and \( a \), then a member of the \( F_1 \) generation has the same chance of inheriting \( A \) as it has of inheriting \( a \). The second law is that **pairs of alleles segregate independently**, so that inheritance of the alleles of gene \( A \) is independent of inheritance of the alleles of gene \( B \). Because of these laws, the outcomes of genetic crosses are predictable (**Figure 3.13**).

When Mendel’s work was rediscovered in 1900, his second law worried the early geneticists because it was soon established that genes reside on chromosomes, and it was realized that all organisms have many more genes than chromosomes.
Chromosomes are inherited as intact units, so it was reasoned that the alleles of some pairs of genes will be inherited together because they are on the same chromosome (Figure 3.14). This is the principle of genetic linkage, and it was quickly shown to be correct, although the results did not turn out exactly as expected. The complete linkage that had been anticipated between many pairs of genes failed to materialize. Pairs of genes were either inherited independently, as expected for genes in different chromosomes, or if they showed linkage, it was only partial linkage: sometimes they were inherited together and sometimes they were not (Figure 3.15). The resolution of this contradiction between prediction and observation was the critical step in the development of genetic mapping techniques.

**Partial linkage is explained by the behavior of chromosomes during meiosis**

The critical breakthrough was achieved by Thomas Hunt Morgan, who made the conceptual leap between partial linkage and the behavior of chromosomes when the nucleus of a cell divides. Cytologists in the late nineteenth century had distinguished two types of nuclear division: mitosis and meiosis. Mitosis is more common: it is the process by which the diploid nucleus of a somatic cell...
divides to produce two daughter nuclei, both of which are diploid (Figure 3.16). Approximately $10^{17}$ mitoses are needed to produce all the cells required during a human lifetime. Before mitosis begins, each chromosome in the nucleus is replicated, but the resulting daughter chromosomes do not immediately break away from one another. To begin with, they remain attached at their centromeres. The daughters do not separate until later in mitosis, when the chromosomes are distributed between the two new nuclei. Obviously it is important that each of the new nuclei receives a complete set of chromosomes, and most of the intricacies of mitosis appear to be devoted to achieving this end.

Mitosis illustrates the basic events occurring during nuclear division, but it is the distinctive features of meiosis that interest us. Meiosis occurs only in reproductive cells and results in a diploid cell giving rise to four haploid gametes, each of which can subsequently fuse with a gamete of the opposite sex during sexual reproduction. The fact that meiosis results in four haploid cells whereas mitosis gives rise to two diploid cells is easy to explain: meiosis involves two nuclear divisions, one after the other, whereas mitosis is just a single nuclear division. This is an important distinction, but the critical difference between mitosis and meiosis is more subtle. Recall that in a diploid cell there are two, separate copies of each chromosome (Chapter 1). We refer to these as pairs of homologous chromosomes. During mitosis, homologous chromosomes remain separate from one another, so each member of the pair replicates and is passed to a daughter nucleus independently of its homolog. In meiosis, however, the pairs of homologous chromosomes are by no means independent. During the initial stage of meiosis, called prophase I, each chromosome lines up with its homolog to form a bivalent (Figure 3.17). This occurs after each chromosome has replicated but before the replicated structures split, so the bivalent in fact contains four chromosome copies, each of which is destined to find its way into one of the four gametes that will be produced at the end of meiosis. Within the bivalent, the chromosome arms (chromatids) can undergo physical breakage and exchange of segments of DNA. The process is called crossing over, or recombination, and was discovered by the Belgian cytologist Janssens in 1909. This was just two years before Morgan started to think about partial linkage.
How did the discovery of crossing over help Morgan explain partial linkage? To understand this, we need to think about the effect that crossing over can have on the inheritance of genes. Let us consider two genes, each of which has two alleles. We will call the first gene A and its alleles A and a, and the second gene B with alleles B and b. Imagine that the two genes are located on chromosome number 2 of Drosophila melanogaster, the species of fruit fly studied by Morgan. We are going to follow the meiosis of a diploid nucleus in which one copy of chromosome 2 has alleles A and B, and the second has a and b. This situation is illustrated in Figure 3.18. Consider the two alternative scenarios:

- A crossover does not occur between genes A and B. In this case, two of the resulting gametes will contain chromosome copies with alleles A and B, and the other two will contain a and b. In other words, two of the gametes have the genotype AB and two have the genotype ab.

- A crossover does occur between genes A and B. This leads to segments of DNA containing gene A being exchanged between homologous chromosomes. The eventual result is that each gamete has a different genotype: one AB, one aB, one Ab, and one ab.

Now think about what would happen if we looked at the results of meiosis in 100 identical cells. If crossovers never occur, then the resulting gametes will have the following genotypes: 200 AB and 200 ab. This is complete linkage: genes A and B behave as a single unit during meiosis. But if (as is more likely) crossovers occur between A and B in some of the nuclei, then the allelic pairs will not be inherited as single units. Let us say that crossovers occur during 40 of the 100 meioses. The following gametes will result: 160 AB, 160 ab, 40 Ab, and 40 aB. The linkage is not complete; it is only partial. As well as the two parental genotypes (AB and ab), we see gametes with recombinant genotypes (Ab and aB).
Figure 3.18 The effect of a crossover on linked genes. The drawing shows a pair of homologous chromosomes, one red and the other blue. A and B are linked genes with alleles A, a, B, and b. On the left is meiosis with no crossover between A and B. Two of the resulting gametes have the genotype AB and the other two are ab. On the right, a crossover occurs between A and B. The four resulting gametes display all four of the possible genotypes—AB, ab, Ab, and aB.

From partial linkage to genetic mapping

Once Morgan understood how partial linkage could be explained by crossing over during meiosis, he was able to devise a way of mapping the relative positions of genes on a chromosome. In fact, the most important work was done not by Morgan himself but by an undergraduate in his laboratory, Arthur Sturtevant. Sturtevant assumed that crossing over was a random event, with an equal chance of it occurring at any position along a pair of lined-up chromatids. If this assumption is correct, then two genes that are close together will be separated by crossovers less frequently than two genes that are more distant from one another. Furthermore, the frequency with which the genes are unlinked by crossovers will be directly proportional to how far apart they are on the chromosome. The recombination frequency is therefore a measure of the distance between two genes. If you work out the recombination frequencies for different pairs of genes, you can construct a map of their relative positions on the chromosome.

The first map that Sturtevant constructed showed the positions of four genes on chromosome 1 of the fruit fly (Figure 3.19). Morgan’s group then set about mapping as many fruit fly genes as possible, and by 1915 they had assigned locations for 85 of them. These genes fall into four linkage groups, corresponding to the four pairs of chromosomes seen in the fruit fly nucleus. The distances between genes are expressed in map units; one map unit is the distance between two genes that recombine with a frequency of 1%. According to this notation, the distance between the genes for white eyes and yellow body, which recombine with a frequency of 1.3%, is 1.3 map units (see Figure 3.19). More recently the name centiMorgan (cM) has begun to replace the map unit. Each of the 85 genes initially mapped by Morgan specified a phenotype, such as eye color or wing or body shape, that could be typed simply by examining the fruit flies obtained as a result of the genetic crosses. The technique is equally efficacious with genes that are typed by biochemical tests and with DNA markers such as RFLPs, SSLPs, and SNPs, whose alleles are identified by PCR or some other type of DNA analysis (Section 3.2). Linkage analysis can therefore be used with many different types of organisms, as we will see in the next section, and the resulting maps can show the positions of many different types of markers.

Before moving on, there is one final issue relating to the basic principles of linkage analysis that we must consider. It turns out that Sturtevant’s assumption

![Diagram of meiosis and crossover](image)

**Figure 3.19 Working out a genetic map from recombination frequencies.** The example is taken from the original experiments carried out with fruit flies by Arthur Sturtevant. All four genes are on chromosome 1 of the fruit fly. Recombination frequencies between the genes are shown, along with their deduced map positions.

<table>
<thead>
<tr>
<th>Genes</th>
<th>Phenotypes</th>
</tr>
</thead>
<tbody>
<tr>
<td>m</td>
<td>Miniature wings</td>
</tr>
<tr>
<td>v</td>
<td>Vermilion eyes</td>
</tr>
<tr>
<td>w</td>
<td>White eyes</td>
</tr>
<tr>
<td>y</td>
<td>Yellow body</td>
</tr>
</tbody>
</table>

**Recombination frequencies**

- Between m and v = 3.0%
- Between m and y = 33.7%
- Between v and w = 29.4%
- Between w and y = 1.3%

**Deduced map positions**

<table>
<thead>
<tr>
<th>y</th>
<th>w</th>
<th>v</th>
<th>m</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1.3</td>
<td>30.7</td>
<td>33.7</td>
</tr>
</tbody>
</table>
about the randomness of crossovers was not entirely justified. Comparisons between genetic maps and the actual positions of markers on DNA molecules, as revealed by physical mapping and DNA sequencing, have shown that some regions of chromosomes, called recombination hotspots, are more likely to be involved in crossovers than others. This means that a genetic map distance does not necessarily indicate the physical distance between two markers (see Figure 3.26). Also, we now realize that a single chromatid can participate in more than one crossover at the same time but that there are limitations on how close together these crossovers can be, leading to more inaccuracies in the mapping procedure. Despite these qualifications, linkage analysis usually makes correct deductions about marker order, and distance estimates are sufficiently accurate to generate genetic maps that are of value as frameworks for genome sequencing projects and for use in techniques such as marker-assisted selection. We will therefore move on to consider how linkage analysis is carried out with different types of organisms.

3.4 LINKAGE ANALYSIS WITH DIFFERENT TYPES OF ORGANISMS

To see how linkage analysis is actually carried out, we need to consider three quite different situations:

- Linkage analysis with species such as fruit flies and mice, with which we can carry out planned breeding experiments
- Linkage analysis with humans, with whom we cannot carry out planned experiments but instead can make use of family pedigrees
- Linkage analysis with bacteria, which do not undergo meiosis

Linkage analysis when planned breeding experiments are possible

The first type of linkage analysis that we will study is the modern counterpart of the method developed by Morgan and his colleagues. The method is based on analysis of the progeny of experimental crosses set up between parents of known genotypes and is, at least in theory, applicable to all eukaryotes. Ethical considerations preclude this approach with humans, and practical problems such as the length of the gestation period and the time taken for the neonate to reach maturity (and hence to participate in subsequent crosses) limit the effectiveness of the method with some animals and plants.

If we return to Figure 3.18, we see that the key to genetic mapping is being able to determine the genotypes of the gametes resulting from meiosis. In a few situations this is possible by directly examining the gametes. For example, the gametes produced by some microbial eukaryotes, including the yeast Saccharomyces cerevisiae, can be grown into colonies of haploid cells. The genotypes of these haploid colonies can then be identified by biochemical tests and by DNA marker typing. Direct genotyping of gametes is also possible with higher eukaryotes if DNA markers are used, as PCR can be carried out with the DNA from individual spermatozoa, enabling RFLPs, SSLPs, and SNPs to be typed. Unfortunately, sperm typing is laborious. Routine linkage analysis with higher eukaryotes is therefore carried out not by examining the gametes directly but by determining the genotypes of the diploid progeny that result from fusion of two gametes, one from each of a pair of parents. In other words, a genetic cross is performed.

The complication with a genetic cross is that the resulting diploid progeny are the product not of one meiosis but of two (one in each parent), and in most organisms, crossover events are equally likely to occur during production of male and female gametes. Somehow we have to be able to disentangle from the genotypes of the diploid progeny the crossover events that occurred in each of these two meioses. This means that the cross has to be set up with care. The standard procedure is to use a test cross. This is illustrated in Figure 3.20, where we have set up a

Figure 3.20 A test cross between alleles displaying dominance and recessiveness. A and B are markers with alleles A, a, B, and b. The resulting progeny are typed by examining their phenotypes. Because the double-homozygous parent (Parent 2) has both recessive alleles, a and b, it effectively makes no contribution to the phenotypes of the progeny. The phenotype of each individual in the F1 generation is therefore the same as the genotype of the gamete from Parent 1 that gave rise to that individual.
test cross to map the two markers we met earlier: A (alleles A and a) and B (alleles B and b), both on chromosome 2 of the fruit fly. The critical feature of a test cross is the genotypes of the two parents:

- One parent is a double heterozygote. This means that all four alleles are present in this parent: its genotype is AB/ab. This notation indicates that one of the pair of homologous chromosomes has alleles A and B, and the other has a and b. Double heterozygotes can be obtained by crossing two pure-breeding strains, for example, AB/AB × ab/ab.

- The second parent is a pure-breeding double homozygote. In this parent, both homologous copies of chromosome 2 are the same: in the example shown in Figure 3.20, both chromosome copies have alleles a and b and the genotype of the parent is ab/ab.

The double heterozygote has the same genotype as the cell whose meiosis we followed in Figure 3.18. Our objective, therefore, is to infer the genotypes of the gametes produced by this parent and to calculate the fraction that are recombinants. Note that all the gametes produced by the second parent (the double homozygote) will have the genotype ab regardless of whether they are parental or recombinant gametes. Alleles a and b are both recessive, so meiosis in this parent is, in effect, invisible when the phenotypes of the progeny are examined. This means that, as shown in Figure 3.20, the phenotypes of the diploid progeny can be unambiguously converted into the genotypes of the gametes from the double-heterozygous parent. The test cross therefore enables us to make a direct examination of a single meiosis and hence to calculate a recombination frequency and map distance for the two markers being studied.

The power of this type of linkage analysis is enhanced if more than two markers are followed in a single cross. This not only generates recombination frequencies more quickly but also enables the relative order of markers on a chromosome to be determined by simple inspection of the data. This is because two recombination events are required to unlink the central marker from the two outer markers in a series of three, whereas either of the two outer markers can be unlinked by just a single recombination (Figure 3.21). A double recombination is less likely than a single one, so unlinking of the central marker will occur relatively infrequently. A set of typical data from a three-point cross is shown in Table 3.2. A test cross has been set up between a triple heterozygote (ABC/abc) and a triple homozygote (abc/abc). The most frequent progeny are those with one of the two parental genotypes, resulting from an absence of recombination events in the region containing the markers A, B, and C. Two other classes of progeny are relatively frequent (51 and 63 progeny in the example shown). Both of these are presumed to arise from a single recombination. Inspection of their genotypes shows that, in the first of these two classes, marker A has become unlinked from B and C, and in the second class, marker B has become unlinked from A and C. The implication is that A and B are the outer markers. This is confirmed by the number of progeny in which marker C has become unlinked from A and B. There are only two of these, showing that a double recombination is needed to produce this genotype. Marker C is therefore between A and B.

### Table 3.2 Set of Typical Data from a Three-Point Test Cross

<table>
<thead>
<tr>
<th>Genotypes of progeny</th>
<th>Number of progeny</th>
<th>Inferred recombination events</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABC/abc, abc/abc</td>
<td>987</td>
<td>None (parental genotype)</td>
</tr>
<tr>
<td>aBC/abc, Abc/abc</td>
<td>51</td>
<td>One, between A and B/C</td>
</tr>
<tr>
<td>AbC/abc, aBc/abc</td>
<td>63</td>
<td>One, between B and A/C</td>
</tr>
<tr>
<td>ABC/abc, aBC/abc</td>
<td>2</td>
<td>Two, one between C and A and one between C and B</td>
</tr>
</tbody>
</table>

**Figure 3.21** Effects of crossovers during a trihybrid cross. Either of the two outer markers can be unlinked by just a single recombination event, but two recombinations are required to unlink the central marker from the two outer markers.
Just one additional point needs to be considered. If, as shown in Figure 3.20 and Table 3.2, markers whose alleles display dominance and recessiveness are examined in a test cross, then the double- or triple-homozygous parent must have alleles for the recessive phenotypes. If, on the other hand, co-dominant markers are used, then the double-homozygous parent can have any combination of homozygous alleles \((AB/AB, Ab/Ab, aB/aB, or ab/ab)\). Figure 3.22, which gives an example of this type of test cross, shows the reason for this. Note that DNA markers typed by PCR display what is, in effect, co-dominance: Figure 3.22 therefore shows a typical scenario encountered when linkage analysis is being carried out with DNA markers.

**Gene mapping by human pedigree analysis**

With humans it is, of course, impossible to preselect the genotypes of parents and set up crosses designed specifically for mapping purposes. Instead, data for the calculation of recombination frequencies have to be obtained by examining the genotypes of the members of successive generations of existing families. This is called **pedigree analysis**. Often, only limited data are available, and their interpretation is often difficult because a human pairing rarely results in a convenient test cross, and often the genotypes of one or more family members are unobtainable because those individuals are dead or unwilling to cooperate.

The problems are illustrated by Figure 3.23. In this example, we are studying a genetic disease present in a family of two parents and six children. Genetic diseases are frequently used as gene markers in humans, the disease state being one allele and the healthy state being a second allele. The pedigree in Figure 3.23A shows us that the mother is affected by the disease, as are four of her children. We know from family accounts that the maternal grandmother also suffered from this disease, but both she and her husband, the maternal grandfather, are now dead. We can include them in the pedigree, with slashes indicating that they are dead, and we cannot obtain any further information on their genotypes. We know that the disease gene is present on the same chromosome as a microsatellite, which we call \(M\), four alleles of which, \(M_1, M_2, M_3,\) and \(M_4\), are present in the living family members. Our aim is to map the position of the disease gene relative to the microsatellite.

To establish a recombination frequency between the disease gene and microsatellite \(M\), we must determine how many of the children are recombinants. If we look at the genotypes of the six children, we see that children 1, 3, and 4 have the disease allele and the microsatellite allele \(M_1\). Children 2 and 5 have the healthy allele and the microsatellite allele \(M_2\). We can therefore construct two alternative hypotheses. One hypothesis is that the two copies of the relevant homologous chromosomes in the mother have the genotypes disease-\(M_1\) and healthy-\(M_2\); therefore, children 1–5 have parental genotypes and child 6 is the one and only recombinant (Figure 3.23B). This would suggest that the disease gene and the microsatellite are relatively closely linked and that crossovers between them occur infrequently. The alternative hypothesis is that the mother’s chromosomes have the genotypes healthy-\(M_1\) and disease-\(M_2\); this would mean that children 1–5 are recombinants and child 6 has the parental genotype. This would mean that the gene and microsatellite are relatively far apart on the chromosome. We cannot determine which of these hypotheses is correct: the data are frustratingly ambiguous.

The most satisfying solution to the problem posed by the pedigree in Figure 3.23 would be to know the genotype of the grandmother. Let us pretend that this is a soap-opera family and that the grandmother is not really dead. To everyone’s surprise, she reappears just in time to save the declining audience ratings. Her genotype for microsatellite \(M\) turns out to be \(M_1M_5\) (Figure 3.23C). This tells us that the chromosome inherited by the mother has the genotype disease-\(M_1\). We can therefore conclude with certainty that Hypothesis 1 is correct and that only child 6 is a recombinant.
Resurrection of key individuals is not usually an option open to real-life geneticists, although DNA can be obtained from old pathology specimens such as slides and Guthrie cards, the latter of which contain blood samples from newborn children. Imperfect pedigrees are analyzed statistically, by use of a measure called the lod score. This stands for logarithm of the odds that the genes are linked, and it is used primarily to determine if the two markers being studied lie on the same chromosome; in other words if the genes are linked or not. A lod score of 3 or more corresponds to odds of 1000:1 and is usually taken as the minimum for confidently concluding that this is the case. If the lod analysis establishes linkage, then additional lod scores can be calculated for each of a range of recombination frequencies, in order to identify the frequency most likely to have given rise to the data obtained by pedigree analysis. Ideally the available data will derive from more than one pedigree, increasing confidence in the result. The analysis is less ambiguous for families with larger numbers of children, and as we saw in Figure 3.23, it is important that the members of at least three generations can be genotyped. For this reason, family collections have been established, such as the one maintained by the Centre d’Études du Polymorphisme Humaine (CEPH) in Paris. The CEPH collection contains cultured cell lines from families in which all four grandparents as well as at least eight second-generation children could be sampled. This collection is available for DNA marker mapping by any researcher who agrees to submit the resulting data to the central CEPH database.
Genetic mapping in bacteria

The final type of genetic mapping that we must consider is the strategy used with bacteria. The main difficulty that geneticists faced when trying to develop genetic mapping techniques for bacteria is that these organisms are normally haploid, and so they do not undergo meiosis. Some other way therefore had to be devised to induce crossovers between homologous segments of bacterial DNA. The answer was to make use of three natural methods that exist for transferring pieces of DNA from one bacterium to another (Figure 3.24):

- In **conjugation**, two bacteria come into physical contact and one bacterium (the donor) transfers DNA to the second bacterium (the recipient). The transferred DNA can be a copy of some or possibly all of the donor cell’s chromosome, or it could be a segment of chromosomal DNA, up to $1 \times 10^6$ base pairs or 1 megabase (Mb) in length, integrated in a plasmid. The latter is called **episome transfer**.

- **Transduction** involves transfer of a small segment of DNA, up to 50 kb or so, from donor to recipient via a bacteriophage.

- In **transformation**, the recipient cell takes up from its environment a fragment of DNA, rarely longer than 50 kb, released from a donor cell.

Biochemical markers are often used, where the dominant or **wild-type** phenotype is possession of a biochemical characteristic (for example, the ability to synthesize tryptophan) and the recessive phenotype is the complementary characteristic (for example, the inability to synthesize tryptophan). The DNA transfer is usually set up between a donor strain that possesses the wild-type allele and a recipient with the recessive allele, and transfer into the recipient strain is monitored by looking for acquisition of the biochemical function specified by the gene being studied. This is illustrated in Figure 3.25A, where we see a functional gene for tryptophan biosynthesis being transferred from a wild-type bacterium (genotype described as $trp^+$) to a recipient that lacks a functional copy of this gene ($trp^-$). The recipient is called a tryptophan **auxotroph**, the term used to describe a mutant bacterium that can survive only if provided with a nutrient (in this case, tryptophan) not required by the wild type. After transfer, two crossovers are needed to integrate the transferred gene into the recipient cell’s chromosome, converting the recipient from $trp^-$ to $trp^+$.

The precise details of the mapping procedure depend on the type of gene transfer that is being used. During conjugation, DNA is transferred from donor to recipient in the same way that a string is pulled through a tube. The relative positions of markers on the DNA molecule can therefore be mapped by determining the times at which the markers appear in the recipient cell. In the example shown in Figure 3.25B, markers A, B, and C are transferred 8, 20, and 30 minutes after the beginning of conjugation, respectively. The entire Escherichia coli chromosome takes approximately 100 minutes to transfer. In contrast, transduction and transformation mapping enable markers that are relatively close together to be

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**Figure 3.24** Three ways of achieving DNA transfer between bacteria.

- **Conjugation** can result in transfer of chromosomal or plasmid DNA from the donor bacterium to the recipient. Conjugation involves physical contact between the two bacteria, with transfer thought to occur through a narrow tube called the **pilus**. **Transduction** is the transfer of a small segment of the donor cell’s DNA via a bacteriophage. **Transformation** is similar to transduction, but naked DNA is transferred. The events illustrated in panels B and C are often accompanied by death of the donor cell. In transduction (B), death occurs when the bacteriophages emerge from the donor cell. In transformation (C), release of DNA from the donor cell is usually a consequence of the cell’s death through natural causes.
mapped, because the transferred DNA segment is short (<50 kb), so the probability of two markers being transferred together depends on how close together they are on the bacterial chromosome (Figure 3.25C).

The limitations of linkage analysis

As we have seen, the data for linkage analysis are obtained by experimental utilization of the natural biological processes that govern inheritance of markers during meiosis and transfer of DNA between non-meiotic species such as bacteria. It is difficult, if not impossible, to modify these processes to improve the quality of the resulting data. Meiosis, for example, is a complex cellular pathway (see Figure 3.17), and there is little that a geneticist can do to alter the pathway in a particular organism to increase the accuracy or degree of detail in a genetic map. This means that linkage analysis has natural limitations that influence the utility of the resulting map. The following are the two most important limitations:

- The resolution of a genetic map depends on the number of crossovers that have been scored. This is not a major problem for microorganisms because they can be obtained in huge numbers, enabling many crossovers to be studied, resulting in a highly detailed genetic map in which the markers are just a few kilobases apart. For example, when the Escherichia coli genome sequencing project began in 1990, the latest genetic map for this organism comprised over 1400 markers, an average of one per 3.3 kb. This was sufficiently detailed to ensure that the genome sequence was assembled correctly. Similarly, the Saccharomyces cerevisiae project was supported by a fine-scale genetic map (approximately 1150 markers, on average one per 10 kb). The problem with humans and most other eukaryotes is that it is simply not possible to obtain large numbers of progeny, so relatively few meioses can be studied and the resolving power of linkage analysis is restricted. This means markers that are several tens of kilobases apart may appear at the same position on the genetic map.

- Genetic maps have limited accuracy. We touched on this point in Section 3.3 when we assessed Sturtevant’s assumption that crossovers occur at random along chromosomes. This assumption is only partly correct because the presence of recombination hotspots means that crossovers are more likely to occur at some points than at others. The effect that this can have on
the accuracy of a genetic map was illustrated in 1992, when the complete sequence for *Saccharomyces cerevisiae* chromosome III was published, enabling the first direct comparison to be made between a genetic map and the actual positions of markers as shown by DNA sequencing (Figure 3.26). There were considerable discrepancies, even to the extent that one pair of genes had been ordered incorrectly by genetic analysis. Bear in mind that *S. cerevisiae* is one of the two eukaryotes (fruit fly is the second) whose genomes have been subjected to intensive genetic mapping. If the yeast genetic map is inaccurate, then how precise are the genetic maps of organisms subjected to less-detailed analysis?

Because of these limitations, other ways of mapping markers onto chromosomes have been developed, with the aim of creating maps that have a greater density of markers with those markers more accurately positioned. These methods do not make use of linkage analysis and hence are not based on conventional genetic techniques. These alternative methods, which collectively make up the approach called physical mapping, are described in the next two sections.

### 3.5 Physical Mapping by Direct Examination of DNA Molecules

A plethora of physical mapping techniques have been developed, but these can conveniently be grouped into two categories depending on the approach used to identify marker locations:

- Methods that involve direct examination of DNA molecules or chromosomes
- Methods that assign markers to DNA fragments whose positions within an intact DNA molecule are known or can be inferred

The simplest direct examination methods are those that enable the positions of restriction sites to be located in a DNA molecule. This process is called restriction mapping.

**Conventional restriction mapping is applicable only to small DNA molecules**

Genetic mapping using RFLPs as DNA markers can locate the positions of polymorphic restriction sites within a genome (Section 3.2), but very few restriction sites are polymorphic, so many sites are not mapped by this technique (Figure 3.27). Could we increase the marker density on a genome map by using an alternative method to locate the positions of some of the nonpolymorphic restriction sites? This is what restriction mapping achieves.

The simplest way to construct a restriction map is to compare the fragment sizes produced when a DNA molecule is digested with two different restriction enzymes that recognize different target sequences. An example using the restriction enzymes EcoRI and BamHI is shown in Figure 3.28. This example illustrates the conventional approach to restriction mapping of a small DNA molecule, which involves carrying out three types of digests:

- First, a sample of the DNA molecule is digested with just one of the enzymes and the sizes of the resulting fragments are measured by agarose gel electrophoresis. Next, another sample of the molecule is digested with the second enzyme, and the resulting fragments are again sized in an agarose gel.
The results so far enable the number of restriction sites for each enzyme to be worked out but do not allow their relative positions to be determined.

- A sample of the DNA molecule is then cut with both enzymes together. In the example shown in Figure 3.28, the fragment sizes produced by this double restriction enable three of the four restriction sites to be mapped.

- A partial restriction is carried out, in which a single enzyme is used but the digestion does not go to completion, because the reaction is incubated only for a short time or a suboptimal incubation temperature is used. Partial restriction leads to a complex set of products, the complete restriction products now being supplemented with partially restricted fragments.
that still contain one or more uncut restriction sites. The sizes of the partially restricted fragments enable the map to be completed.

The method described above will yield an unambiguous map if there are relatively few cut sites for the enzymes being used. However, as the number of cut sites increases, so also do the numbers of single-, double-, and partial-restriction products whose sizes must be measured and compared in order for the map to be constructed. Computer analysis can be brought into play, but problems still eventually arise. A stage will be reached when a digest contains so many fragments that individual bands merge on the agarose gel, increasing the chances of one or more fragments being measured incorrectly or missed entirely. If several fragments have similar sizes, then even if they can all be identified, it may not be possible to assemble them into an unambiguous map. The conventional approach to restriction mapping is therefore more applicable to small rather than large molecules, the upper limit depending on the frequency of the restriction sites in the molecule being mapped. In practice, if a DNA molecule is less than 50 kb in length, it is usually possible to construct a restriction map for a selection of enzymes with six-nucleotide recognition sequences. Fifty kilobases is far below the minimum size for bacterial or eukaryotic chromosomes, although it does cover a few viral and organelle genomes, and whole-genome restriction maps constructed in this way have indeed been important in directing sequencing projects with these small molecules. The approach is equally useful after bacterial or eukaryotic genomic DNA has been cloned, if the cloned fragments are less than 50 kb in length.

**Optical mapping can locate restriction sites in longer DNA molecules**

It is also possible to use methods other than electrophoresis to map restriction sites in DNA molecules. With the technique called optical mapping, restriction sites are located simply by observing the cut DNA molecules under a microscope. DNA in solution takes up a random coil configuration, with groups of molecules tending to clump together in masses. The key to optical mapping is therefore the ability to extend individual DNA molecules into a linear configuration so that the locations of cuts made by a restriction enzyme, as visualized by observation of the molecule, accurately reflect the positions of the restriction sites in the DNA sequence. If parts of the DNA molecule remain as a random coil or are not fully extended, then working out the actual distances between the restriction sites will be much more difficult. In the earliest form of optical mapping, the molecules were extended by a process called gel stretching. Chromosomal DNA was suspended in molten agarose and placed on a microscope slide held at a slight angle so that the agarose flowed slowly along the slide as it cooled and solidified. Under these conditions, the DNA molecules contained within the agarose line up and become extended (Figure 3.29A). The gel also contains the restriction enzyme, which can be activated by adding magnesium ions (all restriction enzymes require magnesium in order to work). The molecules are then visualized by adding a fluorescent dye, such as DAPI (4’,6-diamino-2-phenylindole dihydrochloride), which stains the DNA so that the fibers can be seen when the slide is examined under a high-power fluorescence microscope. The restriction sites in the extended molecules gradually become gaps as the degree of fiber extension is reduced by the natural springiness of the DNA, enabling the relative positions of the cuts to be recorded.

Gel stretching is relatively easy to carry out, but the distortions inherent in viewing the DNA fibers while they are contained in a gel droplet limits the degree of resolution that can be achieved with this approach. An alternative method for stretching out the molecules without the use of a gel is molecular combing. A silicone-coated coverslip is dipped into a solution of DNA, left for 5 minutes (during which time the DNA molecules attach to the coverslip by their ends), and then removed from the solution at a constant speed, typically 0.3 mm s\(^{-1}\) (Figure 3.29B). The force required to pull the DNA molecules through the meniscus causes them to line up. Once in the air, the surface of the coverslip dries, retaining the DNA molecules as an array of parallel fibers. With this method, restriction sites less than 800 bp apart can be visualized.
Optical mapping was first applied to large DNA fragments cloned in BAC vectors (Section 2.3). The feasibility of using this technique with genomic DNA was then established with studies of a 1 Mb chromosome of the malaria parasite *Plasmodium falciparum*, and with the two chromosomes and a plasmid of the bacterium *Deinococcus radiodurans*, which are 2.65, 0.41, and 0.18 Mb, respectively (see Table 8.2). Molecules over 1 Mb in length are difficult to purify and extend without accidental breakage, so most optical maps are built up from the data obtained from a series of overlapping fragments. The 2.65 Mb *D. radiodurans* chromosome, for example, was mapped from 157 fragments. This means that the procedure is labor-intensive as many separate observations have to be made, and the amount of labor increases disproportionately as the length of the starting molecule increases. Recent work has therefore focused on automating the procedure so that restriction sites can be mapped in many fragments in a high-throughput manner. These automated procedures make use of microfluidic devices that extend the molecules and then move them, one by one, past an optical detector. In some systems, a version of molecular combing is used to extend the molecules, but in others the molecules are partially extended by movement through a grid of electrodes and are then fully extended by being pushed, by the solvent flow, into a series of nanochannels that are only just wide enough for the linear molecules to squeeze through (Figure 3.30). Of course, this method will work only if cleavage of the DNA fragment by the restriction enzyme is delayed until the fragment enters the nanochannels. One way of achieving this outcome is to design the microfluidic architecture in such a way that a magnesium ion gradient is established within each nanochannel, so that the restriction enzyme is activated only when it...
enters the channel along with the DNA fragment. The restriction sites are therefore cut within the nanochannels, and the resulting gaps in the DNA fragment are immediately recorded by the detection system. These automated approaches for data generation, along with computer analysis of the resulting data, have greatly extended the scope of optical mapping, with maps of this type now available for a range of plant and animal genomes.

**Optical mapping can be used to map other features in a DNA molecule**

The realization during the 2000s that observation of extended DNA molecules is a feasible method for mapping restriction sites has led to the development of innovative versions of optical mapping, ones that enable markers other than restriction sites to be mapped. These modifications to optical mapping derive, in part, from parallel developments that were occurring in the use of a second technique for physical mapping of DNA molecules, called **fluorescent in situ hybridization (FISH)**.

As in optical mapping, FISH enables the position of a marker on a chromosome or extended DNA molecule to be directly visualized. The difference is that, with FISH, the marker is a DNA sequence contained in the DNA molecule, whose location is visualized by hybridization with a fluorescent DNA probe that is complementary to and hence binds to the marker sequence (Figure 3.31). The technique was first used in the 1980s with **metaphase chromosomes** (Section 7.1). These chromosomes, prepared from nuclei that are undergoing division, are highly condensed, with each chromosome in a set taking up a recognizable appearance, characterized by the position of its centromere and the banding pattern that emerges after the chromosome preparation is stained (see Figure 7.6). This type of FISH can therefore identify the position of a marker relative to the centromere and the chromosome bands, but cannot achieve any degree of high-resolution mapping, as two markers have to be at least 1 Mb apart to be resolved as separate hybridization signals. The main application of metaphase FISH has therefore been in determining on which chromosome a new marker is located and providing a rough idea of its map position, as a preliminary to finer-scale mapping by other methods.

During the 1990s, modified versions of FISH were developed in which the target material was not the metaphase structures but instead mechanically stretched chromosomes or ones prepared from the prophase or interphase stages of nuclear division, when the chromosomes are naturally more extended. Even with these innovations, markers closer than 25 kb cannot be resolved. To improve the resolution of FISH even further, it was therefore necessary to abandon intact chromosomes and instead use purified DNA. This approach, initially called **fiber-FISH**, is essentially a modified version of optical mapping, and is carried out with stretched DNA fragments in microfluidic devices with architectures similar to those described earlier. The advantage of fiber-FISH compared with restriction site mapping is that the probe can be designed to target any desired DNA sequence, so there is no limit to the types of marker that can be detected.

The main challenge presented by this type of optical mapping is ensuring that the probe remains attached to its specific position on the DNA fragment as the fragment is extended and passed through the microfluidic channels and past the detector. With a traditional hybridization probe, the target DNA must be at least partially denatured to expose a single-stranded region to which the probe anneals. The second DNA strand will then compete with the probe and possibly displace it, re-forming the double-stranded molecule. If this occurs prior to passage of the DNA past the detector, then no data will be obtained. One solution to

*Figure 3.31 Fluorescent in situ hybridization.* A sample of dividing cells is dried onto a microscope slide and treated with formamide so that the chromosomes become denatured but do not lose their characteristic metaphase morphologies. The position at which the probe hybridizes to the chromosomal DNA is visualized by detecting the fluorescent signal emitted by the labeled DNA.
this problem is to use a **peptide nucleic acid (PNA)** as the probe. This is a polynucleotide analog in which the sugar–phosphate backbone is replaced by amide bonds (Figure 3.32). The hybridization between a PNA probe and its target on a DNA molecule is more stable than the normal DNA–DNA interaction for two reasons. First, the stability of a DNA–DNA hybrid is weakened, to some extent, by repulsion between the negatively charged sugar–phosphate backbones of the two polynucleotides. The amide backbone of a PNA is uncharged, so this repulsion does not occur. Second, there are two different ways in which a PNA can base-pair to its target. As well as attaching via standard Watson–Crick base pairs, a PNA with a high pyrimidine content can also form **Hoogsteen base pairs** with the target. Hoogsteen base pairs involve the same combinations (A-T and G-C) as Watson–Crick base pairs, but the hydrogen bonding that holds the pairs together involves different groups on the purine and pyrimidine bases (Figure 3.33). This means that a single DNA strand can attach to two PNAs at the same time, one by Watson–Crick pairing and one by Hoogsteen pairing. The resulting **triplex** structure, PNA₂DNA, is more stable than a DNA–DNA hybrid and so is unlikely to break down during the optical mapping procedure.

There are two other innovations of optical mapping that we should briefly consider while we are exploring this method. These innovations are not directly relevant to the identification of marker positions on a genome map, but nonetheless they provide important information on the structure and expression of a genome:

- **GC-rich regions** can be located by partially denaturing the DNA fragments, which can be achieved by raising the temperature or including a chemical denaturant such as formamide in the microfluidic solvent. Because a G-C base pair has three hydrogen bonds, compared to just two for A-T, the GC-rich regions are more likely to remain double-stranded under these conditions. If a double-strand-specific dye is added, then the GC-rich regions will be revealed. In some genomes, GC-rich regions indicate the locations of genes, so the data provided by this method can be useful when a genome sequence is being annotated.

- Patterns of DNA methylation can be assessed if optical mapping is carried out with a restriction enzyme that is unable to cut the DNA when its restriction site is methylated. As we will see in Section 10.3, the addition of methyl groups to certain nucleotides is one way in which a gene can be silenced. By identifying whether a particular region of the genome is methylated or unmethylated, it is therefore possible to infer if the genes in that region are inactive or are being expressed.

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**Figure 3.32** A short stretch of peptide nucleic acid. A peptide nucleic acid has an amide backbone instead of the sugar–phosphate structure found in a standard nucleic acid.

**Figure 3.33** Hoogsteen base pairs. This type of base pairing can form between PNA and DNA strands.
3.6 PHYSICAL MAPPING BY ASSIGNING MARKERS TO DNA FRAGMENTS

The second approach to physical mapping involves assigning markers to genome fragments, on the basis that two markers that occur in the same fragment must be located close to one another in the genome. In this method, each marker is called a sequence-tagged site (STS), an STS simply being a short DNA sequence, generally between 100 and 500 bp in length, that is easily recognizable and occurs only once in the chromosome or genome being studied. To map a set of STS markers, a collection of overlapping DNA fragments from a single chromosome or from the entire genome is needed. In the example shown in Figure 3.34, a fragment collection has been prepared from a single chromosome, with each point along the chromosome represented, on average, five times in the collection. The data from which the map will be derived are obtained by determining which fragments contain which STS markers. This can be done by hybridization analysis, but PCR is generally used because it is quicker and has proven to be more amenable to automation. The chances of two STS markers being present on the same fragment will depend on how close together they are in the genome. If they are very close, then there is a good chance that they will always be on the same fragment; if they are further apart, then sometimes they will be on the same fragment and sometimes they will not. The data can therefore be used to calculate the distance between two markers, in a manner analogous to the way in which map distances are determined by linkage analysis (Section 3.4). Remember that in linkage analysis a map distance is calculated from the frequency at which crossovers occur between two markers. STS mapping is essentially the same, except that the map distance is based on the frequency at which breaks occur between two markers.

This description of STS mapping leaves out some critical questions: What exactly is an STS? How is the DNA fragment collection obtained?

Any unique sequence can be used as an STS

To qualify as an STS, a DNA sequence must satisfy two criteria. The first is that its sequence must be known, so that a PCR assay can be set up to test for the presence or absence of the STS on different DNA fragments. The second requirement is that the STS must have a unique location in the chromosome being studied, or in the genome as a whole if the DNA fragment set covers the entire genome. If the STS sequence occurs at more than one position, then the mapping data will be ambiguous. Care must therefore be taken to ensure that an STS does not include sequences found in repetitive DNA.

Figure 3.34 A fragment collection suitable for STS mapping. The fragments span the entire length of a chromosome, with each point on the chromosome present (on average) in five fragments. The two blue markers are close together on the chromosome map and there is a high probability that they will be found on the same fragment. The two green markers are more distant from one another and so are less likely to be found on the same fragment.
Chapter 3: Mapping Genomes

Section 2.1

DNA fragments for STS mapping can be obtained as radiation hybrids

The second component of an STS mapping procedure is the collection of DNA fragments spanning the chromosome or genome being studied. This collection is sometimes called the mapping reagent, and at present there are two ways in which it can be assembled: as a clone library and as a panel of radiation hybrids. We will consider radiation hybrids first.

A radiation hybrid is a cell or organism that contains fragments of chromosomes from a second organism. The technology was initially developed with human chromosomes, starting in the 1970s when it was discovered that exposure of human cells to X-ray doses of 3000–8000 rad causes the chromosomes to break up randomly into fragments, with larger X-ray doses producing smaller fragments (Figure 3.36A). This treatment is lethal for the human cells, but the chromosome fragments can be propagated if the irradiated cells are subsequently fused with nonirradiated cells of a hamster or other rodent. Fusion is stimulated either chemically, with polyethylene glycol, or by exposure to Sendai virus (Figure 3.36B). Not all of the hamster cells take up chromosome fragments, so a means of identifying the hybrids is needed. The routine selection process is to use a hamster cell line that is unable to make either thymidine kinase (TK) or

![Figure 3.35 One method for preparing cDNA. Most eukaryotic mRNAs have a poly(A) tail at their 3’-end (Section 1.2). This series of A nucleotides is used as the priming site for the first stage of cDNA synthesis, carried out by reverse transcriptase, a DNA polymerase that copies an RNA template (Section 2.1). The primer is a short, synthetic DNA oligonucleotide, typically 20 nucleotides in length and made up entirely of T's, known as an oligo(dT) primer. When the first-strand synthesis has been completed, the preparation is treated with ribonuclease H, which specifically degrades the RNA component of an RNA–DNA hybrid. Under the conditions used, the enzyme does not degrade all of the RNA, instead leaving short segments that prime the second DNA strand-synthesis reaction, this one catalyzed by DNA polymerase I. This polymerase possesses a 5’ → 3’ exonuclease activity (Section 2.1) and so is able to degrade the RNA primers and replace these with DNA, completing synthesis of the second strand of cDNA. These are easy criteria to satisfy, and STS markers can be obtained in many ways; the most common sources are expressed sequence tags (ESTs), SSLPs, and random genomic sequences:

- Expressed sequence tags are short sequences obtained by analysis of complementary DNA (cDNA) clones. Complementary DNA is prepared by converting an mRNA preparation into double-stranded DNA (Figure 3.35). Because the mRNA in a cell is derived from protein-coding genes, cDNAs and the ESTs obtained from them represent the genes that were being expressed in the cell from which the mRNA was prepared. ESTs are looked upon as a rapid means of gaining access to the sequences of important genes, and they are valuable even if their sequences are incomplete. An EST can also be used as an STS, if it comes from a unique gene and not from a member of a gene family in which all the genes have the same or very similar sequences.

- An SSLP, whose use in genetic mapping we examined in Section 3.2, can also act as an STS in physical mapping. SSLPs that have already been mapped by linkage analysis are particularly valuable, as they provide a direct connection between the genetic and physical maps.

- Random genomic sequences are obtained by sequencing random pieces of cloned genomic DNA or simply by downloading sequences that have been deposited in the databases. If these sequences contain SNPs that have already been mapped by linkage analysis, then again a direct connection can be made between the genetic and physical maps.
A clone library can be used as the mapping reagent

A preliminary to the sequencing of a large and complex genome is to break the genome or isolated chromosomes into fragments and to clone each one in a high-capacity vector such as a BAC (Section 2.3). This results in a clone library, a collection of DNA fragments, with an average size of several hundred kilobases. The fragments in the various clones form an overlapping series, which means that as well as supporting the sequencing work, the clone library can also be used as a mapping reagent in STS analysis.

A clone library can be prepared from genomic DNA, in which case it represents the entire genome, or a chromosome-specific library can be made if the starting DNA comes from just one type of chromosome. The latter is possible because individual chromosomes can be separated by flow cytometry. To carry out this technique, dividing cells (ones with condensed chromosomes) are carefully broken open so that a mixture of intact chromosomes is obtained. The chromosomes are then stained with a fluorescent dye. The amount of dye that a chromosome binds depends on its size, so larger chromosomes bind more dye and fluoresce more brightly than smaller ones. The chromosome preparation is diluted and passed through a fine aperture, producing a stream of droplets, each one containing a single chromosome (Figure 3.37). The droplets pass through a detector that measures the amount of fluorescence and hence identifies which droplets contain the particular chromosome being sought. An electric charge is applied to these drops and no others, enabling the droplets containing the desired chromosome hypoxanthine phosphoribosyltransferase (HPRT). Deficiencies in either of these two enzymes are lethal when the cells are grown in a medium containing a mixture of hypoxanthine, aminopterin, and thymidine (HAT medium). After fusion, the cells are placed in HAT medium. Those that grow are hybrid hamster cells that have acquired human DNA fragments that include genes for the human TK and HPRT enzymes, which are synthesized inside the hybrids, enabling these cells to grow in the selective medium. The treatment results in hybrid cells that contain a random selection of human DNA fragments inserted into the hamster chromosomes. Typically the fragments are 5–10 Mb in size, with each cell containing fragments equivalent to 15–35% of the human genome. The collection of cells is called a radiation hybrid panel and can be used as a mapping reagent in STS mapping, provided that the PCR assay used to identify the STS does not amplify the equivalent region of DNA from the hamster genome.

Radiation hybrid mapping was important in constructing the first physical maps of the human genome, with a panel of less than 200 hybrids enabling 41,000 STS markers to be mapped with 100 kb resolution. This means that if two markers are less than 100 kb apart, then they will appear to occupy the same position in the genome. This degree of resolution is much less than for optical mapping, with which it can be possible to distinguish markers less than 1 kb apart, but it is still satisfactory for the initial mapping of an unsequenced genome. Following the success of the approach with the human genome, radiation hybrid mapping was applied to other mammals and to non-mammalian species such as zebrafish and chicken. Some progress has also been made in adapting the technique for use with plants. For example, a barley radiation hybrid panel has been created by irradiating barley protoplasts, to fragment their chromosomes, and then fusing these cells with tobacco protoplasts. A cotton panel was generated by irradiating pollen of one species of cotton, *Gossypium hirsutum*, and then using this pollen to fertilize the related species *Gossypium barbadense*. Similar approaches are proving successful with wheat, a recent study with a panel of 115 radiation hybrids enabling 26,299 SNPs to be mapped onto the D genome with 249 kb resolution.
to be deflected and separated from the rest. What if two different chromosomes have similar sizes, as is the case with human chromosomes 21 and 22? These can usually be separated if the dye used is not one that binds nonspecifically to DNA but instead has a preference for AT- or GC-rich regions. Examples of such dyes are Hoechst 33258 and chromomycin A3, respectively. Two chromosomes that are the same size rarely have identical GC contents and so can be distinguished by the amounts of AT- or GC-specific dye that they bind.

Compared with radiation hybrid panels, clone libraries have one important advantage for STS mapping. This is the fact that the assembly of overlapping clones can be used as the base material for a lengthy, continuous DNA sequence, and the STS data can then be used to anchor this sequence precisely onto the physical map. If the STS markers also include SSLPs or SNPs that have been mapped by linkage analysis, then the DNA sequence, physical map, and genetic map can all be integrated.

**SUMMARY**

- Genome maps provide the framework for sequencing projects because they indicate the positions of genes and other recognizable features and hence enable the accuracy of an assembled DNA sequence to be checked.

- Genome maps are used in methods for identifying the functions of genes involved in disease and for identifying QTLs that control traits such as meat productivity in farm animals.

- In the first genetic maps, the markers were genes whose alleles could be distinguished because they gave rise to easily recognized phenotypes, such as different eye colors, or whose alleles could be distinguished by biochemical tests.

- Today, DNA markers are also extensively used for mapping these markers including restriction fragment length polymorphisms (RFLPs), simple sequence length polymorphisms (SSLPs), and single-nucleotide polymorphisms (SNPs), all of which can be typed quickly and easily by PCR.

- The relative positions of genes and DNA markers on chromosomes are determined by linkage analysis, which enables the recombination frequency between a pair of markers to be calculated, providing the data needed to deduce the relative positions of the markers on the genetic map.

- For many organisms, linkage analysis is carried out by following the inheritance of markers in planned breeding experiments, but this is not possible for humans. Instead, genetic mapping of the human genome depends on examination of marker inheritance in large families, a procedure called pedigree analysis.

- Genetic maps have relatively poor resolution and tend to be inaccurate and must be refined by physical mapping if the map is to be used in a genome sequencing project.

- The positions of restriction sites in a small DNA molecule can be determined by restriction mapping.
Optical mapping enables the positions of restriction sites and some other sequence features to be directly visualized in a long DNA molecule.

The most detailed physical maps are obtained by sequence-tagged site (STS) content mapping, which makes use of a mapping reagent, a collection of overlapping DNA fragments that span an entire chromosome or genome. The mapping reagent can be a library of clones or a radiation hybrid panel.

**SHORT ANSWER QUESTIONS**

1. Describe the past and current use of maps in genome sequencing projects.
2. Clearly explain the differences between a genetic and a physical map of a genome.
3. How has PCR made the analysis of RFLPs much faster and easier? What was required to map RFLPs prior to the utilization of PCR?
4. Explain why SNPs are now the most widely used type of DNA marker, and outline the various methods used to type SNPs.
5. How does the linkage between genes provide a critical component to genetic mapping? Describe how genetic maps of individual chromosomes are obtained in (A) fruit flies and (B) humans.
6. What are Mendel’s two Laws of Genetics? What component of genetic mapping is not covered by Mendel’s Laws?
7. Explain why a double homozygote is used for test crosses in linkage analysis experiments. Why is it preferable that the homozygote alleles be recessive for the traits being tested?
8. Outline the limitations of human pedigree analysis and describe how the impact of these limitations is minimized in actual pedigree studies.
9. Briefly describe the three methods used to obtain maps of bacterial genomes.
10. Describe the basis to optical mapping and explain why optical mapping has become important in genome studies.
11. How are radiation hybrids used in the construction of genome maps?
12. How would a scientist prepare a clone library of DNA from just a single chromosome?

**IN-DEPTH PROBLEMS**

1. What are the ideal features of a DNA marker that will be used to construct a genetic map? To what extent can RFLPs, SSLPs, or SNPs be considered ideal DNA markers?
2. Explore and assess the applications of DNA chip technology in biological research.
3. What features would be desirable for an organism that is to be used for extensive studies of heredity?
4. Will maps ever become entirely unnecessary in genome research?
5. Which is more useful—a genetic or a physical map?
FURTHER READING

Books on the history of genetics

Genetic and DNA markers

Linkage analysis

Restriction and optical mapping

Radiation hybrids
The ultimate objective of a genome project is the complete DNA sequence for the organism being studied. This chapter describes the techniques and research strategies that are used during the sequencing phase of a genome project, when this ultimate objective is being directly addressed. Techniques for sequencing DNA are clearly of central importance in this context, and we begin the chapter with a detailed examination of sequencing methodology. This methodology is of little value, however, unless the short sequences that result from individual sequencing experiments can be linked together in the correct order to give the master sequences of the chromosomes that make up the genome. The second part of this chapter therefore describes the strategies used to ensure that the master sequences are assembled correctly.

Over the years, a number of different methods for DNA sequencing have been developed, and others are likely to become important in the future. The techniques in use today can be divided into two categories:

- The **chain-termination method** (Section 4.1), which was first devised by Fred Sanger and colleagues in the mid-1970s
- **Next-generation sequencing** (Section 4.2), which is a collection of methods, each of which utilizes a **massively parallel** strategy in order to generate millions of sequences at the same time

### 4.1 Chain-termination Sequencing

Chain-termination sequencing was first introduced in the 1970s and gradually became the most popular of the various sequencing methods that were available at that time. All genome sequencing projects completed prior to the mid-2000s were carried out with the chain-termination method, including the Human Genome Project and projects for several other eukaryotes and many types of bacteria and archaea. Nowadays, genome projects rely much more on next-generation techniques, which enable vast amounts of sequence to be obtained much more rapidly, but the chain-termination method is still performed in most molecular biology labs as a means of sequencing short DNA molecules, such as PCR products and small inserts cloned in plasmid or bacteriophage vectors.

**Chain-termination sequencing in outline**

Chain-termination DNA sequencing is based on the principle that single-stranded DNA molecules that differ in length by just a single nucleotide can be separated from one another by **polyacrylamide gel electrophoresis**. If the electrophoresis is carried out in a capillary tube 50–80 cm in length, with a bore of 0.1 mm, then it is possible to resolve a family of molecules representing all lengths up to 1500 nucleotides, with the single-stranded molecules emerging one after another from the end of the capillary (Figure 4.1).

Chain-termination sequencing is carried out with a DNA polymerase, which makes copies of the DNA molecule that is being sequenced. The first step is to anneal a short oligonucleotide to the template DNA. This oligonucleotide
4.1 Chain-Termination Sequencing

Chapter 4: Sequencing Genomes

subsequently acts as the primer for synthesis of a new DNA strand that is complementary to the template (Figure 4.2A). The strand-synthesis reaction, which requires the four deoxynucleotide triphosphates (dNTPs: dATP, dCTP, dGTP, and dTTP) as substrates, would normally continue until several thousand nucleotides have been polymerized. This does not occur in a chain-termination sequencing experiment because, as well as the four deoxynucleotides, small amounts of the four dideoxynucleotide triphosphates (ddNTPs: ddATP, ddCTP, ddGTP, and ddTTP) are added to the reaction. Each of these dideoxynucleotides is labeled with a different fluorescent marker.

The polymerase enzyme does not discriminate between deoxy- and dideoxynucleotides, but once incorporated, a dideoxynucleotide blocks further strand synthesis.

Figure 4.1 Polyacrylamide gel electrophoresis in a capillary system can resolve single-stranded DNA molecules that differ in length by just one nucleotide. (A) The dimensions of a capillary gel used in chain-termination DNA sequencing. (B) Separation of DNA molecules of different lengths during electrophoresis.

Figure 4.2 Chain-termination DNA sequencing. (A) Chain-termination sequencing involves the synthesis of new strands of DNA that are complementary to a single-stranded template. (B) Strand synthesis does not proceed indefinitely because the reaction mixture contains small amounts of the four dideoxynucleotides, which block further elongation because they have a hydrogen atom rather than a hydroxyl group attached to the 3'-carbon. (C) Incorporation of ddATP results in chains that are terminated opposite Ts in the template. This generates the A family of terminated molecules. Incorporation of the other dideoxynucleotides generates the C, G, and T families.
4.1 Chain-Termination Sequencing

Chapter 4: Sequencing Genomes

... elongation because it lacks the 3'-hydroxyl group needed to form a connection with the next nucleotide (Figure 4.2B). Because the normal deoxynucleotides are also present, in larger amounts than the dideoxynucleotides, the strand synthesis does not always terminate close to the primer: in fact, several hundred nucleotides may be polymerized before a dideoxynucleotide is eventually incorporated. The result is a set of new molecules, all of different lengths and each ending in a dideoxynucleotide whose identity indicates the nucleotide, A, C, G, or T, that is present at the equivalent position in the template DNA (Figure 4.2C).

To determine the DNA sequence, all we have to do is identify the dideoxynucleotide at the end of each chain-terminated molecule. This is where the polyacrylamide gel comes into play. The DNA mixture is loaded onto the capillary gel, and electrophoresis is carried out to separate the molecules according to their lengths. After separation, the molecules are run past a fluorescence detector capable of discriminating the labels attached to the dideoxynucleotides (Figure 4.3A). The detector therefore determines whether each molecule ends in A, C, G, or T. The sequence can be printed out for examination by the operator (Figure 4.3B) or entered directly into a storage device for future analysis.

Not all DNA polymerases can be used for sequencing

Any template-dependent DNA polymerase is capable of extending a primer that has been annealed to a single-stranded DNA molecule, but not all polymerases do...
this in a way that is useful for DNA sequencing. Three criteria in particular must be fulfilled by a sequencing enzyme:

- **High processivity.** This refers to the length of polynucleotide that is synthesized before the polymerase terminates through natural causes. A sequencing polymerase must have high processivity so that it does not dissociate from the template before incorporating a dideoxynucleotide.

- **Negligible or zero 5’ → 3’ exonuclease activity.** Most DNA polymerases also have exonuclease activities, meaning that they can degrade DNA polynucleotides as well as synthesize them (Section 2.1). This is a disadvantage in DNA sequencing because removal of nucleotides from the 5’-ends of the newly synthesized strands alters the lengths of these molecules, making it impossible to determine the correct sequence.

- **Negligible or zero 3’ → 5’ exonuclease activity.** This is also desirable so that the polymerase does not remove the dideoxynucleotide at the end of a completed strand. If this happens, then the strand might be further extended. The net result will be that there are few short strands in the reaction mixture, and the sequence close to the primer will be unreadable.

These are stringent requirements that, in the early days of DNA sequencing, were met by using artificially modified enzymes. In the original method for chain-termination sequencing, the Klenow polymerase was used. This is a version of *Escherichia coli* DNA polymerase I from which the 5’ → 3’ exonuclease activity of the standard enzyme has been removed, either by cleaving away the relevant part of the protein or by genetic engineering (Section 2.1). The Klenow polymerase has relatively low processivity, limiting the length of sequence that can be obtained from a single experiment to about 250 bp, and it gives nonspecific products—strands that have terminated naturally rather than by incorporation of a dideoxynucleotide—in the sequencing reaction. To avoid these problems, most sequencing today makes use of the *Taq* DNA polymerase, which has high processivity and no exonuclease activity and so is ideal for chain-termination sequencing, enabling sequences of 750 bp and longer to be obtained in a single experiment.

**Chain-termination sequencing with *Taq* polymerase**

The chain-termination method that uses *Taq* polymerase is called **thermal cycle sequencing.** It is carried out in a similar way to PCR, but just one primer is used and the reaction mixture includes the four dideoxynucleotides (Figure 4.4). Because there is only one primer, only one strand of the starting molecule is copied, and the product accumulates in a linear fashion, not exponentially as is the case in a standard PCR. The presence of the dideoxynucleotides in the reaction mixture causes chain termination, as in the standard methodology, and the family of resulting strands can be analyzed and the sequence read in the usual way.

Thermal cycle sequencing is usually carried out with PCR products or with DNA that has been cloned in a plasmid or phage vector. If a PCR product is being sequenced, then one of the primers from the original PCR can be used in the sequencing reaction. If two separate reactions are carried out, one with each of the two PCR primers, then **forward** and **reverse sequences** are obtained (Figure 4.5A).

**Figure 4.4 Thermal cycle sequencing.** A reaction similar to PCR is carried out, with just one primer and with the four dideoxynucleotides present in the reaction mixture. The result is a set of chain-terminated strands, such as the A family in the part of the reaction shown here. These strands, along with the products of the C, G, and T reactions, are then electrophoresed and imaged in order to read the sequence.
This is an advantage if the PCR product is more than 750 bp and hence too long to be sequenced completely in one experiment. Alternatively, it is possible to extend the sequence in one direction by synthesizing a new primer designed to anneal at a position within the PCR product.

If cloned DNA is being sequenced, then a **universal primer** can be used. This is a primer that is complementary to the vector DNA immediately adjacent to the point into which new DNA is ligated (Figure 4.5B). Any fragment that is cloned in the vector can therefore be sequenced with the same primer. Once again, both forward and reverse universal primers can be used, enabling sequences to be obtained from both ends of the insert, and an internal primer can be used to provide sequence from the middle region of a long insert.

**Strengths and limitations of chain-termination sequencing**

During the late 1970s, two DNA sequencing methods were developed: the chain-termination method, which we have just studied, and **chemical degradation sequencing**. The latter took a radically different approach in which a double-stranded DNA molecule was treated with chemicals that cut the molecule at specific nucleotide positions. The lengths of the resulting fragments were then examined by polyacrylamide gel electrophoresis in order to deduce the sequence of nucleotides in the starting molecule. Both methods were equally popular to begin with, but the chain-termination procedure gained ascendancy, partly because the chemicals used in the chemical degradation method are toxic and therefore hazardous to the health of the researchers doing the sequencing experiments, but mainly because it was easier to automate chain-termination sequencing. Automated sequencers with multiple capillary gels working in parallel can read up to 384 different sequences in a one-hour period, which means that with an average of 750 bp per individual experiment, almost 7 Mb of information can be generated per machine in 24 hours. This, of course, requires round-the-clock technical support, ideally with robotic devices used to prepare the sequencing reactions and to load the reaction products into the sequencers.

If such a factory approach can be established and maintained, then the data needed to sequence an entire genome can be generated in a relatively short time. Chain-termination sequencing carried out in this intensive manner was used in the first genome sequencing projects and proved effective in particular with bacterial genomes, many of which are less than 5 Mb in length. For example, to sequence the *Haemophilus influenzae* genome (Section 4.3), 28,643 chain-termination experiments were carried out, generating 11.6 Mb of sequence. The length of the
Chapter 4: Sequencing Genomes

H. influenzae genome is 1.8 Mb, but no sequencing method is entirely accurate, so it is necessary to sequence each region of a genome multiple times, in order to identify errors present in individual sequence reads (Figure 4.6). With the chain-termination method, to ensure that errors are identified, at least \(5 \times\) sequence depth or coverage is needed, meaning that every nucleotide is present in five different reads. Despite this requirement, chain-termination sequencing was used to obtain the human genome sequence, as well as that of other eukaryotes such as the nematode worm Caenorhabditis elegans and the fruit fly Drosophila melanogaster. The human genome is 3.235 Mb, so for \(5 \times\) sequence depth, a total of \(5 \times 3.235 = 16,175\) Mb of sequence was required, equivalent to over 21.5 million chain-termination sequences averaging 750 bp in length. In fact this target was exceeded: the Human Genome Project generated 23,147 Mb of sequence by the time that the first draft of the genome was published in 2001 (Section 4.4).

Despite these successes, it was realized during the early 2000s that progress in studying genomes would be slow if sequencing remained dependent on the chain-termination method, even when multiple sequencing machines are operated in a factory setting. It would take months or years of effort to obtain enough data to assemble the genome sequence of each new species that was studied. Cost is also an important consideration, especially for genome resequencing, in which the sequences of multiple individuals within a species are obtained. Comparisons between these genome sequences enable variations to be identified that might be associated with characteristics such as inherited disease in humans or adaptation to environmental extremes for a crop plant. One of the goals of personalized medicine is to use individual human genome sequences to make accurate diagnoses of a person’s risk of developing a disease and to use that person’s genetic characteristics to plan effective therapies and treatment regimes. For personalized medicine and other research programs based on genome resequencing to become a reality, cost-effective and rapid ways of sequencing individual genomes are needed.

The chain-termination method is still used routinely to sequence small pieces of DNA, where its speed and ease of use outweigh other issues. For genome projects, chain-termination sequencing has now been almost entirely superseded by next-generation sequencing methods, which are able to generate much larger amounts of sequence data more quickly and at less cost.

4.2 NEXT-GENERATION SEQUENCING

Next-generation sequencing is the term applied to a variety of methods that enable thousands or millions of DNA fragments to be sequenced in parallel in a single experiment. The preparation and use of this sequencing library is the distinctive feature that distinguishes these methods from chain-termination sequencing, which is able to sequence only individual DNA fragments, each fragment obtained by a different PCR or from a different clone. Next-generation methods therefore enable the vast amounts of data needed to assemble an entire genome sequence to be obtained much more rapidly than with the chain-termination approach. At this scale there are also significant cost savings because a single sequencing
4.2 NEXT-GENERATION SEQUENCING

A machine can produce the required data in a single or small number of runs, compared to the multiple runs using factories of sequencers that are needed when the chain-termination method is used.

**Preparation of a sequencing library is the common feature of next-generation methods**

The common feature of the various next-generation sequencing methods is the prior preparation of a library of DNA fragments that have been immobilized on a solid support in such a way that multiple sequencing reactions can be carried out side by side in a **massively parallel array** format (Figure 4.7). The fragments are usually 100–500 bp in length, the precise sizes depending on the lengths of the individual sequences that can be obtained by the next-generation method that is being employed. The most popular way of breaking genomic DNA down into fragments of these sizes is by **sonication**, a technique that uses high-frequency sound waves to make random cuts in DNA molecules. Random breakage is important because each fragment will be sequenced from its ends. With next-generation methods it is not possible to direct the sequencing toward the middle of a fragment, as can be done by designing an internal primer for the chain-termination method. The ends must therefore be randomly distributed throughout the starting DNA molecule in order to ensure that the entire molecule is sequenced.

Two different immobilization methods are commonly used in next-generation sequencing. In the first method, the solid support is a glass slide that has been coated with many copies of a short oligonucleotide (Figure 4.8). **Adaptors**, short pieces of double-stranded DNA whose sequences match that of the oligonucleotide, are ligated to the ends of the DNA fragments, which are then denatured. The resulting single-stranded molecules attach to the glass slide by base pairing between their adaptor sequences and the immobilized oligonucleotides. In the second immobilization method, the solid support is provided by small metallic beads that are coated with the protein **streptavidin**. Once again, the DNA fragments are ligated to adaptors, in this case adaptors that carry a **biotin** label attached to their 5’-ends. Biotin is a small organic molecule that binds strongly to streptavidin, so the fragments become attached to the metallic beads by biotin–streptavidin linkages (Figure 4.9A). The ratio of DNA fragments to beads is set so that, on average, just one fragment becomes attached to each bead. The beads are then shaken in an oil–water mixture in order to generate an emulsion.

**Figure 4.7** A DNA library immobilized on a solid support. In reality, a single massively parallel array will have millions of components.

**Figure 4.8** Immobilization of DNA fragments in a sequencing library by base pairing to oligonucleotides on a glass slide. Base pairing between the immobilized oligonucleotides and the adaptors ligated to the ends of the DNA fragments results in attachment of the fragments to the slide.
conditions set so that there will be just one bead in each aqueous droplet within the emulsion (Figure 4.9B). Each aqueous droplet is then transferred into a different well in a multiple array on a plastic strip.

The final step in preparation of the library is amplification of the immobilized DNA fragments by PCR, to produce a sufficient number of identical copies to be sequenced. The adaptors now play a second role as they provide the annealing sites for the primers for this PCR. The same pair of primers can therefore be used to amplify all the fragments, even though the fragments themselves have many different sequences. With the glass slide method, the PCR products become attached to adjacent oligonucleotides, so each starting fragment is amplified into an immobilized cluster of identical fragments (Figure 4.10A). If metallic beads are used, then the PCR is carried out in the oil emulsion, so that the products of each PCR are retained within their own water droplet, prior to deposition of those droplets into the wells on the plastic strip (Figure 4.10B).

Figure 4.9 Immobilization of DNA fragments by base pairing to metallic beads. (A) Each DNA fragment is attached to a single bead via a streptavidin–biotin linkage. (B) Beads, with their attached DNA fragments, are shaken in an oil–water mixture in such a way that, within the resulting emulsion, each individual water droplet contains a single bead.

Figure 4.10 Amplification of immobilized libraries. (A) With the glass slide method, the fragment copies made by PCR attach to adjacent oligonucleotides, resulting in a cluster of identical immobilized fragments. (B) When metallic beads are used, PCR is carried out in the oil–water emulsion, and the fragment copies are retained in their individual water droplets.
Various next-generation sequencing methods have been devised

Over the last few years, there has been immense competition between different companies striving to develop next-generation sequencing platforms that combine the greatest speed in generating accurate sequence data with the lowest cost. Currently the most popular method is based on **reversible terminator sequencing** which, like the chain-termination method, makes use of modified nucleotides that block strand synthesis when incorporated at the end of a polynucleotide that is being synthesized by a DNA polymerase. The difference is that the termination step is reversible, because the chemical group attached to the 3'-carbon of the modified nucleotide can be removed once the identity of this nucleotide has been confirmed (Figure 4.11). In the simplest method, this removable blocking group is a fluorescent label, a different one for each of the four nucleotides. There are no normal deoxynucleotides present in the reaction mixture, so each step in strand synthesis is accompanied by a pause, during which an optical device detects the fluorescent label, thereby identifying the terminal nucleotide. An enzyme then removes the label, enabling the next terminator nucleotide to be added and the detection process to be repeated. When used in next-generation sequencing, the process is initiated by a primer that anneals to the adaptor sequences attached to the ends of the DNA fragments during library preparation. Every cluster of fragments in the library is therefore sequenced at the same time. The method generates relatively short sequence reads, with a maximum length of 300 bp, but is so massively parallel that up to 2000 Mb of sequence can be obtained per run. This technology is usually referred to as **Illumina sequencing**, named after the company that markets the necessary equipment.

Prior to the introduction of Illumina technology, the most popular next-generation method was based on an approach called **pyrosequencing**. In this method, the reaction mixture contains only deoxynucleotides, and template copying proceeds without artificial termination. Strand synthesis is followed by detecting flashes of chemiluminescence generated by the enzyme sulfurylase from the molecule of pyrophosphate that is released each time DNA polymerase adds a deoxynucleotide to the 3'-end of the growing strand. A flash of chemiluminescence therefore signals the successful copying of one position in the template molecule. Of course, if all four deoxynucleotides were added at once, then flashes of light would be seen all the time and no useful sequence information would be

(A) A reversible terminator nucleotide

(B) Reversible terminator sequencing

**Figure 4.11 Reversible terminator sequencing.** (A) The structure of a reversible terminator nucleotide with a removable fluorescent blocking group attached to the 3'-carbon. (B) After each nucleotide addition, there is a pause while the fluorescent label is detected and the terminal nucleotide is identified. The blocking group is then removed so that the next nucleotide addition can occur.
**Figure 4.12** Pyrosequencing. The strand-synthesis reaction is carried out in the absence of terminating nucleotides. Each deoxynucleotide is added individually in a repetitive series, along with a nucleotidase enzyme that degrades the deoxynucleotide if it is not incorporated into the strand being synthesized. Incorporation is detected by a flash of chemiluminescence induced by pyrophosphate release from the deoxynucleotide. The order in which deoxynucleotides are added to the growing strand can therefore be followed.

**Figure 4.13** The basis to SOLiD sequencing. The mixture of 5-mer oligonucleotides (oligonucleotides that are five nucleotides in length) includes all 1024 possible sequences. The oligonucleotide that is complementary to the template DNA immediately adjacent to the primer hybridizes and is attached to the primer by DNA ligase. The cycle then repeats, resulting in hybridization and ligation of the second oligonucleotide.

The deoxynucleotides are therefore added individually in a repetitive series (e.g. A, then T, then G, then C, then A, then T, etc.) and the pattern of light emissions is used to deduce the order in which nucleotides are incorporated into the growing strand (**Figure 4.12**). Pyrosequencing of fragment libraries immobilized on magnetic beads is the procedure used by the next-generation method called **454 sequencing**, the name once again referring to the company that initially developed this particular technology. In its most advanced format, 454 sequencing can give sequence reads up to 1000 bp in length, with enough reads to generate a total of 700 Mb of DNA sequence per run.

The **ion torrent** method uses a similar approach to pyrosequencing, with a repetitive series of nucleotides flowed over an immobilized fragment library. However, with this method, the detection system is directed at the hydrogen ions that, along with pyrophosphate, are released every time a nucleotide is incorporated into the growing strand. The reaction is carried out with DNA fragments immobilized on acrylamide beads, each bead in a well lined with an **ion-sensitive field effect transistor** (ISFET). The ISFET generates an electronic pulse each time it detects hydrogen ions, these pulses being related to the flow of nucleotides over the well in order to deduce the sequence of the immobilized fragments. Read lengths of up to 400 bp are possible, but the main advantage of this technology is the electronic detection system, which has lower construction and running costs compared with the optical detectors used in the Illumina and 454 platforms.

The fourth of the next-generation sequencing technologies, called **sequencing by oligonucleotide ligation and detection** (SOLiD), takes a radically different approach. The sequence is deduced not by polymerase-directed synthesis of a new DNA strand but by hybridization of a series of oligonucleotides whose sequences are complementary to that of the template. To begin the sequencing process, a primer is attached to the template DNA, using a terminal adaptor sequence as the annealing site, just as in the sequencing-by-synthesis methods. A set of 1024 oligonucleotides, representing each of the possible five-nucleotide sequences, is then added, along with a DNA ligase. The sequence of one of these oligonucleotides will be complementary to the template DNA immediately adjacent to the primer. This oligonucleotide will hybridize and be attached to the primer by DNA ligase (**Figure 4.13**). The process of hybridization–ligation continues for a set number of cycles until 50–75 nucleotides of the template have been covered. SOLiD sequencing is therefore quite straightforward in outline. However, it is computationally intensive because of the way in which the sequences of the oligonucleotides that hybridize to the template are deduced.
Each of these oligonucleotides is labeled with a fluorescent marker, but only four different markers are used in total. This means that the markers divide the 1024 oligonucleotides into four families of 256 sequences. The groupings are not random; instead each family comprises four dibase sets, with each dibase set comprising the 64 oligonucleotides with the same two initial nucleotides. For example, the AT dibase set includes all the oligonucleotides with sequence ATNNN, where N is any nucleotide. Detection of the marker attached to a hybridizing oligonucleotide therefore assigns a color to the terminal pair of nucleotides in that five-nucleotide sequence. Clearly this results in an incomplete sequence (only two of every five nucleotides are read) as well as one that is ambiguous (the same label is used for four different dibase sets). The sequencing process is therefore repeated with a second primer that anneals at a position offset by one nucleotide \((n − 1)\) compared to the first primer \((\text{Figure 4.14})\). This is followed by three further repeats with primers that anneal at the \(n − 2, n − 3,\) and \(n − 4\) positions. The result is that every nucleotide in the template is read twice, and the combination of colors assigned to that nucleotide enables its identity to be determined without ambiguity. Because of this double read, SOLiD sequencing is highly accurate, but the read lengths are shorter than for the other methods, typically 50 or 75 nucleotides depending on the number of hybridization–ligation cycles that are used. The computational issues are less severe if the expected sequence of the template is known, which means that the major application of SOLiD is not \(\textit{de novo}\) sequencing of genomic DNA fragments but rather genome resequencing, aimed at detecting polymorphisms in DNA samples from different individuals.

**Third- and fourth-generation methods enable sequencing in real time**

New DNA sequencing technologies are continually being developed with the aim of enabling genome sequences to be assembled more quickly and at less cost. One limitation of the three sequencing-by-synthesis methods described above is that detection of each nucleotide addition requires a brief delay in the DNA polymerization process. With the reversible terminator method, the delay is caused by the need to remove the 3’-blocking group after each nucleotide addition, and during pyrosequencing and ion torrent sequencing, the delay occurs because each nucleotide is presented individually to the polymerase. This delay increases the period of time needed to complete a sequence read and also decreases the processivity of the polymerase, limiting the lengths of those reads.

The recent focus has therefore been on methods that avoid a delay at the nucleotide detection step and enable a sequence to be read during the normal, unimpeded progression of the polymerase along the template. This is referred to as **third-generation sequencing**, or sequencing in real time. One of the most promising third-generation methods so far developed is **single-molecule real-time sequencing**. A sophisticated optical system called a **zero-mode waveguide** is used to observe the copying of a single DNA template \((\text{Figure 4.15})\). The nucleotide substrates are still labeled with fluorescent markers, but the optical system is so precise that there is no need to use a blocking group to delay the polymerization process to allow the detection to take place. Instead, the label is removed immediately after nucleotide incorporation, so strand synthesis progresses...
without interruption. Read lengths of up to 20,000 bp have been reported. This technology was initially developed by Pacific Biosciences and is often called PacBio sequencing.

The next logical step, fourth-generation sequencing, would be to dispense with the strand-synthesis step and read the sequence of a DNA molecule directly without copying that molecule in any way. This is the aim of nanopore sequencing. This method makes use of a synthetic membrane with small pores just large enough for a DNA molecule to pass through. An electrical current is set up, positive on one side of the membrane and negative on the other, so electrophoresis causes the DNA molecule to approach one of the nanopores. This molecule is double-stranded, but the presence of a helicase enzyme in the vicinity of the nanopore breaks the base pairs, so the DNA unwinds and just one strand passes through the pore (Figure 4.16). The sequence of this strand can be read because each of the four nucleotides has a different shape and so occludes the nanopore in a different way, resulting in a slightly different perturbation of the flow of ions passing through the membrane. These perturbations are measured in order to deduce the sequence of the polynucleotide. Because no synthesis is involved, the length of the sequence is not limited by polymerase processivity, and reads of up to 50 kb have been reported. At present the technology is limited because the accuracy of sequence identification is hampered by the speed at which the polynucleotide passes through the nanopore. Improvements in the detection systems are therefore being sought, along with ways of modifying the nanopore structure so that progress of the polynucleotide is slowed down.

4.3 HOW TO SEQUENCE A GENOME

Having understood the methods available for sequencing DNA, the next question we must address is how the multitude of short sequence reads generated by these methods are assembled into the sequence of an entire genome. Since the 1990s, when the chain-termination method was first automated, the actual generation of sequence data has not been a limiting factor in genome sequencing projects. Instead, the main challenge lies with sequence assembly, the procedure used to convert thousands or millions of short sequence reads into a contiguous genome sequence. The most straightforward approach to sequence assembly is to build...
up the master sequence directly from the short sequences obtained from individual sequencing experiments, simply by examining those sequences for overlaps (Figure 3.1). This is called the **shotgun method**.

The potential of the shotgun method was proven by the *Haemophilus influenzae* sequence

During the early 1990s, there was extensive debate about whether the shotgun method would work in practice. Many molecular biologists were of the opinion that the amount of data handling needed to compare all the sequence reads and identify overlaps, even with the smallest genomes, would be beyond the capabilities of existing computer systems. These doubts were laid to rest in 1995 when the sequence of the 1830 kb genome of the bacterium *Haemophilus influenzae* was published.

The strategy used to obtain the *H. influenzae* genome sequence is shown in Figure 4.17. The first step was to sonicate the genomic DNA in order to break it into fragments. The fragments were then electrophoresed and those in the size range 1.6–2.0 kb purified from the agarose gel and ligated into a plasmid vector. From the resulting library, 19,687 clones were taken at random and 28,643 chain-termination sequencing experiments were carried out. The number of sequencing experiments was greater than the number of plasmids because both ends of some inserts were sequenced. Of these sequencing experiments, 16% were considered to be failures because they resulted in less than 400 bp of sequence. The remaining 24,304 sequences gave a total of 11,631,485 bp, corresponding to six times the length of the *H. influenzae* genome. This amount of redundancy was deemed necessary to ensure complete coverage. Sequence assembly required 30 hours on a computer with 512 megabytes of random access memory (RAM) and resulted in 140 lengthy contiguous sequences, each of these **sequence contigs** representing a different, nonoverlapping portion of the genome.

The next step was to join up pairs of contigs by obtaining sequences from the gaps between them. First, the library was checked to see if there were any clones whose two end-sequences were located in different contigs. If such a clone could be identified, then additional sequencing of its insert would close the sequence gap between the two contigs (Figure 4.18A). In fact, there were 99 clones in this category, so 99 of the gaps could be closed without too much difficulty.

This left 42 gaps, which probably consisted of DNA sequences that were unstable in the cloning vector and therefore not present in the library. To close these physical gaps, a second clone library was prepared, this one with a different type of vector. Rather than using another plasmid, in which the uncloned sequences would probably still be unstable, the second library was prepared in a bacteriophage λ vector (Section 2.3). This new library was probed, one at a time, with 84 oligonucleotides that had sequences identical to the sequences at the ends of the unlinked contigs (Figure 4.18B). The rationale was that if two oligonucleotides hybridized to the same λ clone, then the ends of the contigs from which they were derived must lie within that clone, and sequencing the DNA in the λ clone would therefore close the gap. Twenty-three of the 42 physical gaps were dealt with in this way.

A second strategy for gap closure was to use pairs of oligonucleotides, from the set of 84 described above, as primers for PCR of *H. influenzae* genomic DNA. Some oligonucleotide pairs were selected at random, and those spanning a gap were identified simply from whether or not they gave a PCR product (see Figure 4.17).

**Figure 4.17 Shotgun sequencing of the *Haemophilus influenzae* genome.** *H. influenzae* DNA was sonicated, and fragments with sizes between 1.6 and 2.0 kb were purified from an agarose gel and ligated into a plasmid vector to produce a clone library. End-sequences were obtained from clones taken from this library, and a computer was used to identify overlaps between sequences. This resulted in 140 sequence contigs.
Figure 4.18 Methods used to close the gaps in the initial assembly of the *Haemophilus influenzae* genome sequence. (A) Sequence gaps are ones that can be closed by further sequencing of clones already present in the library. In this example, the end-sequences of contigs 1 and 2 lie within the same plasmid clone, so further sequencing of this DNA insert with internal primers will provide the sequence to close the gap. (B) Physical gaps are stretches of sequence that are not present in the clone library, probably because these regions are unstable in the cloning vector that was used. Two strategies for closing these gaps are shown. On the left, a second clone library, prepared with a bacteriophage λ vector rather than a plasmid vector, is probed with oligonucleotides corresponding to the ends of the contigs. Oligonucleotides 1 and 7 both hybridize to the same clone, whose insert must therefore contain DNA spanning the gap between contigs 1 and 4. On the right, PCR is carried out with pairs of oligonucleotides. Only numbers 1 and 7 give a PCR product, confirming that the contig ends represented by these two oligonucleotides are close together in the genome. The PCR product or the insert from the λ clone can therefore be sequenced to close the gap between contigs 1 and 4.

Many prokaryotic genomes have been sequenced by the shotgun method

The demonstration that a small genome can be sequenced relatively rapidly by the shotgun method led to a sudden plethora of completed microbial genomes. The 580 kb genome of *Mycoplasma genitalium* was completed in 1995, shortly after publication of the *H. influenzae* sequence. Despite the initial doubts...
about the feasibility of the approach, it soon became accepted that the shotgun method was capable of assembling the genome sequences of most, if not all, prokaryotic species. The relatively short lengths of these genomes mean that the computational requirements for finding sequence overlaps are not too great. With the introduction of next-generation sequencing in the 2000s, gap closure became less of an issue, because a single experiment can easily generate sequence reads whose total length is several hundred times that of the genome.

Many prokaryotic genomes have been assembled by de novo sequencing, as described above for H. influenzae, with the genome sequence worked out solely by finding overlaps between individual sequence reads. The genomes of over 6500 prokaryotic species have been sequenced in this way. There are, however, more than 40,000 complete prokaryotic genome sequences in the databases. This is because some species have been sequenced multiple times in order to understand the amount of variation that exists in their genomes (Section 8.2). These resequencing projects make use of a second, less computer-intensive approach to assembly, in which the existing sequence is used as a reference genome for assembly of additional genome sequences from the same species. Rather than looking for overlaps among the sequence reads for the genome that is being assembled, individual reads are simply placed on to the reference sequence by looking for regions of sequence identity or similarity (Figure 4.19).

The reference genome approach can also be used for de novo sequencing if the species whose genome is being sequenced is related to other species whose genomes are already assembled. The rationale is that the two sequences will be sufficiently similar for the known genome to direct assembly of the sequence for the new species. Usually, the reads from the new genome will be assembled into contigs before comparison with the reference, in order to increase the degree of certainty that can be assigned to matches. With this approach, care must be taken to avoid assembly errors that can arise if there are any regions in the new genome where the gene order has been changed by recombination (Figure 4.20). This kind of rearrangement does not alter the sequence of the segment that moves; it merely places that segment at a different position in the genome. This means that the only difference between the sequences will be at the boundaries of the segments that have been rearranged. If these boundaries are not identified, then reliance on the reference genome might lead to the rearrangement not being recognized in the genome being assembled.

Figure 4.19 Using a reference genome during a resequencing project. The reference genome is used to place the reads from the second genome at their correct relative positions.

Figure 4.20 Recombination can cause a problem when a reference genome is used. In this example, the positions of genes C and D are different in the reference genome compared with the genome being sequenced. If the sequence coverage does not include the critical regions at the boundaries for this recombination event, then the fact that the gene order is different in the genome being sequenced will not be recognized.
Shotgun sequencing of eukaryotic genomes requires sophisticated assembly programs

The shotgun approach is also being applied to eukaryotic genomes, but with these organisms sequence assembly is complicated by two factors. The first complication is the size of the read data set that must be assembled. Eukaryotic genomes are much longer than prokaryotic ones (for example, the human genome is 3235 Mb, compared with just 1.83 Mb for *H. influenzae*), so many more sequence reads are required in order to ensure adequate coverage. In fact, the problem as far as assembly is concerned is not the number of reads but the number of pairs of reads, as we must compare pairs in order to identify overlaps. The required data analysis therefore becomes disproportionately more complex as the number of sequence reads increases: for $n$ reads, the number of pairs is given by $2n^2 - 2n$.

The second problem with the shotgun method is that it can lead to errors if the genome contains repetitive DNA sequences. These are sequences, up to several kilobases in length, that are repeated at two or more places in a genome. As we learned in Section 3.1, they cause problems for the shotgun approach because a sequence that lies partly or wholly within one repeat element might accidentally be assigned an overlap with the identical sequence present in a different repeat element (see Figure 3.3A). This could lead to a part of the genome sequence being placed at the incorrect position or left out entirely. Most prokaryotic genomes contain relatively little repetitive DNA, but repeat sequences are common in eukaryotes, and in some species they make up more than 50% of the genome.

The first sequence assemblers—software packages that convert sequence reads into contigs—simply compared pairs of reads, merged those with the longest overlaps, and repeated the process until no further overlaps were present (Figure 4.21). The result is an overlap graph, from which a master sequence can be assembled. In computational parlance, this approach uses a greedy algorithm, one that makes the most logical choice at each step in an iterative process. However, greedy algorithms are notorious for sometimes ending up with a sub-optimal solution, because each step is taken in isolation without consideration of the nature of the problem as a whole. This is the case with the overlap graph method, as the presence of repetitive DNA is likely to lead to errors in assembly, as described above. Construction of an overlap graph is also computationally intensive. The approach is feasible when the chain-termination method is used to sequence a genome but is impossible with the much greater number of short reads obtained by a next-generation sequencing method.

More recently developed sequence assemblers take an alternative approach that is less computationally intensive and hence is applicable to next-generation sequence reads. These assemblers make use of a representation called a De Bruijn graph, which is a mathematical concept for identifying overlaps between strings of symbols. The strings must be of equal length, so when this method is used in sequence assembly, the initial step is to break the sequence reads into smaller segments or *k-mers*, typically 20–30 nucleotides in length. Duplicate *k*-mers are discarded, so this step reduces the size of the data set. Each *k*-mer is then converted into a prefix sequence (the *k*-mer minus its last nucleotide) and a suffix sequence (the *k*-mer minus its first nucleotide) (Figure 4.22A). Overlaps between *k*-mers are then identified by searching for pairs where the
4.3 HOW TO SEQUENCE A GENOME

(A) The prefix and suffix versions of a k-mer

<table>
<thead>
<tr>
<th>k-mer</th>
<th>ATGCAGCTATATAGCGGATG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prefix sequence</td>
<td>ATGCAGCTATATAGCGGATG</td>
</tr>
<tr>
<td>Suffix sequence</td>
<td>TGCACTATATAGCGGATG</td>
</tr>
</tbody>
</table>

(B) Reading a sequence for a De Bruijn graph

TGA ➔ GAC ➔ ACC ➔ CCC ➔ CGC ➔ GCA ➔ CAG ➔ AGT ➔ GTT ➔ TTA

Master sequence TGACGAGTTA

(C) An Eulerian pathway through a De Bruijn graph

TGA ➔ GAC ➔ ACC ➔ CCC ➔ CGC ➔ GCA ➔ CAG ➔ AGT ➔ GTT ➔ TTA

Master sequence TGACGACCTA

suffix of one k-mer is identical to the prefix of the second k-mer. Once all overlaps are identified, the k-mers are linked together as a De Bruijn graph, where each k-mer is depicted as a node and lines (edges) connect k-mers whose suffix and prefix sequences overlap (Figure 4.22B). The master sequence can then be read from the graph. If the sequence contains repetitive DNA, then the De Bruijn graph will be branched rather than linear. If this is the case, then the computer attempts to identify an Eulerian pathway through the graph, which is a pathway that visits each edge just once. If an Eulerian pathway can be identified, then the correct sequence assembly will be achieved, despite the presence of the repeated sequences (Figure 4.22C).

Even if the De Bruijn graph is so complex that an Eulerian pathway cannot be identified, the branched structure is still valuable, as it indicates the regions of the sequence in which there are unresolved repetitive DNA regions. These regions can then be studied in more detail, for example, by preparing a second sequencing library in which the fragments are longer than the longest repeat sequences in the region being studied. The two ends of each of these longer fragments are sequenced, to give paired-end reads. The paired-end reads from any fragments that contain a copy of the repetitive DNA sequence can then be used to resolve the genome assembly on either side of that repeat (Figure 4.23).

Paired-end reads can also be used to identify sequence contigs that are adjacent to one another in the genome. The result is a series of scaffolds, each scaffold comprising a set of sequence contigs separated by gaps that lie between the paired-end reads (Figure 4.24). As more sequences are added in to the data set, longer and longer scaffolds are built up. The accuracy of sequence assembly can be further checked by obtaining pairs of end-sequences from fragments of 100 kb or more that have been cloned in a high-capacity vector such as a bacterial vector.

Figure 4.22 Sequence assembly by use of a De Bruijn graph. (A) A k-mer of 20 nucleotides and its prefix and suffix sequences. (B) An example of sequence assembly by use of a De Bruijn graph. In this example, the k-mers are just three nucleotides in length, which is too short for sequence assembly in the real world but serves to illustrate the principle of the method. Pairs of k-mers with identical suffix and prefix sequences have been identified: GA for the first two k-mers in the series and AC for k-mers 2 and 3. The edges, shown here as arrows, link pairs of k-mers, resulting in a series from which the master sequence can be read. (C) Sequences containing repetitive DNA regions can be assembled correctly from a De Bruijn graph if an Eulerian pathway through the graph can be identified. This is a pathway that visits every edge just once. In this example, the Eulerian pathway follows the edges labeled 1–9 in series, resulting in unambiguous assembly of the master sequence even though that sequence contains a repeat of the motif ACC.

Figure 4.23 Using paired-end reads to resolve sequence assembly of a region that contains repetitive DNA. (A) In this example, the De Bruijn graph is ambiguous but identifies three segments of sequence that could be located downstream of a repeat sequence. (B) The locations of the paired-end reads obtained from a fragment that spans the repeat sequence identify which of the segments is positioned downstream of the repeat sequence.
artificial chromosome (BAC). If a pair of end-sequences does not fall within a single scaffold at their anticipated positions relative to one another, then an error in assembly has occurred.

**More complex genomes can be sequenced by a hierarchical shotgun approach**

The first eukaryotic genomes, including the human genome, were sequenced by a modification of the shotgun approach called **hierarchical shotgun sequencing**. This approach involves a pre-sequencing phase during which the genome is broken into large fragments, typically 300 kb in length, and these fragments are cloned into a high-capacity vector such as a BAC (Section 2.3). Clones that contain overlapping fragments of DNA are then identified, enabling a contiguous series or **clone contig** to be built up (Figure 4.25). The sequence of each clone insert is then assembled by the shotgun method, and the master sequence is built up by joining together the insert sequences in the order dictated by the contig. Repetitive DNA will cause a problem only if two or more copies of the same repeat sequence are present in a single clone insert, and even then it might be possible to identify errors in the assembly by examining the sequences of clones that overlap with the one containing the repeats (Figure 4.26).

The hierarchical approach is still used today in projects attempting to sequence the larger eukaryotic genomes, such as those of barley, wheat, and other plants. The main difficulty with this approach is the time and effort needed to identify clones that contain overlapping inserts. The simplest way to build up an overlapping...
series of cloned DNA fragments is to begin with one clone from a library, identify a second clone whose insert overlaps with the insert in the first clone, then identify a third clone whose insert overlaps with the second clone, and so on. This is the basis of **chromosome walking**, which was the first method devised for assembly of clone contigs. In its original form, the insert from one clone was used as a hybridization probe to screen all the other clones in the library. Clones whose inserts overlap with the probe give positive hybridization signals, and their inserts can be used as new probes to continue the walk (**Figure 4.27**). This method suffers from the drawback that if the probe contains a repeat sequence, then it will hybridize not only to overlapping clones but also to nonoverlapping ones whose inserts contain copies of the repeat. An alternative approach that avoids this problem is to design primers that will amplify a segment at the end of a clone insert and then use these primers in attempted PCR with all the other clones in the library. A second clone that gives a PCR product of the correct size must contain an insert that overlaps with that of the original clone (**Figure 4.28**). To speed up the process even more, rather than performing a PCR with each individual clone, a combinatorial process can be used, in which groups of clones are mixed together in such a way that unambiguous identification of overlapping inserts can still be made. This method is illustrated in **Figure 4.29**.

Even when the screening step is carried out by combinatorial PCR, chromosome walking is a slow process, and it is rarely possible to assemble contigs of more than 15–20 clones by this method. So what alternative methods are there?

The main alternative is to use a **clone fingerprinting** technique. Clone fingerprinting provides information on the physical structure of a cloned DNA fragment. This physical information, or fingerprint, is compared with equivalent data from other clones, enabling those with similarities, possibly indicating
overlaps, to be identified. One or a combination of the following techniques can be used (Figure 4.30):

- **Restriction patterns** can be generated by digesting clones with a variety of restriction enzymes and separating the products in an agarose gel. If two clones contain overlapping inserts, then their restriction fingerprints will

**Figure 4.29** Combinatorial screening of clones in microtiter trays. In this example, a library of 960 clones has to be screened by PCR. Rather than carrying out 960 individual PCRs, the clones are grouped as shown and just 296 PCRs are performed. In most cases, the results enable positive clones to be identified unambiguously. In fact, if there are few positive clones, then sometimes they can be identified by just the row and column PCRs. For example, if positive PCRs are obtained with tray 2 row A, tray 6 row D, tray 2 column 7, and tray 6 column 9, then it can be concluded that there are two positive clones, one in tray 2 well A7 and one in tray 6 well D9. The well PCRs are needed if there are two or more positive clones in the same tray.

**Figure 4.30** Four clone fingerprinting techniques.

(A) Restriction fingerprint

(B) Repetitive DNA fingerprint

(C) Repetitive DNA PCR

(D) STS content mapping
have bands in common, as both will contain fragments derived from the overlap region.

- **Repetitive DNA fingerprints** can be prepared by analyzing a set of restriction fragments by Southern hybridization with probes specific for one or more types of repeat sequence. As with restriction fingerprints, overlaps are identified by looking for two clones that have some hybridizing bands in common.

- **Repetitive DNA PCR**, or interspersed repeat element PCR (IRE-PCR), uses primers that anneal within repeat sequences and so amplify the *single-copy DNA* between two neighboring repeats. Because repeat sequences are not evenly spaced in a genome, the sizes of the products obtained after repetitive DNA PCR can be used as a fingerprint in comparisons with other clones in order to identify potential overlaps.

- **Sequence-tagged site (STS) content mapping** is particularly useful because it can result in a clone contig that is anchored onto a physical map of STS locations. PCRs directed at individual STS markers (*Section 3.6*) are carried out with each member of a clone library. If it is presumed that the STS has a single copy in the genome, then all clones that give PCR products must contain overlapping inserts.

As with chromosome walking, efficient application of these fingerprinting techniques requires combinatorial screening of gridded clones, ideally with computerized methodology for analyzing the resulting data.

**What is a genome sequence and do we always need one?**

In this chapter we have examined how genome sequences are obtained. We have not, however, asked ourselves what we mean by the term “genome sequence”. This is what we must do now.

We have already noted that for most species there is no such thing as *the* genome sequence. The individual members of a species have their own personal variations, and the variable positions often make up a substantial fraction of the genome as a whole. The human genome, for example, contains some 10 million single-nucleotide polymorphisms (SNPs), positions that are variable in different people. This is an average of one SNP for every 325 bp of the genome. This means that a genome sequence is, at best, just the genome of a single representative of a species. In fact, for some of the first species to be studied, the genome sequence was a conglomerate of different genomes, as separate sequencing libraries were prepared for different chromosomes, and the DNA for those libraries did not always come from the same individual. Even if a single source is used, there is the complication that each member of a diploid species has two copies of its genome, and those two copies might differ at various SNP positions. These *heterozygosities* will appear as ambiguities during the assembly process (*Figure 4.31*), and usually one of the variations will be chosen to represent that position in the genome sequence.

It is also important to recognize that a complete sequence—one in which every nucleotide is known, with no errors, and every segment placed at its correct position—is currently unattainable for a eukaryotic genome. The sequence referred to as the *finished* genome will typically still have some unsequenced gaps between contigs and an average of up to one error per $10^4$ nucleotides. In contrast, a *draft* genome sequence has a greater error rate, more gaps, and possibly some ambiguity about the order and/or orientation of some sequence contigs. Moving down the scale, we have partially assembled genomes, which might be works in progress, and, in extreme cases, unassembled collections of sequence reads.

```
...ATGAGCATCGATGCA
CAGCAGATTGAGCTAC...
```
To avoid confusion over the meaning of terms such as “finished” and “draft”, a number of statistics have been devised to provide a more accurate measure of the degree of completeness of a genome sequence. One of these is the **N50 size**, which can be applied to either contigs or scaffolds. A contig N50 size is derived as follows:

- First, the total length of all the contigs added together is calculated.
- The contigs are then ordered by length from longest to shortest.
- Beginning with the longest contig, the individual lengths are added together until the combined length is equal to half the total length of all the contigs.
- The N50 value is the length of the last and hence shortest contig whose addition makes the combined length greater than 50% of the total.

A higher N50 value therefore indicates a more complete assembly. A different value, the **NG50 size**, is based not on the total length of contigs or scaffolds but on the actual genome size. The NG50 value therefore enables direct comparisons to be made between genome assemblies for different species.

Finally, we must appreciate that a genome sequence, however complete, is not always needed. All parts of the genome are fascinating, but the degree of fascination for any component is in the eye of the beholder. For example, for some research questions, the critical issue is the sequence of the **exome**, which is the complete set of exons in the genome and hence the part that codes for proteins. Sequence variations in the exomes of different individuals might reveal protein polymorphisms that underlie cancer or some other disorder. The human exome comprises approximately 48 Mb of DNA, about 1.5% of the total genome. Sequencing the exome would therefore be a less challenging task than sequencing the entire genome. With next-generation methods, sequencing can be directed at the exome, or any other part of the genome, by including **target enrichment** during preparation of the sequencing library. A large set of oligonucleotides, each typically 150 nucleotides in length, is synthesized, the sequences of these oligonucleotides corresponding to the sequences of the part of the genome that is being targeted (Figure 4.32). The oligonucleotides then act as **baits** that hybridize to,

![Figure 4.32 Target enrichment](image.png)

**(A)** DNA capture

![DNA capture diagram](image.png)

**(B)** Comparison between sequencing before and after capture

![Comparison diagram](image.png)

*Figure 4.32 Target enrichment.* (A) Baits are used to capture DNA fragments representing segments of the genome that are of interest. (B) Only the captured DNA fragments are sequenced.
and hence capture, the DNA fragments representing the segments of interest. If the baits are attached to magnetic beads, the captured DNA fragments can be collected with a magnet, and nontargeted parts of the genome, which remain in solution, are discarded. The captured DNA is then released from the baits and used to prepare the sequencing library.

4.4 A SURVEY OF EUKARYOTIC GENOME SEQUENCING PROJECTS

To complete this chapter, we will examine the genome projects for four eukaryotic species, to explore how the challenges inherent in sequencing and assembling a genome have been dealt with as the technologies have gradually advanced.

The Human Genome Project: genome sequencing in the heroic age

The Human Genome Project (HGP) was established in the late 1980s as a loose but organized collaboration between geneticists in all parts of the world, with the objective of obtaining, by 2005, a finished sequence of at least 95% of the euchromatin component of the human genome, the euchromatin being the part in which most of the genes are located (Section 10.1). Prior to the beginning of the HGP, complete sequences had been obtained only for viral and organelle genomes, the longest being the chloroplast genomes of tobacco and other plants, which are 155 kb. The massive conceptual leap to the 3235 Mb human genome was looked on as mad by some biologists. Although comprehensive genetic maps had been constructed for fruit flies and a few other organisms, the problems inherent in analysis of human pedigrees (Section 3.4) and the relative paucity of polymorphic genetic markers meant that most geneticists doubted whether a map of the human genome could ever be achieved, and it was assumed that without a map it would be impossible to assemble the genome sequence.

The initial goal that the HGP set itself was therefore a genetic map with a density of one marker per 1 Mb, although it was thought that a density of one per 2–5 Mb might be the realistic limit. The initial breakthrough came from the discovery of restriction fragment length polymorphisms (RFLPs), which were the first highly polymorphic DNA markers to be recognized in animal genomes. In 1987 the first human RFLP map was published, comprising 393 RFLPs and 10 additional polymorphic markers, with an average marker density of one per 10 Mb. By 1994, the genetic map had been extended to 7000 markers, most of which were simple sequence length polymorphisms (SSLPs), with a density of one marker per 0.7 Mb. Physical mapping did not lag far behind, with publication in 1995 of a radiation hybrid map of 15,088 STS markers, with an average density of one per 199 kb. This map was later supplemented with an additional 20,104 STS markers, most of these being expressed sequence tags (ESTs) and hence positioning protein-coding genes on the physical map. The combined STS maps included positions for almost 7000 polymorphic SSLPs that had also been mapped onto the genome by genetic means. As a result, the physical and genetic maps could be directly compared, and clone contig maps that included STS data could be anchored onto both maps. The net result was a comprehensive, integrated map that was then used as the framework for the DNA sequencing phase of the Human Genome Project.

The sequencing phase made use of a library of 300,000 BAC clones whose positions on the genome map were known and which could therefore be assembled into a series of clone contigs. The strategy was therefore to carry out shotgun sequencing of individual BAC inserts, by the chain-termination method, and from these inserts to build up contiguous blocks of sequence that could be anchored onto the genome map. Just as the sequencing phase of the HGP was beginning, a second group of researchers started to explore the possibility of generating the final sequence by the shotgun method, without recourse to the laborious assembly of clone contigs. The possibility that the HGP might not in fact provide the first human genome sequence stimulated the organizers of the project to bring forward their
planned dates for completion of a working draft. The first draft sequence of an entire human chromosome (number 22) was published in December 1999, and the draft sequence of chromosome 21 appeared a few months later. Finally, on June 26, 2000, accompanied by the President of the United States, the leaders of the two projects, Francis Collins and Craig Venter, jointly announced completion of their genome drafts, which appeared in print eight months later.

The draft sequence obtained by the clone contig method covered just 90% of the genome, with the missing 320 Mb lying predominantly in constitutive heterochromatin. This component of the genome is made up of chromosomal regions in which the DNA is very tightly packaged and in which there are few if any genes (Section 10.1). Within the 90% of the genome that was covered, each part had been sequenced at least four times, providing an acceptable level of accuracy, but only 25% had been sequenced the 8–10 times necessary before the work is considered to be finished. Furthermore, this draft sequence had approximately 150,000 gaps, and it was recognized that some segments had probably not been ordered correctly. The International Human Genome Sequencing Consortium, which managed the final phase of the project, set as its goal a finished sequence of at least 95% of the euchromatin, with an error rate of less than one in $10^4$ nucleotides, and all except the most refractory gaps filled. Achieving this goal required further sequencing of 46,000 BAC, PAC, fosmid, and cosmid clones. The first finished chromosome sequences began to appear in 2004, with the entire genome sequence being considered complete a year later.

The HGP was a monumental undertaking that stretched the limits of the mapping and sequencing technologies that were available in the 1980s and 1990s. As well as the sequence itself, the project laid the groundwork for future genome initiatives. In particular, the success of the shotgun approach showed that it would not always be necessary to have a comprehensive map of a eukaryotic genome in order to perform the sequence assembly. It was also recognized that, having sequenced one copy of the human genome, sequencing additional copies would be less difficult, as the initial sequence could be used as a reference for assembly of subsequent versions. We will now look at one particularly interesting way in which the human genome was used as a reference for assembly of a second genome.

The Neanderthal genome: assembly of an extinct genome by use of the human sequence as a reference

Neanderthals are extinct hominins who lived in Europe and parts of Asia between 200,000 and 30,000 years ago. Their preserved skeletons show many features similar to those of our own species, *Homo sapiens*, leading paleontologists to classify Neanderthals as another member of the *Homo* genus, *Homo neanderthalensis*, our closest relatives in the fossil world. Many of the distinctive features of Neanderthals are thought to be adaptations to the cold climates of the Ice Ages, such as an enlarged nasal cavity, which might have helped to warm the air before it entered the lungs (Figure 4.33).

If the last Neanderthal died out 30,000 years ago, then how can we obtain the sequence of its genome? Ancient DNA has provided an answer. It has been known for some years that DNA molecules can survive the death of the organism in which they are contained, being recoverable centuries and possibly millennia later as short, degraded fragments preserved in bones and other biological remains. Unfortunately, the study of ancient DNA has been plagued with controversies. In the early 1990s there were many reports of ancient human DNA being detected in bones and other archaeological specimens, but often it turned out that what had been sequenced was not ancient DNA at all but contaminating modern DNA from the archaeologist who had excavated the specimen or from the molecular biologist who carried out the DNA extraction. The worldwide success of the film *Jurassic Park* led to reports of DNA in insects preserved in amber and even in dinosaur bones, but all these claims are now known to be incorrect. Many biologists started to wonder if ancient DNA existed at all, but gradually it has become...
clear that if the work is carried out with extreme care, it is sometimes possible to extract authentic ancient DNA from specimens up to about one million years old. This is old enough to include Neanderthals.

Although they are different species, it was anticipated that the Neanderthal and \textit{H. sapiens} genomes would be sufficiently similar for the human genome to act as a reference for assembly of the Neanderthal sequence reads. It would not therefore be necessary to attempt a \textit{de novo} assembly of the Neanderthal sequence. There would, however, be one problem not usually encountered during a genome assembly. DNA degrades over time, and some of these degradation processes result in what ancient DNA researchers call \textbf{miscoding lesions}. A miscoding lesion is a chemical change that results in a nucleotide being read incorrectly during a sequencing experiment. The commonest miscoding lesion is removal of the amino group from a cytosine base, which is promoted by the presence of water and gives rise to uracil (\textbf{Figure 4.34A}). If this miscoding lesion occurs in an ancient DNA molecule, then the C will mistakenly be read as a T. These sequence errors must be distinguished from genuine positions at which the Neanderthal genome differs from the \textit{H. sapiens} reference. For the Neanderthal genome sequence to be accurate, it would therefore be necessary to achieve high coverage, so that each nucleotide in the genome is present in as many sequence reads as possible (\textbf{Figure 4.34B}). Those reads will be derived from different fragments of ancient DNA, and it would be extremely unlikely for all of those fragments to have a miscoding lesion at the same position. Genuine differences between the Neanderthal and \textit{H. sapiens} sequences would therefore be signaled by a SNP that is present in each of the sequence reads covering a particular nucleotide. A miscoding lesion, on the other hand, would only occur in one of those reads.

The first complete Neanderthal genome sequence was obtained from a toe bone of an adult female, dated to approximately 50,000 years ago and recovered from a cave site in the Altai Mountains of Siberia. Five libraries were prepared and sequenced, from which over 2,278,000,000 Neanderthal sequence reads were obtained, by the Illumina method. The average length of these reads was only 75 bp, a second problem with ancient DNA being that the polynucleotides degrade into short fragments over time, limiting the lengths of individual sequence reads. Nevertheless, it was possible to assemble the reads into a genome sequence with an average of 52× coverage, sufficient for a high degree of sequence accuracy.

What has the sequence told us about Neanderthals and their relationships with present-day humans? To find the answer to that question, you will have to wait until \textbf{Section 18.3}, where we examine in more detail the fast-moving field of \textbf{paleogenomics}: the study of the genomes of extinct species.

\textbf{The giant panda genome: shotgun sequencing based entirely on next-generation data}

The giant panda, \textit{Ailuropoda melanoleuca}, is a critically endangered species with possibly as few as 1500 wild individuals left alive in various mountainous regions of western China (\textbf{Figure 4.35}). Its genome is about 2400 Mb, somewhat shorter than the human genome, due to a lower repetitive DNA content; the number of genes is about the same. The giant panda genome was the first to be sequenced entirely from next-generation data. Previously, eukaryotic genomes that were

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure434}
\caption{\textbf{Miscoding lesions cause problems when sequencing ancient DNA.} (A) Deamination of cytosine gives uracil. When a DNA sequence is read, this miscoding lesion results in the C being read as a T. (B) High coverage is needed to distinguish miscoding lesions from genuine sequence variations. In this example, there are two possible miscoding lesions. One can be identified with confidence as a miscoding lesion because it is present in just one of four sequence reads. The second possible miscoding lesion is present in a part of the genome that is covered by just a single ancient DNA sequence read. It is impossible to tell if this C → T anomaly is a miscoding lesion or a genuine sequence variation.}
\end{figure}
small enough for shotgun assembly were sequenced by a hybrid approach employing both chain-termination sequencing and a next-generation method. Contigs of short sequence reads were generated by the next-generation method. Chain-termination sequencing was then used to obtain paired-end reads from large cloned fragments, and these end reads directed assembly of the contigs into scaffolds. The drawback with this approach was the additional time and expense needed to prepare the clone library and sequence the clones individually to obtain the paired-end reads. A strategy that dispensed with the chain-termination component, and enabled the entire genome to be sequenced solely from next-generation reads, would therefore be quicker and less costly.

To sequence the giant panda genome, a series of sequencing libraries was prepared, containing fragments with average lengths of 150 bp, 500 bp, 2 kb, 5 kb, and 10 kb. The libraries were sequenced by the Illumina method, generating a total of 176,000 Mb of sequence. The genome was then built up in a stepwise manner (Figure 4.36):

- Short reads of less than 500 bp were assembled, using a De Bruijn algorithm to identify overlaps. The resulting contigs had a total length of 2000 Mb with 39× coverage but an N50 size of only 1.5 kb, indicating that, at this stage of the assembly process, the individual contigs were very short.

- Paired-end reads from the fragments in each library were then used to assemble the contigs into scaffolds. These reads were obtained by a modification of the standard Illumina procedure that enables both ends of each immobilized fragment to be sequenced. The resulting scaffolds had an N50 size of 1.3 Mb and a total sequence length of 2300 Mb. The increase in assembly length from 2000 to 2300 Mb was due to the presence of gaps between the contigs in the individual scaffolds.

- Gaps within scaffolds were at least partially closed by identifying paired-end reads that had just one read anchored in a contig and the other end unanchored in a gap. The unanchored ends were used to seed additional assembly of short reads. This stage increased the contig N50 size to 40 kb, with the remaining gaps (largely comprising highly repetitive DNA) making up only 2.4% of the total scaffold length. This final assembly had an average coverage of 65×, with 95% of the genome covered by 20 reads or more.

![Figure 4.36](image)

**Figure 4.36** Sequencing the giant panda genome. (A) Short reads of less than 500 bp were assembled to give contigs. (B) Paired-end reads were used to assemble the contigs into scaffolds. (C) Gaps between contigs were partially closed by short read assembly around the free ends of paired-end reads that had just one read anchored in a contig.
Analysis of the giant panda genome revealed some interesting information regarding its very specialized diet. One of the reasons why pandas have such a narrow geographical range is because their diet is made up primarily of bamboo. The panda genome included most of the genes associated with a carnivorous diet, suggesting that pandas could also digest meat. However, genes for a taste receptor contained mutations that might have made meat unpleasant to the panda. Eating bamboo requires the ability to break down cellulose, and no genes that would enable the panda to do this could be identified in its genome. One possibility is that the panda’s *microbiome*—the collection of bacteria in its digestive system—provides this essential dietary function.

The initial giant panda genome has subsequently been used as a reference for assembly of the genome sequences for 34 additional individuals, representing 2% of the total population. This resequencing project identified three distinct panda populations, pointing the way to planned breeding programs aimed at maintaining the genetic diversity of the species as a whole. Comparisons of the genome sequences also suggested that although the population numbers have been affected by changes in the climate, the main factor responsible for the decline in the number of giant pandas has been human destruction of their environment.

### The barley genome: the concept of gene space

Barley is the world’s fourth most important crop in terms of annual global production. Most of the barley that is grown is used to feed farm animals, but about a quarter is consumed by humans, directly or indirectly in food products or as alcoholic beverages prepared from barley malt. Barley is potentially a rich source of dietary fiber, and crop breeding programs aimed at increasing its fiber content are underway. Attempts are also being made to understand the adaptability of barley to extreme environments, with the hope that varieties can be developed that are able to remain productive when the climate changes. Those changes might result in increased demand for crops able to grow in dry environments, so breeding drought tolerance into barley is an important goal of current breeding programs.

Breeding programs are possible without extensive knowledge of a plant’s genome, but knowing the identities and map positions of key genes and quantitative trait loci (QTLs) makes a directed breeding program much more effective, and much more likely to have a successful outcome. The barley genome is 5100 Mb in length, with 80% of the genome made up of repetitive DNA, greatly complicating the assembly of sequence reads. At present, not even a draft genome sequence has been completed. Can an alternative description of the barley genome be devised, one that satisfies, at least to some extent, the need to identify genes and map positions in the DNA sequence? Barley researchers have come up with the concept of *gene space*, which is, in essence, the sequences of the vast majority of barley genes anchored on to a detailed genome map. The key stages in development of the barley gene space were as follows (Figure 4.37):

- A detailed physical map was constructed by STS content analysis of 571,000 BAC clones. These clones were assembled into 9265 contigs covering 4980 Mb of the genome, over 97% of the total. The clone contigs had an N50 length of 904 kb, long enough for individual genes to be contained entirely within a single contig.

- From the BAC library, the inserts from 5341 clones known to contain genes, as well as 937 chosen at random, were sequenced, and the sequences were placed on the physical map. Additional sequences were obtained from the ends of the inserts in 304,523 BAC clones. This work provided a sequence-enriched map comprising 1136 kb of anchored sequence, with much of that sequence containing genes.

- Genomic DNA was used to prepare two Illumina sequencing libraries, with fragment sizes of 300 bp and 2.5 kb. *De novo* assembly of the reads
generated contigs totaling 1900 Mb of sequence, but many of these contigs were short and made up entirely of repetitive DNA. Of the 376,261 contigs that were longer than 1 kb and lacked repeat sequences, 112,989 had overlaps with components of the sequence-enriched map. This increased the amount of sequence contained in the map to over 300 Mb.

- Data from genetic mapping were added to the sequence-enriched map. The sequence contigs were searched for SNPs and other short sequences whose map positions had previously been identified by linkage analysis. As a result, 4556 contigs could be positioned on the genetic map.

- Finally, genes were located within the sequence-enriched map. RNA was prepared from plants at eight different stages of development, from initial germination of the seedling through flowering and formation of seeds. The RNA was converted into cDNA, and a sequencing library was prepared. The resulting RNA-sequencing or RNA-seq data represent the exome, so when they are assembled, these reads give the sequences of the exonic parts of the genes. These exon sequences were then added to the map by searching for them within the sequenced parts of the clone contigs. An additional 28,592 cDNA sequences that had previously been obtained were also mapped in this way. The RNA-seq and cDNA sequences identified 26,159 genes, of which 24,154 could be positioned on the sequence-enriched map.

The barley gene space therefore comprises the sequences of the exons of 26,159 genes, with 24,154 of these positioned on the combined physical and genetic maps, along with additional sequence data containing SNPs and other DNA markers whose positions on the physical and/or genetic maps are also known.
Although not a draft genome sequence, the gene space provides the essential information on gene sequences and marker locations needed for studies aimed at identifying genes specifying important characteristics and for the design of breeding programs that make use of those genes.

**SUMMARY**

- Chain-termination sequencing is used to sequence short DNA molecules such as PCR products.
- Next-generation sequencing is a collection of methods that enable thousands or millions of DNA fragments to be sequenced in parallel in a single experiment.
- Next-generation methods begin with preparation of a library of DNA fragments that have been immobilized on a solid support in such a way that multiple sequencing reactions can be carried out side by side in a massively parallel array format. Various sequencing methods are applied to the library.
- Third- and fourth-generation sequencing provide advantages over next-generation methods and may replace the latter in the medium term.
- When a genome is being sequenced the major challenge is assembling all the mini-sequences, obtained from the multiple sequencing experiments, in the correct order.
- With a small bacterial genome, sequence assembly is possible by the shotgun method, which simply involves examining sequence reads for overlaps.
- Shotgun sequencing of eukaryotic genomes requires more complex assembly procedures such as the use of De Bruijn graphs.
- More complex genomes can be sequenced by the hierarchical shotgun approach, which makes use of a clone contig comprising a series of clones, in a high-capacity vector such as a BAC, that contain overlapping fragments that have been anchored onto a physical and/or genetic map of the genome under study.
- The completeness of a genome sequence is described by its N50 size.
- A draft of the human genome sequence was completed in 2000. The sequence of the Neanderthal genome is also known.
- The giant panda genome was the first to be sequenced entirely by shotgun assembly of next-generation sequencing data.

**SHORT ANSWER QUESTIONS**

1. Explain how the inclusion of dideoxynucleotides in a strand-synthesis reaction enables a DNA sequence to be read.

2. DNA polymerases used for chain-termination DNA sequencing should have which three properties?

3. Compare the strengths and weaknesses of chain-termination and next-generation sequencing methods. What are the applications of chain-termination sequencing in modern genomics research?

4. How is a sequencing library prepared during a next-generation sequencing project?

5. Describe how the DNA sequence is obtained by (A) reversible terminator sequencing, (B) pyrosequencing, (C) the ion torrent method, and (D) SOLiD sequencing.
6. What are the advantages of third- and fourth-generation sequencing methods?
7. Outline the approach that was used in order to sequence the *Haemophilus influenzae* genome.
8. What factors complicate the application of shotgun sequencing to a eukaryotic genome?
9. Describe how sequence reads are assembled into contiguous sequences.
10. Describe the hierarchical shotgun method for genome sequencing.
11. What methods can be used to perform clone fingerprinting?
12. Compare the approaches used to obtain the first sequences of (A) the human genome and (B) the giant panda genome.

**IN-DEPTH PROBLEMS**

1. In the late 1970s, two methods for DNA sequencing—the chain-termination and chemical degradation methods—were developed. Initially, both methods were used extensively, but gradually the chain-termination method became more popular. Why did the chemical degradation method go out of favor?

2. You have isolated a new species of bacterium whose genome is a single DNA molecule of approximately 2.6 Mb. Write a detailed project plan to show how you would obtain the genome sequence for this bacterium.

3. A 122 bp DNA molecule is randomly broken into overlapping fragments and those fragments are sequenced. The resulting sequence reads are as follows:
   - CGTAGCTAGCTAGCGATT
   - GATTAGTTCCGGCCCATTCG
   - GCTGTAGCATGTTTTCGC
   - TTCGCTCAGCATCGGATT
   - ATTTAGTTGACGTAGCA
   - CATTCCGGATGCTATCTCT
   - GTTGACGCTACGGCGGG
   - TCGTAGCTAGCTAGCGAT
   - ATGCTATCTCATCTGATT
   - ATTTAGTTCGCCCCATTCGC
   - ATTTAGTTCGGTACGATCG
   - ATGCATCGTAGCTAGCTAG
   - CTCAGCATCGGATTTAGTT
   - CGATGCTATCTCATCTGAT
   - CGCATACGGCGGGGGGAT

   Is it possible to reconstruct the sequence of the original molecule by searching for overlaps between pairs of reads? If not, then what problem has arisen and how might this problem be solved?

4. Critically evaluate the strengths and weaknesses of the hierarchical shotgun approach as a means of sequencing a large eukaryotic genome.
5. A pharmaceutical company has invested a great deal of time and money to sequence the gene for a genetic disease. The company is now studying the gene and its protein product and is working to develop drugs to treat the disease. Does the company have the right (in your opinion) to patent the gene sequence? Justify your answer.

**FURTHER READING**

**Chain-termination sequencing**


**Next-generation sequencing and third- and fourth-generation methods**


**Shotgun sequencing**


**Sequence assembly**


**Hierarchical shotgun sequencing**


**Iconic sequencing projects**


A genome sequence is not an end in itself. Major challenges still have to be met in locating the genes and other interesting features within the genome sequence and assigning functions to those genes whose roles are unknown. These challenges can be addressed by a combination of computer analysis and experimentation, but a complete description of a genome is rarely possible. Even though the finished version of the human genome sequence has been available since 2004, there are still uncertainties regarding the number of genes in the genome, and many of the identified genes have unknown functions. The development of new methods for understanding genome sequences is one of the central goals of genomics research.

This chapter describes the methods used in genome annotation, the process by which genes are located in a genome sequence. In Chapter 6, we will explore the various ways in which functions can be assigned to unknown genes.

5.1 Genome Annotation by Computer Analysis of the DNA Sequence

Once an assembled genome sequence has been obtained, various methods can be employed to locate the genes that are present. These methods can be divided into those that involve simply inspecting the sequence, by eye or more frequently by computer, and those methods that locate genes by experimental analysis.

Sequence inspection can be used to locate genes because genes are not random series of nucleotides but instead have distinctive features. At present we do not fully understand the nature of all these specific features, and sequence inspection is therefore not a foolproof way to locate genes, but it is still a powerful tool and is usually the first method applied to analysis of a new genome sequence. The computer techniques form part of the methodology called bioinformatics, and it is with these that we begin.

The coding regions of genes are open reading frames

Genes that code for proteins comprise open reading frames (ORFs) consisting of a series of codons that specify the amino acid sequence of the protein that the gene codes for (Figure 5.1). The ORF begins with an initiation codon, usually (but not always) ATG, and ends with a termination codon, either TAA, TAG, or TGA (Section 1.3). ORF scanning or ab initio gene prediction, which involves searching a DNA sequence for ORFs that begin with an ATG and end with a termination triplet, is therefore one way of looking for genes. The analysis is complicated by the fact that each DNA sequence has six reading frames, three in one direction and three in the reverse direction on the complementary strand (Figure 5.2), but computers are quite capable of scanning all six reading frames for ORFs. How effective is this as a means of gene location?

The key to the success of ORF scanning is the frequency with which termination codons appear in the DNA sequence. If the DNA has a random sequence and a GC content of 50%, then each of the three termination codons (TAA, TAG, and TGA) will appear, on average, once every $4^3 = 64$ bp. If the GC content is greater than 50%, then the termination codons, being AT-rich, will occur less frequently, but
Figure 5.2 A double-stranded DNA molecule has six reading frames. Both strands are read in the 5' → 3' direction. Each strand has three reading frames, depending on which nucleotide is chosen as the starting position.

Figure 5.3 ORF scanning is an effective way of locating genes in a bacterial genome. The diagram shows 4522 bp of the lactose operon of *Escherichia coli* with all ORFs longer than 50 codons marked. The sequence contains two real genes, *lacZ* and *lacY*, indicated by the red lines. These real genes cannot be mistaken because they are much longer than the spurious ORFs, shown in yellow.

Simple ORFs are less effective with genomes of higher eukaryotes

Although ORF scans work well for bacterial genomes, they are less effective for locating genes in DNA sequences from higher eukaryotes. This is partly because there is substantially more space between the real genes in a eukaryotic genome (for example, approximately 62% of the human genome is intergenic), increasing the chances of finding spurious ORFs. But the main problem with the human genome, and the genomes of higher eukaryotes in general, is that their genes are often split by introns (Section 1.2) and so do not appear as continuous ORFs in the DNA sequence. Many exons are shorter than 100 codons, some consisting of fewer

one will still be expected every 100–200 bp. This means that random DNA should not show many ORFs longer than 50 codons in length, especially if the presence of a starting ATG triplet is used as part of the definition of an ORF. Most genes, on the other hand, are longer than 50 codons: the average lengths are 300–350 codons for bacterial genes and approximately 450 codons for humans. ORF scanning, in its simplest form, therefore takes a figure of, say, 100 codons as the shortest length of a putative gene and records positive hits for all ORFs longer than this.

How well does this strategy work in practice? With bacterial genomes, simple ORF scanning is an effective way of locating most of the genes in a DNA sequence. This is illustrated by Figure 5.3, which shows a segment of the *E. coli* genome with all ORFs longer than 50 codons highlighted. The real genes in the sequence cannot be mistaken because they are much longer than 50 codons in length. With bacteria, the analysis is further simplified by the fact that the genes are very closely spaced and hence there is relatively little intergenic DNA in the genome (only 11% for *E. coli*; see Section 11.2). If we assume that the real genes do not overlap, which is true for most bacterial genes, then a short, spurious ORF would have to be located in an intergenic region in order for it to be mistaken for a real gene. This means that if the intergenic component of a genome is small, then there is a reduced chance of making mistakes in interpreting the results of a simple ORF scan.
than 50 codons, and continuing the reading frame into an intron usually leads to a
termination sequence that appears to close the ORF (Figure 5.4). In other words,
the genes of a higher eukaryote do not appear in the genome sequence as long
ORFs, and simple ORF scanning cannot locate them.

Solving the problem posed by introns is the main challenge for bioinformatics
writers new software programs for ORF location. Three modifications to the
basic procedure for ORF scanning have been adopted:

- **Codon bias** is taken into account. Codon bias refers to the fact that not all
codons are used with equal frequency in the genes of a particular organism. For example, leucine is specified by six codons in the genetic code (TTA, TTG, CTT, CTC, CTA, and CTG; see Figure 1.26), but in human genes
leucine is most frequently coded by CTG and is only rarely specified by TTA or CTA. Similarly, of the four valine codons, human genes use GTG four times more frequently than GTA. The biological reason for codon bias is not understood, but all organisms have a bias, which is different in different species. Real exons are expected to display codon bias, whereas chance series of triplets do not. The codon bias of the organism being studied is therefore written into the ORF-scanning software.

- **Exon–intron boundaries** can be searched for, as these have distinctive
sequence features, although unfortunately the distinctiveness of these
sequences is not so great as to make their location a trivial task. The sequence of the upstream exon–intron boundary is usually described as 5’-AG GTAAGT-3’, with the arrow indicating the precise boundary point. However, only the GT immediately after the arrow is invariable: elsewhere in the sequence, nucleotides other than the ones shown are quite often found. In other words, the sequence is a **consensus**, by which we mean that the sequence shows the most frequent nucleotide at each position in all of the upstream exon–intron boundaries that are known, but in any particular boundary sequence, one or more of these positions might have a different nucleotide (Figure 5.5). The downstream intron–exon boundary is even less well defined: 5’-PyPyPyPyPyPyNCAG-3’, where Py means one of the pyrimidine nucleotides (T or C) and N is any nucleotide.

- **Upstream regulatory sequences** can be used to locate the regions where
genes begin. These regulatory sequences, like exon–intron boundaries, have distinctive sequence features that they possess in order to carry out their role as recognition signals for the DNA-binding proteins involved in
gene expression (Section 12.2). Unfortunately, as with exon–intron boundaries, the regulatory sequences are variable, more so in eukaryotes than in prokaryotes, and in eukaryotes not all genes have the same collection of regulatory sequences. Using these to locate genes is therefore problematic.

These three extensions of simple ORF scanning, despite their limitations, are generally applicable to the genomes of all higher eukaryotes. Additional strategies are also possible with individual organisms, depending on the special features of their genomes. For example, vertebrate genomes contain **CpG islands** upstream of many genes. These are sequences of approximately 1 kb in which the GC content is greater than the average for the genome as a whole. Some 40–50% of human genes are associated with an upstream CpG island. These sequences are distinctive, and
when one is located in vertebrate DNA, a strong assumption can be made that a gene begins in the region immediately downstream.

*Ab initio* gene prediction with eukaryotic genomes remains an inefficient process, despite the increasing sophistication of computer programs that have been developed for this task. For most genomes, the starts and ends of genes can be predicted with almost 100% accuracy, but the accuracy in identifying exon–intron boundaries is much lower, usually only 60–70%. These figures assume that there is some *a priori* knowledge of parameters such as codon bias. If the genome is completely unstudied, then the accuracy of gene prediction will be lower, even though most gene prediction software includes a machine learning function, so the computer becomes trained to recognize appropriate patterns of codon usage as it gradually builds up the genome annotation.

### Locating genes for noncoding RNA

ORF scanning is appropriate for protein-coding genes, but what about those genes for noncoding RNAs such as rRNA and tRNA (*Section 1.2*)? These genes do not comprise open reading frames and hence will not be located by the methods described above. Noncoding RNA molecules do, however, have their own distinctive features, which can be used as an aid in their discovery in a genome sequence. The most important of these features is the ability to fold into a secondary structure, such as the *cloverleaf* adopted by tRNA molecules (*Figure 5.6A*). These secondary structures are held together by base pairing not between two separate polynucleotides, as in the DNA double helix, but by *intramolecular base pairing* between different parts of the same polynucleotide. In order for intramolecular base pairs to form, the nucleotide sequences in the two parts of the molecule must be complementary, and to produce a complex structure such as the cloverleaf, the components of these pairs of complementary sequences must be arranged in a characteristic order within the RNA sequence (*Figure 5.6B*). These features provide a wealth of information that can be used to locate tRNA genes in a genome sequence, and programs designed for this specific purpose are usually very successful.

As well as tRNAs, rRNAs and some of the short noncoding RNAs (*Section 1.2*) also adopt secondary structures that have sufficient complexity to enable their genes to be identified without too much difficulty. Other noncoding RNA genes are less easy to locate because the RNAs take up structures that involve relatively little base pairing or the base pairing is not in a regular pattern. Three approaches are used for location of the genes for these RNAs:

- Although some noncoding RNAs do not adopt complex secondary structures, most contain one or more *stem-loops* (or *hairpins*), which result from the simplest type of intramolecular base pairing (*Figure 5.7*). Programs that scan DNA sequences for such structures therefore identify regions where noncoding RNA genes might be present. These programs incorporate thermodynamic rules that enable the stability of a stem–loop to be estimated, taking into account features such as the size of the loop, the number of base pairs in the stem, and the proportion of G-C base pairs (recall that these are more stable than A-T pairs because they are held together by three rather than two hydrogen bonds; see *Figure 1.9*). A putative stem–loop structure with an estimated stability above a chosen limit is considered a possible indicator of the presence of a noncoding RNA gene.

#### Figure 5.6 The distinctive features of tRNAs aid in location of the genes for these noncoding RNAs. (A) All tRNAs fold into the cloverleaf structure, which is held together by intramolecular base pairing in the four highlighted regions. (B) The DNA sequence for the gene for one of the *Escherichia coli* tRNAs specific for the amino acid leucine is shown. Highlighted segments correspond to the regions of intramolecular base pairing shown in part A. The sequence constraints imposed by the need for these segments to be able to base-pair to one another provide features that can be searched for by computer programs designed to locate tRNA genes.
• As with protein-coding genes, a search can be made for regulatory sequences associated with genes for noncoding RNAs. These regulatory sequences are different than those for protein-coding genes and may be present within a noncoding RNA gene as well as upstream of it.

• In compact genomes, attention is directed toward regions that remain after a comprehensive search for protein-coding genes. Often these empty spaces are not empty at all, and a careful examination will reveal the presence of one or more noncoding RNA genes.

Homology searches and comparative genomics give an extra dimension to gene prediction

The limitations of ab initio gene prediction can be offset to a certain extent by the use of a homology search to test whether a series of triplets is a real exon or a chance sequence. In this analysis, the DNA databases are searched to compare the test sequence with genes that have already been sequenced. If the test sequence is part of a gene that has already been sequenced by someone else, then an identical match will be found, but this is not the point of a homology search. Instead the intention is to determine if an entirely new sequence is similar to any known genes, because if it is, then there is a chance that the test and match sequences are homologous, meaning that they represent genes that are evolutionarily related. The main use of homology searching is to assign functions to newly discovered genes, and we will therefore return to it when we deal with this aspect of genome analysis in the next chapter (Section 6.1). The technique is also central to gene prediction because it enables the authenticity of tentative exon sequences located by ORF scanning to be tested. If the tentative exon sequence gives one or more positive matches after a homology search then it is probably a real exon, but if it gives no match then its authenticity must remain in doubt until it is assessed by one or other of the experiment-based genome annotation techniques.

A more precise version of homology searching is possible when genome sequences are available for two or more related species. Related species have genomes that share similarities inherited from their common ancestor, overlaid with species-specific differences that have arisen since the species began to evolve independently (Figure 5.8). Because of natural selection, the sequence similarities between related genomes are greatest within the genes and lowest in the intergenic regions. Therefore, when related genomes are compared, homologous genes are easily identified because they have high sequence similarity, and any ORF that does not have a clear homolog in the second genome can be discounted as almost certainly being a chance sequence and not a genuine gene. This value of this type of analysis, called comparative genomics, was illustrated during annotation of the genome of the yeast Saccharomyces cerevisiae. Complete or partial sequences were available not only for this yeast but also for various other members of the

![Figure 5.7 A typical RNA stem–loop structure.](image)

![Figure 5.6A Gene organization](image)

![Figure 5.8 Related species have similar genomes.](image)
Chapter 5: Genome Annotation

5.2 Genome Annotation by Analysis of Gene Transcripts

Saccharomycetes, including *Saccharomyces paradoxus*, *Saccharomyces mikatae*, and *Saccharomyces bayanus*, the species most closely related to *S. cerevisiae*. Comparisons between these genomes confirmed the authenticity of a number of *S. cerevisiae* ORFs and also enabled almost 500 putative ORFs to be removed from the *S. cerevisiae* catalog on the grounds that they have no equivalents in the related genomes. The analysis is made even more powerful by the **synteny**, or conservation of gene order, displayed by the genomes of these related yeasts. Although each genome has undergone its own species-specific rearrangements, there are still substantial regions where the gene order in the *S. cerevisiae* genome is the same as in one or more of the related genomes. This makes it very easy to identify homologous genes, but more importantly, it enables a spurious ORF, especially a short one, to be discarded with great confidence, because its expected location in a related genome can be searched in detail to ensure that no equivalent is present (Figure 5.9).

Synteny is also displayed by other related groups of species. A second important example occurs among the grasses, including commercially important cereals such as barley, maize, sorghum, sugar cane, foxtail millet, and rice. The rice genome, which at 430 Mb is the smallest of this group, was sequenced in 2005. The wild grass *Brachypodium distachyon* has an even smaller genome, of 270 Mb, and this also has been sequenced and shown to be syntenic with the larger cereal genomes. The synteny within this family has been important in annotating the genomes of those cereals that have been sequenced and has also been utilized in establishment of resources such as the barley gene space (Section 4.4). Of the 24,154 barley genes whose sequences were obtained by RNA-seq and cDNA sequencing, 3743 were assigned their positions in the barley gene space not by identification of overlaps with sequences present in the sequence-enriched map, but by inferring the map positions of those genes by comparison with the equivalent parts of the *Brachypodium*, *rice*, and sorghum genomes.

**5.2 GENOME ANNOTATION BY ANALYSIS OF GENE TRANSCRIPTS**

The second approach to genome annotation makes use of experimental techniques to locate genes within a genome sequence. These methods are not usually based on direct examination of DNA molecules but instead rely on detection of the RNA molecules that are transcribed from genes. All genes are transcribed into RNA, and if the gene is **discontinuous**, then the **primary transcript** is subsequently processed to remove the introns and link up the exons (Section 12.4). Techniques that map the positions of transcribed sequences in a DNA fragment can therefore be used to locate exons and entire genes. One problem to be kept in mind is that the transcript is usually longer than the coding part of the gene because it begins several tens of nucleotides upstream of the initiation codon and continues several tens or hundreds of nucleotides downstream of the termination codon (Figure 5.10). Because of these upstream and downstream **untranslated regions** (UTRs), transcript analysis does not give a precise definition of the start and end of the coding region of a gene, but it does tell you that a gene is present in a particular region and it can locate the exon–intron boundaries. Often this is sufficient information to enable the coding region to be delineated.

**Figure 5.9** Using comparisons with syntenic genomes to test the authenticity of a short ORF. In this example, the ORF is present in three of the four related genomes and hence is likely to be a genuine gene.

**Figure 5.10** A transcript is longer than a gene.
Hybridization tests can determine if a fragment contains transcribed sequences

The simplest procedures for studying transcribed sequences are based on hybridization analysis. RNA molecules can be separated by specialized forms of agarose gel electrophoresis, transferred to a nitrocellulose or nylon membrane, and examined by the process called northern hybridization. This differs from Southern hybridization (Section 2.1) only in the precise conditions under which the transfer is carried out, and the fact that it was not invented by a Dr. Northern and so does not have a capital N. If a northern blot of cellular RNA is probed with a labeled fragment of the genome, then RNAs transcribed from genes within that fragment will be detected (Figure 5.11). Northern hybridization is therefore, theoretically, a means of determining the number of genes present in a DNA fragment and the size of each coding region. There are two weaknesses with this approach:

- Some individual genes give rise to two or more transcripts of different lengths because some of their exons are optional and may or may not be retained in the mature RNA (Section 12.4). If this is the case, then a fragment that contains just one gene could detect two or more hybridizing bands in the northern blot. A similar problem can occur if the gene is a member of a multigene family (Section 7.3).

- With many species, it is not practical to make an mRNA preparation from an entire organism, so the extract is obtained from a single organ or tissue. Any genes not expressed in that organ or tissue will not be represented in the RNA population and so will not be detected when the RNA is probed with the DNA fragment being studied. Even if the whole organism is used, not all genes will give hybridization signals, because many are expressed only at a particular developmental stage and others are weakly expressed, meaning that their RNA products are present in amounts too low to be detected by hybridization analysis.

A second type of hybridization analysis avoids the problems with poorly expressed and tissue-specific genes by searching not for RNAs but for related sequences in the DNAs of other organisms. This approach, like homology searching, is based on the fact that homologous genes in related organisms have similar sequences, whereas the intergenic DNA is usually quite different. If a DNA fragment from one species is used to probe a Southern transfer of DNAs from related species, and one or more hybridization signals are obtained, then it is likely that the probe contains one or more genes (Figure 5.12). This is called zoo-blotting.

**Figure 5.11 Northern hybridization.** An RNA extract is electrophoresed under denaturing conditions in an agarose gel. After ethidium bromide staining, two bands are seen. These are the two largest rRNA molecules (Section 1.2), which are abundant in most cells. The smaller tRNAs, which are also abundant, are not seen because they are so short that they run out the bottom of the gel, and in most cells, none of the mRNAs is abundant enough to form a band visible after ethidium bromide staining. The gel is blotted onto a nylon membrane and, in this example, probed with a radioactively labeled DNA fragment. A single band is visible on the autoradiograph, showing that the DNA fragment used as the probe contains part or all of one transcribed sequence.

**Figure 5.12 Zoo-blotting.** The objective is to determine if a fragment of human DNA hybridizes to DNAs from related species. Samples of human, chimpanzee, cow, and rabbit DNAs are therefore prepared, restricted, and electrophoresed in an agarose gel. Southern hybridization is then carried out with a human DNA fragment as the probe. A positive hybridization signal is seen with each of the animal DNAs, suggesting that the human DNA fragment contains an expressed gene. Note that the hybridizing restriction fragments from the cow and rabbit DNAs are smaller than the hybridizing fragments in the human and chimpanzee samples. This indicates that the restriction map around the transcribed sequence is different in cows and rabbits but does not affect the conclusion that a homologous gene is present in all four species.
**Methods are available for precise mapping of the ends of transcripts**

Northern hybridization and zoo-blotting can identify a DNA fragment that contains a gene, but those methods do not enable the gene to be positioned within that fragment with any degree of accuracy. More precise mapping of the transcript onto the DNA sequence requires a different approach. One possibility is a special type of PCR that uses RNA rather than DNA as the starting material. The first step in this type of PCR is to convert the RNA into single-stranded cDNA with reverse transcriptase (see Figure 3.35), after which the cDNA is amplified with Taq polymerase in the same way as in a normal PCR. These methods go under the collective name of **reverse transcriptase polymerase chain reaction (RT-PCR)**, but the particular version that interests us at present is **rapid amplification of cDNA ends (RACE)**. In the simplest form of this method, one of the primers is specific for an internal region of the gene being studied, close to its start site. This primer attaches to the mRNA for the gene and directs the first reverse transcriptase-catalyzed stage of the process, during which a cDNA copy of the start of the mRNA is made, the 3′-end of this cDNA corresponding exactly with the 5′-end of the mRNA (Figure 5.13). The 3′-end of the cDNA is then extended by treatment with terminal deoxynucleotidyl transferase to give a short poly(A) tail. The second primer anneals to this poly(A) sequence and, during the first round of the normal PCR, converts the single-stranded cDNA into a double-stranded molecule, which is subsequently amplified as the PCR proceeds. The sequence of this amplified molecule will reveal the position of the start of the clone.

Other approaches for precise transcript mapping involve **heteroduplex analysis**. This method requires a single-stranded version of the gene being studied, which can be obtained by cloning in a vector based on **M13 bacteriophage**. The replication process of M13 involves synthesis of phage particles that contain a single-stranded copy of the phage genome. DNA that has been cloned in an M13 vector can therefore be obtained as single-stranded DNA by purification from the recombinant phages. When mixed with an appropriate RNA preparation, the transcribed sequence in the single-stranded DNA hybridizes with the equivalent mRNA, forming a double-stranded heteroduplex. In the example shown in Figure 5.14, a restriction fragment spanning the start of an mRNA has been cloned. Some of the cloned fragment participates in the heteroduplex, but the rest does not. The single-stranded regions can be digested by treatment with a single-strand-specific nuclease such as S1. The size of the heteroduplex is determined by degrading the RNA component with alkali and electrophoresing the resulting single-stranded DNA in an agarose or polyacrylamide gel. This size measurement is then used to position the start of the transcript relative to the restriction site at the end of the cloned fragment.

**Exon–intron boundaries can also be located with precision**

Heteroduplex analysis can also be used to locate exon–intron boundaries. The method is almost the same as that shown in Figure 5.14 with the exception that

**Figure 5.13** RACE, or rapid amplification of cDNA ends. The RNA being studied is converted into a partial cDNA by extension of a DNA primer that anneals at an internal position close to the 5′-end of the molecule. The 3′-end of the cDNA is further extended by treatment with terminal deoxynucleotidyl transferase in the presence of dATP, which results in a series of As being added to the cDNA. This series of As acts as the annealing site for the anchor primer. Extension of the anchor primer leads to a double-stranded DNA molecule, which can now be amplified by standard PCR. This is 5′-RACE, so called because it results in amplification of the 5′-end of the starting RNA. A similar method, 3′-RACE, can be used if the 3′-end sequence is desired.
the cloned restriction fragment spans the exon–intron boundary being mapped rather than the start of the transcript.

A second method for finding exons in a genome sequence is called exon trapping. This requires a special type of vector that contains a minigene consisting of two exons flanking an intron sequence, the first exon being preceded by the sequence signals needed to initiate transcription in a eukaryotic cell (Figure 5.15). To use the vector, the piece of DNA to be studied is inserted into a restriction site located within the vector’s intron region. The vector is then introduced into a suitable eukaryotic cell line, where it is transcribed and the RNA produced from it is spliced. The result is that any exon contained in the genomic fragment becomes attached between the upstream and downstream exons from the minigene. RT-PCR with primers annealing within the two minigene exons is now used to amplify a DNA fragment, which is sequenced. As the minigene sequence is already known, the nucleotide positions at which the inserted exon starts and ends can be determined, precisely delineating this exon.

5.3 ANNOTATION BY GENOMEWIDE RNA MAPPING

The transcript analysis methods that we have studied so far are designed for the mapping of individual genes onto short sequences of DNA. As there are tens of thousands of genes in the average vertebrate or plant genome, reliance on these methods for genome annotation would make RNA mapping a lengthy and highly tedious process. We therefore need to explore alternative methods that enable multiple transcripts to be mapped simultaneously.
Tiling arrays enable transcripts to be mapped onto chromosomes or entire genomes

The first methods for multiple, parallel transcript mapping made use of tiling arrays. A tiling array is a special type of DNA chip. We learned in Section 3.2 that a DNA chip is a small piece of glass or silicon onto which many different oligonucleotides have been immobilized in an ordered array, and we saw how, by hybridizing a DNA sample to the chip, the SNPs within that DNA could be typed (see Figure 3.9). In a tiling array, the oligonucleotides form a series that covers the length of a chromosome sequence or the sequence of an entire genome, the series either overlapping or having small gaps between adjacent oligonucleotides (Figure 5.16). To squeeze all the necessary oligonucleotides onto the chip, a high-density array is needed. In the traditional technology, each oligonucleotide was synthesized separately and then spotted onto the chip at its appropriate position. This method is suitable for preparing low-density arrays for typing a relatively small number of SNPs, but it does not enable a high-density tiling array to be prepared. To achieve high density, the oligonucleotides must be synthesized directly on the surface of the chip. This presents a challenge, as the normal synthesis method involves adding nucleotides one-by-one to the growing end of an oligonucleotide, with the sequence determined by the order in which the nucleotide substrates are added to the reaction mixture. If used for synthesis on a chip, this method would result in every oligonucleotide having the same sequence. Instead, modified nucleotide substrates are added that have to be light-activated before they will attach to the end of a growing oligonucleotide. The nucleotides are added one after another to the chip surface, with photolithography used to direct pulses of light onto individual positions in the array. Only the oligonucleotides that are light-activated will be extended by the nucleotide that is present at any particular step (Figure 5.17). This method enables the highest density arrays, with up to 300,000 oligonucleotides per square centimeter, to be prepared.

The tiling array is hybridized not to DNA, as would be the case with SNP typing, but to a labeled sample of RNA from the organism whose genome is being annotated. Those positions on the array that hybridize will be ones containing oligonucleotides that hybridize to molecules in the RNA samples, and hence they will reveal the positions in the genome of transcribed sequences. The hybridization data will not accurately locate each gene, for two reasons. First, as described above, the transcript might be longer than the gene, so oligonucleotides whose positions lie in the upstream and downstream UTRs will also give signals. Second, the accuracy of mapping depends on the lengths of the oligonucleotides and of the overlaps or gaps between them in the array. In the examples of array design shown in Figure 5.16, the accuracy is ±30 for the overlapping array and ±70 nucleotides for the gapped array.

The source of the RNA preparation also has to be chosen with care to ensure that the sample contains transcripts of as many genes as possible. With a higher eukaryote such as humans, it is very difficult, if not impossible, to obtain a sample that contains transcripts of every gene in the genome, because not all genes are expressed in a single organ or cell type, and even in one cell type the gene expression pattern varies over time. In fact, the major use of tiling arrays is in understanding these cell-specific and time-dependent patterns of gene expression. It is therefore necessary to test RNA samples from many different tissues in order to build up a complete genome annotation by this, or any other, genomewide method for transcript mapping. The RNA that is prepared can also be fractionated...
in various ways so that only certain types of genes are targeted by the tiling array. The most commonly used fractionation procedure is to preselect RNAs that carry a poly(A) tail at their 3’-ends. The tail is a series of up to 250 adenine nucleotides added to the end of a eukaryotic mRNA after transcription (Section 1.2). An affinity chromatography column containing oligo(dT)–cellulose (cellulose beads to which short oligonucleotides of thymidine have been attached) can therefore be used to purify the polyadenylated mRNA fraction from a eukaryotic RNA sample (Figure 5.18).

**Transcript sequences can be directly mapped onto a genome**

The most direct way of using transcript analysis in genome annotation is to sequence those transcripts and then to use the sequence data to search the genome for the genes from which the RNAs are transcribed. On a small scale, this approach is possible by sequencing the inserts in libraries of cloned cDNAs. The utility of cDNA library sequencing depends on two factors. The first concerns the frequency of the desired cDNAs in the library. Most libraries are dominated by identical clones representing genes that are expressed at the highest level in the cells from which the RNA was obtained. In contrast, clones representing rare transcripts, from genes with low expression levels, will be infrequent in the library and it might be necessary to screen many clones before the desired one is identified. If the genes in, for example, a single bacterial artificial chromosome (BAC) clone are being mapped, then one way around this problem is to use cDNA capture or cDNA selection to enrich the library for the desired clones. The BAC fragment is repeatedly hybridized to the pool of cDNAs, with nonhybridized cDNAs washed away and discarded. Because the cDNA pool contains so many different sequences, it is generally not possible to discard all the irrelevant clones by these repeated hybridizations, but it is possible to increase significantly the frequency of those clones that specifically hybridize to the DNA fragment. This reduces the size of the library that must subsequently be screened under stringent conditions to identify the desired clones.

The second factor that determines the success or failure of cDNA sequencing in genome annotation is the completeness of the individual cDNA molecules. Usually, cDNAs are made by copying RNA molecules into single-stranded DNA with reverse transcriptase and then converting the single-stranded DNA into double-stranded DNA with a DNA polymerase (see Figure 3.35). There is always a chance that one or other of the strand-synthesis reactions will not proceed to completion, resulting in a truncated cDNA. The presence of intramolecular base pairs in the RNA can also lead to incomplete copying. The sequences of truncated cDNAs can be used to locate genes in a DNA sequence, but they may lack the sequences needed to delineate the start and end points of the gene or the exact positions of exon–intron boundaries.

Since the development of next-generation sequencing, methods for genome annotation using RNA-seq data have become increasingly attractive. RNA-seq is simply the application of Illumina or some other high-throughput sequencing method to a library that has been prepared from cDNA rather than directly from DNA. The sequence reads therefore correspond to segments of the transcripts in the original RNA sample. Those reads can be mapped directly onto a genome sequence, in a manner identical to the way in which DNA sequence reads are mapped onto a reference genome during a genome resequencing project (Section 4.3). The only difference is that the RNA reads do not give lengthy scaffolds but instead form clusters that map specifically onto the transcribed parts of the genome (Figure 5.19A). An alternative strategy that achieves the same result is to apply a de novo assembly method to the collection of RNA-seq reads and then map the assembled contigs onto the reference genome (Figure 5.19B). The advantage with the latter approach is that many genes are members of multigene families, the members of which display sequence similarity (Section 7.3). If individual, short RNA reads are mapped directly onto the reference genome, then some might be identical to segments of two or more members of a multigene
family, complicating the mapping process. If, on the other hand, the complete transcript sequence is determined prior to mapping, then the members of a gene family are easily distinguished.

RNA-seq mapping is becoming increasingly important in genome annotation and in studies of gene expression patterns in different tissues. The analysis is, however, computationally intensive, and methods that still make use of RNA-seq data but take shortcuts to gene mapping are constantly being sought. One such method is cap analysis gene expression (CAGE). The basis to this method is the presence of the cap structure at the 5'-end of a eukaryotic mRNA. The cap is a 7-methylguanine nucleotide that is added to the start of the mRNA after transcription, becoming attached to the first nucleotide of the transcript by an unusual 5'-5' phosphodiester bond (Section 1.2). The cap is utilized in the CAGE method in the following way. The first stage of the standard cDNA synthesis process is carried out, creating an RNA–DNA hybrid (Figure 5.20). The cap structure is still attached to the RNA component of the hybrid, at least for those molecules that are either full-length copies or truncated fragments that include the start of the RNA. Because the 7-methylguanine within the cap structure is attached to the next nucleotide by a 5'-5' bond, both its 2'- and 3'-carbons retain their hydroxyl groups. These two hydroxyls form a diol structure. Addition of an oxidizing agent specifically breaks the bond between the 2'- and 3'-carbons of the diol, creating an open-chain sugar structure that can be covalently attached to the small organic molecule called biotin. This enables the capped cDNAs to be captured, through use of magnetic beads coated with avidin, a protein from egg white that has a high binding affinity for biotin. Following capture, the RNA component of the hybrids is degraded and the second cDNA strand is synthesized. In low-throughput versions of CAGE, the cDNA is now sequenced and the sequence, corresponding to the 5'-end of the transcript, is mapped. Innovations designed for RNA-seq extend the CAGE method a little further by ensuring that the end fragments that are sequenced are all very short. Prior to second-strand synthesis, a short oligonucleotide containing a recognition sequence for the restriction enzyme EcoP151 is ligated to the start of the cDNA (Figure 5.21). EcoP151 is a type III restriction endonuclease (Section 2.1) that makes a cut not within the recognition site but 27 base pairs away from that site. After second-strand synthesis, the enzyme is added. The cDNA is cut, releasing a 27 bp tag fragment representing the 5'-end of the original mRNA. All the tags from all the cDNAs in the sample are collected, sequenced, and mapped onto the reference genome. Although CAGE is methodologically complex, the computational phase of the procedure is less intensive than mapping of reads from complete RNA-seq libraries. CAGE therefore holds promise as a means of rapidly identifying the start regions of genes in a genome sequence.
5.4 Genome Browsers

The most convenient way to display the results of a genome annotation project is in a graphical format, with the DNA sequence forming the x-axis and the positions of genes and other interesting features marked at their appropriate map positions. A genome browser is a software package that enables genome annotation data to be displayed in this way. The display might be quite complex, because as well as indicating the positions of ORFs that have been identified with high confidence, the browser might also display short ORFs of questionable authenticity in all six reading frames, noncoding RNA genes, the starts and ends of transcripts, positions of mapped DNA markers, locations and identities of repetitive DNA sequences, etc. The software will enable the map to be displayed at different levels of magnification, so the entire length of a chromosome can be viewed, or the operator can zoom in to a level at which individual nucleotides are distinguished. Most browsers also incorporate a search function so that particular genes, markers, or map positions can be quickly located.

Many genome browsers are used online, with the draft and final annotations of a newly sequenced genome accessible to other researchers. One of the imperatives of genomics research is that data must be made publicly available. Databases for the curation of DNA sequences have been established for many years. The most important is GenBank, which is maintained by the National Center for Biotechnology Information (NCBI), part of the US National Institutes of Health. Online genome browsers fulfill the same purpose for genome annotations. Two of the most widely used are Ensembl, which is maintained by the European Bioinformatics Institute and the UK Sanger Institute, and the UCSC Genome Browser of the University of California, Santa Cruz (Figure 5.22). Both Ensembl and the UCSC Genome Browser hold annotations of the human genome along with those of several other vertebrates and invertebrates. There are also more specialized online genome browsers, such as the Plant GDB, which holds plant genome annotations.

**Figure 5.21** Rapid sequencing of cDNA ends following CAGE. The single-stranded cDNA resulting from the CAGE process shown in Figure 5.20 is ligated to an oligonucleotide containing an EcoP151 restriction site. Following second-strand synthesis, treatment with EcoP151 releases a tag fragment. Collection of all the tag fragments followed by RNA-seq gives the 5′-terminal sequences for all the mRNAs in the original sample.

**Figure 5.22** An example of the information provided by an online genome browser. This example shows the annotation for human chromosome 1 between nucleotide positions 16,532,000 and 16,534,500, as displayed by the UCSC Genome Browser (redrawn slightly). The user can configure the browser to show many different features of genome annotation. This configuration shows the locations of protein-coding genes, with exons as boxes and introns as lines, various types of repeated DNA sequence, tRNA genes, SNPs, and STS markers. In the bottom panel the GC content is indicated, based on a sliding window of 5 bp.
Chapter 5: Genome Annotation

**SUMMARY**

- Genome annotation is the process by which genes are located in a genome sequence.
- Protein-coding genes can be identified by searching for open reading frames, though this is complicated in eukaryotes by the presence of introns, whose boundary sequences are variable and which therefore cannot be identified accurately.
- Genes for noncoding RNAs can be located by searching for their characteristic features, primarily the ability of the RNAs to fold into secondary structures based on the formation of base-paired stem–loops.
- Genes can also be located by homology analysis, which uses the presence of an equivalent gene in a second genome as evidence that a putative gene in the test genome is genuine.
- Experimental methods for gene location are based on detection of RNA molecules transcribed from the genome. These techniques include transcript mapping by reverse-transcriptase PCR (RT-PCR) or heteroduplex analysis.
- Exon–intron boundaries can be identified experimentally by the procedure called exon trapping.
- Tiling arrays are used for the genomewide mapping of transcript positions.
- RNA sequencing using next-generation methods, including cap analysis gene expression, is becoming increasingly important in the identification of the transcribed parts of a genome.
- Genome annotations are displayed using a genome browser.

**SHORT ANSWER QUESTIONS**

1. Why is it relatively easy to identify ORFs in prokaryotic genomes by computer analysis?
2. Describe how ORF scans are used to search for genes in eukaryotic genome sequences.
3. What is meant by the term “codon bias”?
4. Outline the structural features of functional RNA molecules, such as tRNA and rRNA, that can be searched for in a genome sequence to identify the genes encoding these RNA molecules.
5. Define the term “homologous”, as used when comparing gene sequences.
6. Give examples of the use of comparative genomics in genome annotation.
7. What are the two limitations that arise when northern analysis is used to determine the number of genes present in a DNA fragment?
8. Describe how rapid amplification of cDNA ends (RACE) is used to map the transcription initiation site of a gene.
9. How is heteroduplex analysis used to map transcripts?
11. What is a tiling array, and how are tiling arrays used in genome annotation?
12. Explain how the computational challenges of RNA-seq are reduced by the method called cap analysis gene expression (CAGE).
IN-DEPTH PROBLEMS

1. To what extent do you believe it will be possible in future years to use bioinformatics to obtain a complete description of the locations and functions of the protein-coding genes in a eukaryotic genome sequence?

2. Devise a hypothesis to explain the codon biases that occur in the genomes of various organisms. Can your hypothesis be tested?

3. “Comparative genomics has an important role to play in the study of disease genes.” Evaluate this statement.

4. Studies with tiling arrays have revealed that many transcripts are synthesized from the sequences in the intergenic spaces lying between the open reading frames in the human genome. Use the internet (and possibly other chapters of this book) to explore the nature and possible functions of these RNAs.

5. Use the Ensembl Bacteria genome browser (http://bacteria.ensembl.org/index.html) to locate the position of the β-galactosidase gene in the Escherichia coli genome. Draw (or export) a map of the genes in the 30 kb region centered on the β-galactosidase gene.

FURTHER READING

Gene location by computer analysis

Comparative genomics

Experimental methods for gene location

Genomewide RNA mapping

URLs for genome browsers
UCSC Genome Browser. https://genome.ucsc.edu/
Plant GDB. http://www.plantgdb.org/
Once a new gene has been located in a genome sequence, the question of its function has to be addressed. This is turning out to be an important area of genomics research, because completed sequencing projects have revealed that we know rather less than we thought about the content of individual genomes. *Escherichia coli* and *Saccharomyces cerevisiae*, for example, were studied intensively by conventional genetic analysis before the advent of sequencing projects, and geneticists were at one time fairly confident that most of the genes in these species had been identified. The genome sequences revealed that in fact there are large gaps in our knowledge. Of the 4288 protein-coding genes in the initial annotation of the *E. coli* genome sequence, only one-third were described as well characterized and 38% had no attributed function. The figures were very similar for *S. cerevisiae*. Methods that enable functions to be assigned to genes are therefore of critical importance in understanding a genome sequence.

As with genome annotation, attempts to determine the functions of unknown genes are made by computer analysis and by experimental studies. We begin with the computer methods.

### 6.1 COMPUTER ANALYSIS OF GENE FUNCTION

We have already seen that computer analysis plays an important role in locating genes in DNA sequences and that one of the most powerful tools available for this purpose is homology searching, which locates genes by comparing the DNA sequence under study with all the other DNA sequences in a database. The basis of homology searching is that related genes have similar sequences and so a new gene can be discovered by virtue of its similarity to an equivalent, already sequenced gene from a different organism. Now we will look more closely at homology analysis and see how it can be used to assign a function to a new gene.

**Homology reflects evolutionary relationships**

Homologous genes are ones that share a common evolutionary ancestor, revealed by sequence similarities between the genes. These similarities form the data on which molecular phylogenies are based, as we will see in Section 18.4. Homologous genes fall into two categories (Figure 6.1):

- **Orthologous** genes are homologs present in different organisms whose common ancestor predates the split between the species. Orthologous genes usually have the same or very similar functions. For example, the myoglobin genes of humans and chimpanzees are orthologs.

![Figure 6.1 Orthologous and paralogous genes.](image)
Section 7.3

Paralogous genes are present in the same organism, often as members of a recognized multigene family (Section 7.3); their common ancestor may or may not predate the species in which the genes are now found. For example, the myoglobin and β-globin genes of humans are paralogs: they originated by duplication of an ancestral gene some 550 million years ago (Section 18.2).

Two homologous genes do not usually have identical nucleotide sequences, because the two genes undergo different random changes by mutation, but they have similar sequences because these random changes have operated on the same starting sequence, the common ancestral gene. Homology searching makes use of these sequence similarities. The basis of the analysis is that if a newly sequenced gene turns out to be similar to a previously sequenced gene, then an evolutionary relationship can be inferred and the function of the new gene is likely to be the same, or at least similar, to the function of the known gene.

It is important not to confuse the words homology and similarity. It is incorrect to describe a pair of related genes as 80% homologous if their sequences have 80% nucleotide identity (Figure 6.2). Two genes are either evolutionarily related or they are not; there are no in-between situations, and it is therefore meaningless to ascribe a percentage value to homology.

Homology analysis can provide information on the function of a gene

A homology search can be conducted with a DNA sequence, but usually a tentative gene sequence is converted into an amino acid sequence before the search is carried out. One reason for this is that there are 20 different amino acids in proteins but only four nucleotides in DNA, so genes that are unrelated usually appear to be more different from one another when their amino acid sequences are compared (Figure 6.3). A homology search is therefore less likely to give spurious results if the amino acid sequence is used.

A homology search program begins by making alignments between the query sequence and sequences from the databases. For each alignment, a score is calculated from which the operator can gauge the likelihood that the query and test sequences are homologs. There are two ways of generating the score:

- The simplest programs count the number of positions at which the same amino acid is present in both sequences. This number, when converted into a percentage, gives the degree of identity between two sequences.
- More sophisticated programs use the chemical relatedness between non-identical amino acids to assign a score to each position in the alignment, a higher score for identical or closely related amino acids (for example, leucine and isoleucine, or aspartic acid and asparagine) and a lower score for less closely related amino acids (for example, cysteine and tyrosine, or phenylalanine and serine). This analysis determines the degree of similarity between a pair of sequences.

To achieve the highest possible score, the algorithm introduces gaps at various positions in one or both sequences, up to limits set by the operator. This approach

![Figure 6.2](image_url)

**Figure 6.2** Two DNA sequences with 80% sequence identity.

| Sequence 1 | GGTGAGGTTATCATCCACATCTGACTACCTCATTGGAGAGCGAGCACTG |
| Sequence 2 | GGTGAGGTTATCATCCACATCTGACTACCTCATTGGAGAGCGAGCACTG |
| Identities | *** *** *** ** *****  ********* ***  *** ********* |

![Figure 6.3](image_url)

**Figure 6.3** Lack of homology between two sequences is often more apparent when comparisons are made at the amino acid level. Two nucleotide sequences are pictured, with nucleotides that are identical in the two sequences shown in green and nonidentities shown in pink. The two nucleotide sequences are 76% identical, as indicated by the asterisks. This might be taken as evidence that the sequences are homologous. However, when the nucleotide sequences are translated into amino acids, the identity decreases to 28%. Identical amino acids are shown in gold and nonidentities shown in red. The comparison between the amino acid sequences suggests that the genes are not homologous and that the similarity at the nucleotide level was fortuitous. The amino acid sequences have been written with one-letter abbreviations (see Table 1.2).
parallels processes thought to occur during the evolution of genes, when blocks of nucleotides coding for individual or adjacent amino acids may have been inserted into or deleted from genes.

The practicalities of homology searching are not at all daunting. Several software programs exist for this type of analysis, the most popular of which is BLAST (Basic Local Alignment Search Tool). The analysis can be carried out simply by logging on to the website for one of the DNA databases and entering the sequence into the online search tool. The standard BLAST program is efficient at identifying homologous genes that have more than 40% sequence similarity but is less effective at recognizing evolutionary relationships if the similarity is lower than this amount. The modified version called PSI-BLAST (position-specific iterated BLAST) identifies more distantly related sequences by combining the homologous sequences from a standard BLAST search into a profile, the features of which are used to identify additional homologous sequences that were not detected in the initial search.

Homology searching with BLAST and similar programs has gained immense importance in genomics research, but its limitations must be recognized. A growing problem is the presence in the databases of genes whose stated functions are incorrect. If one of these genes is identified as a homolog of the query sequence, then the incorrect function will be passed on to this new sequence, adding to the problem. There are also several cases where homologous genes have quite different biological functions: an example is the crystallins of the eye lens, some of which are homologous to metabolic enzymes. Homology between a query sequence and a crystallin therefore does not mean that the query sequence is a crystallin, and similarly, an apparently clear homology between a query sequence and a metabolic enzyme might not mean that the query sequence is a metabolic enzyme.

**Identification of protein domains can help to assign function to an unknown gene**

What if a homology search with the DNA or amino acid sequence of an unknown gene fails to reveal any matches in the databases? All is not lost, as it may be possible to deduce at least some part of the function of the gene by searching the amino acid sequence for motifs that encode protein **domains** of known function. A protein domain is a segment of a protein that possesses a characteristic tertiary structure that provides the protein with a particular biochemical function. An example of a protein domain is the **Cys2His2 zinc finger**, which comprises a series of 12 or so amino acids, including two cysteines and two histidines, that form a segment of β-sheet followed by an α-helix. These two structures, which form the finger projecting from the surface of the protein, hold between them a bound zinc atom, coordinated to the two cysteines and two histidines (Figure 6.4). Zinc fingers are DNA-binding structures (Section 11.2), so identification in an unknown gene of an amino acid sequence that can encode a zinc finger indicates that the gene codes for a DNA-binding protein. As well as function, the amino acid sequence can also provide information on the subcellular location of a protein. This can be inferred by searching for motifs called **sorting sequences**, which direct proteins to organelles such as the nucleus or mitochondria or might specify that the protein is secreted from the cell.

As with homology searching, identification of conserved sequence motifs in an unknown gene can be performed online, by use of search tools at protein structure databases such as PROSITE, which is maintained by the Swiss Institute of Bioinformatics. The results of such an analysis must, however, be interpreted with care. The presence of a shared domain indicates that two proteins can perform a similar biochemical activity, but that does not necessarily mean that the proteins have similar overall functions. This point is illustrated by the **Tudor domain** family of proteins. As the name implies, these proteins are related in that each possesses one or more copies of the Tudor domain, a five-stranded β-sheet structure encoded by a sequence of approximately 60 amino acids. The Tudor domain binds...
to methylated arginine and/or lysine amino acids contained in other proteins. This is a specific biochemical activity, but it is associated with a variety of different protein functions. The first Tudor domain protein to be discovered, coded by the Drosophila melanogaster gene called tudor, is involved in synthesis of piwi-interacting RNAs (piRNAs) in the developing oocyte. These are short noncoding RNAs that bind to piwi proteins, forming complexes that regulate gene expression during various developmental processes (Section 12.1). Other Tudor domain proteins, in a range of species, are also involved in piRNA synthesis, but other members of the family are associated with RNA splicing (Section 12.4), the RNA interference pathway (Section 12.3), the response to DNA damage (Section 16.2), and histone modification (Section 10.2). In all of these processes, the Tudor domain protein is thought to exert its effect through attachment to methylated arginine and/or lysine amino acids in target proteins. Identification of a Tudor domain sequence in an unknown gene therefore enables a specific biochemical activity to be identified, but on its own this does not enable the actual function of the gene to be assigned, beyond placing that function among the variety of roles performed by known Tudor domain proteins.

Annotation of gene function requires a common terminology

An easily overlooked but nonetheless vital aspect of genome annotation is the need to have an agreed and consistent terminology with which to describe the functions of different genes. Consistency is needed for two reasons. First, a rigorous comparison can be made between the genome annotations of two different species only if the same terminology has been used to name the genes in those two genomes. If the annotations use different terms for the same gene function, then attempts to identify the similarities between the two genomes will give inaccurate results, because the computer making the comparison will fail to recognize the co-identity of the genes. If the terminology is based on a hierarchical classification system, then a second advantage is that a gene whose function has been only partially deduced—for example, by identification of one or more protein domains—can still be given a meaningful descriptor that indicates what is known about its function.

The first comprehensive protein classification system predates the advent of genome sequencing but provides the hierarchical structure that is needed for genome annotation. This system, which applies only to enzymes, was first agreed upon by the International Union of Biochemistry and Molecular Biology in 1961. In this classification, enzymes are initially divided into six broad groups:

- EC (Enzyme Commission) group 1, oxidoreductases
- EC 2, transferases
- EC 3, hydrolases
- EC 4, lyases
- EC 5, isomerases
- EC 6, ligases

Each of these groups is further subdivided in such a way that each individual enzyme has its own four-part EC number. For example, EC 3.2.1.2 is a hydrolase (EC 3) that breaks glycosidic bonds (EC 3.2) in which the linkage between the sugar and the second molecule includes an oxygen or sulfur atom (EC 3.2.1), specifically the (\(\alpha 1\rightarrow 4\)) O-glycosidic bonds between glucose units in starch, glycogen, and related polysaccharides, in such a way as to release maltose disaccharide units from the nonreducing ends of the polymers (EC 3.2.1.2) (Figure 6.5). The enzyme activity is therefore described in a specific and hierarchical fashion, one that is much more informative than \(\beta\)-amylase, which is the common name for
this enzyme. It has been estimated that if a homology search reveals 40% amino acid sequence similarity between an unknown gene and a database entry, then the function of the unknown gene can be assigned as far as the first three parts of the EC number (EC 3.2.1). This degree of similarity indicates that the catalytic mechanism of the protein specified by the unknown gene is the same as that of the database match. If the similarity is 60% or higher, then both the query and match are likely to use the same substrate, so the fourth digit of the EC number can be assigned.

A second scheme for describing gene functions, the gene ontology (GO) nomenclature, was initially designed for annotation of the Drosophila genome but has subsequently been applied to many other species. The GO system, which can be applied to any protein, not just to enzymes, is not so much a classification scheme as a detailed set of standardized words and phrases that are used to describe a protein’s molecular function, the biological process to which it contributes, and its location in the cell. For example, the molecular function for β-amylase is described as “Catalysis of the reaction: (1,4-α-D-glucosyl)(n + 1) + H₂O = (1,4-α-D-glucosyl)(n − 1) + α-maltose. This reaction is the hydrolysis of 1,4-α-glucosidic linkages in polysaccharides so as to remove successive maltose units from the nonreducing ends of the chains.” This description is accompanied by an information modeling device called a directed acyclic graph (DAG), which gives a hierarchical categorization of the molecular function, similar to the way an enzyme function is categorized within an EC number (Figure 6.6).

Because the GO vocabulary is standardized, GO descriptions can be searched using a computer, which not only enables homologous genes in different genomes to be identified, but also makes it possible to identify groups of genes with similar functions in a single genome or across a set of genomes. The DAG enables the search to be conducted at different levels, so in the case of β-amylase the search can be for “β-amylase activity” and hence directed specifically at β-amylase homologs, or the search term could be “hydrolase activity, hydrolyzing O-glycosyl” so that a broader group of related hydrolase enzymes are also identified.

6.2 ASSIGNING FUNCTION BY GENE INACTIVATION AND OVEREXPRESSION

Computer methods for assigning functions to unknown genes are becoming increasingly sophisticated, but the bioinformatics approach has limitations and cannot identify the function of every new gene that is discovered in a genome. Experimental methods are therefore needed to complement and extend the results of computer analysis.
Functional analysis by gene inactivation

Devising experimental methods for functional analysis of new genes is proving to be one of the biggest challenges in genomics research. Most molecular biologists would agree that the methodologies and strategies currently in use are not entirely adequate for assigning functions to the vast numbers of unknown genes being discovered by sequencing projects. The problem is that the objective—to plot a course from gene to function—is the reverse of the route conventionally taken by genetic analysis, in which the starting point is an observable characteristic, or phenotype, and the objective is to identify the underlying gene or genes. The problem we are currently addressing takes us in the opposite direction: starting with a new gene and, ideally, arriving at identification of the associated phenotype.

In conventional genetic analysis, the genetic basis of a phenotype is usually studied by searching for mutant organisms in which the phenotype has become altered. The mutants might be obtained experimentally, for example, by treating a population of organisms (such as a culture of bacteria) with ultraviolet radiation or a mutagenic chemical, or the mutants might be present in a natural population. The gene or genes that have been altered in the mutant organism are then studied by genetic crosses (Section 3.4), which can locate the position of a gene in a genome and also determine if the gene is the same as one that has already been characterized. The gene can then be studied further by molecular biology techniques such as cloning and sequencing.

The general principle of this conventional analysis is that the genes responsible for a phenotype can be identified by determining which genes are inactivated in organisms that display a mutant version of the phenotype. If the starting point is the gene, rather than the phenotype, then the equivalent strategy would be to mutate the gene and identify the phenotypic change that results. This is the basis of most of the techniques used to assign functions to unknown genes.

Individual genes can be inactivated by homologous recombination

The easiest way to inactivate a specific gene is to disrupt it with an unrelated segment of DNA (Figure 6.7). This can be achieved by homologous recombination between the chromosomal copy of the gene and a second piece of DNA that shares some sequence identity with the target gene. Homologous and other types of recombination are complex events, which we will examine in detail in Section 17.1. For present purposes, it is enough to know that if two DNA molecules have similar sequences, then recombination can result in segments of the molecules being exchanged.

How is gene inactivation carried out in practice? We will consider two examples, the first with S. cerevisiae. After completing the genome sequence in 1996, yeast molecular biologists embarked on a coordinated, international effort to determine the functions of as many of the unknown genes as possible. Much of this work made use of the technique shown in Figure 6.8. The central component is the deletion cassette, which carries a gene for antibiotic resistance. This gene is not a normal component of the yeast genome but it will work if transferred into a yeast chromosome, giving rise to a transformed yeast cell that is resistant to the antibiotic geneticin. Before the deletion cassette is used, new segments of DNA are attached as tails to either end. These segments have sequences identical to parts of the yeast gene to be inactivated. After the modified cassette is introduced into a yeast cell, homologous recombination occurs between the DNA tails and the chromosomal copy of the yeast gene, replacing the latter with the antibiotic-resistance gene. Cells that have undergone the replacement are therefore selected by plating the culture onto agar medium containing geneticin. The resulting colonies lack the target gene activity, and their phenotypes can be examined to gain some insight into the function of the gene.

This method of gene inactivation is straightforward to carry out but time-consuming if it needs to be applied independently to each gene being studied. In the yeast project, this was an important consideration, as only 60% of the 6274
open reading frames (ORFs) greater than 100 codons in length could be assigned functions either from results of previous genetic analysis of yeast or by homology searching. The remaining 40%, over 2500 genes in total, had to be subjected to experimental analysis in order to assign functions. A high-throughput version of the gene inactivation method was therefore devised, called the barcode deletion strategy. This strategy uses a modified version of the basic deletion cassette system, with the difference that the cassette also includes two 20-nucleotide barcode sequences, different for each deletion, that act as tags for that particular mutant (Figure 6.9). Each barcode is flanked by the same pair of sequences and so can be amplified by a single PCR. This means that groups of mutated yeast strains, each with a different inactivated gene, can be mixed together and their phenotypes can be screened in a single experiment. For example, to identify genes required for growth in a glucose-rich medium, a collection of mutants would be mixed together and cultured under these conditions. After incubation, DNA is prepared from the culture and the barcoding PCR is carried out. The result is a mixture of PCR products, each representing a different barcode, where the relative abundance of each barcode indicates the abundance of each mutant after growth in glucose-rich medium. Barcodes that are absent or present only at low abundance indicate mutants whose inactivated genes were needed for growth under these conditions.

The second approach to gene inactivation that we will study is used with mice rather than yeast. The mouse is a popular model organism for humans because the mouse genome is similar to the human genome, containing many of the same genes. Functional analysis of unknown human genes is therefore being carried out largely by inactivating the equivalent genes in the mouse, as these experiments would be ethically unthinkable with humans. The procedure makes use of homologous recombination in a manner identical to that described for yeast, and once again the result is a cell in which the target gene has been inactivated. The problem is that we do not want just one mutated cell, we want a whole mutant mouse, as only with the complete organism can we make a full assessment of the effect of the gene inactivation on the phenotype. To achieve this, it is necessary to use a special type of mouse cell: an embryonic stem or ES cell. Unlike most mouse cells, ES cells are totipotent, meaning that they are not committed to a single developmental pathway and can therefore give rise to all types of differentiated cells. The engineered ES cell is therefore injected into a mouse embryo, which continues to develop and eventually gives rise to a chimera, a mouse whose cells are a mixture of mutant ones, derived from the engineered ES cells, and non-mutant ones, derived from all the other cells in the embryo. This is still not quite what we want, so the chimeric mice are allowed to mate with one another. Some of the offspring result from fusion of two mutant gametes and will therefore be nonchimeric, as every one of their cells will carry the inactivated gene. These are

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**Figure 6.8 Use of a yeast deletion cassette.** The deletion cassette consists of an antibiotic-resistance gene preceded by the promoter sequence needed for expression in yeast and flanked by two restriction sites. The start and end segments of the target gene are inserted into the restriction sites, and the vector is introduced into yeast cells. Recombination between the gene segments in the vector and the chromosomal copy of the target gene results in disruption of the latter. Cells in which the disruption has occurred are identifiable because they now express the antibiotic-resistance gene and so will grow on an agar medium containing geneticin. The gene designation KanR is an abbreviation for kanamycin resistance. Kanamycin is the family name of the group of antibiotics that includes geneticin.

**Figure 6.9 A deletion cassette used in the barcode strategy.** The two molecular barcodes are 20-nucleotide sequences, different for each cassette, that can be amplified by PCR. During homologous recombination, the barcodes are inserted into the yeast genome along with the kanamycin-resistance gene. The barcodes therefore provide specific tags for each individual gene deletion.
knockout mice, and with luck their phenotypes will provide the desired information on the function of the gene being studied. This works well for many gene inactivations, but some are lethal and so cannot be studied in a homozygous knockout mouse. Instead, a heterozygous mouse is obtained, the product of fusion between one normal and one mutant gamete, in the hope that the phenotypic effect of the gene inactivation will be apparent even though the mouse still has one correct copy of the gene being studied.

Gene inactivation without homologous recombination

Homologous recombination is not the only way to disrupt a gene in order to study its function. One alternative is to use transposon tagging, in which inactivation is achieved by insertion of a transposable element, or transposon, into the gene. Most genomes contain transposable elements (Section 9.2) and although the bulk of these are inactive, there are usually a few that retain their ability to move to new positions in the genome. Under normal circumstances, transposition is a relatively rare event, but it is sometimes possible to use recombinant DNA techniques to make modified transposons that change their position in response to an external stimulus. One way of doing this, involving the yeast transposon Ty1, is shown in Figure 6.10. Transposon tagging is also important in analysis of the fruit fly genome, by use of the endogenous Drosophila transposon called the P element. The weakness with transposon tagging is that it is difficult to target individual genes, because transposition is more or less a random event and it is impossible to predict where a transposon will end up after it has jumped. If the intention is to inactivate a particular gene, then it is necessary to induce a substantial number of transpositions and then to screen all the resulting organisms to find one with the correct insertion. Transposon tagging is therefore more applicable to global studies of genome function, in which genes are inactivated at random and groups of genes with similar functions are identified by examining the progeny for interesting phenotype changes.

A completely different approach to gene inactivation is provided by RNA interference, or RNAi, one of a series of natural processes by which short RNA molecules influence gene expression in living cells (Section 12.3). When used in genomics research, RNAi provides a means of silencing the expression of a target gene, not by disrupting the gene itself but by destroying its mRNA. This is accomplished by introducing short double-stranded RNA molecules, whose sequences match that of the mRNA being targeted, into the cell. The double-stranded RNAs are broken down into shorter molecules, which induce degradation of the mRNA (Figure 6.11).
RNA interference was initially shown to work effectively in the worm *Caenorhabditis elegans*, the genome of which has been completely sequenced and which is looked on as an important model organism for higher eukaryotes (Section 14.3). Virtually all of the 20,000 predicted genes in the *C. elegans* genome have been individually silenced by RNA interference. The key step in any RNAi experiment is introducing into the test organism the double-stranded RNA molecule that will give rise to the single-stranded interfering RNAs. With *C. elegans* this can be achieved by feeding the RNA to the worms. *C. elegans* eats bacteria, including *Escherichia coli*, and is often grown on a lawn of bacteria on an agar plate. If the bacteria contain a cloned gene that directs expression of a double-stranded RNA with the same sequence as a *C. elegans* gene, then after ingestion, the RNAi pathway begins to operate. Alternatively, the double-stranded RNA can be directly microinjected into the worm, but this is more time-consuming.

RNA interference occurs naturally in most eukaryotes, but its general application as a means of studying gene function has been hindered by three issues:

- RNAi does not always result in complete silencing of the target gene. Often the silencing is incomplete and is referred to as knockdown rather than knockout. Depending on the degree of silencing, it might or might not be possible to assess the effect of a gene knockdown on phenotype.

- The interfering RNAs are so short that off-target effects are possible. These occur when the interfering RNAs bind to mRNAs other than the targets, resulting in silencing of more than one gene.

- In mammals, the artificial introduction of double-stranded RNA often results in activation of signaling proteins called interferons, which stimulate an antiviral defense process that is displayed by both cultured cells and whole organisms. This interferon response results in phenotypic changes that can mask the specific change occurring due to silencing of the target gene. Some mammalian cells, such as mouse oocytes, lack the interferon response, but with most mammalian systems special strategies have to be devised when RNAi is being used for gene knockdown.

A second type of natural gene inactivation process, which avoids many of the problems associated with RNAi, makes use of a *programmable nuclease*. This is a nuclease that can be directed to a specific site in a genome and hence can be programmed to make a double-stranded cut in a selected gene (Figure 6.12). The cut stimulates a natural repair process, called nonhomologous end-joining (NHEJ) in eukaryotes, which joins the DNA strands together again. However, NHEJ is error-prone and usually results in a short insertion or deletion occurring at the repair site. If the repair is within a gene, then the change in nucleotide sequence will inactivate the gene. The inactivation is complete, and hence a true knockout, and is permanent, as opposed to an RNAi knockdown, which must be maintained by the continual presence of the interfering RNAs. A programmable nuclease is therefore an ideal system for analysis of gene function, but does it work in practice? Several systems have been explored, but the one that has gained prominence makes use of the Cas9 endonuclease, which is directed to its target site by a 20-nucleotide guide RNA. The binding site for the guide RNA must be immediately upstream of a 5′-NGG-3′ or 5′-NAG-3′ sequence (where N is any nucleotide), giving a 23-bp target that is cleaved by the endonuclease (Figure 6.13). Because the target sequence is precisely known, specificity can be assured, and off-target effects avoided, by ensuring that this sequence is unique in the genome being studied.

The Cas9 endonuclease is a component of the prokaryotic immune system called clustered regularly interspaced short palindromic repeats (CRISPR;

**Figure 6.12** Gene inactivation with a programmable nuclease. The cut made by the nuclease is repaired by nonhomologous end-joining, which is error-prone and likely to insert or delete a few base pairs of DNA at the repair site, disrupting the target gene.
Section 8.2. A eukaryotic cell therefore has to be engineered to synthesize the nuclease prior to performing the programmed gene inactivation. One approach is to use an adeno-associated virus vector to clone the Cas9 gene from Streptococcus pyogenes into the cell line being studied and then to introduce one or more guide RNA sequences via a second cloning experiment. Alternatively, both the endonuclease gene and the guide RNA sequence can be introduced together.

Gene overexpression can also be used to assess function

So far we have concentrated on techniques that result in inactivation of the gene being studied (loss of function). The complementary approach is to engineer an organism in which the test gene is much more active than normal (gain of function) and to determine what changes, if any, this has on the phenotype. The results of these experiments must be treated with caution because of the need to distinguish between a phenotype change that is due to the specific function of an overexpressed gene and a less specific phenotype change that reflects the abnormality of the situation where a single gene product is being synthesized in excessive amounts, possibly in tissues in which the gene is normally inactive. Despite this qualification, overexpression has provided some important information on gene function.

To overexpress a gene, a special type of cloning vector must be used, one designed to ensure that the cloned gene directs the synthesis of as much protein as possible. The vector is therefore multiplicity, meaning that it multiplies inside the host organism to 40–200 copies per cell, so there are many copies of the test gene. The vector must also contain a highly active promoter sequence (Section 12.2) so that each copy of the test gene is converted into large quantities of mRNA, again ensuring that as much protein as possible is made. In the example shown in Figure 6.14, the cloning vector contains a highly active promoter that is expressed only in the liver, so each transgenic mouse overexpresses the test gene in its liver. This approach has been used with genes whose sequences suggested that they code for proteins that are secreted into the bloodstream. After synthesis in the liver of the transgenic mouse, the test protein is secreted and the phenotype of the transgenic mouse is examined in the search for clues regarding the function of the cloned gene. An interesting discovery was made when it was realized that one mouse containing a human transgene had bones that were significantly more dense than those of normal mice. This was important for two reasons: first, it enabled the relevant gene to be identified as one involved in bone synthesis, and second, the discovery of a protein that increases bone density has implications for the development of treatments for human osteoporosis, a fragile-bone disease.
The phenotypic effect of gene inactivation or overexpression may be difficult to discern

The critical aspect of a gene inactivation or overexpression experiment is the need to identify a phenotypic change, the nature of which gives a clue to the function of the manipulated gene. This can be much more difficult than it sounds. With any organism, the range of phenotypes that must be examined is immense. Even with a unicellular organism such as yeast, the list is quite lengthy (Table 6.1A), and

<table>
<thead>
<tr>
<th>Phenotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>(A) Screens of all Saccharomyces cerevisiae genes</td>
</tr>
<tr>
<td>DNA synthesis and the cell cycle</td>
</tr>
<tr>
<td>RNA synthesis and processing</td>
</tr>
<tr>
<td>Protein synthesis</td>
</tr>
<tr>
<td>Stress responses</td>
</tr>
<tr>
<td>Cell wall synthesis and morphogenesis</td>
</tr>
<tr>
<td>Transport of biochemicals within the cell</td>
</tr>
<tr>
<td>Energy and carbohydrate metabolism</td>
</tr>
<tr>
<td>Lipid metabolism</td>
</tr>
<tr>
<td>DNA repair and recombination</td>
</tr>
<tr>
<td>Development</td>
</tr>
<tr>
<td>Meiosis</td>
</tr>
<tr>
<td>Chromosome structure</td>
</tr>
<tr>
<td>Cell architecture</td>
</tr>
<tr>
<td>Secretion and protein trafficking</td>
</tr>
<tr>
<td>(B) Screens of genes involved in early embryogenesis of Caenorhabditis elegans</td>
</tr>
<tr>
<td>Sterility/impaired fertility in parent</td>
</tr>
<tr>
<td>Osmotic integrity</td>
</tr>
<tr>
<td>Polar body extrusion</td>
</tr>
<tr>
<td>Passage through meiosis</td>
</tr>
<tr>
<td>Entry into interphase</td>
</tr>
<tr>
<td>Cortical dynamics</td>
</tr>
<tr>
<td>Pronuclear/nuclear appearance</td>
</tr>
<tr>
<td>Centrosome attachment</td>
</tr>
<tr>
<td>Pronuclear migration</td>
</tr>
<tr>
<td>Spindle assembly</td>
</tr>
<tr>
<td>Spindle elongation/integrity</td>
</tr>
<tr>
<td>Sister chromatid separation</td>
</tr>
<tr>
<td>Nuclear appearance</td>
</tr>
<tr>
<td>Chromosome segregation</td>
</tr>
<tr>
<td>Cytokinesis</td>
</tr>
<tr>
<td>Asymmetry of division</td>
</tr>
<tr>
<td>Pace of cell division</td>
</tr>
<tr>
<td>General pace of development</td>
</tr>
<tr>
<td>Severe pleiotropic defects</td>
</tr>
<tr>
<td>Integrity of membrane-bound organelles</td>
</tr>
<tr>
<td>Egg size</td>
</tr>
<tr>
<td>Aberrant cytoplasmic structures</td>
</tr>
<tr>
<td>Complex combination of defects</td>
</tr>
</tbody>
</table>
Chapter 6: Identifying Gene Functions

with multicellular eukaryotes it is much more so (Table 6.1B). In higher organisms some phenotypes (such as behavioral ones) are difficult, if not impossible, to assess in a comprehensive fashion. Furthermore, the effect of gene inactivation can be very subtle and may not be recognized when the phenotype is examined. A good example of the problems that occur was provided by the longest gene on yeast chromosome III, which, at 2167 codons and with typical yeast codon bias, simply had to be a functional gene rather than a spurious ORF. Inactivation of this gene had no apparent effect: the mutant yeast cells appeared to have an identical phenotype to normal yeast. For some time it was thought that perhaps this gene was dispensable, its protein product either involved in some completely nonessential function or having a function that is duplicated by a second gene. Eventually it was shown that the mutants die when they are grown at low pH in the presence of glucose and acetic acid, which normal yeasts can tolerate. This observation enabled the gene to be linked with the processes by which yeast cells secrete unwanted compounds, such as acetate, out of the cell. This is definitely an essential function, but this essentiality was difficult to track down from the phenotype tests.

Even when the most careful screens are carried out, many gene inactivations appear to give no discernible phenotypic change. Almost 5000 of the 6692 genes in the yeast genome can be individually inactivated without causing the cells to die, and inactivation of many of these 5000 genes has no detectable effect on the metabolic properties of the cell under normal growth conditions. The phenotypic effects of these genes only become apparent, if at all, when the cells are grown under a range of different conditions or when groups of genes that contribute to the same phenotype are co-inactivated. In the human genome, there appears to be a subset of several hundred genes that are nonessential, both copies of which can be inactivated, due to natural mutation, without any discernible effect on the health of the individual. These observations suggest that a complete functional annotation of the genomes of many species will not be achievable by approaches that are based solely on gene inactivation or overexpression.

6.3 UNDERSTANDING GENE FUNCTION BY STUDIES OF EXPRESSION PATTERN AND PROTEIN PRODUCT

Gene inactivation and overexpression are the primary techniques used by genome researchers to determine the function of a new gene, but they are not the only procedures that can be used to provide information on gene activity. Additional insights into gene function can be obtained by identifying in which tissues, and at what times, a gene is expressed and by direct examination of the protein coded by the gene.

Reporter genes and immunocytochemistry can be used to locate where and when genes are expressed

Clues to the function of a gene can often be obtained by determining where and when the gene is active. If gene expression is restricted to a particular organ or tissue of a multicellular organism, or to a single set of cells within an organ or tissue, then this positional information can be used to infer the general role of the gene product. The same is true of information relating to the developmental stage at which a gene is expressed. This type of analysis has proved particularly useful in understanding the activities of genes involved in the earliest stages of development in Drosophila (Section 14.3) and is increasingly being used to unravel the genetics of mammalian development. It is also applicable to those unicellular organisms, such as yeast, that have distinctive developmental stages in their life cycle.

Determining the pattern of gene expression within an organism is possible with a **reporter gene**. This is a gene whose expression can be monitored in a convenient way, ideally by visual examination (Table 6.2): for example, cells that
express the reporter gene may become blue, fluoresce, or give off some other visible signal. For the reporter gene to give a reliable indication of where and when a test gene is expressed, the reporter must be subject to the same regulatory signals as the test gene. This is achieved by replacing the ORF of the test gene with the ORF of the reporter gene (Figure 6.15). Most of the regulatory signals that control gene expression are contained in the region of DNA upstream of the ORF, so the reporter gene should now display the same expression pattern as the test gene. The expression pattern can therefore be determined by examining the organism for the reporter signal.

As well as knowing in which cells a gene is expressed, it is often useful to locate the position within the cell where the protein coded by the gene is found. For example, key data regarding gene function can be obtained by showing that the protein product is located in mitochondria, in the nucleus, or on the cell surface. Reporter genes cannot help here because the DNA sequence upstream of the gene—the sequence to which the reporter gene is attached—is not involved in targeting the protein product to its correct intracellular location. Instead, the amino acid sequence of the protein itself contains the targeting information. Therefore, the only way to determine where the protein is located is to search for it directly. This can be done by **immunocytochemistry**, which makes use of an antibody that is specific for the protein of interest and so binds to this protein and no other. The antibody is labeled so that its position in the cell, and hence the position of the target protein, can be visualized (Figure 6.16). Fluorescent labeling and confocal microscopy are used for low-resolution studies; alternatively, high-resolution immunocytochemistry can be carried out by electron microscopy with an electron-dense label such as colloidal gold.

**Directed mutagenesis can be used to probe gene function in detail**

Inactivation and overexpression can determine the general function of a gene, but they cannot provide detailed information on the activity of a protein coded by a gene. For example, it might be suspected that part of a gene codes for an amino acid sequence that directs its protein product to a particular compartment in the cell or is responsible for the ability of the protein to respond to a chemical or physical signal. To test these hypotheses, it would be necessary to delete or alter the relevant part of the gene sequence but to leave the bulk unmodified so that the protein is still synthesized and retains the major part of its activity. The various procedures of **site-directed** or **in vitro mutagenesis** can be used to make these subtle changes. These are important techniques whose applications lie not only with the study of gene activity but also in the area of **protein engineering**, where the objective is to create novel proteins with properties that are better suited for use in industrial or clinical settings.

**Figure 6.15 A reporter gene.** The open reading frame of the reporter gene replaces the open reading frame of the gene being studied. The result is that the reporter gene is placed under control of the regulatory sequences that usually dictate the expression pattern of the test gene.

**Figure 6.16 Immunocytochemistry.** The cell is treated with an antibody that is labeled with a red fluorescent marker. Examination of the cell shows that the fluorescent signal is associated with the inner mitochondrial membrane. A working hypothesis would therefore be that the target protein is involved in electron transport and oxidative phosphorylation, as these are the main biochemical functions of the inner mitochondrial membrane.
For many years, biologists have made use of conventional **mutagenesis**, in which an organism is exposed to a mutagenic chemical in order to induce mutations in its genome. These mutations occur randomly at unspecified positions in a DNA molecule, so large numbers of mutated organisms must be screened to find a mutation of interest. Even with microbes, which can be screened in huge numbers, the best that can be hoped for is a range of mutations in the correct gene, one of which might affect an interesting part of the protein being studied. Site-directed mutagenesis offers a means of making much more specific mutations. The most important of these methods are as follows:

- **Oligonucleotide-directed mutagenesis**, in which an oligonucleotide containing a single base-pair mismatch, corresponding to the desired mutation, is annealed to a single-stranded version of the relevant gene, which is usually obtained by cloning the gene with a M13 bacteriophage vector. If a DNA polymerase is added to the single-stranded DNA, the oligonucleotide primes a strand-synthesis reaction that continues all the way around the circular template molecule (Figure 6.17A). After introduction into *E. coli*, DNA replication produces numerous copies of this recombinant DNA molecule. Half of these are copies of the original strand of DNA, and half are copies of the strand that contains the mutated sequence. All of these double-stranded molecules direct synthesis of M13 phage particles, so about half the phages released from the infected bacteria carry a copy of the mutated molecule. The phages are plated onto solid agar so that plaques are produced, and the mutant ones are identified by hybridization probing with the original oligonucleotide (Figure 6.17B).

- **Artificial gene synthesis** involves constructing the gene in the test tube, placing mutations at all the desired positions. The gene is constructed by synthesizing a series of partially overlapping oligonucleotides, each typically about 150 nucleotides in length. The gene is then assembled by filling in the gaps between the overlaps with DNA polymerase.

- PCR can also be used to create mutations in cloned genes, though like oligonucleotide-directed mutagenesis, only one mutation can be created per experiment. The method shown in Figure 6.18 involves two PCRs, each with one normal primer (which forms a fully base-paired hybrid with the template DNA) and one mutagenic primer (which contains a single base-pair mismatch). The mutation is therefore initially present in two PCR
products, each corresponding to one half of the starting DNA molecule. The two PCR products are then mixed together and a final PCR cycle is carried out to construct the full-length, mutated DNA molecule.

After mutagenesis, the mutated gene can be placed back in its original host by homologous recombination, as described in Section 6.2, or transferred to an *E. coli* vector designed for synthesis of protein from cloned DNA, so that a sample of the mutated protein can be obtained. If homologous recombination is used, then we must have a way of knowing which cells have taken up a copy of the mutated gene. Even with yeast, this will be only a fraction of the total. Normally we would solve this problem by placing a marker gene (for example, one coding for antibiotic resistance) next to the mutated gene and looking for cells that take on the phenotype conferred by this marker. In most cases, cells that insert the marker gene into their genome also insert the closely attached mutated gene and so are the ones we want. The problem is that in a site-directed mutagenesis experiment we must be sure that any change in the activity of the gene being studied is the result of the specific mutation that was introduced into the gene, rather than the indirect result of changing its environment in the genome by inserting a marker gene next to it. The answer is to use a more complex, two-step gene replacement (Figure 6.19). In this procedure the target gene is first replaced with the marker gene on its own, and the cells in which this recombination takes place are identified by selecting for the marker gene phenotype. These cells are then used in the second stage of the gene replacement, when the marker gene is replaced by the mutated gene, success now being monitored by looking for cells that have lost the marker gene phenotype. These cells contain the mutated gene, and their phenotypes can be examined to determine the effect of the directed mutation on activity of the protein product.

### 6.4 USING CONVENTIONAL GENETIC ANALYSIS TO IDENTIFY GENE FUNCTION

We noted at the start of Section 6.2 that functional annotation of a genome represents the reverse of the conventional approach to genetics, because functional annotation starts with a gene and attempts to discover its function, whereas
conventional genetics starts with a phenotype and attempts to discover the gene or genes responsible for that phenotype. The conventional approach is sometimes called forward genetics, with reverse genetics comprising the methods that start with the gene. So far, we have only studied the use of reverse genetics in functional annotation of a genome, but that does not mean that forward genetics is no longer important. Quite the opposite is true. Forward genetics is still central to many areas of genomics research, in particular as a means of identifying human genes responsible for inherited diseases.

Identification of human genes responsible for inherited diseases

An inherited disease is one that is caused by a defect in the genome and which can be passed from parents to offspring. There are over 6000 monogenic inherited diseases known in the human population, each resulting from a defect in a single gene. The frequencies of these diseases vary enormously: the commonest types, such as inherited breast cancer and cystic fibrosis, occur once in a few hundred or few thousand births, but there are also very rare diseases with just a few occurrences every year (Table 6.3). Inherited diseases also affect other animals, in particular those with low genetic diversity due to artificial breeding, such as some breeds of pedigree dogs.

Forward genetics is able to identify the gene that confers a phenotype even if very little is known about that phenotype. In fact, all that is necessary is to establish that inheritance of the phenotype follows a simple Mendelian pattern (Section 3.3) and hence is specified by a single gene. If this is the case for an inherited disease, then DNA samples can be gathered from members of affected families and pedigree analysis carried out to determine the linkage between the disease gene and mapped DNA markers (Section 3.4). To illustrate the methodology in greater detail, we will examine the way in which pedigree analysis, coupled with reverse genetics, was used to identify a gene that confers susceptibility to inherited breast cancer.

The first pedigree studies of inherited breast cancer aimed to locate the position of the causative gene relative to restriction fragment length polymorphisms (RFLPs) (Section 3.2) that had already been mapped onto the human genome. This work showed that, in families with a high incidence of breast cancer, a significant number of the women who suffered from the disease possessed the same allele of the RFLP called D17S74. This observation suggested that the breast cancer gene must lie close to D17S74 on the human genome. The rationale is that recombination is unlikely between two markers that are close together, so the alleles of those two markers will be inherited together (Figure 6.20). This is called

<table>
<thead>
<tr>
<th>Disease</th>
<th>Symptoms</th>
<th>Frequency (births per year)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inherited breast cancer</td>
<td>Cancer</td>
<td>1 in 300 females</td>
</tr>
<tr>
<td>Cystic fibrosis</td>
<td>Lung disease</td>
<td>1 in 2000</td>
</tr>
<tr>
<td>Huntington’s chorea</td>
<td>Neurodegeneration</td>
<td>1 in 2000</td>
</tr>
<tr>
<td>Duchenne muscular dystrophy</td>
<td>Progressive muscle weakness</td>
<td>1 in 3000 males</td>
</tr>
<tr>
<td>Hemophilia A</td>
<td>Blood disorder</td>
<td>1 in 4000 males</td>
</tr>
<tr>
<td>Sickle cell anemia</td>
<td>Blood disorder</td>
<td>1 in 10,000</td>
</tr>
<tr>
<td>Phenylketonuria</td>
<td>Mental retardation</td>
<td>1 in 12,000</td>
</tr>
<tr>
<td>β-Thalassemia</td>
<td>Blood disorder</td>
<td>1 in 20,000</td>
</tr>
<tr>
<td>Retinoblastoma</td>
<td>Cancer of the eye</td>
<td>1 in 20,000</td>
</tr>
<tr>
<td>Hemophilia B</td>
<td>Blood disorder</td>
<td>1 in 25,000 males</td>
</tr>
<tr>
<td>Tay–Sachs disease</td>
<td>Blindness, loss of motor control</td>
<td>1 in 200,000</td>
</tr>
</tbody>
</table>
linkage disequilibrium. In our example, the defective allele of the disease gene is linked to one allele of D17S74, and the nondefective allele of the gene is linked to the second allele of D17S74. As the RFLP had previously been mapped to the long arm of chromosome 17, it could be concluded that the breast cancer gene must also be located in this part of the genome, probably within the chromosomal region designated q21 (Figure 6.21). A more specific map position was then deduced from additional pedigree analyses, which examined linkage between the breast cancer gene and short tandem repeats (STRs) known to be present in the q21 region. These analyses placed the breast cancer gene between two STRs called D17S1321 and D17S1325, which are approximately 600 kb apart.

Further pedigree analysis, with increasing numbers of families, could conceivably have resulted in an even more specific location for the breast cancer gene, but once a human gene has been placed within a genomic region of 1 Mb or less, it is usually possible for reverse genetics methods to complete the identification. The genome annotation shows that the region between D17S1321 and D17S1325 contains over 60 genes, any of which could be the breast cancer gene. The expression patterns of these candidate genes were studied, with the expectation that the breast cancer gene would be expressed in breast and ovarian tissue, as ovarian cancer is frequently associated with inherited breast cancer. Genes with the expected expression profile were then used in BLAST searches of other mammalian genomes, on the basis that a human gene that is important enough to cause disease when mutated would be expected to have homologs in a range of other mammals. Finally, the sequences of those genes still looked on as candidates were examined in women with and without inherited breast cancer to see if the genes from affected individuals contained mutations that might explain why they have the disease. When these analyses were complete, the most likely candidate was an approximately 100 kb gene, comprising 22 exons and coding for an 1863 amino acid protein. This gene, subsequently named BRCA1, was expressed in breast and ovary tissue and had homologs in mice, rats, rabbits, sheep, and pigs but not in chickens. Critically, the alleles of this gene in five susceptible families contained mutations likely to lead to a nonfunctioning protein. Subsequent studies have shown that the protein specified by BRCA1 is involved in transcription regulation and DNA repair and also acts as a tumor suppressor gene, inhibiting abnormal cell division.

Genowide association studies can also identify genes for diseases and other traits

Although pedigree analysis can be used to identify the gene responsible for a monogenic disease, it is less successful with the many human diseases that have more complex genetic backgrounds. Several types of cancer, as well as disorders such as coronary heart disease and osteoporosis, are polygenic, meaning that they are controlled not just by one gene but by many genes working together. Most polygenic traits are quantitative, so affected individuals display different degrees of susceptibility, depending on the particular combination of alleles that they possess. This means that two members of a family that display the disease can have different genotypes. Under these circumstances, the disease phenotype will not follow a simple Mendelian inheritance pattern, and it becomes impossible to use pedigree data to link the disease to DNA markers with any degree of certainty.
A genomewide association study (GWAS) is an alternative approach to gene identification that can work with polygenic traits. Rather than linking DNA markers with individual genes, a GWAS attempts to identify all of the markers, from all over the genome, that are associated with the disease. The positions of these markers will reveal the positions of candidate genes that might form part of the polygenic trait. A GWAS therefore requires both a large cohort of individuals, including people with and without the disease, and a large panel of DNA markers that will be typed in DNA samples from those individuals. The DNA samples are often obtained from biobanks, which are collections of biological material such as blood samples, donated following informed consent by patients and volunteers, with each sample accompanied by a detailed description of that individual's disease status. The DNA markers are invariably single-nucleotide polymorphisms (SNPs), because of the huge number of SNPs whose precise positions in the human genome are known and because of the ease with which multiple SNPs can be typed by chip technology (Section 3.2). One of the first GWAS projects studied age-related macular degeneration, which causes vision problems in elderly people. By typing 226,204 SNPs in 96 affected individuals and 50 controls, two SNPs that displayed strong association with the disease were identified. Both SNPs are located in an intron within the gene for complement factor B, a protein involved in control of the inflammatory response. This is one of five genes, on three different chromosomes, now thought to be involved in age-related macular degeneration.

GWAS projects have gradually become more ambitious, with highly complex traits such as hypertension (high blood pressure) now being addressed. These studies use cohort sizes of tens of thousands and may type a million or more SNPs, the larger numbers enabling a greater degree of resolution so that associations with multiple genetic loci can be identified in a single screen. Over 60 genes have been associated with hypertension as a result of a series of GWAS projects carried out in different parts of the world. The GWAS approach has also been applied to species other than humans, including crop plants, where it is proving valuable in identifying genes responsible for complex traits such as heading date (the time taken for the plant to reach the flowering stage) and the number and weight of seeds that are produced.

SUMMARY

- Gene functions can tentatively be assigned by homology analysis, because homologous genes are evolutionarily related and often, but not always, have similar functions.
- Identification of conserved sequence motifs can also help to determine the function of a gene.
- Most experimental techniques for the functional analysis of genes involve examining the effect of gene inactivation on the phenotype of the organism.
- Gene inactivation can be achieved by homologous recombination with a defective copy of the gene.
- Inactivation is also achieved by insertion of a transposon into the gene, by RNA interference, and by use of the CRISPR technology.
- Gene overexpression can also be used to assess function.
- With both inactivation and overexpression experiments it may be difficult to discern a phenotypic change, and the precise function of the gene may remain elusive.
- The cellular location of a protein can be determined by expression of a reporter gene or by immunocytochemistry.
More detailed studies of gene function can be carried out by site-directed mutagenesis.

Candidates for human disease genes can be identified by pedigree analysis and by genomewide association studies.

**SHORT ANSWER QUESTIONS**

1. What is the difference between orthologous and paralogous genes?
2. Describe how a BLAST search is carried out and explain why errors are sometimes made when assigning gene function using this approach.
3. How is identification of protein domains used in the functional analysis of a genome sequence?
4. Outline the key features of the EC and GO systems for the classification of gene functions.
5. Discuss the role of homologous recombination in studies of gene function.
6. Describe how gene inactivation is achieved by transposon tagging.
7. What is RNA interference and how is it used in studies of gene function?
8. How is the CRISPR system used to inactivate eukaryotic genes?
9. Outline the applications of gene overexpression in studies of gene function.
10. Compare the strengths and weaknesses of gene inactivation and overexpression experiments in the functional annotation of a genome sequence.
11. Describe how immunocytochemistry and directed mutagenesis are used to understand gene functions.
12. Outline the possible approaches that can be used to identify the gene responsible for an inherited disease.

**IN-DEPTH PROBLEMS**

1. Perform a BLAST search (https://blast.ncbi.nlm.nih.gov/Blast.cgi) with the following amino acid sequence:
   GLSDGEWQLNLNVWKEADLHGHQEVILLFKGMETLKFDFK
   FKLSEKGEMLVILKHGNTVETALEILKKKALELFKNIAAKTELGLG.
   What protein has this amino acid sequence? Are the homologous sequences identified for this search mostly orthologs or paralogs?
2. An important protein domain has the following amino acid sequence:
   KRARTATYQTELEKEHFHNRYLRRRIEIAHALCLSERQIKIWFQ
   NRRMKWKKDN
   Identify the domain and describe its function.
3. Gene inactivation studies have suggested that at least some genes in a genome are redundant, meaning that they have the same function as a second gene and so can be inactivated without affecting the phenotype of the organism. What evolutionary questions are raised by genetic redundancy? What are the possible answers to these questions?
4. Explore the natural role of RNA interference in living organisms.
5. Gene overexpression has so far provided limited but important information on the function of unknown genes. Assess the overall potential of this approach in functional analysis.
**FURTHER READING**

Assigning function by computer analysis


RNA interference studies


CRISPR


Other methods for gene inactivation


Overexpression, immunocytochemistry, and directed mutagenesis


Identifying gene function by conventional genetics


In the next three chapters, we will survey the anatomies of the various types of genome that are found on our planet. There are three chapters because there are three types of genome to consider:

- **Eukaryotic nuclear genomes** (this chapter), of which the human genome is the one of greatest interest to us
- **Genomes of prokaryotes and of eukaryotic organelles** (Chapter 8), which we will consider together because eukaryotic organelles are descended from ancient prokaryotes
- **Viral genomes and mobile genetic elements** (Chapter 9), grouped together because some mobile elements are related to viral genomes

### 7.1 Nuclear Genomes Are Contained in Chromosomes

The nuclear genome is split into a set of linear DNA molecules, each contained in a chromosome. No exceptions to this pattern are known: all eukaryotes that have been studied have at least one chromosome, and the DNA molecules are always linear. The only variability lies with chromosome number, which appears to be unrelated to the biological features of the organism. For example, yeast has 16 chromosomes, four times as many as the fruit fly. The ant *Myrmecia pilosula* has just one chromosome, and the Indian muntjac deer has only four. Nor is chromosome number linked to genome size: some salamanders have genomes 30 times bigger than the human version but split into half the number of chromosomes. These comparisons are interesting but at present do not tell us anything useful about the genomes themselves; they are more a reflection of the nonuniformity of the evolutionary events that have shaped genome architecture in different organisms.

**Chromosomes are much shorter than the DNA molecules they contain**

During cell division, each human chromosome adopts a compact structure that is just a few micrometers in length. In contrast, the shortest of the 24 DNA molecules...
that make up the human genome is 1.6 cm in length, and the longest is 8.5 cm. The average length is over 4 cm. A highly organized packaging system is therefore needed to fit a DNA molecule into its chromosome. We must understand this packaging system before we start to think about how genomes function, because the way in which DNA is packaged has an influence on the processes involved in expression of individual genes (Chapter 10).

The important breakthroughs in understanding DNA packaging were made in the early 1970s by a combination of biochemical analysis and electron microscopy. It was already known that nuclear DNA is associated with DNA-binding proteins called histones, but the exact nature of the association had not been delineated. In 1973–1974, several groups carried out nuclease protection experiments on chromatin (DNA–histone complexes) that had been gently extracted from nuclei by methods designed to retain as much of the chromatin structure as possible. In a nuclease protection experiment, the complex is treated with an enzyme that cuts the DNA at positions that are not protected by attachment to a protein. The sizes of the resulting DNA fragments indicate the positioning of the protein complexes on the original DNA molecule (Figure 7.1). After limited nuclease treatment of purified chromatin, the bulk of the DNA fragments have lengths of approximately 200 bp and multiples thereof, suggesting a regular spacing of histone proteins along the DNA.

In 1974, these biochemical results were supplemented by electron micrographs of purified chromatin, which enabled the regular spacing inferred by the protection experiments to be visualized as beads of protein on the string of DNA (Figure 7.2A). Further biochemical analysis indicated that each bead, or nucleosome, contains eight histone protein molecules, two each of histones H2A, H2B, H3, and H4. Structural studies have shown that these eight proteins

---

**Figure 7.1** Nuclease protection analysis of chromatin from human nuclei.
Chromatin is gently purified from nuclei and treated with a nuclease enzyme. On the left, the nuclease treatment is carried out under limiting conditions so that the DNA is cut, on average, just once in each of the linker regions between the bound proteins. After removal of the protein, the DNA fragments are analyzed by agarose gel electrophoresis and found to be 200 bp in length or multiples thereof. On the right, the nuclease treatment proceeds to completion, so all the DNA in the linker regions is digested. The remaining DNA fragments are all 146 bp in length. The results show that, in this form of chromatin, protein complexes are spaced along the DNA at regular intervals, one for each 200 bp, with 146 bp of DNA closely attached to each protein complex.

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**Figure 7.2** Nucleosomes. (A) Electron micrograph of a purified chromatin strand showing the beads-on-a-string structure. (B) Model for the beads-on-a-string structure, in which each bead is a barrel-shaped nucleosome with the DNA wound twice around the outside. Each nucleosome is made up of eight proteins: a central tetramer of two histone H3 and two histone H4 subunits, plus a pair of H2A–H2B dimers, one above and one below the central tetramer (see Figure 7.4). (C) The precise position of the linker histone relative to the nucleosome is not known, but as shown here, the linker histone may act as a clamp, preventing DNA from detaching from the outside of the nucleosome. (A, Courtesy of Barbara Hamkalo, University of California, Irvine.)
form a barrel-shaped **core octamer** with the DNA wound twice around the outside (Figure 7.2B). Between 140 and 150 bp of DNA (depending on the species) is associated with the nucleosome particle, and each nucleosome is separated by 50–70 bp of linker DNA, giving the repeat length of 190–220 bp previously shown by the nuclease protection experiments.

As well as the proteins of the core octamer, there is a group of additional histones, all closely related to one another and collectively called **linker histones**. In humans these include histones H1.0–H1.5, H1oo, H1t, and H1x. A single linker histone is attached to each nucleosome, to form the **chromatosome**, but the precise positioning of this linker histone is not known. Structural studies support the traditional model in which the linker histone acts as a clamp, preventing the coiled DNA from detaching from the nucleosome (Figure 7.2C). However, other results suggest that, at least in some organisms, the linker histone is not located on the extreme surface of the nucleosome–DNA assembly, as would be expected if it really were a clamp, but instead is inserted between the core octamer and the DNA.

The beads-on-a-string structure shown in Figure 7.2A is thought to represent an unpacked form of chromatin that might occur only infrequently in living nuclei. Very gentle cell-breakage techniques developed in the mid-1970s revealed a more condensed version of the complex, approximately 30 nm in width, called the **30 nm fiber**. The exact way in which nucleosomes associate to form the 30 nm fiber is not known, but several models have been proposed, two of which are shown in Figure 7.3. The individual nucleosomes within the 30 nm fiber may be held together by interactions between the linker histones, or the attachments may involve the core histones, whose protein tails extend outside the nucleosome (Figure 7.4). The latter hypothesis is attractive because chemical modification of these tails results in the 30 nm fiber opening up, enabling genes contained within it to be activated (Section 10.2).

**Special features of metaphase chromosomes**

The 30 nm fiber is probably the major type of chromatin in the nucleus during **interphase**, the period between nuclear divisions. When the nucleus divides, the DNA adopts a more compact form of packaging, resulting in the highly condensed **metaphase chromosomes** that can be seen with the light microscope and have the appearance generally associated with the word chromosome (Figure 7.5). The metaphase chromosomes form at a stage in the **cell cycle** after DNA replication has taken place, and so each one contains two copies of its chromosomal.
DNA molecule. The two copies are held together at the **centromere**, which has a specific position within each chromosome. The arms of the chromosome, which are called **chromatids** and have terminal structures called **telomeres**, are of different lengths in different chromosomes. Individual chromosomes can therefore be recognized because of the lengths of their chromatids and the location of the centromere relative to the telomeres. Further distinguishing features are revealed when chromosomes are stained. There are a number of different staining techniques (**Table 7.1**), each resulting in a banding pattern that is characteristic for a particular chromosome. This means that the set of chromosomes possessed by an organism can be represented as a **karyogram**, in which the banded appearance of each one is depicted. The human karyogram is shown in **Figure 7.6**.

The human karyogram is typical of that of the great majority of eukaryotes, but some organisms display unusual features not displayed by the human genome. These include the following:

- **Microchromosomes** are found in birds and some fish, reptiles, and amphibians. They are relatively short in length, less than 20 Mb, but often they are rich in genes. The chicken genome, for example, is split into 38 autosomes and two sex chromosomes, the latter called Z and W. Of the 38 autosomes, only five are classified as **macrochromosomes**, which are longer than 50 Mb and hence comparable in size to human chromosomes. The other 33 autosomes include five of 20–50 Mb, usually referred to as intermediate chromosomes, and 28 microchromosomes, the majority of which are less than 10 Mb, with several less than 1 Mb. Within the microchromosomes there are 13–42 genes/Mb of DNA, compared with just 9–16 genes/Mb of DNA in the macrochromosomes. The gene density in the microchromosomes is therefore some 2–3 times greater than that in the macrochromosomes.

- **B chromosomes** are additional chromosomes possessed by some individuals in a population, but not all. They are common in plants and are also known in fungi, insects, and animals. B chromosomes appear to be fragmentary versions of normal chromosomes that result from unusual events during nuclear division. Some contain genes, often for rRNAs, but it is not clear whether these genes are active. The presence of B chromosomes can affect the biological characteristics of the organism, particularly in plants where they are associated with reduced viability. It is presumed that B chromosomes are gradually lost from cell lineages as a result of irregularities in their inheritance pattern.

- **Holocentric chromosomes** do not have a single centromere but instead have multiple structures that act as centromeres spread along their length. The nematode worm *Caenorhabditis elegans* has holocentric chromosomes.

<table>
<thead>
<tr>
<th>Technique</th>
<th>Procedure</th>
<th>Banding pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>G-bandung</td>
<td>Mild proteolysis followed by staining with Giemsa</td>
<td>Dark bands are AT-rich; pale bands are GC-rich</td>
</tr>
<tr>
<td>R-bandung</td>
<td>Heat denaturation followed by staining with Giemsa</td>
<td>Dark bands are GC-rich; pale bands are AT-rich</td>
</tr>
<tr>
<td>Q-bandung</td>
<td>Stain with quinacline</td>
<td>Dark bands are AT-rich; pale bands are GC-rich</td>
</tr>
<tr>
<td>C-bandung</td>
<td>Denature with barium hydroxide and then stain with Giemsa</td>
<td>Dark bands contain constitutive heterochromatin (see Section 10.1)</td>
</tr>
</tbody>
</table>
DNA–protein interactions in centromeres and telomeres

The DNA contained within centromeres and telomeres and the proteins attached to this DNA have special features related to the particular functions of these structures.

The initial breakthroughs in understanding the nucleotide sequence of centromeric DNA in higher eukaryotes were made with the plant *Arabidopsis thaliana*, whose amenity to genetic analysis enabled the positions of the centromeres on the DNA sequence to be located with some precision. During the *Arabidopsis* genome sequencing project, which was completed in 2000, a special effort was made to
sequence these centromeric regions, which are often excluded from genome sequences because of problems in obtaining an accurate reading through the highly repetitive structures that characterize these regions. *Arabidopsis* centromeres span 0.4–3.0 Mb of DNA and each one is made up largely of 178–180 bp repeat sequences. Before the *Arabidopsis* sequences were obtained, it was thought that these repeat sequences were by far the principal component of centromeric DNA. However, *Arabidopsis* centromeres also contain multiple copies of a variety of repeats that are also found elsewhere in the genome, and we now know that the same is true for the centromeric DNA of most species. In humans, for example, the centromeric DNA is 1–5 Mb, of which 1–4 Mb is made of 171 bp *alpheid* DNA repeats and the remainder comprises other families of repeated sequences up to 70 bp in length.

*Arabidopsis* and humans display the basic pattern for centromeric DNA as seen in virtually all eukaryotes. We call these **regional centromeres**, to indicate that each centromere covers a region of the chromosomal DNA. An interesting variation occurs in the yeast *Saccharomyces cerevisiae*, which has short **point centromeres**, which do not contain repetitive DNA and instead are defined by a single-copy sequence, approximately 120 bp in length. This sequence is made up of two short elements, called CDEI and CDEII, which flank a longer element called CDEIII (Figure 7.7). The sequence of CDEII is variable, though always very rich in A and T nucleotides, whereas both CDEI and CDEIII are highly conserved, meaning that their sequences are very similar in all 16 yeast chromosomes. Mutations in CDEII rarely affect the function of the centromere, but a mutation in CDEI or CDEIII usually prevents the centromere from forming. The short, nonrepetitive nature of the yeast centromeric DNA has enabled progress to be made in understanding how the DNA interacts with proteins to form a functional centromere. A key role is played by a special chromosomal protein called Cse4, which is similar in structure to histone H3 and replaces H3 in a centromeric nucleosome. The exact composition of this nucleosome is unclear. It could be an octamer, just like the standard nucleosome but with H3 replaced with Cse4, but alternative possibilities are a hemisome (one molecule each of Cse4 and histones H2A, H2B, and H4) or a tetrasome of two Cse4 molecules and two copies of H4. Two other proteins, Cbf1 and Cbf3, form interactions with the CDEI and CDEIII sequences, respectively, and also bind to at least some of the 20 or so additional proteins that form the **kinetochore**, the structure that acts as the attachment point for the microtubules that draw the divided chromosomes into the daughter nuclei (Figure 7.8). This model of the yeast point centromere appears also to apply to the regional centromeres of other eukaryotes, in which the protein CENP-A replaces histone H3. However, a regional centromere contains many nucleosomes, and it is not clear whether all of them are modified in the same way.

The second important part of the chromosome is the terminal region or telomere. Telomeres are important because they mark the ends of chromosomes and

---

**Figure 7.7** *Saccharomyces cerevisiae* centromeric DNA. CDEI is 9 bp in length, CDEII is 80–90 bp, and CDEIII is 11 bp. Additional sequences flanking the region shown here are looked on as part of the centromeric DNA, whose full length is approximately 120 bp.

**Figure 7.8** The yeast centromere. (A) DNA–protein interactions in the yeast centromere. The diagram is purely schematic, as the precise positioning of the proteins and of the DNA components is unknown. (B) The role of kinetochores during nuclear division. During the anaphase period of nuclear division, individual chromosomes are drawn apart by the contraction of microtubules attached to the kinetochores.
therefore enable the cell to distinguish a real end from an unnatural end caused by chromosome breakage. This is an essential requirement because the cell must repair the latter but not the former. Telomeric DNA is made up of hundreds of copies of a repeated motif, 5′-TTAGGG-3′ in humans, with a short extension of the 3′-terminus of the double-stranded DNA molecule (Figure 7.9). A series of telomere-binding proteins, including TRF1, TRF2, and POT1, bind to the telomere repeat sequence and together with other proteins make a structure called a shelterin. This structure protects the telomeres from degradation by nuclease enzymes and mediates the enzymatic activity that maintains the length of each telomere during DNA replication.

### 7.2 HOW ARE THE GENES ARRANGED IN A NUCLEAR GENOME?

Now that we understand the physical structure of the eukaryotic nuclear genome, we can move on to consider the genetic features: which genes are present and how those genes are arranged.

**Genes are not evenly distributed within a genome**

Much of what we know about the identities and relative positions of genes in eukaryotic genomes has been obtained by annotation of genome sequences by the bioinformatic and experimental methods that we studied in Chapters 5 and 6. Prior to the genome sequencing era, linkage analysis had enabled some genes to be mapped and their functions to be assigned, at least for well-studied species such as yeast, fruit flies, and humans. DNA sequences were also available for many individual genes and for short segments of individual chromosomes, such as the 65 kb stretch of human chromosome 11 that contains the β-globin gene cluster (see Figure 7.19). Geneticists at that time, therefore, had a broad but approximate understanding of gene distribution within, for example, the human genome, along with a detailed knowledge of the structures of those individual genes that had been cloned and sequenced.

One outcome of the pre-genome sequencing studies was a growing awareness that genes are not arranged evenly along the length of a eukaryotic chromosome. This hypothesis was supported by two lines of evidence. The first of these relates to the banding patterns produced when chromosomes are stained. The dyes used in these procedures (see Table 7.1) bind to DNA molecules, but in most cases they have preferences for certain base pairs. Giemsa, for example, has a greater affinity for DNA regions that are rich in A and T nucleotides. The dark G-bands in the human karyogram (see Figure 7.6) were therefore thought to be AT-rich regions of the genome. The base composition of the human genome as a whole is 59.7% A + T, so the dark G-bands must have AT contents substantially greater than 60%. Cytogeneticists therefore predicted that there would be fewer genes in dark G-bands because genes generally have AT contents of 45–50%.

The second line of evidence pointing to uneven gene distribution derived from the isochore model of genome organization. According to this model, first proposed in the early 1980s, eukaryotic genomes are mosaics of segments of DNA, each at least 300 kb in length, with each segment having a uniform base composition that differs from that of the adjacent segments. Support for the isochore model came from experiments in which genomic DNA was broken into fragments of approximately 100 kb and treated with dyes that bind specifically to AT- or GC-rich regions, and the pieces then separated by isopycnic or buoyant density centrifugation. In this method, the DNA fragments are pipetted onto the surface of a high-density solution, such as 8 M cesium chloride, which is then centrifuged at a very high speed: at least 450,000 g for several hours. The centrifugal force carries some of the CsCl molecules toward the bottom of the centrifuge tube, forming a density gradient. Each DNA fragment migrates to the position in the tube where the density of the CsCl solution equals its own buoyant density. The latter
is influenced by a number of factors, including conformation (linear, circular, and supercoiled versions of the same DNA molecule have different buoyant densities), but if all the molecules being studied are linear, then their buoyant densities are determined mainly by their GC content, according to the formula

\[
\text{% GC content} = \frac{\text{buoyant density (g cm}^{-3}) - 1.660}{0.098} \times 100
\]

When this experiment was carried out with human DNA, five fractions were seen, each identified as a different isochore type with a distinctive base composition: two lower density AT-rich isochores, called L1 and L2, and three higher density GC-rich classes called H1, H2, and H3 (Figure 7.10). The last of these, H3, is the least abundant in the human genome, making up only 3% of the total, but it contains over 25% of the genes, a clear indication that genes are not distributed evenly through the human genome.

The first eukaryotic genome sequences confirmed that genes are unevenly distributed along the lengths of individual chromosomes. For example, the average gene density in the Arabidopsis thaliana genome is 25 genes/100 kb, but even outside the centromeres and telomeres, where there are very few genes, the density varies from 1 to 38 genes/100 kb, as illustrated in Figure 7.11 for the largest of the plant’s five chromosomes. The same is true for the human genome: several chromosomes contain gene deserts in which the density is very low over regions as long as several megabase pairs. The distribution of protein-coding genes between different human chromosomes is also very uneven, ranging from 3.16 genes/Mb for chromosome 13 to 22.61 genes/Mb for chromosome 19. To a certain extent, the interpretation of G-bands as regions of low gene content has been confirmed, as these bands often contain just a few long genes, with multiple introns, occasionally with a single gene spanning an entire band. The isochore model, however, has fared less well. Although the original proponents of the model still argue that isochores can be detected in most if not all eukaryotic genomes, examination of genome sequences suggests that the isochore theory oversimplifies what is, in reality, a much more complex pattern of variations in base composition along the length of a eukaryotic chromosome. Mammalian genomes, for example, do contain discrete regions within which the GC content is homogeneous, but these homogeneous domains make up only two-thirds of the genome as a whole, while the remainder has mixed GC composition with no recognizable domain structure. The majority of the homogeneous domains are less than 100 kb, with only 2% of these domains, covering less than 28% of the genome, made of segments of >300 kb, the supposed lengths of isochores. So the isochore theory might be a misconception, but it has been a useful misconception, as it played an important role in stimulating molecular biologists of the pre-sequence era to think about genome structure.

### A segment of the human genome

The variations in gene density that occur along the length of a eukaryotic chromosome mean that it is difficult to identify regions in which the organization of the genes can be looked on as typical of the genome as a whole. Despite this difficulty,
it is clear that the overall pattern of gene organization varies greatly between different eukaryotes, and we need to understand these differences because they reflect important distinctions between the genetic features and evolutionary histories of these genomes. To begin to address this issue, we will look in detail at a small part of the human genome.

The segment that we will examine (Figure 7.12) comes from midway along the long arm of human chromosome 1. It is 200 kb in length and runs from nucleotide position 55,000,000 to position 55,200,000. The segment contains the following:

- All or part of three protein-coding genes:
  - The end of the **BSND** gene, which starts at position 54,998,944. **BSND** codes for a chloride channel protein. This is a membrane-bound protein that forms a pore through which various ions, including chloride, can enter and leave the cell.
  - **PCSK9** codes for proprotein convertase subtilisin/kexin type 9, a protein made in liver, intestine, and kidney tissues that is involved in breakdown of low-density lipoproteins, thereby playing an important role in the metabolism of cholesterol.
  - The start of **USP24**, specifying ubiquitin-specific peptidase 24, a protease that removes **ubiquitin** side chains from proteins that have been modified by **ubiquitination**. Ubiquitin is a small regulatory protein whose addition to or removal from a protein controls that protein’s location in the cell and eventual degradation (Section 13.3). **USP24** ends at position 55,215,366, so most of it is contained in the segment shown in Figure 7.12.

  Note that each of these three genes is discontinuous: there are three introns in **BSND**, 11 in **PCSK9**, and 73 in **USP24**.

- A vast number of **interspersed repeat** sequences. These are sequences that recur at many places in the genome. There are four main types of interspersed repeats, called **SINEs** (short interspersed nuclear elements), **LINEs** (long interspersed nuclear elements), **LTR** (long terminal repeat) elements, and **DNA transposons** (Section 9.2). Multiple copies of each type are seen in this short segment of the genome, in both the intergenic regions and in the introns of the protein-coding genes.

The most striking feature of this 200 kb segment of the human genome is the relatively small amount of space taken up by the coding parts of the genes. When

**Figure 7.12** A 200 kb segment of the human genome. The segment runs from nucleotide position 55,000,000 to position 55,200,000 of chromosome 1. Within the genes, exons are shown as green boxes and introns are shown as gray boxes. (Data from the UCSC Genome Browser, hg38 assembly.)
added together, the total length of exons (the parts of genes that contain the biological information) is 10,664 bp, equivalent to 5.33% of the 200 kb segment. In fact, this segment is rather rich in genes: all the exons in the human genome make up only 48 Mb, just 1.5% of the total. In contrast, 44% of the genome is taken up by interspersed repeats (Figure 7.13).

The yeast genome is very compact

How extensive are the differences in gene organization among eukaryotes? There are certainly very substantial differences in genome size, with the smallest eukaryotic genomes being less than 10 Mb in length and the largest over 100,000 Mb. As can be seen in Figure 7.14 and Table 7.2, this size range coincides to a certain extent with the complexity of the organism: the simplest eukaryotes such as fungi have the smallest genomes, and higher eukaryotes such as vertebrates and flowering plants have the largest ones. This might appear to make sense, as one would expect the complexity of an organism to be related to the number of genes in its genome, so that higher eukaryotes would need larger genomes to accommodate the extra genes. However, the correlation is far from precise. The human genome is 3235 Mb and, according to the most recent annotation, contains 20,441 protein-coding genes. The genome of the yeast *Saccharomyces cerevisiae*, which at 12.2 Mb is 0.004 times the size of the human nuclear genome, would therefore be expected to contain 0.004 × 20,441 genes, which is just 82. In fact the *S. cerevisiae* genome contains 6692 protein-coding genes.

For many years the lack of precise correlation between the complexity of an organism and the size of its genome was looked on as a bit of a puzzle, the so-called C-value paradox. In fact, the answer is quite simple: space is saved in the genomes of less complex organisms because the genes are more closely packed together. The *S. cerevisiae* genome illustrates this point, as we can see in Figure 7.15, where a typical 200 kb segment of the yeast genome is displayed. This segment comes from chromosome IV, which is the largest of the 16 yeast chromosomes but still only 1.53 Mb in length, reflecting the much smaller size of the yeast genome as a whole. The 200 kb segment, running from nucleotide position 250,000 to position 450,000, therefore comprises 13% of the length of chromosome IV, and in fact it ends adjacent to the centromere of this chromosome. When we compare this segment with the 200 kb stretch of the human genome that we previously examined (Figure 17.5A and B; see also Figure 17.12), three differences immediately become apparent:
7.2 HOW ARE THE GENES ARRANGED IN A NUCLEAR GENOME?

- The gene density in the yeast genome is much higher than that for humans. This segment of chromosome IV contains 104 genes thought to code for proteins, four that specify transfer RNAs, and one small nucleolar RNA gene.

- Relatively few of the yeast genes are discontinuous. In this segment of chromosome IV there are four introns, one in each of four protein-coding genes. In the entire yeast genome there are only 344 discontinuous genes, and the vast majority of these genes have just one intron each.

- This part of chromosome IV contains just two interspersed repeats. Both are truncated LTR elements (Section 9.2), one called a delta sequence and the other a tau sequence. Interspersed repeats make up only 3.4% of the yeast genome. The most prevalent types are full-length LTR elements (about 50 copies in total depending on the particular strain of \textit{S. cerevisiae}) and truncated LTR sequences (300–400 copies).

  The genetic organization of the yeast genome is clearly much more compact than that of the human version. The genes themselves are much shorter, having fewer introns, and the spaces between the genes are relatively short, with much less space taken up by interspersed repeats and other noncoding sequences.

**Gene organization in other eukaryotes**

The hypothesis that less complex eukaryotes have more compact genomes holds when other species are examined. Next we will examine a 200 kb segment of the fruit fly genome. If we agree that a fruit fly is more complex than a yeast cell but less complex than a human, then we would expect the organization of the fruit fly genome to be intermediate between that of yeast and humans. This is what we see in Figure 7.15C. Again we have chosen the largest of the chromosomes in the karyogram, our segment running from positions 5,300,000 to 5,500,000 on
the left arm of chromosome 3. There are eight genes in this region, six of which are discontinuous, with some of the introns similar in length to those in human genes. There are only two interspersed repeat sequences, both of which are LTR elements. The fruit fly genome also contains SINEs, LINEs, and DNA transposons, but there are none in this particular segment. The picture is similar when the entire genome sequences of the three organisms are compared (Table 7.3).

The gene density in the fruit fly genome is intermediate between that of yeast and humans, and the average fruit fly gene has many more introns than the average yeast gene but fewer introns than the average human gene.

The comparison between yeast, fruit fly, and human genomes also holds true when we consider interspersed repeats (see Table 7.3). These make up 3.4% of the yeast genome, about 12% of the fruit fly genome, and 44% of the human genome. It is beginning to become clear that interspersed repeats play an intriguing role in dictating the compactness or otherwise of a genome. This is strikingly illustrated by the maize genome, which at 2500 Mb is relatively small for a flowering plant. The 200 kb segment shown in Figure 7.15D has nine genes, seven of which contain one or more short introns. Instead of the genes, the dominant feature of this genome

<table>
<thead>
<tr>
<th>Species</th>
<th>Genome size (Mb)</th>
</tr>
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<tbody>
<tr>
<td><em>Encephalitozoon intestinalis</em></td>
<td>2.3</td>
</tr>
<tr>
<td><em>Saccharomyces cerevisiae</em></td>
<td>12.2</td>
</tr>
<tr>
<td><em>Aspergillus nidulans</em></td>
<td>31</td>
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<tr>
<td><strong>Protozoa</strong></td>
<td></td>
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<td><em>Plasmodium falciparum</em></td>
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<td><em>Dictyostelium discoideum</em></td>
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<td><em>Neospora caninum</em></td>
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<tr>
<td><em>Amoeba dubia</em></td>
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</tr>
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<td><strong>Invertebrates</strong></td>
<td></td>
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<tr>
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</tr>
<tr>
<td><em>Drosophila melanogaster</em></td>
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<td><em>Bombyx mori</em> (silkworm)</td>
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<tr>
<td><em>Strongylocentrotus purpuratus</em> (sea urchin)</td>
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</tr>
<tr>
<td><em>Laupala sp. (cricket)</em></td>
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<tr>
<td><em>Locusta migratoria</em> (locust)</td>
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<tr>
<td><em>Takifugu rubripes</em> (pufferfish)</td>
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<tr>
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</tr>
<tr>
<td><em>Homo sapiens</em></td>
<td>3235</td>
</tr>
<tr>
<td><em>Protopterus aethiopicus</em> (marbled lungfish)</td>
<td>143,000</td>
</tr>
<tr>
<td><strong>Plants</strong></td>
<td></td>
</tr>
<tr>
<td><em>Arabidopsis thaliana</em> (vetch)</td>
<td>135</td>
</tr>
<tr>
<td><em>Oryza sativa</em> (rice)</td>
<td>430</td>
</tr>
<tr>
<td><em>Zea mays</em> (maize)</td>
<td>2500</td>
</tr>
<tr>
<td><em>Pisum sativum</em> (pea)</td>
<td>4300</td>
</tr>
<tr>
<td><em>Triticum aestivum</em> (wheat)</td>
<td>16,500</td>
</tr>
<tr>
<td><em>Paris japonica</em> (canopy plant)</td>
<td>165,000</td>
</tr>
</tbody>
</table>
segment is the interspersed repeats, which have been described as forming a sea within which islands of genes are located. The interspersed repeats are mainly of the LTR element type, which comprise a large part of the intergenic regions of the segment, and on their own are estimated to make up approximately 50% of the maize genome. It is becoming clear that one or more families of interspersed repeats have undergone a massive proliferation in the genomes of certain species. This may provide an explanation for the most puzzling aspect of the C-value paradox, which is not the general increase in genome size that is seen in increasingly complex organisms but the fact that similar organisms can differ greatly in genome size. A good example is provided by *Amoeba dubia*, which, being a protozoan, might be expected to have a genome of less than 100 kb, similar to other protozoa such as *Dictyostelium discoideum* (see Table 7.2). In fact the *Amoeba* genome is over 200,000 Mb. Similarly, we might guess that the genomes of crickets would be similar in size to those of other insects, but crickets have genomes of approximately 2000 Mb, 11 times that of the fruit fly.

### 7.3 HOW MANY GENES ARE THERE AND WHAT ARE THEIR FUNCTIONS?

In Chapters 5 and 6 we examined the methods used to locate genes in a genome sequence and to assign functions to those genes. We learnt that although there are a variety of bioinformatics and experimental methods for gene identification
and functional analysis, a complete genome annotation is, at present, difficult or impossible to achieve with any eukaryotic species. This means that we do not know precisely how many genes are present in a genome, and we cannot give a full description of the functions of the proteins specified by that genome. We can, however, make reasonable estimates of the gene numbers for many species whose genomes have been sequenced, and we can make cautious extrapolations from those genes whose functions have been identified to the functional capacity of the genome as a whole.

**Gene numbers can be misleading**

The current annotation of the human genome recognizes 20,441 protein-coding genes and 22,219 genes for noncoding RNAs. During recent years, the trend has been for the number of protein-coding genes to decrease as questionable ORFs are gradually discarded, with some estimates suggesting that there might be as few as 19,000 in the human genome. The number of noncoding RNA genes, on the other hand, has increased substantially in recent years as different types of noncoding RNA are discovered. This area of genomics research is very fluid at the moment, as we will discover when we study the compositions of transcriptomes in Section 12.1. Future changes in the accepted numbers of noncoding RNA genes, in any organism, are likely to be a balance between the discovery of new genes whose products are genuinely functional and the removal of sequences initially identified as genes but whose RNA transcripts are subsequently interpreted as junk.

Our expectation might be that humans, being the most sophisticated species on the planet, would have more genes than any other organism. An initial comparison between the numbers of protein-coding genes in different species supports this prejudice (Table 7.4). Yeast has just 6692 genes, the fruit fly has fewer than 14,000, and chickens have 15,508. But the correlation begins to go awry when we look more carefully at the figures. Humans and other primates are by no means the most complex organisms in terms of gene counts: *Arabidopsis thaliana* has over 27,000 protein-coding genes and rice has over 35,000. We might ascribe the higher gene numbers of plants to the need for plants to code for proteins involved in photosynthesis, but this would be a misassumption, because the photosynthetic capability of plants is outweighed in this regard by the many unique specializations of mammals and other higher vertebrates. In any case, the higher gene content of plant genomes is neither the only nor the most striking anomaly revealed by the numbers in Table 7.4. The genome of the microscopic worm called *Caenorhabditis elegans*, whose adult body comprises just over 1000 cells, contains 20,362 protein-coding genes, which is almost certainly more than the actual number of functional protein-coding genes in the human genome.

These gene number comparisons lead us into an important aspect of genome biology. Before the human genome was sequenced, it was anticipated that there

<table>
<thead>
<tr>
<th>Species</th>
<th>Protein-coding genes</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Saccharomyces cerevisiae</em> (budding yeast)</td>
<td>6692</td>
</tr>
<tr>
<td><em>Schizosaccharomyces pombe</em> (fission yeast)</td>
<td>5145</td>
</tr>
<tr>
<td><em>Caenorhabditis elegans</em> (nematode worm)</td>
<td>20,362</td>
</tr>
<tr>
<td><em>Arabidopsis thaliana</em> (plant)</td>
<td>27,416</td>
</tr>
<tr>
<td><em>Drosophila melanogaster</em> (fruit fly)</td>
<td>13,918</td>
</tr>
<tr>
<td><em>Oryza sativa</em> (rice)</td>
<td>35,679</td>
</tr>
<tr>
<td><em>Gallus gallus</em> (chicken)</td>
<td>15,508</td>
</tr>
<tr>
<td><em>Homo sapiens</em> (human)</td>
<td>20,441</td>
</tr>
</tbody>
</table>

Data from Ensembl release 85, Ensembl Plants release 32, and Ensembl Fungi release 32.
Chapter 7: Eukaryotic Nuclear Genomes

would be 80,000–100,000 protein-coding genes, this number remaining in vogue up to a few months before the draft sequence was completed in 2000. This early estimate was high because it was based on the supposition that, in most cases, a single gene specifies a single mRNA and a single protein. According to this model, the number of genes in the human genome should be similar to the number of proteins in human cells, leading to the estimates of 80,000–100,000. The discovery that the actual number of protein-coding genes is much lower indicates that it is possible for an individual gene to specify more than one protein. This is the case for many of the discontinuous genes in the human genome (Section 1.2). When introns were first discovered, it was thought that a discontinuous gene would have just one splicing pathway, in which all of the exons are joined together to give a single mRNA. We now know that many discontinuous genes have alternative splicing pathways, which means that their pre-mRNAs can be processed in a variety of ways, to give a series of mRNAs made up of different combinations of exons. Each of these genes can therefore direct synthesis of related but different proteins.

An example of a human gene with two splicing pathways, one followed in the thyroid and a second in nervous tissue, is shown in Figure 7.16. Alternative splicing is relatively common in vertebrates, with 75% of all human protein-coding genes, representing 95% of those with two or more introns, undergoing alternative splicing, giving rise to an average of four different spliced mRNAs per gene. Alternative splicing also occurs in lower eukaryotes, but it is less prevalent. In C. elegans, for example, only about 25% of the protein-coding genes have alternative splicing pathways, with an average of 2.2 variants per gene.

Because of alternative splicing, the question “How many genes are there?” has no real biological significance, as the number of genes does not indicate the number of proteins that can be synthesized and hence is not a measure of the coding capacity of a genome. A better measure of the biological complexity of an organism is provided by categorizing the genes, including the splice variants, according to function. Now the problem becomes lack of completeness, because of the difficulty in identifying functions, even for a relatively simple organism such as Saccharomyces cerevisiae. It is quite probable that certain categories of genes are underrepresented in the existing annotations, because those genes have functions that are particularly difficult to identify. With these qualifications in mind, we will compare the gene catalogs of different species.

Gene catalogs reveal the distinctive features of different organisms

The functions of more than half of the 20,000 human protein-coding genes are known or can be inferred with a reasonable degree of certainty. These functions are described by the Gene Ontology (GO) nomenclature (Section 6.1), which means that groups of genes with related activities can be identified. The GO system enables function to be described in different ways (for example, molecular function, biological process), and groups of genes can be further subclassified in a hierarchical manner. To illustrate the degree of sophistication that can be attained, we will explore the human gene catalog as described in terms of
molecular function (Figure 7.17). We see that, at the highest level of classification for this ontology term, 5570 genes code for proteins that are involved in binding, which is defined as noncovalent interaction between a protein and specific sites on another molecule, and another 5090 genes code for proteins with catalytic activity (enzymes). If we drill down to the next level of the GO hierarchy, we discover that 2854 of the binding proteins attach to other proteins and 2350 bind to nucleic acids. Among the proteins with catalytic activity, the largest group is hydrolases with 2134 genes, followed by transferases with 1542 genes. Back at the upper level of molecular function, the next largest group of genes (after those involved in binding and catalytic activity) codes for proteins with receptor activity. These specify the cell-surface and cytoplasmic receptor proteins that respond to extracellular signals such as the presence of hormones and growth factors. There is also a group of 718 genes involved in signal transduction, which form the pathways leading from the receptor proteins to the genes and enzymes whose activities must be changed in response to the extracellular signals. Another 1014 genes in the human catalog code for proteins with transporter...
activity, responsible for the controlled movement of molecules and ions across the outer cell membrane and into and out of organelles such as the mitochondria, and a further 927 are structural proteins, forming part of, for example, the cell cytoskeleton and the extracellular matrix.

Similar analyses of the human gene catalog can be carried out for other high-level GO categories such as biological process. These functional analyses are interesting with regard to the descriptions they provide of the biological and biochemical capabilities of the human genome, but on its own the catalog of a single species has limited information content. These catalogs become more revealing when comparisons are made between species, as then it is possible to understand the genomic basis for some of the distinctive features of different organisms. These studies suggest that all eukaryotes possess the same basic set of genes but that more complex species have a greater number of genes in each category. This point is illustrated by comparing the molecular function categorizations for human, fruit fly, yeast, *C. elegans*, and *A. thaliana* (Figure 7.18). Humans have the greatest number of genes in five of the nine categories; the exceptions are antioxidant activity, catalytic activity, translation regulator, and transporter activity, where *Arabidopsis* comes out on top. This is understandable in view of the photosynthetic capability of *Arabidopsis*, which requires an additional set of genes not present in the other four genomes included in this comparison. These extra enzymes are needed not only to carry out the Calvin cycle and other unique biochemical pathways of photosynthesis but also to deal with the oxidants produced as a byproduct of photosynthetic activity and to transport ions and molecules into and out of the chloroplasts, which are absent in the other four species. Other interesting features emerge: notably, that humans are the only species in which binding is the category containing the greatest number of genes. This indicates that a greater ability to form protein–protein and protein–DNA interactions might explain, in part, the biological sophistication of humans and other vertebrates.

**Figure 7.18 Comparison of the gene catalogs of different species.** The gene catalogs of humans, *Drosophila melanogaster*, *Saccharomyces cerevisiae*, *Caenorhabditis elegans*, and *Arabidopsis thaliana* are shown, categorized according to the GO term molecular function. (Data from PANTHER 11.0.)
It is also interesting to note that *C. elegans* has a relatively high number of genes involved in receptor activity and signal transduction, twice as many as fruit flies and over half the number present in the human genome. Probing down into the subcategories of these functions, it becomes clear that the human and fruit fly genomes specify a broader range of receptor proteins, whereas *C. elegans* relies on a large set of receptors of a single type (G-protein receptors). In contrast, the three genomes code for similar sets of transporter proteins, suggesting that the ability to control the movement of metabolites across their membranes was fully mature at a relatively early stage in eukaryote evolution. Comparisons between gene catalogs therefore help us to understand not only the coding capabilities of different genomes but also when those capabilities evolved.

The GO system is not the only possible way of categorizing a gene catalog. An interesting alternative is to base the classification not on the functions of genes but on the structures of the proteins that they specify. A protein molecule is constructed from a series of **domains**, each of which has a particular biochemical function. Examples are the **zinc finger**, which is one of several domains that enable a protein to bind to a DNA molecule (Section 11.2), and the death domain, made up of six α-helices, which is present in many proteins involved in apoptosis. Each domain has a characteristic amino acid sequence, usually not exactly the same sequence in every example of that domain but close enough for the presence of a particular domain to be recognizable by examining the amino acid sequence of the protein. The amino acid sequence of a protein is specified by the nucleotide sequence of its gene, so the domains present in a protein can be determined from the nucleotide sequence of the gene that codes for that protein. The genes in a genome can therefore be categorized according to the protein domains that they specify. This method has the advantage that it can be applied to genes whose overall functions are not known and hence can encompass a larger proportion of the genes present in a genome. It shows that a vertebrate genome specifies a number of protein domains that are rare or absent from the genomes of the other organisms. These domains include several involved in activities such as cell adhesion, electrical coupling between cells, and growth of nerve cells (Table 7.5). These functions are interesting because they are ones that we look on as conferring the distinctive features of vertebrates compared with other types of eukaryotes.

**Families of genes**

Since the earliest days of DNA sequencing, it has been known that **multigene families**—groups of genes of identical or similar sequence—are common features of many genomes. For example, every eukaryote that has been studied (as well as

### TABLE 7.5 EXAMPLES OF PROTEIN DOMAINS SPECIFIED BY DIFFERENT GENOMES

<table>
<thead>
<tr>
<th>Domain</th>
<th>Function</th>
<th>Number of genes in the genome that contain the domain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Human</td>
</tr>
<tr>
<td>Zinc finger, Cys&lt;sub&gt;2&lt;/sub&gt;His&lt;sub&gt;2&lt;/sub&gt; type</td>
<td>DNA binding</td>
<td>2474</td>
</tr>
<tr>
<td>Zinc finger, GATA type</td>
<td>DNA binding</td>
<td>44</td>
</tr>
<tr>
<td>Homeobox</td>
<td>Gene regulation during development</td>
<td>827</td>
</tr>
<tr>
<td>Death</td>
<td>Programmed cell death</td>
<td>118</td>
</tr>
<tr>
<td>Connexin</td>
<td>Electrical coupling between cells</td>
<td>70</td>
</tr>
<tr>
<td>Ephrin</td>
<td>Nerve cell growth</td>
<td>15</td>
</tr>
</tbody>
</table>

Data from InterPro 58.0.
all but the simplest bacteria) has multiple copies of the genes for ribosomal RNAs. This is illustrated by the human genome, which contains several thousand genes for the 5S rRNA, many of these located in a single cluster on chromosome 1. There are also several hundred copies of a repeat unit containing the 28S, 5.8S, and 18S rRNA genes, with major groups of this unit on each of chromosomes 13, 14, 15, 21, and 22 (see Figure 7.6). Ribosomal RNAs are components of the protein-synthesizing particles called ribosomes, and it is presumed that their genes are present in multiple copies because there is a heavy demand for rRNA synthesis during cell division, when several tens of thousands of new ribosomes must be assembled.

The rRNA genes are examples of simple or classical multigene families, in which all the members have identical or nearly identical sequences. These families are believed to have arisen by gene duplication, with the sequences of the individual members kept identical by an evolutionary process that, as yet, has not been fully described (Section 18.2). Other multigene families, more common in higher eukaryotes than in lower eukaryotes, are classified as complex because the individual members, although similar in sequence, are sufficiently different for the gene products to have distinctive properties. The mammalian globin genes are one of the best examples of this type of multigene family. Globins are the blood proteins that combine to make hemoglobin, each molecule of hemoglobin comprising two α-type and two β-type globins. In humans, the α-type globins are coded by a small multigene family on chromosome 16 and the β-type globins are coded by a second family on chromosome 11 (Figure 7.19). These genes were among the first to be sequenced, back in the late 1970s. The sequence data showed that the genes in each family are similar to one another but by no means identical. In fact, the nucleotide sequences of the two most different genes in the β-type cluster, coding for the β- and ε-globins, display only 79.1% identity. Although this is similar enough for both proteins to be β-type globins, it is sufficiently different for them to have distinctive biochemical properties. Similar variations are seen in the α-cluster.

Why are the members of the globin gene families so different from one another? The answer was revealed when the expression patterns of the individual genes were studied. It was discovered that the genes are expressed at different stages in human development: for example, in the β-type cluster, ε is expressed in the early embryo, Gγ and Aγ (whose protein products differ by just one amino acid) in the fetus, and δ and β in the adult (see Figure 7.19). The biochemical properties of the different β-globin proteins reflect the slight changes in the physiological role that hemoglobin plays during the course of human development. For example, the developing fetus must obtain oxygen from its mother. This means that oxygen must be transferred from the mother’s hemoglobin molecules to those of the fetus. For this to be possible, fetal hemoglobin must have a greater affinity for oxygen compared to adult hemoglobin, so when the two types of hemoglobin mix in the placenta, the fetus is able to steal oxygen from its mother. The nucleotide sequences of the Gγ and Aγ genes specify globin proteins with the higher degree of oxygen affinity that is needed to allow this to happen.

In some multigene families, the individual members are clustered, as with the globin genes, but in others the genes are dispersed around the genome. An example of a dispersed family is provided by the five human genes for aldolase, an enzyme involved in energy generation, which are located on chromosomes 3, 9, 10, 16, and 17. The important point is that, even though dispersed, the members of the multigene family have sequence similarities that point to a common origin. The answer was revealed when the expression patterns of the individual genes were studied. It was discovered that the genes are expressed at different stages in human development: for example, in the β-type cluster, ε is expressed in the early embryo, Gγ and Aγ (whose protein products differ by just one amino acid) in the fetus, and δ and β in the adult (see Figure 7.19). The biochemical properties of the different β-globin proteins reflect the slight changes in the physiological role that hemoglobin plays during the course of human development. For example, the developing fetus must obtain oxygen from its mother. This means that oxygen must be transferred from the mother’s hemoglobin molecules to those of the fetus. For this to be possible, fetal hemoglobin must have a greater affinity for oxygen compared to adult hemoglobin, so when the two types of hemoglobin mix in the placenta, the fetus is able to steal oxygen from its mother. The nucleotide sequences of the Gγ and Aγ genes specify globin proteins with the higher degree of oxygen affinity that is needed to allow this to happen.
evolutionary origin. When these sequence comparisons are made, it is sometimes possible to see relationships not only within a single gene family but also between different families. All of the genes in the \( \alpha \)- and \( \beta \)-globin families, for example, have some sequence similarity and are thought to have evolved from a single ancestral globin gene. We therefore refer to these two multigene families as comprising a single globin gene superfamily, and from the similarities between the individual genes we can chart the duplication events that have given rise to the series of genes that we see today (Section 18.2).

**Pseudogenes and other evolutionary relics**

As well as the functional genes that are expressed at different developmental stages, the human globin gene clusters also contain five pseudogenes. These are the sequences labeled \( \psi_{A1}, \psi_{A2}, \psi_{\alpha}, \psi_{\beta} \) in the \( \alpha \)-globin cluster and \( \psi_{\beta} \) in the \( \beta \)-globin cluster (see Figure 7.19). Two of the five genes in the dispersed aldolase family are pseudogenes; these are located on chromosomes 3 and 10. What are these pseudogenes?

A pseudogene is a sequence of nucleotides that resembles a genuine gene but does not specify a functional RNA or protein. Pseudogenes are derived from genuine genes and can therefore be looked on as an evolutionary relic, an indication that genomes are continually undergoing change. In some cases, a gene loses its function and becomes a pseudogene simply because its nucleotide sequence changes by mutation. Many mutations have only minor effects on the activity of a gene but some are more important, and it is quite possible for a single nucleotide change to result in a gene becoming completely nonfunctional. Once a pseudogene has become nonfunctional, it will degrade through accumulation of more mutations, and eventually it will no longer be recognizable as a gene relic. Pseudogenes that arise in this way are called conventional or nonprocessed pseudogenes. They fall into two broad classes:

- **A duplicated pseudogene** arises when a member of a multigene family becomes inactivated by mutation. This event is usually not deleterious to the organism because the other genes in the family are still active and the function specified by the pseudogene is not lost. Comparisons between different genomes have revealed many instances where a pseudogene in one species is the homolog of a functional gene in a second species. For example, the \( \delta \)-globin gene, which is active in humans, is a pseudogene in mice. The implication is that the \( \delta \)-globin gene became inactivated by a mutation that occurred at some point during the evolutionary lineage leading to mice, after this lineage diverged from the one leading to humans.

- **Unitary pseudogenes** also arise from mutation, but in this case the gene is not a member of a family, so the resulting loss of function is not compensated by the activity of other genes. Unitary pseudogenes are rare, because the loss of function will usually be lethal, which means that cells that experience such a mutation will die and not contribute to the subsequent evolutionary lineage. Those unitary pseudogenes that appear in a genome are therefore ones whose loss of function could be tolerated. There are probably less than 50 unitary pseudogenes in the human genome, the best-known example being the l-gulono-\( \gamma \)-lactone oxidase pseudogene. The functional version of this gene enables many mammals to synthesize ascorbic acid, but in the Haplorhini group of primates the gene is a pseudogene, which means that haplorhines, including humans, must obtain ascorbic acid (otherwise known as vitamin C) from their diets.

Other pseudogenes arise by a process not involving mutation. These are called processed pseudogenes, and they result from an abnormal adjunct to gene expression. A processed pseudogene is derived from the mRNA copy of a gene by synthesis of a cDNA copy, which subsequently reinserts into the genome (Figure 7.20). Because a processed pseudogene is a copy of an mRNA molecule,
it does not contain any introns that were present in its parent gene. It also lacks the nucleotide sequences immediately upstream of the parent gene, which is the region in which the signals used to switch on expression of the parent gene are located. The absence of these signals means that a processed pseudogene is inactive. Additionally, genomes also contain other evolutionary relics in the form of truncated genes, which lack a greater or lesser stretch from one end of the complete gene, and gene fragments, which are short isolated regions from within a gene (Figure 7.21).

In recent years, there has been increasing debate about the possibility that some sequences identified as pseudogenes do in fact have a functional role of some kind. Some pseudogenes are transcribed into RNA and a smaller number, just over 100 for the human genome, also direct synthesis of a protein. Expression is not, in itself, evidence of a function, because it is conceivable that a nonprocessed pseudogene could be transcribed and/or translated simply because its upstream signals and open reading frame have not yet decayed to the stage where expression is impossible. Before a pseudogene is reassigned as a functional sequence, it is necessary to prove that the expression product plays some active role in the cell. There are suggestions that this might be the case for at least a few pseudogenes. An example in humans is PTENP1, which is a nonprocessed pseudogene derived from the gene for the PTEN phosphatase, an enzyme involved in one of the signal transduction pathways that control cell division. Expression of the PTEN gene is regulated in part by miRNAs that attach to the PTEN RNA and promote its degradation (Section 12.3). The transcripts from the PTENP1 pseudogene also bind some of these miRNAs, reducing their abundance in the cell and ensuring that the PTEN gene is not completely silenced (Figure 7.22A). In experimental systems, reducing the level of PTENP1 transcription leads to silencing of PTEN, which in turn results in an increased cell division rate. These results correlate with the observation that the PTENP1 pseudogene is deleted in some types of colon cancer, implying that in these cells the absence of PTENP1 transcripts results in PTEN silencing and the uncontrolled division that gives rise to the cancerous state (Figure 7.22B). The system as a whole would appear to provide strong evidence that the PTENP1 sequence is not a genuine pseudogene but instead plays an important regulatory function. However, if a gene is providing a

![Figure 7.21](image_url)  
**Figure 7.21** A truncated gene and a gene fragment.

![Figure 7.22](image_url)  
**Figure 7.22** A possible function for the PTENP1 pseudogene. (A) In normal tissue, the binding of miRNAs to PTENP1 transcripts is thought to prevent silencing of PTEN. (B) In some cancers, PTENP1 is deleted. The absence of the miRNA binding sites normally provided by the PTENP1 transcripts might lead to additional silencing of PTEN mRNAs, resulting in a loss of control over cell division.
useful function, then we expect natural selection to be acting in a positive way on that gene, and evolutionary studies have failed to find evidence for this being the case with PTENP1. Similar evolutionary studies have also failed to reveal indicators of positive selection for most of the human pseudogenes that give rise to protein products. Questions remain, therefore, about the importance of the apparent roles played by those pseudogenes that are still expressed.

7.4 THE REPETITIVE DNA CONTENT OF EUKARYOTIC NUCLEAR GENOMES

Our examination of the genomes of humans and other eukaryotes showed us that large parts of these DNA sequences are made up of repetitive elements (see Figures 7.12 and 7.15). Repetitive DNA can be divided into two categories (Figure 7.23): interspersed repeats, whose individual repeat units are distributed around the genome in an apparently random fashion, and tandemly repeated DNA, whose repeat units are placed next to each other in an array.

**Tandemly repeated DNA is found at centromeres and elsewhere in eukaryotic chromosomes**

Tandemly repeated DNA is also called satellite DNA because DNA fragments containing tandemly repeated sequences form satellite bands when genomic DNA is fractionated by density gradient centrifugation (see Section 7.2). For example, when broken into fragments 50–100 kb in length, human DNA forms a main band (buoyant density 1.701 g cm$^{-3}$) and three satellite bands (1.687, 1.693, and 1.697 g cm$^{-3}$). The main band contains DNA fragments made up mostly of single-copy sequences with GC compositions close to 40.3%, the average value for the human genome. The satellite bands contain fragments of repetitive DNA, and hence have GC contents and buoyant densities that are atypical of the genome as a whole (Figure 7.24). This repetitive DNA is made up of long series of tandem repeats, possibly hundreds of kilobases in length. A single genome can contain several different types of satellite DNA, each with a different repeat unit; these units range from less than 5 to more than 200 bp in length. The three satellite bands in human DNA include at least four different repeat types.

We have already encountered one type of human satellite DNA: the alphoid DNA repeats found in the centromere regions of chromosomes (Section 7.1). Although some satellite DNA is scattered around the genome, most is located in the centromeres, where it may play a structural role, possibly as a binding site for one or more of the special centromeric proteins.

**Minisatellites and microsatellites**

Although not appearing in satellite bands on density gradients, two other types of tandemly repeated DNA are also classed as satellite DNA. These are minisatellites and microsatellites. Minisatellites form clusters up to 20 kb in length, with repeat units up to 25 bp in length; microsatellite clusters are shorter, usually less than 150 bp, and the repeat unit is usually 13 bp or less.

Minisatellite DNA is a second type of repetitive DNA that we are already familiar with because of its association with structural features of chromosomes.
Telomeric DNA, which in humans comprises hundreds of copies of the motif 5′-TTAGGG-3′ (see Figure 7.9), is an example of a minisatellite. We know a certain amount about how telomeric DNA is formed, and we know that it has an important function in DNA replication (Section 15.4). In addition to telomeric minisatellites, some eukaryotic genomes contain various other clusters of minisatellite DNA, many, although not all, located near the ends of chromosomes. The functions of these other minisatellite sequences have not been identified.

Microsatellites are also examples of tandemly repeated DNA. The most common type of human microsatellite is dinucleotide repeats, with approximately 1.5 million copies in the genome as a whole, most frequently repeats of the motif AT. There are also over 1 million trinucleotide repeats. As with interspersed repeats, it is not clear whether microsatellites have a function. It is known that they arise through an error in the process responsible for copying of the genome during cell division, called slippage (Section 16.1), and they might simply be unavoidable products of genome replication.

Although their function, if any, is unknown, microsatellites have proved very useful to geneticists. Many microsatellites are variable, meaning that the number of repeat units in the array is not the same in all members of a species. This is because additional slippage sometimes occurs when a microsatellite is copied during DNA replication, leading to insertion or, less frequently, deletion of one or more of the repeat units. No two humans alive today, except monozygotic twins, triplets, etc., have exactly the same combination of microsatellite length variants: if enough microsatellites are examined, then a unique genetic profile can be established for each person. Genetic profiling is well-known as a tool in forensic science (Figure 7.25), but identification of criminals is a fairly trivial application of microsatellite variability. More sophisticated methodology makes use of the fact that a person’s genetic profile is inherited partly from the mother and partly from the father. This means that microsatellites can be used to establish kinship relationships and population affinities, not only for humans but also for other animals and for plants.

**Interspersed repeats**

Tandemly repeated DNA sequences are thought to have arisen by expansion of a progenitor sequence, either by replication slippage, as described for microsatellites, or by DNA recombination processes. Both of these events result in a series of linked repeats rather than individual repeat units scattered around the genome. Interspersed repeats must therefore have arisen by a different mechanism, one that can result in a copy of a repeat unit appearing in the genome at a position distant from the location of the original sequence. The most frequent way in which this occurs is by transposition, and most interspersed repeats have inherent transpositional activity. Transposition is also a feature of some viral genomes, which are able to insert into the genome of the infected cell and then move from place to place within that genome. Some interspersed repeats are clearly descended from transposable viruses, and because of this relationship we will postpone discussion of these and the other types of interspersed repeats until Chapter 9, after we have looked in detail at the features of viral genomes.
SUMMARY

- The eukaryotic nuclear genome is split into a set of linear DNA molecules, each of which is contained in a chromosome.
- Within a chromosome, the DNA is packaged by association with histone proteins to form nucleosomes, which interact with one another to give the 30 nm fiber and higher orders of chromatin structure.
- The most compact organization results in the metaphase chromosomes that can be observed by light microscopy of dividing cells and which take up characteristic banding patterns after staining.
- The centromeres, which are visible in metaphase chromosomes, contain special proteins that make up the kinetochore, the attachment point for the microtubules that draw the divided chromosomes into the daughter nuclei.
- Telomeres, the structures that maintain the chromosome ends, contain repetitive DNA and special binding proteins.
- Genes are not evenly spread along vertebrate chromosomes; some chromosomes have gene deserts where the gene density is very low.
- The coding parts of genes make up only a small part of the human genome, less than 1.5%, with 44% of the genome made up of various types of repetitive DNA sequence. In contrast, the S. cerevisiae genome is much more compact, with only 3.4% of the DNA taken up by repeat sequences. In general, larger genomes are less compact, explaining why organisms with similar numbers of genes can have genomes of very different sizes.
- Humans have 20,441 protein-coding genes, about the same number as the nematode worm Caenorhabditis elegans. However, the human genome specifies more proteins than that of C. elegans because of alternative splicing.
- Comparisons of gene catalogs listing the functions of the genes in a genome suggest that all eukaryotes possess the same basic set of genes, but that more complex species have a greater number of genes in different functional categories.
- Many genes are organized into multigene families whose members have similar or identical sequences, and in some families, such as the vertebrate globin genes, the members are expressed at different developmental stages.
- Eukaryotic nuclear genomes also contain evolutionary relics such as pseudogenes and gene fragments.
- The repetitive DNA content of a eukaryotic nuclear genome comprises interspersed repeats, much of which has transpositional activity, and tandemly repeated DNA, which includes the satellite DNA found at centromeres, mini-satellites such as telomeric DNA, and microsatellites.

SHORT ANSWER QUESTIONS

1. What does the treatment of eukaryotic chromatin with nucleases reveal about the packaging of eukaryotic DNA?
2. Describe how the nucleosomes are arranged in the 30 nm fiber.
3. List the special features of (A) microchromosomes and (B) B chromosomes.
4. What did researchers find when they sequenced the centromeres of Arabidopsis? Why was this finding surprising?
5. Explain why it is important that chromosomes have telomeres at their ends.

6. What differences in gene distribution and repetitive DNA content are seen when yeast and human chromosomes are compared?

7. The human genome contains many fewer genes than were predicted by many researchers. Why were these initial predictions so high?

8. What aspects of the biology of the different species are revealed when the human, fruit fly, yeast, C. elegans, and A. thaliana gene catalogs are compared?

9. Describe the organization of the human globin gene families and indicate the functions of each of the genes in these families.

10. Distinguish between the two types of nonprocessed pseudogene.

11. Describe the events that give rise to a processed pseudogene.

12. What types of repetitive DNA are present in the human genome?

IN-DEPTH PROBLEMS

1. What impact is DNA packaging likely to have on the expression of individual genes?

2. Defend or attack the isochore model.

3. Discuss possible functions for the intergenic component of the human genome.

4. To what extent is it possible to describe the typical features of a eukaryotic genome?

5. What would be the implications for genome evolution if some pseudogenes retained their functions or acquired new functions?

FURTHER READING

Chromosome structure


Gene distribution


Key papers and databases on eukaryotic genome structure and content


Genetic features


Online resources


KEGG (Kyoto Encyclopedia of Genes and Genomes). http://www.genome.jp/kegg/ A collection of databases including details of the structures and contents of all sequenced genomes.

Prokaryotes are organisms whose cells lack extensive internal compartments. There are two very different groups of prokaryotes, distinguished from one another by characteristic genetic and biochemical features:

- The bacteria, which include most of the commonly encountered prokaryotes including the Gram-negatives (such as *Escherichia coli*), the Gram-positives (such as *Bacillus subtilis*), the cyanobacteria (such as *Anabaena*), and many more
- The archaea, which are less well studied and were once thought to be extremophiles, living only in inhospitable environments such as hot springs and acidic streams. Now we know that they are much more common and are present in many nonextreme environments, including the human gut.

In this chapter we will examine the genomes of prokaryotes, and also of eukaryotic mitochondria and chloroplasts which, as they are descended from bacteria, have genomes that display many prokaryotic features. Because of the relatively small sizes of prokaryotic genomes, over 40,000 complete sequences, representing 6500 species of bacteria and archaea, have been obtained. As a result, we are beginning to understand a great deal about the anatomies of prokaryotic genomes, and in some respects we know more about these organisms than we do about eukaryotes. The picture that is emerging is one of immense variability among the prokaryotes as a whole and in some cases even between closely related species.

### 8.1 PHYSICAL FEATURES OF PROKARYOTIC GENOMES

Prokaryotic genomes are very different from eukaryotic ones, in particular with regard to the physical organization of the genome within the cell. Although the word chromosome is used to describe the DNA-protein structures present in prokaryotic cells, this is a misnomer, as the structure has few similarities with a eukaryotic chromosome.

The traditional view has been that in a typical prokaryote the genome is contained in a single circular DNA molecule, localized within the nucleoid, which is the lightly staining region of the otherwise featureless prokaryotic cell (Figure 8.1). This is certainly true for *E. coli* and many of the other commonly studied bacteria. However, as we will see, our growing knowledge of prokaryotic genomes is leading us to question several of the preconceptions that became established during the pre-genome era of microbiology. These preconceptions relate both to the physical structure of the prokaryotic genome and to its genetic organization.

The traditional view of the prokaryotic chromosome

As with eukaryotic chromosomes, a prokaryotic genome has to squeeze into a relatively tiny space (the circular *E. coli* chromosome has a circumference of 1.6 mm, whereas an *E. coli* cell is just 1.0 × 2.0 μm), and as with eukaryotes, this is achieved with the help of DNA-binding proteins that package the genome in an organized fashion.
Most of what we know about the organization of DNA in the nucleoid comes from studies of *E. coli*. The first feature to be recognized was that the circular *E. coli* genome is **supercoiled**. Supercoiling occurs when additional turns are introduced into the DNA double helix (positive supercoiling) or if turns are removed (negative supercoiling). With a linear molecule, the torsional stress introduced by over- or underwinding is immediately released by rotation of the ends of the DNA molecule, but a circular molecule, having no ends, cannot reduce the strain in this way. Instead the circular molecule responds by winding around itself to form a more compact structure (**Figure 8.2**). Supercoiling is therefore an ideal way to package a circular molecule into a small space. Evidence that supercoiling is involved in packaging the circular *E. coli* genome was first obtained in the 1970s from examination of isolated nucleoids and was subsequently confirmed as a feature of DNA in living cells in 1981. In *E. coli*, the supercoiling is thought to be generated and controlled by the DNA topoisomerase called DNA gyrase, which we will look at in more detail in **Section 15.1** when we examine the roles of these enzymes in DNA replication.

Studies of isolated nucleoids suggest that the *E. coli* DNA molecule does not have unlimited freedom to rotate once a break is introduced. The most likely explanation is that the bacterial DNA is attached to proteins that restrict its ability to relax, so that rotation at a break site results in loss of supercoiling from only a small segment of the molecule (**Figure 8.3**). The strongest evidence for this domain model has come from experiments that exploit the ability of trimethylpsoralen to distinguish between supercoiled and relaxed DNA. When it is photoactivated by a pulse of light of wavelength 360 nm, trimethylpsoralen binds to double-stranded DNA at a rate that is directly proportional to the degree of torsional stress possessed by the molecule. The degree of supercoiling can therefore be assayed by measuring the amount of trimethylpsoralen that binds to a molecule in unit time. After *E. coli*
cells have been irradiated to introduce single-strand breaks into their DNA molecules, the amount of trimethylpsoralen binding is indirectly proportional to the radiation dose (Figure 8.4). This is the response predicted by the domain model, in which the overall supercoiling of the molecule is gradually relaxed as greater doses of radiation cause breaks within an increasing number of domains. In contrast, if the E. coli nucleoid were not organized into domains, then a single break in the DNA molecule would lead to complete loss of supercoiling; irradiation would therefore have an all-or-nothing effect on trimethylpsoralen binding.

The current model has the E. coli DNA attached to a protein core from which supercoiled loops radiate out into the cell. Each loop contains 10–100 kb of supercoiled DNA, the amount of DNA that becomes unwound after a single break. The protein component of the nucleoid is made up of a variety of nucleoid-associated proteins, which are present in most bacteria that have been studied, have some amino acid sequence similarity with the eukaryotic histone H2B. Each HU protein is a dimer of two subunits, either two HUα, two HUβ, or a heterodimer comprising one of each type of monomer. When crystals of HU are prepared for diffraction studies, some of the subunits form octamers, around which DNA could be wound to give a structure resembling a nucleosome. However, this type of association has not been observed in vivo and there is no evidence that HU proteins act in this way in the nucleoid. They may simply induce bends in the DNA in order to facilitate formation of the supercoiled loops. A packaging role has also been proposed for the histone-like nucleoid structuring protein (H-NS): it binds specifically to AT-rich regions, which are thought to be present at the boundaries of the supercoiled loops. H-NS might therefore be a component of the nucleoid core.

The preceding discussion refers specifically to the E. coli chromosome, which we look on as typical of bacterial chromosomes in general. But we must be careful to make a distinction between the bacterial chromosome and that of the second group of prokaryotes, the archaea. Although a single group of archaea, the Crenarchaeota, which includes many marine species, has nucleoid proteins similar to the bacterial ones that we have discussed above, another group, the Euryarchaeota, possesses proteins that are much more similar to histones. In some species, these form a tetramer that associates with approximately 60 bp of DNA to form a structure similar to a eukaryotic nucleosome. In other species, the histone-like proteins form larger multimers of various sizes that are able to bind longer segments of DNA. Currently we have very little information on the archaeal nucleoid, but the assumption is that in the Euryarchaeota these histone-like proteins play a central role in DNA packaging.

Some bacteria have linear or multipartite genomes

The E. coli genome, as described above, is a single circular DNA molecule. This is also the case with the vast majority of bacterial and archaeal chromosomes that have been studied, but an increasing number of linear versions are being found. The first of these, for Borrelia burgdorferi, the organism that causes Lyme disease, was described in 1989, and during the following years similar discoveries were made for Streptomyces coelicolor and Agrobacterium tumefaciens. Linear molecules have free ends, which must be distinguishable from DNA breaks, so these chromosomes require terminal structures equivalent to the telomeres of eukaryotic chromosomes (Section 7.1). In Borrelia and Agrobacterium, the real chromosome ends are distinguishable because a covalent linkage is formed between the 5′- and 3′-ends of the polynucleotides in the DNA double helix, and in Streptomyces, the ends appear to be marked by special binding proteins.

A second and more widespread variation on the E. coli theme is the presence in some prokaryotes of multipartite genomes, which are genomes that are divided into two or more DNA molecules. With these multipartite genomes, a problem often arises in distinguishing a genuine part of the genome from a
8.1 Physical Features of Prokaryotic Genomes

Chapter 8: Genomes of Prokaryotes and Eukaryotic Organelles

A plasmid is a small piece of DNA, often but not always circular, that coexists with the main chromosome in a bacterial cell (Figure 8.5). Some types of plasmid are able to integrate into the main genome, but others are thought to be permanently independent. Their replication process is distinct from that of the main chromosome, and some can reach copy numbers of a thousand or more in a single cell. When the bacterium divides, the plasmids are partitioned between the daughter cells by a process different from the one that results in each daughter receiving its copy of the main chromosome. Plasmids carry genes that are not usually present in the main chromosome, but in many cases these genes are nonessential to the bacterium, coding for characteristics such as antibiotic resistance, which the bacterium does not need if the environmental conditions are amenable (Table 8.1). As well as this apparent dispensability, many plasmids are able to transfer from one cell to another, and the same plasmids are sometimes found in bacteria that belong to different species. These various features of plasmids suggest that they are independent entities and that in most cases the plasmid content of a prokaryotic cell should not be included in the definition of its genome.

With a bacterium such as *E. coli* K12, which has a 4.64 Mb chromosome and can harbor various combinations of plasmids, none of which is more than a few kilobases in size and all of which are dispensable, it is acceptable to define the main chromosome as the genome. With other prokaryotes it is not so easy (Table 8.2). *Vibrio cholerae* O1 El Tor, the pathogenic bacterium that causes cholera, has two circular DNA molecules, one of 2.96 Mb and the other of 1.07 Mb, with 73% of the organism’s 4113 genes on the larger of these. It would appear obvious that these two DNA molecules together constitute the *Vibrio* genome, but closer examination reveals that most of the genes for central cellular activities such as genome expression and energy generation, as well as the genes that confer pathogenicity, are located on the larger molecule. The smaller molecule contains many essential genes but also has certain features that are considered characteristic of plasmids, notably the presence of an integron, a set of genes and other DNA sequences that enable plasmids to capture genes from bacteriophages and other plasmids. It therefore appears possible that the smaller genome is a megaplasmid that was acquired by the ancestor to *Vibrio* at some period in the bacterium’s evolutionary past. *Deinococcus radiodurans* R1, whose genome is of particular interest because it contains many genes that help this bacterium resist the harmful effects of radiation, is constructed on similar lines, with essential genes distributed among two circular chromosomes and two plasmids. However, the *Vibrio* and *Deinococcus* genomes are relatively noncomplex compared with *Borrelia burgdorferi* B31, whose linear chromosome of 911 kb, carrying 875 genes, is accompanied by up to 19 linear and circular plasmids, which together contribute another 504 kb and another 478 genes. Although the functions of most of these genes are unknown, those that have been identified include several that would normally be considered dispensable, such as genes for membrane proteins and purine biosynthesis. The implication is that at least some of the *Borrelia* plasmids are essential components of the genome, leading to the possibility that some

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**Table 8.1 Features of Typical Plasmids**

<table>
<thead>
<tr>
<th>Type of plasmid</th>
<th>Gene functions</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resistance</td>
<td>Antibiotic resistance</td>
<td>RbK of <em>Escherichia coli</em> and other bacteria</td>
</tr>
<tr>
<td>Fertility</td>
<td>Conjugation and DNA transfer between bacteria</td>
<td>F of <em>E. coli</em></td>
</tr>
<tr>
<td>Killer</td>
<td>Synthesis of toxins that kill other bacteria</td>
<td>Col of <em>E. coli</em>, for colicin production</td>
</tr>
<tr>
<td>Degradative</td>
<td>Enzymes for metabolism of unusual molecules</td>
<td>TOL of <em>Pseudomonas putida</em>, for toluene metabolism</td>
</tr>
<tr>
<td>Virulence</td>
<td>Pathogenicity</td>
<td>Ti of <em>Agrobacterium tumefaciens</em>, conferring the ability to cause crown gall disease in dicotyledonous plants</td>
</tr>
</tbody>
</table>
prokaryotes have highly multipartite genomes, comprising a number of separate DNA molecules, more akin to what we see in the eukaryotic nucleus rather than the typical prokaryotic arrangement.

The complications posed by bacteria such as *Vibrio* and *Deinococcus* have prompted microbial geneticists to invent a new term, *chromid*, to describe a plasmid that carries essential genes. This means that we now distinguish between three, rather than just two, types of DNA molecule that might be found in a bacterium (Figure 8.6):

- One or more bacterial chromosomes, carrying essential genes and located in the nucleoid
8.2 Genetic Features of Prokaryotic Genomes

- Genuine plasmids, which are distinct from a bacterial chromosome because of their special plasmid partitioning system and whose genes are nonessential to the bacterium.

- Chromids, which use a plasmid partitioning system but carry genes that the bacterium needs to survive.

According to this nomenclature, *V. cholerae* has one chromosome and one chromid, and *D. radiodurans* has two chromosomes and two chromids.

8.2 Genetic Features of Prokaryotic Genomes

Genome annotation by sequence inspection is much easier for prokaryotes compared with eukaryotes (Section 5.1), and for most of the prokaryotic genomes that have been sequenced, we have reasonably accurate estimates of the number of genes and fairly comprehensive lists of gene functions. The results of these studies have been surprising and have forced microbiologists to reconsider the meaning of species when applied to prokaryotes. We will examine these evolutionary issues later in this chapter. First, we must look at the way in which the genes are organized in a prokaryotic genome.

Gene organization in the *E. coli* K12 genome

We are already familiar with the notion that bacterial genomes have compact genetic organizations with very little space between genes, as this was an important part of our discussion of the strengths and weaknesses of open reading frame (ORF) scanning as a means of identifying the genes in a genome sequence (see Figure 5.3). To reemphasize this point, the complete circular gene map of the *E. coli* K12 genome is shown in Figure 8.7. There is intergenic DNA in the *E. coli* genome, but it accounts for only 11% of the total and it is distributed around the genome in small segments that do not show up when the map is drawn at this
scale. In this regard, *E. coli* is typical of all prokaryotes whose genomes have so far been sequenced: prokaryotic genomes have very little wasted space. There are theories that this compact organization is beneficial to prokaryotes, for example, by enabling the genome to be replicated relatively quickly, but these ideas have never been supported by hard experimental evidence.

Let us now look more closely at the *E. coli* K12 genome. A typical 50 kb segment is shown in Figure 8.8. When we compare this segment with a typical part of the human genome (see Figure 7.12), it is immediately obvious that in the *E. coli* segment there are more genes and much less space between them, with 43 genes taking up 85.9% of the segment. Some genes have virtually no space between them: *thrA* and *thrB*, for example, are separated by a single nucleotide, and *thrC* begins at the nucleotide immediately following the last nucleotide of *thrB*. These three genes are an example of an operon, a group of genes involved in a single biochemical pathway (in this case, synthesis of the amino acid threonine) and expressed in conjunction with one another. In general, prokaryotic genes are shorter than their eukaryotic counterparts: the average length of a bacterial gene is about two-thirds that of a eukaryotic gene, even after the introns have been removed from the latter. Bacterial genes appear to be slightly longer than archaeal ones.

Two other features of prokaryotic genomes can be deduced from Figure 8.8. The first is the infrequency of repetitive sequences. Most prokaryotic genomes do not have anything equivalent to the high-copy-number interspersed repeat families found in eukaryotic genomes. They do, however, possess certain sequences that might be repeated elsewhere in the genome, such as the insertion sequences IS1 and IS186 that can be seen in the 50 kb segment shown in Figure 8.8. These are examples of transposable elements, sequences that have the ability to move around the genome and, in the case of insertion elements, to transfer from one organism to another, even sometimes between two different species. The positions of the IS1 and IS186 elements shown in Figure 8.8 refer only to the particular *E. coli* isolate from which this sequence was obtained: if a different isolate is examined, then the insertion sequences could well be in different positions or might be entirely absent from the genome. Several families of transposable elements are known in prokaryotic genomes, and we will examine their structures when we study mobile genetic elements in more detail in Section 9.2. Many prokaryotic genomes also contain at least a few nontransposable repeat sequences. The two most important classes are as follows:

- **Repetitive extragenic palindromic (REP) sequences**, most of which are 20–35 bp in length and occur singly or in arrays. Many REP sequences are transcribed into short RNA molecules that can fold into complex stem-loop structures, which in some species might play a role in gene regulation.

- **Clustered regularly interspaced short palindromic repeats (CRISPRs)**, which we have already met as the source of the programmable nuclease that forms the basis of one of the gene inactivation procedures used to assign functions to eukaryotic genes (Section 6.2). CRISPRs are 20–50 bp sequences found in tandem arrays, with each pair of repeats separated by a spacer of similar length but with a unique sequence. Some spacer sequences resemble segments of bacteriophage genomes, leading to the suggestion that CRISPRs represent a prokaryotic immune system, in which transcripts of the spacers act as guide RNAs that bind to invading phage genomes, enabling a Cas endonuclease (whose gene is usually located adjacent to a CRISPR array) to cut and hence inactivate the phage DNA.

The number of transposable and nontransposable repeat sequences present in different prokaryotic genomes varies enormously. Usually they take up less
than 1% of the genome sequence, but there are exceptions. The genome of the meningitis bacterium Neisseria meningitidis Z2491 has over 3700 copies of 15 different types of repeat sequence, collectively making up almost 11% of the 2.18 Mb genome.

The second feature of prokaryotic genomes that can be deduced from Figure 8.8 is the scarcity of introns. E. coli K12 has no discontinuous genes at all, and introns are uncommon among other bacteria and archaea. Those that have been discovered belong to the group I and II types, which are quite different from the introns present in eukaryotic pre-mRNA. Unlike pre-mRNA introns, the group I and II types can fold into complex base-paired structures that have the ability to self-splice, meaning that they can remove themselves from RNA transcripts without the aid of catalytic proteins. At least some are also able to move from one position to another in a genome. Because they are autocatalytic, the insertion of one of these introns into a gene does not affect the ability of that gene to be expressed. Once the gene is transcribed, the intron self-splices, leaving a functional copy of the mRNA. Prokaryotic introns might therefore be looked on as a special type of transposable element, one that targets gene sequences rather than intergenic regions as insertion sites.

Operons are characteristic features of prokaryotic genomes

One characteristic feature of prokaryotic genomes illustrated by E. coli K12 is the presence of operons. An operon is a group of genes that are located adjacent to one another in the genome, with perhaps just one or two nucleotides between the end of one gene and the start of the next. All the genes in an operon are expressed as a single unit. A typical E. coli example is the lactose operon, the first operon to be discovered, which contains three genes involved in conversion of the disaccharide lactose into its monosaccharide units, glucose and galactose (Figure 8.9A). Monosaccharides are substrates for the energy-generating glycolytic pathway, so the function of the genes in the lactose operon is to convert lactose into a form that

![Figure 8.9 Two operons of Escherichia coli.](image)

- **(A)** The lactose operon. The three genes are called lacZ, lacY, and lacA; the first two are separated by 52 bp and the second two by 64 bp. All three genes are expressed together. lacZ codes for the lactose permease that transports lactose into the cell, and lacZ and lacA code for enzymes that split lactose into its component sugars, galactose and glucose.
- **(B)** The tryptophan operon, which contains five genes coding for enzymes involved in the multistep biochemical pathway that converts chorismic acid into the amino acid tryptophan. The genes in the tryptophan operon are closer together than those in the lactose operon: trpE and trpD overlap by 1 bp, as do trpB and trpA; trpD and trpC are separated by 4 bp, and trpC and trpB are separated by 12 bp.
can be utilized by *E. coli* as an energy source. Lactose is not a common component of *E. coli*’s natural environment, so most of the time the operon is not expressed and the enzymes for lactose utilization are not made by the bacterium. When lactose becomes available, the operon is switched on; all three genes are expressed together, resulting in coordinated synthesis of the lactose-utilizing enzymes. This is the classic example of gene regulation in bacteria.

The lactose operon is an example of an **inducible operon**, one that is switched on by a substrate for the enzymes coded by the genes in the operon. Other operons are **repressible**, controlled by a product of the pathway catalyzed by the gene products. An example is the tryptophan operon, which contains five genes that specify the set of enzymes needed to synthesize this amino acid from a precursor called chorismic acid (Figure 8.9B). The regulatory molecule for this operon is tryptophan. When tryptophan levels are low, the operon is expressed so more enzymes are made and more tryptophan can be synthesized. When the levels of tryptophan have been replenished, the operon is switched off.

Altogether there are 850 operons in the *E. coli* K12 genome: 450 of these contain two genes each, and the longest contains 18 genes. Operons are a common feature of many prokaryotic genomes, with over 2000 in some species, but they are not universally frequent. *Lactobacillus helveticus* H10 has 2052 genes but only 35 operons, the longest of which contains six genes. *Pseudomonas syringae* DC3000 has 5619 genes and 25 operons, one of which contains 18 genes, and the marine bacterium *Rhodopirellula baltica* has no operons at all, with each of its 7325 genes forming an individual transcription unit. Operons were once thought to be exclusively a feature of prokaryotic genomes, but we now know that they are not entirely absent in eukaryotes. Closely spaced clusters of genes that are transcribed as a single unit are relatively common in the *Caenorhabditis elegans* genome, and some examples are also known in *Drosophila melanogaster*.

**Prokaryotic genome sizes and numbers of genes vary according to biological complexity**

There is some overlap in size between the largest prokaryotic and smallest eukaryotic genomes, but on the whole, prokaryotic genomes are much smaller (Table 8.3). For example, the *E. coli* K12 genome is just 4.64 Mb, two-fifths the size of the yeast genome, and has only 4315 genes. Most prokaryotic genomes

| TABLE 8.3 GENOME SIZES AND NUMBERS OF GENES FOR VARIOUS PROKARYOTES |
|-----------------------------|---------------------|---------------------|
| Species                     | Size of genome (Mb) | Number of genes     |
| **Bacteria**                |                     |                     |
| *Nasuia deltocephalinicola* NAS-ALF | 0.11                | 169                 |
| *Mycoplasma genitalium* G37  | 0.58                | 559                 |
| *Streptococcus pneumoniae* R6| 2.00                | 2228                |
| *Vibrio cholerae* O1 El Tor  | 4.03                | 4113                |
| *Mycobacterium tuberculosis* H37Rv | 4.41              | 4096                |
| *Escherichia coli* K12     | 4.64                | 4315                |
| *Pseudomonas aeruginosa* PA01 | 6.26              | 5807                |
| *Sorangium cellulosum* So0157-2 | 14.78          | 10,473              |
| **Archaea**                 |                     |                     |
| *Methanocaldocolus jannaschii* DSM2661 | 1.74            | 1875                |
| *Archaeoglobus fulgidus* DSM4304 | 2.18             | 2515                |

Data from Ensembl Bacteria release 32.
are less than 5 Mb in size, but the overall range among sequenced genomes is from just 112 kb for *Nasuia deltocephalinicola* NAS-ALF to 14.8 Mb for *Sorangium cellulosum* So0157-2.

The compact organization of the *E. coli* K12 genome, with the genes making up 89% of the genome sequence, is typical of other prokaryotic genomes: the average gene density is 87%, with most genomes in the range 85–90%. This means that genome size is proportional to the number of genes. The numbers of genes therefore vary over an extensive range, with these numbers reflecting the nature of the ecological niches within which different species of prokaryotes live. The largest genomes tend to belong to free-living species that are found in the soil, the environment that is generally looked on as providing the broadest range of physical and biological conditions, to which the genomes of these species must be able to respond. *S. cellulosum* provides a good example. Its 10,400 protein-coding genes include some specifying enzymes that enable this bacterium to break down cellulose into sugars and others coding for enzymes that synthesize antibacterial and antifungal compounds that help it to compete in the complex soil ecosystem. There are also genes for proteins involved in cell-to-cell communication, which enable the bacteria to migrate together in swarms and to associate into a multicellular fruiting body that produces resistant spores. At the other end of the scale, many of the smallest genomes belong to species that are obligate parasites. *N. deltocephalinicola*, for example, is an endosymbiont of leafhoppers, living inside specialized structures within the insect’s abdomen. The bacteria provide the leafhopper with two amino acids that *Nasuia* can synthesize but insects must obtain from their diet. In return, the bacteria receive various nutrients from the insect, which means that *Nasuia* is able to dispense with many of the enzymes needed by free-living bacteria for the synthesis of metabolites and for energy generation. As a consequence, the *Nasuia* genome can be reduced down to just 137 protein-coding genes, the majority of which are involved in essential functions such as DNA replication, transcription, and translation.

Comparisons between the genomes of different prokaryotes have led to speculation about the smallest number of genes needed to specify a free-living cell. One of the first genomes to be sequenced was that of *Mycoplasma genitalium* G37, a genuinely free-living organism with just 476 protein-coding genes. Experiments in which increasing numbers of *Mycoplasma* genes were inactivated by mutation suggested that 382 of these genes were essential. However, this is the minimal gene set needed for an *M. genitalium* bacterium, and similar inactivation studies with other genomes have shown that the number of essential genes that are identified in this way is species-specific. With some species the minimal gene set is greater than 382, and with a few the number is smaller. In one project a set of just 230 genes was shown to be sufficient for growth of *Salmonella typhimurium* LT2, provided that the mutated bacteria were grown in a rich culture medium from which they could obtain nutrients such as amino acids. By providing many compounds that in its natural environment a bacterium would synthesize for itself, the required gene set can be reduced to a catalog similar to that of *Nasuia* and other symbiotic species.

**Genome sizes and numbers of genes vary within individual species**

Genome projects have confused our understanding of what constitutes a species in the prokaryotic world. This has always been a problem in microbiology because the standard biological definitions of species have been difficult to apply to microorganisms. The early taxonomists such as Linnaeus described species in morphological terms, all members of one species having the same or very similar structural features. This form of classification was in vogue until the early twentieth century and was first applied to microorganisms in the 1880s by Robert Koch and others, who used staining and biochemical tests to distinguish between bacterial species. However, it was recognized that this type of classification was imprecise because many of the resulting species were made up of a variety of types with
The pan-genome concept has been widely applied to various bacteria. For example, the pan-genome of Helicobacter pylori, which is a bacterium that infects the human stomach, includes both pathogenic and non-pathogenic strains. The pan-genome of H. pylori is estimated to contain around 2700 genes, with 1800 genes being present in all strains (the core genome) and 900 genes being present in some but not all strains (the accessory genome). This indicates a high level of genetic variation among different strains of H. pylori, which could contribute to the disease-causing potential of the bacterium.

In contrast, the core genome of Escherichia coli (E. coli) K12, a common laboratory strain, contains only about 500 genes, while the accessory genome contains around 1400 genes. This suggests that E. coli K12 is a genetically simpler organism compared to H. pylori. However, the pan-genome concept also highlights the potential for horizontal gene transfer, where genes from different bacteria can be exchanged, allowing for the rapid evolution of new pathogens.

The pan-genome concept is particularly useful in the study of bacterial species that show significant genetic diversity, such as Salmonella enterica and related species. The pan-genome of Salmonella, which includes both human and animal strains, is estimated to contain around 4000 genes, with about 2500 genes being present in all strains (the core genome) and 1500 genes being present in some but not all strains (the accessory genome). This suggests that Salmonella has a high level of genetic diversity, which could contribute to its ability to cause a wide range of diseases.

The pan-genome concept has also been applied to the study of viruses, such as the human papillomavirus (HPV), which is associated with a range of diseases, including cervical cancer. The pan-genome of HPV includes a large number of genes, with about 600 genes being present in all strains (the core genome) and around 3000 genes being present in some but not all strains (the accessory genome). This suggests that HPV has a high level of genetic diversity, which could contribute to its ability to cause a range of diseases.

The pan-genome concept is also useful in the study of other bacteria, such as Mycobacterium tuberculosis, which is the causative agent of tuberculosis. The pan-genome of M. tuberculosis includes around 3000 genes, with about 1500 genes being present in all strains (the core genome) and around 1500 genes being present in some but not all strains (the accessory genome). This suggests that M. tuberculosis has a high level of genetic diversity, which could contribute to its ability to cause a range of diseases.

The pan-genome concept is particularly useful in the study of bacterial species that show significant genetic diversity, such as Salmonella enterica and related species. The pan-genome of Salmonella, which includes both human and animal strains, is estimated to contain around 4000 genes, with about 2500 genes being present in all strains (the core genome) and 1500 genes being present in some but not all strains (the accessory genome). This suggests that Salmonella has a high level of genetic diversity, which could contribute to its ability to cause a wide range of diseases.

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the remaining 900 were accessory genes. Of the latter, 260 were singleton genes found in just one strain, and the remainder were present in two or more strains but not, by definition, in each of the eight strains.

It was clear from analysis of the *S. agalactiae* pan-genome that the numbers of genes in the core and accessory genomes would change as additional strains are added to the data set. The size of the core genome would be expected to go down as strains were discovered that lacked one or more genes previously assigned to the core set. Conversely, the number of accessory genes would increase as strain-specific genes from each new genome were added to this set. This prediction has turned out to be correct for some, but not all, species. The *E. coli* pan-genome, for example, continues to grow, even though several hundred strains have now been sequenced, and is predicted to contain more than 60,000 genes (Figure 8.11).

Note, however, that the size of the core *E. coli* genome has stabilized and has not changed substantially since a figure of 3188 genes was reached before the 100th genome was sequenced. Because the number of genes is still increasing, *E. coli* is looked on as having an open pan-genome. In contrast, some species have closed pan-genomes, ones that are no longer increasing in size as new strains are sequenced. An example is *Bacillus anthracis*, whose pan-genome comprises 2985 genes, of which 2893 form the core. It has been suggested that the relatively small number of accessory genes possessed by a species with a closed pan-genome reflects a more limited ecological range compared to a species with an open pan-genome, whose vast array of accessory genes presumably enables the species to colonize a larger variety of ecological niches.

**Figure 8.11** The pan-genome of *Escherichia coli*. The graph shows the numbers of genes in the pan-genome and core genome of *E. coli* plotted against the gradually increasing number of strains whose genomes have been sequenced. The pan-genome is defined as open because its size has not yet stabilized. In contrast, the core genome, defined in this example as comprising genes present in >95% of all strains, has stabilized at 3188 genes. The number of genes in the pan-genome is probably an overestimate, because these data include draft genome assemblies likely to contain ORFs that will subsequently be identified as spurious. It is believed that the complete pan-genome of *E. coli* contains about 60,000 genes. (From Land M, Hauser L, Jun SR et al [2015] *Func Integr Genomics* 15:141–161. Courtesy of Springer Science + Business Media under CC BY.)

**Distinctions between prokaryotic species are further blurred by lateral gene transfer**

It has been known since the 1940s that plasmids and occasionally chromosomal genes can move between bacteria by conjugation, transduction via a bacteriophage, or simple uptake of DNA fragments from the environment. These lateral gene transfer processes have been extensively studied, at least with model species such as *E. coli*, because they form the basis of techniques for gene mapping in bacteria (Section 3.4). This early work revealed that, under some circumstances, genes could be transferred between bacteria of different species, for example, between *E. coli* and *S. typhimurium*. The notion that the same genes would occasionally be found in different prokaryotic species was therefore established by the beginning of the genomics era. The extent of lateral gene transfer, as revealed by comparisons between the first few prokaryotic genomes to be sequenced, was nonetheless a major surprise.

Lateral gene transfer is revealed by an inconsistency between the apparent evolutionary relationships of a pair of species, as inferred by studies of individual genes compared with studies of the genomes as a whole. This is because the transfer of a gene from one species to another will have occurred much more recently
in evolutionary terms than the original divergence of the two species from a common ancestor. The copies of the transferred gene in the two species will therefore have relatively similar sequences, as there has been insufficient time for the sequences to diverge greatly by mutation. Comparisons based on the entire genome sequences will therefore place the two species at different positions on the prokaryotic evolutionary tree, whereas comparisons between the gene sequences will place them close together (Figure 8.12). By this type of analysis, many examples of lateral gene transfer have been identified, including some between bacteria and archaea. One outcome of these discoveries has been a greater focus on the processes by which bacteria take up DNA from the environment and the realization not only that DNA uptake is more widespread than originally thought, but also that many species have proteins in their cell membranes whose specific role is to capture DNA fragments from the environment and transport those fragments into the cell.

The impact of lateral gene transfer on the evolution of prokaryotes is still not fully explored. The spread of antibiotic resistance genes through bacterial metapopulations, in hospitals and more broadly in the environment, is clearly only one small facet of lateral gene transfer, albeit the one that has greatest relevance in human society. In some cases, lateral gene transfer appears to have been responsible for a major change in the characteristics of a genus. Almost 25% of the 1952 genes of the thermophilic bacterium *Thermotoga maritima* appear to have been obtained from archaea, these genes possibly forming the basis of this bacterium’s ability to tolerate high temperatures. Transfer of approximately 1000 genes from bacteria to an anaerobic ancestor of the haloarchaea is thought to have enabled these organisms to evolve a tolerance to oxygen and adopt an aerobic lifestyle, albeit as extremophiles in brine pools and other high-salt environments.

Several examples are also known where the smaller-scale transfer of genes between different species appears to have provided the recipient with a novel metabolic capability. The methylaspartate cycle of haloarchaea appears to have evolved in this way. Like many organisms, haloarchaea use acetate as a source of carbon for biosynthesis of products such as amino acids and nucleotides. Unlike most other organisms, some species of haloarchaea use a novel metabolic pathway called the methylaspartate cycle, which is better suited to a high-salt environment, for the first stage of this process, when acetyl-CoA is converted into malate. The methylaspartate cycle combines components of two other pathways: the glyoxylate cycle, which operates in a variety of bacteria and archaea, and the
ethylmalonyl-CoA pathway, which has a more restricted distribution in genera such as *Rhodobacter* and *Methylobacterium*. The two pathways do not operate together in the same species, so evolution of the novel methylaspartate cycle in haloarchoaæa required acquisition of genes from at least two sources. Other examples of metabolic innovation via lateral gene transfer include a variation of the acetyl-CoA synthesis pathway in *Methanosarcina*, by transfer of two genes from a cellulose-degrading *Clostridium* species, and the ability of *Thermosipho* to synthesize vitamin B₁₂ from glutamate, resulting from transfer of 31 genes from members of the phylum Firmicutes.

**Metagenomes describe the members of a community**

In the conventional approach to prokaryotic genomics, a sequencing project is directed at the genome of a single species. A pure culture of that species is prepared, the DNA is extracted, and the resulting next-generation sequencing reads are assembled to give the genome sequence. This approach has one major limitation. For many years, microbiologists have been aware that the artificial culture conditions used to isolate bacteria and archaea from their natural habitats do not suit all species, and many will not grow under these conditions and hence will remain undetected. If a species cannot be grown in culture, then its genome cannot be sequenced, at least not by the conventional approach.

**Metagenomics** addresses this problem by obtaining DNA sequences from all the genomes in a particular habitat, for example, from seawater or from soil. DNA is prepared directly from the environmental sample without any attempt to isolate individual species from that sample. The resulting sequence reads therefore derive from many different genomes, including those of species that cannot be cultured. Assembling the vast mixture of reads into individual genome sequences is a challenge, but is possible if a sufficiently large number of reads is obtained, unless the sample is extremely complex in terms of the numbers of species that are present. The species present in a particular habitat can be identified and the relative abundance of each species can be assessed from the relative numbers of reads that are obtained for each genome. The resulting genome assemblies include species that no microbiologist has ever seen and which are known to science only from their genome sequences. The metabolic capabilities of an unknown species can, however, be inferred from its genome sequence, and an assessment can be made of the contribution that the species makes to its ecosystem, for example, in cycling of nutrients.

In one of the first metagenomic studies, over 1 Mb of sequence was obtained from bacterial DNA from 1500 liters of surface water from the Sargasso Sea. The sequence included segments of the genomes of over 1800 species, of which 148 were totally new. Similar studies have been carried out with samples from sites that have become contaminated with petroleum or acid mine drainage, to assess how the microbial community responds to and helps remediate a polluted environment, and from agricultural soils, with the aim of understanding how microbial activity influences the growth and productivity of crops. But the greatest efforts in metagenomics are being applied to studies of the human *microbiome*. These are the microorganisms that live on or within the human body. Initial estimates are that the entire microbiome of a healthy adult includes 10,000 different species, with perhaps 1000 of these present in the gut. Most of the species are harmless, and pathogens make a significant contribution to the microbiome only when an individual has a specific infection. For many years, the microbiome has been looked on as unimportant, but increasing evidence suggests that at least some of the species carry out useful activities. In the digestive tract, it appears that bacteria break down some types of carbohydrate into metabolites that can be further digested by intestinal cells. Without the bacterial activity, the human host could not use these carbohydrates as nutrients. The aims of the various metagenomic studies of the human microbiome are to catalog the genera present in different parts of the human ecosystem (such as the gut, the respiratory tract, the genitourinary tract, and the skin) to establish how,
if at all, these catalogs vary in different people and in different parts of the world, and to understand how the microbiome influences human health and changes in response to disease.

### 8.3 EUKARYOTIC ORGANELLAR GENOMES

Now we return to the eukaryotic world to examine the genomes present in mitochondria and chloroplasts. The possibility that some genes might be located outside the nucleus—**extrachromosomal genes**, as they were initially called—was first raised in the 1950s as a means of explaining the unusual inheritance patterns of certain genes in the fungus *Neurospora crassa*, the yeast *Saccharomyces cerevisiae*, and the photosynthetic alga *Chlamydomonas reinhardtii*. Electron microscopic and biochemical studies at about the same time provided hints that DNA molecules might be present in mitochondria and chloroplasts. Eventually, in the early 1960s, these various lines of evidence were brought together and the existence of **mitochondrial** and **chloroplast genomes**, independent of and distinct from the eukaryotic nuclear genome, was accepted.

**The endosymbiont theory explains the origin of organellar genomes**

The discovery of organellar genomes led to many speculations about their origins. Today most biologists accept that the **endosymbiont theory** is correct, at least in outline, even though it was considered quite unorthodox when it was first proposed in the 1960s. The endosymbiont theory is based on the observation that gene expression processes occurring in organelles are similar in many respects to equivalent processes in bacteria. In addition, when nucleotide sequences are compared, organellar genes are found to be more similar to equivalent genes from bacteria than they are to eukaryotic nuclear genes. The endosymbiont theory therefore holds that mitochondria and chloroplasts are the relics of free-living bacteria that formed a symbiotic association with the precursor of the eukaryotic cell, way back at the very earliest stages of evolution (**Figure 8.13**).

Support for the endosymbiont theory has come from the discovery of organisms that appear to exhibit stages of endosymbiosis less advanced than seen with mitochondria and chloroplasts. For example, the algae known as glaucophytes possess photosynthetic structures, called **cyanelles**, that are different from chloroplasts and instead resemble ingested cyanobacteria (**Figure 8.14**). Each cyanelle has an external layer of peptidoglycan, thought to be a remnant of the cyanobacterial cell wall, and their light-harvesting proteins resemble those used in free-living cyanobacteria rather than the equivalent structures present in chloroplasts. Modern-day precursors of mitochondrial endosymbiosis have been more difficult to find but one possibility is *Pelomyxa*, a type of amoeba that lacks mitochondria but instead contains symbiotic bacteria, though it is by no means certain that these bacteria provide the amoeba with energy.

If mitochondria and chloroplasts were once free-living bacteria, then since the endosymbiosis was set up, there must have been a transfer of genes from the organelle into the nucleus. We do not understand how this occurred, or indeed whether there was a mass transfer of many genes at once or a gradual trickle from one site to the other. But we do know that DNA transfer from organelle to nucleus, and between organelles, still occurs. This was discovered in the early 1980s, when the first partial sequences of chloroplast genomes were obtained. It was found that in some plants the chloroplast genome contains segments of DNA, often including entire genes, that are copies of parts of the mitochondrial genome. The implication is that this so-called **promiscuous DNA** has been transferred from one organelle to the other. We now know that this is not the only type of transfer that can occur. The *Arabidopsis* mitochondrial genome contains various segments of nuclear DNA as well as 16 fragments of the chloroplast genome, including six tRNA genes that have retained their activity after transfer to the mitochondrion. The nuclear genome of this plant includes several short segments of the chloroplast and mitochondrial genomes as well as a 270 kb
piece of mitochondrial DNA located within the centromeric region of chromosome 2. The transfer of mitochondrial DNA to vertebrate nuclear genomes has also been documented.

As well as the indications that promiscuous DNA provides of the possible transfer of DNA between genomes, there is also one striking example of an endosymbiosis in which the relationship between host and organelle is less developed than is the case with mitochondria and chloroplasts. *Paulinella* is an amoeba with photosynthetic organelles called chromatophores. As with cyanelles, photosynthesis in chromatophores resembles the cyanobacterial processes more closely than the equivalent events in chloroplasts. However, chromatophores, unlike cyanelles, retain a miniature version of a cyanobacterium genome, 1.02 Mb in size and specifying 867 protein-coding genes. This is substantially larger than the typical chloroplast or cyanelle genome, which is less than 0.2 Mb and codes for only 200 genes. The process of genome reduction therefore appears to have reached only an intermediate stage in *Paulinella*. Examination of the chromatophore gene catalog shows that the endosymbiont has lost the ability to make amino acids and some other metabolites, as the genes for these entire pathways are absent from the chromatophore genome. In contrast, the chromatophore retains genes for synthesis of all the proteins and enzymes needed to carry out photosynthesis, as well as genes for DNA replication, transcription, and translation. The chromatophore therefore has the typical features of a symbiont: it is dependent on its host for provision of metabolites that it can no longer make, but it remains autonomous with regard to energy generation and to replication and expression of its genome. In contrast, the genome of an organelle has become so reduced in size and gene content that the organelle is unable to generate energy or replicate and express its genome without the aid of proteins and enzymes coded by nuclear genes.

**Most organellar genomes are circular**

Almost all eukaryotes have mitochondrial genomes, and most photosynthetic eukaryotes have chloroplast genomes. Initially, it was thought that virtually all organellar genomes were circular DNA molecules. Electron microscopy had revealed both circular and linear DNA in some organelles, but it was assumed that the linear molecules were simply fragments of circular genomes that had been broken during preparation for electron microscopy. We still believe that most mitochondrial and chloroplast genomes are circular, but we now recognize that there is a great deal of variability in different organisms. In many eukaryotes, the circular genomes coexist in the organelles with linear versions and, in the case of chloroplasts, with smaller circles that contain subcomponents of the genome as a whole. The latter pattern reaches its extreme in the marine algae called dinoflagellates, whose chloroplast genomes are split into many small circles, each containing just a single gene. We also now realize that the mitochondrial genomes of some microbial eukaryotes (such as *Paramaecium*, *Chlamydomonas*, and several yeasts) are always linear.

Copy numbers for organellar genomes are not particularly well understood. Each human mitochondrion contains about 10 identical molecules, which means that there are about 8000 per cell, but in *S. cerevisiae* the total number is probably smaller, perhaps less than 100 per cell. Photosynthetic microorganisms such as *Chlamydomonas* have 80–90 chloroplast genomes per organelle and approximately 1000 genomes per cell, about one-tenth the number of genomes present in a leaf cell from a higher plant. One mystery, which dates back to the 1950s and has never been satisfactorily solved, is that when organellar genes are studied in genetic crosses, the results suggest that there is just one copy of a mitochondrial or chloroplast genome per cell. This is clearly not the case, and indicates that our understanding of the transmission of organellar genomes from parent to offspring is less than perfect.

Mitochondrial genome sizes are variable ([Table 8.4](#)) and are unrelated to the complexity of the organism. Most multicellular animals have small mitochondrial genomes with a compact genetic organization, where the genes are close together.
with little space between them. The human mitochondrial genome (Figure 8.15), at 16,569 bp, is typical of this type. Most lower eukaryotes such as *S. cerevisiae* (Figure 8.16), as well as flowering plants, have larger and less compact mitochondrial genomes, with a number of the genes containing introns. Chloroplast genomes have less variable sizes (Table 8.4), and most have a structure similar to that shown in Figure 8.17 for the rice chloroplast genome.

### The gene catalogs of organellar genomes

Organellar genomes are much smaller than their nuclear counterparts and we therefore anticipate that their gene contents are much more limited, which is indeed the case. Again, mitochondrial genomes display the greater variability, with gene contents ranging from three for the malaria parasite *Plasmodium falciparum* to 93 for the protozoan *Reclinomonas americana* (Table 8.5). All but the smallest mitochondrial genomes contain genes for noncoding rRNAs, and

**Table 8.4 Sizes of Mitochondrial and Chloroplast Genomes**

<table>
<thead>
<tr>
<th>Species</th>
<th>Type of organism</th>
<th>Genome size (kb)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Plasmodium falciparum</em></td>
<td>Protozoan (malaria parasite)</td>
<td>6</td>
</tr>
<tr>
<td><em>Chlamydomonas reinhardtii</em></td>
<td>Green alga</td>
<td>16</td>
</tr>
<tr>
<td><em>Mus musculus</em></td>
<td>Vertebrate (mouse)</td>
<td>16</td>
</tr>
<tr>
<td><em>Homo sapiens</em></td>
<td>Vertebrate (human)</td>
<td>17</td>
</tr>
<tr>
<td><em>Metridium senile</em></td>
<td>Invertebrate (sea anemone)</td>
<td>17</td>
</tr>
<tr>
<td><em>Chondrus crispus</em></td>
<td>Red alga</td>
<td>26</td>
</tr>
<tr>
<td><em>Aspergillus nidulans</em></td>
<td>Ascomycete fungus</td>
<td>33</td>
</tr>
<tr>
<td><em>Reclinomonas americana</em></td>
<td>Protozoa</td>
<td>69</td>
</tr>
<tr>
<td><em>Saccharomyces cerevisiae</em></td>
<td>Yeast</td>
<td>79</td>
</tr>
<tr>
<td><em>Brassica oleracea</em></td>
<td>Flowering plant (cabbage)</td>
<td>360</td>
</tr>
<tr>
<td><em>Arabidopsis thaliana</em></td>
<td>Flowering plant (vetch)</td>
<td>367</td>
</tr>
<tr>
<td><em>Zea mays</em></td>
<td>Flowering plant (maize)</td>
<td>681</td>
</tr>
<tr>
<td><em>Cucumis sativus</em></td>
<td>Flowering plant (cucumber)</td>
<td>1556</td>
</tr>
</tbody>
</table>

**Chloroplast genomes**

<table>
<thead>
<tr>
<th>Species</th>
<th>Type of organism</th>
<th>Genome size (kb)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Bigelowiella natans</em></td>
<td>Chlorarachniophyte alga</td>
<td>69</td>
</tr>
<tr>
<td><em>Marchantia polymorpha</em></td>
<td>Liverwort</td>
<td>121</td>
</tr>
<tr>
<td><em>Pisum sativum</em></td>
<td>Flowering plant (pea)</td>
<td>122</td>
</tr>
<tr>
<td><em>Oryza sativa</em></td>
<td>Flowering plant (rice)</td>
<td>135</td>
</tr>
<tr>
<td><em>Nicotiana tabacum</em></td>
<td>Flowering plant (tobacco)</td>
<td>156</td>
</tr>
<tr>
<td><em>Chlamydomonas reinhardtii</em></td>
<td>Green alga</td>
<td>204</td>
</tr>
<tr>
<td><em>Floydiella terrestris</em></td>
<td>Green alga</td>
<td>521</td>
</tr>
</tbody>
</table>

Data from NCBI Genome Database.

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**Figure 8.15 The human mitochondrial genome**. The human mitochondrial genome is small and compact, with little wasted space, so much so that genes ATP6 and ATP8 overlap. Abbreviations: ATP6 and ATP8, genes for ATPase subunits 6 and 8; COI, COII, and COIII, genes for cytochrome c oxidase subunits I, II, and III; Cytb, gene for apocytochrome b; ND1–ND6, genes for nicotinamide adenine dinucleotide (NADH) hydrogenase subunits 1–6.
all specify at least some of the protein components of the respiratory chain, the latter being the main biochemical feature of the mitochondrion. The more gene-rich genomes also code for tRNAs, ribosomal proteins, and proteins involved in transcription, translation, and transport of other proteins into the mitochondrion from the surrounding cytoplasm (Table 8.5). Most chloroplast genomes appear to possess the same set of 200 or so genes, again coding for rRNAs and tRNAs, as well as ribosomal proteins and proteins involved in photosynthesis (see Figure 8.17).

A general feature of organellar genomes emerges from Table 8.5. These genomes specify some of the proteins found in the organelle but not all of them. The other proteins are coded by nuclear genes, synthesized in the cytoplasm, and transported into the organelle. If the cell has mechanisms for transporting proteins into mitochondria and chloroplasts, then why not have all the organellar proteins specified by the nuclear genome? We do not yet have a convincing answer to this question, although it has been suggested that at least some of the proteins coded by organellar genomes are extremely hydrophobic and cannot be transported through the membranes that surround mitochondria and chloroplasts, and so cannot be moved into the organelle from the cytoplasm. The only way the cell can get them into the organelle is to make them there in the first place.

**SUMMARY**

- Prokaryotes comprise two distinct types of organism, the bacteria and the archaea.
- The bacterial genome is localized within the nucleoid, the lightly staining region of the otherwise featureless prokaryotic cell. The DNA is attached to a core of binding proteins from which supercoiled loops radiate out into the cell.
Figure 8.17 The rice chloroplast genome. Only those genes with known functions are shown. A number of the genes contain introns, which are not indicated on this map. These discontinuous genes include several for tRNAs, which is why the tRNA genes are of different lengths even though the tRNAs that they specify are all of similar size.

**TABLE 8.5 GENE CONTENTS OF MITOCHONDRIAL GENOMES**

<table>
<thead>
<tr>
<th>Feature</th>
<th>P. falciparum</th>
<th>C. reinhardtii</th>
<th>H. sapiens</th>
<th>S. cerevisiae</th>
<th>A. thaliana</th>
<th>R. americana</th>
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<tr>
<td><strong>Protein-coding genes</strong></td>
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<td></td>
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<td></td>
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<tr>
<td>Respiratory complex</td>
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<td>7</td>
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<td>7</td>
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<td>0</td>
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<td>Transport proteins</td>
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<td>0</td>
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<td>0</td>
<td>0</td>
<td>0</td>
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<td>Translation factor</td>
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<td>0</td>
<td>0</td>
<td>0</td>
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<td>Other</td>
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<td>0</td>
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<td>Total protein-coding genes</td>
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<td>Total noncoding RNA genes</td>
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<td><strong>All genes</strong></td>
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<tr>
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<td>17 kb</td>
<td>79 kb</td>
<td>367 kb</td>
<td>69 kb</td>
</tr>
</tbody>
</table>

Data from NCBI Genome Database. *C. reinhardtii* has a large number of fragmentary ribosomal RNA genes that together specify complete copies of the two standard mitochondrial rRNAs.
• The *E. coli* genome is a single, circular DNA molecule but some prokaryotes have linear genomes and some have multipartite genomes made up of two or more circular and/or linear molecules. In the more complex cases it can be difficult to distinguish which molecules are genuine parts of the genome and which are dispensable plasmids.

• Prokaryotic genomes are very compact, with little repetitive DNA.

• Many genes are organized into operons, the members of which are expressed together and which may have a functional relationship.

• Gene number is related to biological complexity. The largest genomes belong to free-living species found in the soil, and the smallest genomes belong to species that are obligate parasites.

• Genome sizes and gene numbers vary within individual prokaryotic species. The core genome is the set of genes possessed by all members of the species and the accessory genome is the collection of additional genes present in different strains and isolates of that species.

• Many examples of lateral gene transfer between different prokaryotic species have been identified, including some between bacteria and archaea.

• Metagenomics, the study of all the genomes in a habitat such as seawater, is showing that a substantial proportion of the species that are present have never been identified.

• The genomes in the mitochondria and chloroplasts of eukaryotic cells are descended from free-living bacteria that formed a symbiotic association with the precursor of the eukaryotic cell, and hence these genomes have prokaryotic features.

• Most mitochondrial and chloroplast genomes are circular, possibly multipartite, with copy numbers of several thousand per cell.

• Mitochondrial genomes vary in size from 5 to 1500 kb and contain 3–93 genes, including genes for mitochondrial rRNAs, tRNAs, and proteins such as components of the respiratory complex.

• Chloroplast genomes are 60–525 kb with a similar set of some 200 genes, the majority coding for functional RNAs and photosynthetic proteins.

**SHORT ANSWER QUESTIONS**

1. Outline the differences between a eukaryotic chromosome and the *E. coli* chromosome.

2. What experimental evidence suggests that the *E. coli* chromosome is organized into supercoiled domains and is attached to proteins that restrict its ability to relax?

3. What similarities, if any, are there between *E. coli* HU proteins and eukaryotic histone proteins?

4. The *E. coli* genome is a single, circular DNA molecule. What other types of genome structure are found amongst prokaryotes?

5. Describe how the genes and other sequence features are organized in a typical prokaryotic genome. When prokaryotic and mammalian genomes are compared, what differences are seen in the gene density, number of introns, and repetitive DNA content?

6. List the key features of operons and assess the overall importance of the operon as a component of prokaryotic genomic organization.
7. Discuss the factors that influence the number of genes possessed by a prokaryote.

8. Distinguish between the terms core genome and accessory genome.

9. What impact has lateral gene transfer had on the gene content of prokaryotic genomes?

10. Describe the novel information on biology that has been obtained by metagenomic studies of environments such as seawater.

11. Outline the key features of the endosymbiont theory for the origin of mitochondria and chloroplasts.

12. Compare the gene contents of the mitochondria and chloroplasts of different species.

IN-DEPTHPROBLEMS

1. Should the traditional view of the prokaryotic genome as a single, circular DNA molecule be abandoned? If so, what new definition of the prokaryotic genome should be adopted?

2. Speculate on the identities of the approximately 230 genes that constitute the minimum set for a free-living cell.

3. Can the concept of prokaryotic species survive the discoveries that are being made by genome sequencing?

4. Is a definitive test of the endosymbiont theory possible?

5. Why do organelle genomes exist?

FURTHER READING

Prokaryotic nucleoids


Iconic prokaryotic genome sequences


Prokaryotic gene numbers


**Lateral gene transfer**


**Metagenomics**


**Organelle genomes**


**Online resources**


ODB (Operon DataBase). http://operondb.jp/
Viruses are the last and simplest form of life whose genomes we will investigate. In fact, viruses are so simple in biological terms that we have to ask ourselves if they can really be thought of as living organisms. Doubts arise partly because viruses are constructed along lines different from all other forms of life—viruses are not cells—and partly because of the nature of the viral life cycle. Viruses are obligate parasites of the most extreme kind: they reproduce only within a host cell, and in order to replicate and express their genomes, they must subvert at least part of the host’s genetic machinery to their own ends. Some viruses possess genes coding for their own DNA polymerase and RNA polymerase enzymes, but many depend on the host enzymes for genome replication and transcription. All viruses make use of the host’s ribosomes and translation apparatus for synthesis of the polypeptides that make up the protein coats of their progeny. This means that viral genes must be matched to the host genetic system. Viruses are therefore quite specific for particular organisms, and individual types cannot infect a broad spectrum of species.

In this chapter we will also consider the mobile genetic elements that make up a substantial part of the repetitive component of eukaryotic and prokaryotic genomes. We link these elements with viral genomes because it has become clear in recent years that at least some of these repetitive sequences are derived from viruses and are, in effect, viral genomes that have lost the ability to escape from their host cell.

9.1 THE GENOMES OF BACTERIOPHAGES AND EUKARYOTIC VIRUSES

There are a multitude of different types of virus, but the ones that have received most attention from geneticists are those that infect bacteria. These are called bacteriophages, and they have been studied in great detail since the 1930s, when the early molecular biologists, notably Max Delbrück, chose phages as convenient model organisms with which to study genes. We will follow the lead taken by Delbrück and use bacteriophages as the starting point for our investigation of viral genomes.

Bacteriophage genomes have diverse structures and organizations

Bacteriophages are constructed from two basic components: protein and nucleic acid. The protein forms a coat, or capsid, within which the nucleic acid genome is contained. There are three basic capsid structures (Figure 9.1):

- **Icosahedral**, in which the individual polypeptide subunits (protomers) are arranged into a three-dimensional geometric structure that surrounds the nucleic acid. Examples are MS2 phage, which infects *Escherichia coli*, and PM2, which infects *Pseudomonas aeruginosa*.

- **Filamentous**, or helical, in which the protomers are arranged in a helix, producing a rod-shaped structure. The *E. coli* phage called M13 is an example.
9.1 The Genomes of Bacteriophages and Their Viruses

Chapter 9: Viral Genomes and Mobile Genetic Elements

• Head-and-tail, a combination of an icosahedral head, containing the nucleic acid, attached to a filamentous tail and possibly additional structures that facilitate entry of the nucleic acid into the host cell. This is a common structure possessed by, for example, the E. coli phages T4 and λ and by phage SPO1 of Bacillus subtilis.

The term “nucleic acid” has to be used when referring to phage genomes because in some cases these molecules are made of RNA. Viruses are the one form of life that contradicts the conclusion of Avery and his colleagues and of Hershey and Chase that the genetic material is DNA (Section 1.1). Phages and other viruses also break another rule: their genomes, whether of DNA or RNA, can be single-stranded as well as double-stranded. A whole range of different genome structures is known among the phages, as summarized in Table 9.1. With most types of phage there is a single DNA or RNA molecule that comprises the entire genome. However, this is not always the case and a few RNA phages have segmented genomes, meaning that their genes are carried by a number of different RNA molecules. The sizes of phage genomes vary enormously, from about 1.6 kb for the smallest phages to over 150 kb for large ones such as T2, T4, and T6.

Bacteriophage genomes, being relatively small, were among the first to be studied comprehensively by the rapid and efficient DNA sequencing methods that were developed in the late 1970s. The number of genes varies from just four, in the case of MS2, to 278 for the more complex head-and-tail phages (see Table 9.1). The smaller phage genomes of course contain relatively few genes, but these can be organized in a very complex manner. Phage φX174, for example, manages to pack into its genome extra biological information, as several of its genes overlap (Figure 9.2). These overlapping genes share nucleotide sequences (gene B, for example, is contained entirely within gene A) but code for different gene products, as the transcripts are translated from different start

Table 9.1 Features of some Typical Bacteriophages and Their Genomes

<table>
<thead>
<tr>
<th>Phage</th>
<th>Host</th>
<th>Capsid structure</th>
<th>Genome structure</th>
<th>Genome size (kb)</th>
<th>Number of genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>λ</td>
<td>Enterobacteria</td>
<td>Head-and-tail</td>
<td>Linear dsDNA</td>
<td>48.5</td>
<td>73</td>
</tr>
<tr>
<td>M13</td>
<td>Enterobacteria</td>
<td>Filamentous</td>
<td>Circular ssDNA</td>
<td>6.4</td>
<td>10</td>
</tr>
<tr>
<td>MS2</td>
<td>Enterobacteria</td>
<td>Icosahedral</td>
<td>Linear ssRNA</td>
<td>3.6</td>
<td>4</td>
</tr>
<tr>
<td>φ6</td>
<td>Pseudomonas</td>
<td>Icosahedral</td>
<td>Segmented linear dsRNA</td>
<td>2.9, 4.0, 6.4</td>
<td>13</td>
</tr>
<tr>
<td>φX174</td>
<td>Enterobacteria</td>
<td>Icosahedral</td>
<td>Circular ssDNA</td>
<td>5.4</td>
<td>11</td>
</tr>
<tr>
<td>PM2</td>
<td>Pseudoalteromonas</td>
<td>Icosahedral</td>
<td>Linear ssDNA</td>
<td>10.0</td>
<td>22</td>
</tr>
<tr>
<td>SPO1</td>
<td>Bacillus</td>
<td>Head-and-tail</td>
<td>Linear ssDNA</td>
<td>133</td>
<td>204</td>
</tr>
<tr>
<td>T4</td>
<td>Enterobacteria</td>
<td>Head-and-tail</td>
<td>Linear ssDNA</td>
<td>169</td>
<td>278</td>
</tr>
<tr>
<td>T7</td>
<td>Enterobacteria</td>
<td>Head-and-tail</td>
<td>Linear ssDNA</td>
<td>39.9</td>
<td>60</td>
</tr>
</tbody>
</table>

Data from NCBI Genome Database. The genome structure is that in the phage capsid; some genomes exist in different forms within the host cell.

ds, double-stranded; ss, single-stranded.

Figure 9.2 The φX174 genome contains overlapping genes. The genome is made of single-stranded DNA. The expanded region shows the start and end of the overlap between genes E and D. Two other overlapping genes, A* and K, are not shown on this map.
positions and, in most cases, in different reading frames. Overlapping genes are not uncommon in viruses. The larger phage genomes contain more genes, reflecting the more complex capsid structures of these phages and a dependence on a greater number of phage-encoded enzymes during the infection cycle. The T4 genome, for example, includes some 40 genes involved solely in construction of the phage capsid. Despite their complexity, even these large phages still require at least some host-encoded proteins and RNAs in order to carry through their infection cycles.

**Replication strategies for bacteriophage genomes**

Bacteriophages are classified into two groups according to their life cycle: **lytic** and **lysogenic**. The fundamental difference between these groups is that a lytic phage kills its host bacterium very soon after the initial infection, whereas a lysogenic phage can remain quiescent within its host for a substantial period of time, even throughout numerous generations of the host cell. These two life cycles are typified by two *E. coli* phages: the lytic (or virulent) T4 and the lysogenic (or temperate) λ.

The T series of *E. coli* phages (T1–T7) were the first to become available to molecular geneticists and have been the subject of much study. Their lytic infection cycle was first investigated in 1939 by Emory Ellis and Max Delbrück, who added T4 phages to a culture of *E. coli*, waited 3 minutes for the phages to attach to the bacteria, and then measured the number of infected cells over a period of 60 minutes. Their results (Figure 9.3A) showed that there is no change in the number of infected cells during the first 22 minutes of infection, this latent period being the time needed for the phages to reproduce within their hosts. After 22 minutes, the number of infected cells started to increase, showing that lysis of the original hosts had occurred and the new phages that had been produced were now infecting other cells in the culture. The molecular events occurring at the different stages of this one-step growth curve are shown in Figure 9.3B. The initial event is attachment of the phage particle to a receptor protein on the outside of the bacterium. Different types of phage have different receptors: for example, for T4, the receptor is a protein called OmpC (Omp stands for outer membrane protein), which is a type of porin, a protein that forms a channel through the outer cell membrane and facilitates the uptake of nutrients. After attachment, the phage injects its DNA genome into the cell through its tail structure. Immediately after entry of the phage DNA, the synthesis of host DNA, RNA, and protein stops and transcription of the phage genome begins. Within 5 minutes, the bacterial DNA molecule has depolymerized and the resulting nucleotides are being utilized in replication of the T4 genome. After 12 minutes, new phage capsid proteins start to appear and the first complete phage particles are assembled. Finally, at the end of the latent period, the cell bursts and the new phages are released. A typical infection cycle produces 200–300 T4 phages per cell, all of which can go on to infect other bacteria.

Most phages can follow the lytic infection cycle but some, such as λ, can also pursue a lysogenic cycle. In Section 2.3, when we looked at the use of λ phages as cloning vectors, we discovered that during a lysogenic cycle the phage genome becomes integrated into the host DNA. This occurs immediately after entry of the phage DNA into the cell and results in a quiescent form of the bacteriophage, called the prophage (Figure 9.4A). Integration occurs by site-specific recombination (Section 17.2) between identical 15 bp sequences present in the λ and *E. coli* genomes. Note that this means the λ genome always integrates at the same position within the *E. coli* DNA molecule. The integrated prophage can be retained in the host DNA molecule for many cell generations, being replicated along with the host DNA and passed to daughter cells during cell division.

![Figure 9.3](image_url) **Figure 9.3** The lytic infection cycle. (A) The one-step growth curve, as revealed by the experiment conducted by Ellis and Delbrück. (B) Molecular events occurring during the lytic infection cycle.
Chapter 9: Viral Genomes and Mobile Genetic Elements

**Figure 9.4** The lysogenic infection cycle, as followed by bacteriophage λ.

After induction, the infection cycle is similar to the lytic mode.

the bacterial genome and passed with it to the daughter cells. However, a switch to the lytic mode of infection occurs if the prophage is induced by any one of several chemical or physical stimuli. Each of these appears to be linked to DNA damage and possibly therefore signals the imminent death of the host by natural causes. In response to these stimuli, a second recombination event excises the phage genome from the host DNA, phage DNA replication begins, and phage coat proteins are synthesized (Figure 9.4B). Eventually, the cell bursts and new λ phages are released. Lysogeny adds an additional level of complexity to the phage life cycle and ensures that the phage is able to adopt the particular infection strategy best suited to the prevailing conditions.

**Structures and replication strategies for eukaryotic viral genomes**

The capsids of eukaryotic viruses are either icosahedral or filamentous: the head-and-tail structure is unique to bacteriophages. One distinct feature of eukaryotic viruses, especially those with animal hosts, is that the capsid may be surrounded by a lipid membrane, forming an additional component to the viral structure (Figure 9.5). This membrane is derived from the host when the new virus particle leaves the cell and may subsequently be modified by insertion of virus-specific proteins.

Eukaryotic viral genomes display a great variety of structures (Table 9.2). They may be DNA or RNA, single- or double-stranded (or partly double-stranded with single-stranded regions), linear or circular, segmented or nonsegmented. For reasons that no one has ever understood, the vast majority of plant viruses have RNA genomes. Genome sizes cover approximately the same range as seen with phages, although the largest viral genomes (for example, vaccinia virus at 195 kb) are rather bigger than the largest phage genomes.

Although most eukaryotic viruses follow only the lytic infection cycle, few take over the host cell’s genetic machinery to the extent that a bacteriophage does. Many viruses coexist with their host cells for long periods, possibly years, with the host cell functions ceasing only toward the end of the infection cycle, when the viral progeny that have been stored in the cell are released. Other viruses continuously synthesize new virus particles that are extruded from the cell. These long-term infections can occur even if the viral genome does not integrate into the host DNA. Other eukaryotic viruses have life cycles that are more similar to those of lysogenic

<table>
<thead>
<tr>
<th>Virus</th>
<th>Host</th>
<th>Genome structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenovirus</td>
<td>Mammals</td>
<td>Linear dsDNA</td>
</tr>
<tr>
<td>Hepatitis virus</td>
<td>Mammals</td>
<td>Circular, partially ssDNA and partially dsDNA</td>
</tr>
<tr>
<td>Influenza A virus</td>
<td>Mammals</td>
<td>Segmented linear ssRNA</td>
</tr>
<tr>
<td>Parvovirus</td>
<td>Mammals</td>
<td>Linear ssDNA</td>
</tr>
<tr>
<td>Poliovirus</td>
<td>Mammals</td>
<td>Linear ssRNA</td>
</tr>
<tr>
<td>Reovirus</td>
<td>Mammals</td>
<td>Segmented linear dsRNA</td>
</tr>
<tr>
<td>Retroviruses</td>
<td>Mammals, birds</td>
<td>Linear ssRNA</td>
</tr>
<tr>
<td>Tobacco mosaic virus</td>
<td>Plants</td>
<td>Linear ssRNA</td>
</tr>
<tr>
<td>Vaccinia virus</td>
<td>Mammals</td>
<td>Linear dsDNA</td>
</tr>
</tbody>
</table>

The genome structure is that in the viral capsid; some genomes exist in different forms within the host cell. ds, double-stranded; ss, single-stranded.
bacteriophages. A number of DNA and RNA viruses are able to integrate into the genomes of their hosts, sometimes with drastic effects on the host cell. The viral retroelements are examples of integrative eukaryotic viruses. Their replication pathways include a novel step in which an RNA version of the genome is converted into DNA. There are two kinds of viral retroelements: retroviruses, whose capsids contain the RNA version of the genome, and pararetroviruses, whose encapsidated genome is made of DNA. The ability of viral retroelements to convert RNA into DNA was confirmed independently in 1970 by Howard Temin and by David Baltimore. Working with cells infected with retroviruses, both Temin and Baltimore isolated the enzyme, now called reverse transcriptase, which is capable of making a DNA copy of an RNA template (and is of immense utility in the experimental study of genomes; see Section 2.1). The typical retroviral genome is a single-stranded RNA molecule, 7–12 kb in length. After entry into the cell, the genome is copied into double-stranded DNA by a few molecules of reverse transcriptase that the virus carries in its capsid. The double-stranded version of the genome then integrates into the host DNA (Figure 9.6). Unlike λ, the retroviral genome has no sequence similarity with its insertion site in the host DNA. Integration of the viral genome into the host DNA is a prerequisite for expression of the retroviral genes. There are three of these, called gag, pol, and env (Figure 9.7). Each codes for a polypeptide that is cleaved, after translation, into two or more functional gene products. These products include viral coat proteins (from env) and reverse transcriptase (from pol). The protein products combine with full-length RNA transcripts of the retroviral genome to produce new virus particles.

The causative agents of HIV/AIDS (human immunodeficiency virus infection and acquired immune deficiency syndrome) were shown to be retroviruses in 1983–1984. The first human immunodeficiency virus was isolated independently by two groups, led by Luc Montagnier and Robert Gallo. This virus is called HIV-1 and is responsible for the most prevalent and pathogenic form of HIV/AIDS. A related virus, HIV-2, discovered by Montagnier in 1985, is less widespread and causes a milder form of the disease. The human immunodeficiency viruses attack certain types of lymphocyte in the bloodstream, thereby depressing the immune response of the host. These lymphocytes carry on their surfaces multiple copies of a protein called CD4, which acts as a receptor for the virus. An HIV particle binds to a CD4 protein and then enters the lymphocyte after fusion between its lipid envelope and the cell membrane.

Some retroviruses cause cancer

The human immunodeficiency viruses are not the only retroviruses capable of causing diseases. Several retroviruses can induce cell transformation, possibly leading to cancer. Cell transformation involves changes in cell morphology and physiology. In cell cultures, transformation results in a loss of control over growth,
so that transformed cells grow as a disorganized mass, rather than as a monolayer (Figure 9.8). In whole animals, cell transformation is thought to underlie the development of tumors.

There appear to be two distinct ways in which retroviruses can cause cell transformation. With some retroviruses, such as the leukemia viruses, cell transformation is a natural consequence of infection, although it may be induced only after a long latent period during which the integrated provirus lies quiescent within the host genome. Other retroviruses cause cell transformation because of abnormalities in their genome structures. These viruses carry cellular genes that they have captured by some undefined process. With at least one transforming retrovirus (Rous sarcoma virus), this cellular gene is in addition to the standard retroviral genes (Figure 9.9A). With others, the cellular gene replaces part of the retroviral gene complement (Figure 9.9B). In the latter case the retrovirus may be defective, meaning that it is unable to replicate and produce new viruses, as it has lost genes coding for vital replication enzymes and/or capsid proteins. These defective retroviruses are not always inactive, as they can make use of proteins provided by other retroviruses in the same cell (Figure 9.10). The ability of a transforming retrovirus to cause cell transformation lies with the nature of the
cellular gene that has been captured. Often this captured gene (called a v-onc, with onc standing for oncogene) codes for a protein involved in cell proliferation. The normal cellular version of the gene is subject to strict regulation and is expressed only in limited quantities when needed. It is thought that expression of the v-onc gene follows a different, less controlled pattern, either because of changes in the gene structure or because of the influence of expression signals within the retrovirus. One result of this altered expression pattern could be a loss of control over cell division, leading to the transformed state.

Genomes at the edge of life
Viruses occupy the boundary between the living and nonliving worlds. At the very edge of this boundary, or perhaps beyond it, reside a variety of nucleic acid molecules that might or might not be classified as genomes. The satellite RNAs or virusoids are examples. These are RNA molecules, some 320–400 nucleotides in length, which do not encode their own capsid proteins, instead moving from cell to cell within the capsids of helper viruses. The distinction between the two groups is that a satellite virus shares the capsid with the genome of the helper virus, whereas a virusoid RNA molecule becomes encapsidated on its own. They are generally looked on as parasites of their helper viruses, although there appear to be at least a few cases where the helper cannot replicate without the satellite RNA or virusoid, suggesting that at least some of the relationships are symbiotic. Satellite RNAs and virusoids are both found predominantly in plants, as is a more extreme group called the viroids. These are RNA molecules, 240–475 nucleotides in length, which contain no genes and never become encapsidated, spreading from cell to cell as naked RNA. They include some economically important pathogens, such as the citrus exocortis viroid, which reduces the growth of citrus fruit trees. Viroid and virusoid molecules are circular and single-stranded and are replicated by enzymes coded by the genome of the host or helper virus. The replication process results in a series of RNAs joined head to tail, and with some viroids and virusoids these are cleaved by a self-catalyzed reaction in which the RNA molecule acts as an enzyme (Figure 9.11).

Nucleic acid molecules that replicate within plant cells can perhaps be looked on as genomes even if they contain no genes. The same cannot be said for prions, as these infectious, disease-causing particles contain no nucleic acid. Prions are responsible for scrapie in sheep and goats, and their transmission to cattle has led to the new disease called bovine spongiform encephalopathy (BSE). Whether their further transmission to humans causes a variant form of Creutzfeldt–Jakob disease (CJD) is controversial but accepted by many biologists. At first prions were thought to be viruses, but it is now clear that they are made solely of protein. The normal version of the prion protein, called PrP\textsuperscript{C}, is coded by a mammalian nuclear gene and synthesized in the brain, although its function is unknown. PrP\textsuperscript{C} is easily digested by proteases whereas the infectious version, PrP\textsuperscript{Sc}, has a more highly β-sheeted structure that is resistant to proteases and forms fibrillar aggregates that are seen in infected tissues. Once inside a cell, PrP\textsuperscript{Sc} molecules are able to convert newly synthesized PrP\textsuperscript{C} proteins into the infectious form, by a mechanism that is not yet understood, resulting in the disease state. Transfer of one or more of these PrP\textsuperscript{Sc} proteins to a new animal results in accumulation of new PrP\textsuperscript{Sc} proteins in the brain of that animal, transmitting the disease (Figure 9.12). Infectious proteins with similar properties are known in lower eukaryotes, examples being the Ure3 and Psi‘ prions of Saccharomyces cerevisiae. It is clear, however, that prions are gene products rather than genetic material, and despite their infectious properties, which led to the initial confusion regarding their status, they are unrelated to viruses or to subviral particles such as viroids and virusoids.

Figure 9.11 Self-catalyzed cleavage of linked genomes during replication of viroids and virusoids. (A) The replication pathway. (B) The hammerhead structure, which forms at each cleavage site and has enzymatic activity. N indicates any nucleotide.

Figure 9.12 Mode of action of a prion. A normal, healthy sheep has PrP\textsuperscript{C} proteins in its brain. Infection with PrP\textsuperscript{Sc} molecules leads to conversion of newly synthesized PrP\textsuperscript{C} proteins into PrP\textsuperscript{Sc}, leading to the disease state, known as scrapie in sheep.
Chapter 9: Viral Genomes and Mobile Genetic Elements

9.2 MOBILE GENETIC ELEMENTS

In Chapters 7 and 8 we learned that eukaryotic genomes, and to a lesser extent those of prokaryotes, contain interspersed repeats, some with copy numbers of several thousand per genome. For many interspersed repeats, the genomewide distribution pattern is set up by transposition, the process by which a segment of DNA can move from one position to another in a genome. These movable segments are called transposable elements or transposons. Some types move by a conservative process, which involves the excision of the sequence from its original position followed by its reinsertion elsewhere. Conservative transposition therefore results in the transposon simply changing its position in the genome without increasing its copy number (Figure 9.13). Replicative transposition, on the other hand, results in an increase in copy number, because during this process the original element remains in place while a copy is inserted at the new position. This replicative process can therefore lead to a proliferation of the transposon at interspersed positions around the genome.

Both types of transposition involve recombination, and we will therefore deal with the details of the processes when we study recombination and related types of genome rearrangement in Section 17.3. What interests us here is the variety of structures displayed by the transposable elements found in eukaryotic and prokaryotic genomes and the link that exists between these elements and viral genomes.

RNA transposons with long terminal repeats are related to viral retroelements

Replicative transposons can be further subdivided into those that transpose via an RNA intermediate and those that do not. The process that involves an RNA intermediate, which is called retrotransposition, begins with synthesis of an RNA copy of the retrotransposon by the normal process of transcription (Figure 9.14). The transcript is then copied into double-stranded DNA, which initially exists as an independent molecule outside of the genome. Finally, the DNA copy of the transposon integrates into the genome, possibly back into the same chromosome occupied by the original unit or possibly into a different chromosome. The end result is that there are now two copies of the transposon, at different points in the genome. If we compare the mechanism for retrotransposition with that for replication of a viral retroelement, as shown in Figure 9.6, then we see that the two processes are very similar. The one significant difference is that the RNA molecule that initiates the process is transcribed from an endogenous genomic sequence during retrotransposition and from an exogenous viral genome during replication of a viral retroelement. This close similarity alerts us to the relationships that exist between these two types of elements.

RNA transposons, or retroelements, are common features of eukaryotic genomes but are much less common, and less well studied, in prokaryotes. The eukaryotic versions can be broadly classified into two types: those that possess long terminal repeats (LTRs) and those that do not. Long terminal repeats, which play a central role in the process by which the RNA copy of an LTR element is reverse-transcribed into double-stranded DNA (Section 17.3), are also possessed by viral retroelements (see Figure 9.7). It is now clear that these viruses are one member of a superfamily of elements that also includes endogenous LTR transposons.
The first of the endogenous elements to be discovered was the Ty sequence of yeast, which is 6.3 kb in length and has a copy number of about 50 in most Saccharomyces cerevisiae genomes. There are several types of Ty element in yeast genomes, the most abundant of which, Ty1, is similar to the copia retroelement of the fruit fly. These elements are therefore now called the Ty1/copia family. If we compare the structure of a viral retroelement (Figure 9.15A) with that of a Ty1/copia retroelement (Figure 9.15B), we see clear family relationships. Each Ty1/copia element contains two genes, called TyA and TyB in yeast, which are similar to the gag and pol genes of a viral retroelement. In particular, TyB codes for a polyprotein that includes the reverse transcriptase that plays the central role in transposition of a Ty1/copia element. Note, however, that the Ty1/copia element lacks an equivalent of the viral env gene, the one that codes for the viral coat proteins. This means that Ty1/copia retroelements cannot form infectious virus particles and therefore cannot escape from their host cell. They do, however, form virus-like particles (VLPs) consisting of the RNA and DNA copies of the retroelements attached to core proteins derived from the TyA polyprotein. In contrast, the members of a second family of LTR retroelements, called Ty3/gypsy (again after the yeast and fruit fly versions), do have an equivalent of the env gene (Figure 9.15C), and at least some of these can form infectious viruses. Although classed as endogenous transposons, these infectious versions should be looked upon as viral retroelements.

Yeast genomes also contain 300–400 additional copies of the 330 bp LTRs of Ty elements. These solo sequences probably arise by homologous recombination between the two LTRs of a Ty element, which could excise the bulk of the element and leave a single LTR (Figure 9.16). This excision event is probably unrelated to transposition of a Ty element, which occurs by the RNA-mediated process shown in Figure 9.14. The most common of these solo LTRs are called delta sequences, which derive from Ty1/copia elements. Sigma elements, which are solo LTRs from Ty3/gypsy retrotransposons, have copy numbers of 20–30 per genome.

LTR retroelements make up substantial parts of many eukaryotic genomes and are particularly abundant in the larger plant genomes, especially those of grasses such as maize (see Figure 7.15D). They also make up an important component of invertebrate and some vertebrate genomes, but in the genomes of humans and other mammals most of the LTR elements appear to be decayed viral retroelements rather than true transposons. These sequences are called endogenous retroviruses (ERVs) and they make up approximately 9% of the human genome (Table 9.3). Human ERVs are 6–11 kb in length and have copies

### Table 9.3 Transposable Elements in the Human Genome

<table>
<thead>
<tr>
<th>Class</th>
<th>Family</th>
<th>Fraction of genome (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SINE</td>
<td>Alu</td>
<td>10.5</td>
</tr>
<tr>
<td></td>
<td>MIR</td>
<td>2.5</td>
</tr>
<tr>
<td></td>
<td>MIR3</td>
<td>0.4</td>
</tr>
<tr>
<td>LINE</td>
<td>LINE-1</td>
<td>17.5</td>
</tr>
<tr>
<td></td>
<td>LINE-2</td>
<td>3.4</td>
</tr>
<tr>
<td></td>
<td>LINE-3</td>
<td>0.3</td>
</tr>
<tr>
<td>LTR retroelements</td>
<td>ERV1</td>
<td>2.9</td>
</tr>
<tr>
<td></td>
<td>ERVL</td>
<td>5.8</td>
</tr>
<tr>
<td>DNA transposons</td>
<td>TcMar</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td>hAT</td>
<td>2.2</td>
</tr>
</tbody>
</table>

Data from RepeatMasker analysis of hg38 assembly.

LINE, long interspersed nuclear element; SINE, short interspersed nuclear element; LTR, long terminal repeat; ERV, endogenous retrovirus.
of the *gag*, *pol*, and *env* genes. Although most contain mutations or deletions that inactivate one or more of these genes, a few members of the human ERV group HERV-K have functional sequences. By comparing the positions of the HERV-K elements in the genomes of different individuals, it has been inferred that at least some of the HERV-K family are active retroposons. There is also evidence that the RNA copies of some HERV-K elements can be packaged into virus-like particles with the ability to move from cell to cell. These discoveries have prompted studies into possible roles for HERV-K elements in human diseases. HERV-K transcripts and protein products are detectable in the brains of patients suffering from amyotrophic lateral sclerosis (ALS), a neurodegenerative disorder also called Lou Gehrig’s disease, after the famous American baseball player who died of ALS in 1941. Detection of HERV-K products does not prove that these are responsible for ALS, but a possible association is suggested by the demonstration that expression of the HERV-K *env* gene in mouse brains leads to breakdown of motor neuron function and symptoms similar to ALS. It is also possible that HERV-K elements are involved in susceptibility to autoimmune diseases such as rheumatoid arthritis.

**Some RNA transposons lack long terminal repeats**

Not all types of RNA transposons have LTR elements. In mammals the most important types of non-LTR retroelements, or retroposons, are the **LINEs** (long interspersed nuclear elements) and **SINEs** (short interspersed nuclear elements). SINEs have the highest copy number for any type of interspersed repetitive DNA in the human genome, with over 1.7 million copies comprising almost 14% of the genome as a whole (Table 9.3). LINEs are less frequent, with just over 1 million copies, but as they are longer, they make up a larger fraction of the genome (over 20%). The abundance of LINEs and SINEs in the human genome is underlined by their frequency in the 200 kb segment that we looked at in Section 7.2 (see Figure 7.12).

There are three families of LINEs in the human genome, of which one group, **LINE-1**, is both the most frequent and the only type that is able to transpose, as the LINE-2 and LINE-3 families are made up of inactive relics. A full-length LINE-1 element is 6.1 kb and has two genes, one of which codes for a polyprotein similar to the product of the viral *pol* gene (Figure 9.17A). There are no LTRs, but the 3′-end of the LINE is marked by a series of A-T base pairs, giving what is usually referred to as a poly(A) sequence, though of course it is a poly(T) sequence on the other strand of the DNA. Not all copies of LINE-1 are full-length because the reverse transcriptase coded by LINEs does not always make a complete DNA copy of the initial RNA transcript, meaning that part of the 3′-end of the LINE may be lost. This truncation event is so common that only 1% of the LINE-1 elements in the human genome are full-length versions and the average size of all the copies is 900 bp. SINEs are much shorter than LINEs, just 100–400 bp and not containing any genes, which means that SINEs do not make their own reverse transcriptase enzymes (Figure 9.17B). Instead they borrow reverse transcriptases that have been synthesized by LINEs. The commonest SINE in primate genomes is **Alu**, which has a copy number of approximately 1.2 million in humans. An Alu element comprises two halves, each half made up of a similar 120 bp sequence, with a 31–32 bp insertion in the right half (Figure 9.18). The mouse genome has a related element, called B1, which is 130 bp in length and equivalent to half of an Alu sequence. Some Alu elements are actively copied into RNA, providing the opportunity for proliferation of the element. Alu is derived from the gene for the 7SL RNA, a noncoding RNA involved in movement of proteins around the cell. The first Alu element may have arisen by the accidental reverse transcription of a 7SL RNA molecule and integration of the DNA copy into the human genome. Other SINEs are derived from tRNA genes which, like the gene for the 7SL RNA, are transcribed by RNA polymerase III in eukaryotic cells, suggesting that some feature of the transcripts synthesized by this polymerase make these molecules prone to occasional conversion into retroposons.
Although transposition of LINEs and SINEs is a rare event, LINE-1 transposition has been observed in cultured human and mouse cells, and the recent insertions of LINE-1, Alu, and other SINEs into protein-coding sequences is thought to have led to gene inactivations that have given rise to inherited human disorders. This was recognized with a small number of hemophilia patients, whose factor VIII gene was disrupted by a LINE-1 sequence that prevented synthesis of this important blood clotting protein. Since this initial discovery, LINE-1 insertion has been implicated as the causative factor in examples of over 25 diseases, and increasing numbers of examples involving SINEs are being discovered.

A few non-LTR retroelements are known in prokaryotic genomes, but these have much lower copy numbers than eukaryotic RNA transposons. The prokaryotic versions have a broad distribution among bacteria and archaea, but the distribution is uneven: some strains of *E. coli* possess retroelements and others do not. The commonest type of bacterial retroelement is the *retron*, which is a 2 kb sequence that includes a reverse transcriptase gene. A second part of the retron specifies a 70–80 nucleotide RNA which is copied into DNA by the reverse transcriptase. The 5′-end of this single-stranded DNA then forms a 2′–5′ phosphodiester bond with a guanine nucleotide within the RNA, giving rise to an RNA–DNA hybrid that adopts a base-paired secondary structure. Whether or not this structure has any function is still being debated, but there is evidence that synthesis of the retron reverse transcriptase enhances the ability of *Salmonella typhimurium* to colonize the human gut and that the presence of retron sequences increases the pathogenicity of *Vibrio cholerae*.

**DNA transposons are common in prokaryotic genomes**

Not all transposons require an RNA intermediate. Those called DNA transposons are able to transpose in a more direct DNA-to-DNA manner. DNA transposons are an important component of many prokaryotic genomes. The insertion sequences IS1 and IS186, present in the 50 kb segment of *E. coli* DNA that we examined in Section 8.2 (see Figure 8.8), are examples of DNA transposons. Their copy numbers vary in different species and different strains, but a single *E. coli* genome will usually contain 30–50 insertion sequences of various types. An insertion sequence (IS) is 0.7–2.5 kb in length, with most of its sequence taken up by one or two genes that specify the transposase enzyme that catalyzes its transposition (Figure 9.19A). There is a pair of inverted repeats at either end of each IS element, up to about 50 bp in length depending on the type of IS, and insertion of the element into the target DNA creates a pair of short (usually 4–15 bp) direct repeats in the host genome. IS elements can transpose either replicatively or conservatively.

IS elements are also components of a second type of DNA transposon first characterized in *E. coli* and now known to be common in many prokaryotes. These composite transposons are made up of a pair of IS elements flanking a segment of DNA, usually containing one or more genes, often ones coding for antibiotic resistance (Figure 9.19B). Tn10, for example, carries a gene for tetracycline resistance, and Tn5 and Tn903 both carry a gene for resistance to kanamycin. Some composite transposons have identical IS elements at either end, while others have one element of one type and one of another. In some cases the IS elements are oriented as direct repeats, and sometimes they appear as inverted repeats. These variations do not appear to affect the transposition mechanism for a composite transposon, which is conservative in nature and catalyzed by the transposase coded by one or both of the IS elements.

Various other classes of DNA transposon are known in prokaryotes. Two additional important types from *E. coli* are as follows:

- **Tn3-type or unit transposons** have their own transposase gene and so do not require flanking IS elements in order to transpose (Figure 9.19C). Tn3 elements transpose replicatively.

- **Transposable phages** are bacterial viruses that transpose replicatively as part of their normal infection cycle (Figure 9.19D).
Prokaryotic genomes also contain IS fragments that have lost the ability to transpose. These are called **miniature inverted repeat transposable elements (MITEs)**. MITE is a general term for the truncated relic of a DNA transposon, and these sequences were first identified in plants.

**DNA transposons are less common in eukaryotic genomes**

About 3.7% of the human genome is made up of DNA transposons of various types (Table 9.3), all with terminal inverted repeats and all containing a gene for a transposase enzyme that catalyzes the transposition event. However, the vast majority of these elements are inactive, either because the transposase gene is nonfunctional or because sequences at the ends of the transposon, which are essential for active transposition, are missing or mutated.

Active DNA transposons are more common in plants and include the **Ac/Ds transposon** and the **Spm element**, both of which are found in maize. The Ac/Ds elements were the first transposons to be discovered, by Barbara McClintock in the 1950s. Her conclusions—that some genes are mobile and can move from one position to another in a chromosome—were based on exquisite genetic experiments, as the molecular basis of transposition was not understood until the late 1970s. An interesting feature of these plant transposons is that they work together in family groups. For example, the Ac element codes for an active transposase that recognizes both Ac elements and Ds sequences. The latter are versions of Ac that have internal deletions that remove part of the transposase gene, meaning that a Ds element cannot make its own transposase and can move only through the activity of the transposase synthesized by a full-length Ac element (Figure 9.20). Similarly, full-length Spm elements are accompanied by deleted versions that transpose through use of the transposase enzymes coded by the intact elements. The activity of Ac elements is apparent during the normal life cycle of a maize plant: transposition in somatic cells results in changes in gene expression which are manifested in, for example, variegated pigmentation in maize kernels (Figure 9.21).
McClintock’s realization that the maize genome contains transposable elements resulted from her studies into the genetic basis of the different color patterns displayed by kernels. The P element, a DNA transposon in *Drosophila melanogaster*, was similarly discovered from studies of an unusual genetic event that, as it turns out, arises from transposition. This event is called **hybrid dysgenesis** and occurs when females from laboratory strains of *D. melanogaster* are crossed with males from wild populations. The offspring resulting from such crosses are sterile and have chromosomal abnormalities along with a variety of other genetic malfunctions. The explanation is that the genomes of wild fruit flies contain inactive versions of **P elements**, which are typical DNA transposons comprising a transposase gene flanked by inverted terminal repeats, but that laboratory strains lack these P elements. After crossing, the elements inherited from the wild flies become active in the fertilized eggs, transposing into various new positions and causing the gene disruptions that characterize hybrid dysgenesis (**Figure 9.22**). Exactly why this activation occurs is not known, but a more interesting question is why the genomes of wild populations of *D. melanogaster* contain P elements whereas laboratory strains do not. Most of the laboratory strains are descended from flies collected by Thomas Hunt Morgan some 90 years ago and used by Morgan and his colleagues in the first gene mapping experiments (**Section 3.3**). It appears that wild populations at that time lacked P elements, which have somehow proliferated in wild genomes during the last 90 years. The inability of wild and laboratory flies to produce viable offspring means that these two populations fail one of the main criteria used to identify biological species: the ability of all individuals to mate productively. This raises the intriguing possibility that speciation, at least in some organisms, might be driven by differential proliferation of transposable elements within the genomes of members of different populations.

**Figure 9.21** Variegated pigmentation in maize kernels caused by transposition in somatic cells. The highly colored forms of *Zea mays* are popularly known as Indian corn. (Courtesy of Lena Struwe, Rutgers University.)

**Figure 9.22** Hybrid dysgenesis. Crosses between male lab flies and female wild flies give normal progeny, but when the male partner is a wild fly, the offspring are sterile. One possible explanation of hybrid dysgenesis is that the cytoplasm of flies with P elements (P⁺ in this diagram) contains a repressor that prevents P element transposition. The fertilized egg resulting from a cross between a female P⁺ fly and male P⁻ fly will contain this repressor and so the progeny are normal. However, the repressor will not be carried in the sperm from a male P⁺ fly, so the fertilized egg from a cross between a male P⁺ and a female P⁻ fly will lack the repressor, allowing P element transposition to occur and resulting in progeny displaying hybrid dysgenesis.
SUMMARY

- Early studies of viruses focused largely on the bacteriophages, the viruses that infect bacteria.
- Bacteriophages are constructed of protein and nucleic acid, the protein forming a capsid that encloses the genome.
- There are three basic types of capsid structure but many types of genome organization. Different phages have single- or double-stranded DNA or RNA genomes and the genome may be contained in a single molecule or divided into two or more segments.
- Bacteriophages follow two distinct infection cycles. All phages can infect via the lytic cycle, which results in the immediate synthesis of new bacteriophages, usually accompanied by death of the host cell. Some phages can also follow the lysogenic cycle, during which a copy of the phage genome becomes inserted into the host DNA, where it may remain in quiescent form for many generations.
- Eukaryotic viruses are equally diverse in terms of genome organization but display just two capsid structures.
- Most eukaryotic viruses follow a lytic infection cycle but this does not always result in the immediate death of the host cell. A number of DNA and RNA viruses can integrate their genomes into eukaryotic chromosomes in a manner similar to a lysogenic bacteriophage.
- The viral retroelements, which include HIV, the causative agent of HIV/AIDS, are examples of integrative RNA viruses.
- Satellite RNAs and virusoids are different types of infective RNA molecule that contain no genes and depend on other viruses for their transmission. Viroids are small infective RNA molecules that self-replicate, and prions are infective proteins.
- Some mobile genetic elements, which are DNA sequences that can transpose within a genome but cannot escape from the cell, are related to RNA viruses. These elements transpose via an RNA intermediate in a pathway similar to the infection process of viral retroelements.
- The Ty1/copia and Ty3/gypsy retrotransposons, and the endogenous retroviruses of mammals, are the mobile elements most closely related to RNA viruses.
- Mammalian genomes also contain other types of retrotransposon, called LINEs and SINEs, most of which have lost their ability to transpose.
- DNA transposons do not make use of an RNA intermediate in their transposition pathway. These transposons are common in bacteria, within which they are responsible for the spread of genes coding for antibiotic resistance.
- DNA transposons are less widespread in eukaryotes but include some important examples, such as the Ac/Ds transposon of maize, the first transposon of any kind to be studied in detail, and the P element of Drosophila melanogaster, which is responsible for the hybrid dysgenesis that occurs when female laboratory fruit flies are crossed with wild male flies.

SHORT ANSWER QUESTIONS

1. How are viruses different from cells? Is it appropriate to look on viruses as living organisms?
2. Outline the key differences between viral and cellular genomes.
3. Using examples, explain what is meant by overlapping genes, as found in some viral genomes.
4. How long does it take a lytic bacteriophage to lyse a host cell following the initial infection? What is the timeline for the lytic infection cycle of T4 phage?
5. Describe the differences between the capsids of bacteriophages and eukaryotic viruses.
6. List the key stages in the life cycle of a retrovirus.
7. What is a transposon?
8. Describe the characteristic features of the LTR retroelements present in the human genome.
9. Discuss the properties and types of retroposons present in the human genome.
10. What are the general properties of composite transposons?
11. What are the important features of the DNA transposons found in plants?
12. Describe the basis to hybrid dysgenesis in fruit flies.

**IN-DEPTH PROBLEMS**

1. To what extent can viruses be considered a form of life?
2. Bacteriophages with small genomes (for example, φX174) are able to replicate very successfully in their hosts. Why then should other bacteriophages, such as T4, have large and complicated genomes?
3. Some bacteriophages, such as T4, modify the host RNA polymerase after infection so that this polymerase no longer recognizes *E. coli* genes but transcribes bacteriophage genes instead. How might this modification be carried out?
4. Genetic elements that reproduce within or along with a host genome, but confer no benefit on the host, are sometimes called selfish DNA. Discuss this concept, in particular as it applies to transposons.
5. Why do LTR retroelements have long terminal repeats?

**FURTHER READING**

**Classic papers on bacteriophage genetics**


**Bacteriophage genome sequences**


**Eukaryotic viruses**


**Edge of life**


**RNA transposons**


**DNA transposons**


**Online resource**

RepeatMasker. http://www.repeatmasker.org/ *The “Genome Analysis and Downloads” component enables the repetitive DNA content of various genomes to be viewed.*
In order for the cell to utilize the biological information contained within its genome, groups of genes, each gene representing a single unit of information, have to be expressed in a coordinated manner. This coordinated gene expression determines the makeup of the transcriptome, which in turn specifies the nature of the proteome and defines the activities that the cell is able to carry out. In Part III of Genomes, we will examine the events that result in the transfer of biological information from genome to proteome. Our knowledge of these events was initially gained through studies of individual genes, often as naked DNA in test-tube experiments. These experiments provided an interpretation of gene expression that in recent years has been embellished by more sophisticated studies that have taken greater account of the fact that, in reality, it is the genome that is expressed, not individual genes, and this expression occurs in living cells rather than in a test tube.

We begin our investigation of genome expression, here in Chapter 10, by examining the substantial and important impact that the nuclear environment has on utilization of the biological information contained in the genomes of eukaryotes. The accessibility of that information is dependent on the way in which the DNA is packaged into chromatin and is responsive to processes that can silence or inactivate part or all of a chromosome. Chapter 11 then describes the critical role played by DNA-binding proteins in genome expression and explains how the structures of these proteins enable them to recognize their specific binding sites. The composition of the transcriptome, and the role of the transcriptome in the overall process of genome expression, is dealt with in Chapter 12, and Chapter 13 covers the equivalent issues regarding the composition and role of the proteome. As you read Chapters 10–13, you will discover that control over the composition of the transcriptome and the proteome can be exerted at various stages during the overall chain of events that make up genome expression. These regulatory threads will be drawn together in Chapter 14, where we examine how the genome acts within the context of cell and organism by responding to extracellular signals and by driving the biochemical changes that underlie differentiation and development.

10.1 INSIDE THE NUCLEUS

When we look at a genome sequence written out as a series of A, C, G, and T nucleotides or use a genome browser to examine a segment of a chromosome (as in
Figure 5.22, for example), there is a tendency to imagine that all parts of the genome are readily accessible to the DNA-binding proteins that are responsible for its expression. In reality, the situation is very different. The DNA in the nucleus of a eukaryotic cell or the nucleoid of a prokaryote is attached to a variety of proteins that are not directly involved in genome expression and must be displaced in order for RNA polymerase and other expression proteins to gain access to the genes. We know very little about these events in prokaryotes, a reflection of our generally poor knowledge about the physical organization of the prokaryotic genome (Section 8.1), but we are beginning to understand how the packaging of DNA into chromatin (Section 7.1) influences genome expression in eukaryotes. This is an exciting area of molecular biology, with recent research indicating that histones and other packaging proteins are not simply inert structures around which the DNA is wound, but instead are active participants in the processes that determine which parts of the genome are expressed in an individual cell. Many of the discoveries in this area have been driven by new insights into the substructure of the nucleus, so we begin with this topic.

The nucleus has an ordered internal structure

The internal architecture of the nucleus was first examined by light and electron microscopy. With conventional techniques, the inside of the nucleus appears to be relatively unstructured, made up of lighter and darker regions with just one distinct feature, the nucleolus, which is the center for synthesis and processing of rRNA molecules and which appears as a dark area when nuclei are observed with the electron microscope (Figure 10.1). These early microscopy studies therefore suggested that the nucleus has little internal organization, a typical black box in common parlance. In recent years this interpretation has been overthrown, and we now appreciate that the nucleus has an ordered structure that is related to the variety of biochemical activities that it must carry out. Indeed, the inside of the nucleus is just as complex as the cytoplasm of the cell. The only difference is that, in contrast to the cytoplasm, the functional compartments within the nucleus are not individually enclosed by membranes and so are not visible when the cell is observed by conventional light or electron microscopy techniques.

The revised picture of nuclear structure first began to emerge when techniques were developed for labeling different types of nuclear protein with fluorescent markers. This can be achieved by ligating the coding sequence for green fluorescent protein (GFP) to the gene for the protein being studied. Standard cloning techniques are then used to insert the modified gene into the host genome, leading to a recombinant cell that synthesizes a fluorescent version of the protein. Observation of the cell with a fluorescence microscope now reveals the distribution of the labeled protein within the nucleus. For example, nucleoli can be visualized by labeling the fibrillarin protein with GFP (Figure 10.2A), fibrillarin being a component of the small nucleolar ribonucleoproteins that are involved in rRNA processing. Labeling of proteins involved in mRNA splicing (Section 12.4) has shown that this activity is also localized into distinct regions, called speckles (Figure 10.2B), although these are more widely distributed and less well defined than the nucleoli. Other structures, such as Cajal bodies (visible
in Figure 10.2A), which are probably involved in synthesis of small nuclear and small nucleolar RNAs (Section 12.1), can also be seen after fluorescent labeling.

Another new microscopy technique, called fluorescence recovery after photobleaching (FRAP), enables the movement of fluorescently labeled proteins within the nucleus to be visualized. A small area of the nucleus is photobleached by exposure to a tightly focused pulse from a high-energy laser. The laser pulse inactivates the fluorescent signal in the exposed area, leaving a region that appears bleached in the microscopic image. This bleached area gradually retrieves its fluorescent signal, not by a reversal of the bleaching effect but by migration into the bleached region of fluorescent proteins from the unexposed area of the nucleus (Figure 10.3). Rapid reappearance of the fluorescent signal in the bleached area therefore indicates that the tagged proteins are highly mobile, whereas a slow recovery indicates that the proteins are relatively static. Studies of this type have shown that the migration of nuclear proteins does not occur as rapidly as would be expected if their movement were totally unhindered, which is entirely expected in view of the large amounts of DNA and RNA in the nucleus, but that it is still possible for a protein to traverse the entire diameter of a nucleus in a matter of minutes. Proteins involved in genome expression therefore have the freedom needed to move from one activity site to another, as dictated by the changing requirements of the cell. In particular, the linker histones (Section 7.1) continually detach and reattach to their binding sites on the genome. This discovery is important because it emphasizes that the DNA–protein complexes that make up chromatin are dynamic, an observation that has considerable relevance to genome expression, as we will see later in this chapter.

**The DNA content of a nondividing nucleus displays different degrees of packaging**

In Section 7.1 we learned that chromatin is the complex of genomic DNA and chromosomal proteins present in the eukaryotic nucleus. Chromatin structure is hierarchical, ranging from the two lowest levels of DNA packaging, the nucleosome and the 30 nm chromatin fiber (see Figures 7.2 and 7.3), to the metaphase chromosomes, which represent the most compact form of chromatin in eukaryotes and occur only during nuclear division. After division, the chromosomes become less compact and cannot be distinguished as individual structures unless special techniques are used, such as chromosome painting, which we will discuss later. When nondividing nuclei are examined by conventional electron microscopy, all that can be seen, other than the nucleoli, is a mixture of light and dark areas (Figure 10.4). The light areas are called eu chromatin and are thought to be made up of DNA that has a relatively open conformation, either as the 30 nm fiber or possibly simply as nucleosomes in the beads-on-a-string format (see Figure 7.2A). Euchromatin comprises those parts of the chromosomal DNA that contain active genes, ones that are being transcribed into RNA. A less compact packaging is adopted by these regions so that RNA polymerase and other proteins involved in transcription can access these genes.

The dark areas are called heterochromatin and contain DNA that is still in a relatively compact organization, although less compact than in the metaphase structure. These regions are transcriptionally silent, containing few if any active genes. Two types of heterochromatin are recognized:
Chapter 10: Accessing the Genome

• **Constitutive heterochromatin** is a permanent feature of all cells and represents DNA that contains no genes and so can always be retained in a compact organization. This fraction includes centromeric and telomeric DNA, as well as certain regions of some other chromosomes. For example, most of the human Y chromosome is made of constitutive heterochromatin (see Figure 7.6).

• **Facultative heterochromatin** is not a permanent feature but is seen in some cells some of the time. Facultative heterochromatin contains genes that are inactive in some cells or at some periods of the cell cycle. When these genes are inactive, their DNA regions are compacted into heterochromatin.

The nuclear matrix is thought to provide attachment points for chromosomal DNA

The exact organization of the euchromatin regions of the nucleus is not known, but with the electron microscope it is possible to see loops of DNA, each loop between 4 and 200 kb in length. The boundaries of these loops are believed to be attached to a network of fibers called the **nuclear matrix**, equivalent to the cytoskeleton that extends throughout the cytoplasm. However, this is a controversial area of cell biology. It is known that there is a thin network of filaments on the internal side of the nuclear membrane, forming a **nuclear lamina** that is attached to proteins that are embedded in the nuclear membrane. What is questioned is whether this network extends into the internal regions of the nucleus, forming a matrix that permeates the entire nucleoplasm.

The existence of a nuclear matrix was suggested by examination of mammalian nuclei that had been prepared in a special way. After dissolution of membranes by soaking in a mild, nonionic detergent such as one of the Tween compounds, followed by treatment with a deoxyribonuclease to degrade the nuclear DNA and salt extraction to remove the chemically basic histone proteins, a complex network of protein and RNA fibrils was revealed, apparently extending throughout the entire nucleus (Figure 10.5). The structural proteins making up this matrix are lamins and lamins, and they are accompanied by enzymes and other proteins involved in genome replication, such as DNA polymerases and DNA topoisomerases. These experiments might appear to give clear evidence for the existence of the nuclear matrix, but could the structures seen in these electron micrographs be an artifact resulting from the preparation method? Doubts were first raised when intact nuclei were examined by **immunofluorescence microscopy**. Like the methods based on green fluorescent protein that we discussed earlier, immunofluorescence microscopy utilizes a fluorescent marker to visualize the location of particular proteins within a cell. The difference is that visualization is achieved not by engineering the cell to synthesize a GFP fusion of the protein of interest, but instead by treating a tissue section with fluorescently labeled antibodies that bind specifically to that protein. We might anticipate that when this technique is used with antibodies specific for proteins of the nuclear matrix, the latter will be

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**Figure 10.5 Evidence for the existence of a nuclear matrix.** This transmission electron micrograph shows the nuclear matrix of a cultured human HeLa cell. Cells were treated with a nonionic detergent to remove membranes, digested with a deoxyribonuclease to degrade most of the DNA, and extracted with ammonium sulfate to remove histones and other chromatin-associated proteins. (From Penman S, Fulton A, Capco D et al. [1982] Symp. Quant. Biol. 46:1013–1028. With permission from Cold Spring Harbor Laboratory Press.)
revealed as a network of fluorescence. This is not what has been seen: instead, the matrix proteins appear to form distinct spots rather than fibers. Questions regarding the nature and importance of the nuclear matrix are also raised by the apparent absence of a matrix in certain types of cell, including some cancer cell lines.

Perhaps the most persuasive evidence for the existence of a fibrous matrix in the nuclei of most cells is the presence in chromosomal DNA of nucleotide sequences that bind to the matrix proteins. These segments have been given various names in the past but are now commonly called scaffold/matrix attachment regions (S/MARs). Individual S/MARs are 100–1000 bp in length, and although they do not contain any diagnostic sequences, they are AT-rich and can be recognized in genome sequences. There appear to be over 7350 S/MARs in the Drosophila melanogaster genome and 1358 on Arabidopsis thaliana chromosome 4. There are probably 80,000–90,000 in the human genome. Their positions coincide with, or are close to, replication origins and to transcription start points for protein-coding genes, in particular those with a high rate of transcription.

Pairs of S/MARs are thought to delineate each of the euchromatin loops that have been observed with the electron microscope (Figure 10.6). Some S/MARs appear to be used in all cells and are thought to contribute to the underlying structural organization of the nucleus. Other S/MARs are used only in particular cell types, and some form or break their attachments with the matrix proteins in response to extracellular signals such as the presence of a hormone or growth factor. The implication is that the attachments made by these S/MARs are involved in determining and possibly controlling the ability of transcription proteins to gain access to the genes contained in the chromatin loops, by merging adjacent loops to form larger ones when the genes contained in those loops are active and by condensing active loops into smaller ones when their genes are switched off.

Each chromosome has its own territory within the nucleus

Initially it was thought that chromosomes are distributed randomly within a eukaryotic nucleus. We now know that this view is incorrect and that each chromosome occupies its own space or territory. These can be visualized by chromosome painting, which is a version of fluorescent in situ hybridization (FISH; Section 3.5) in which the hybridization probe is a mixture of DNA molecules, with those specific for different regions of a single chromosome all carrying the same fluorescent label. Each chromosome can therefore be painted a different color. When applied to interphase nuclei, chromosome painting reveals territories occupied by individual chromosomes (Figure 10.7). These territories take up the

Figure 10.6 A scheme for organization of euchromatin in the nucleus. Loops of euchromatin DNA, 4–200 kb in length and predominantly in the form of the 30 nm chromatin fiber, are shown attached to the nuclear matrix by AT-rich regions called scaffold/matrix attachment regions (S/MARs).

Figure 10.7 Chromosome territories. (A) Human chromosomes in an interphase nucleus, visualized by hybridization labeling with probes of different colors. The black areas are nucleoli. (B) Interpretation of the image in (A), giving the identity of each chromosome. (From Speicher MR & Carter NP [2005] Nat. Rev. Genet. 6:782–792. With permission from Macmillan Publishing Ltd.)
majority of the space within the nucleus but are separated from one another by **nonchromatin regions**, within which the enzymes and other proteins involved in expression of the genome are located.

Chromosome territories appear to be fairly static within an individual nucleus. This has been concluded from experiments in which CENP-B proteins, which are components of centromeres (Section 7.1), are labeled with green fluorescent protein and the locations of these proteins, and hence of the centromeres, observed over time. On the whole, individual centromeres remain stationary throughout the cell cycle, though there are occasional bursts of relatively slow movement. Although fairly static during the lifetime of a cell, most studies suggest that the relative positioning of territories is not retained after cell division, as different patterns are observed in the nuclei of daughter cells. There may, however, be certain constraints on territory locations, as it has been known for several years that chromosome **translocations**, which result in a segment of one chromosome becoming attached to another chromosome, are more frequent between certain pairs than others. For example, a translocation between human chromosomes 9 and 22, resulting in the abnormal product called the **Philadelphia chromosome**, is a common cause of chronic myeloid leukemia (Figure 10.8). The repeated occurrence of the same translocation suggests that the territories of the interacting pair of chromosomes are frequently close to one another in the nucleus. There is also evidence that, at least in some organisms, certain chromosomes preferentially occupy territories close to the periphery of the nucleus. Relatively little genome expression occurs in this region, and it is often here that those chromosomes containing few active genes are found, for example, the macrochromosomes of the chicken genome (Section 7.1).

The positioning of active genes within individual chromosome territories is a further topic of debate. At one time it was thought that active genes were located on the surface of a territory, adjacent to the nonchromatin region and hence within easy reach of the enzymes and proteins involved in gene transcription. This view is now being questioned, partly as a result of experiments that have shown that RNA transcripts are distributed within territories as well as on their surfaces. More refined microscopic examination has shown that channels run through chromosome territories, linking different parts of the nonchromatin regions and providing a means by which the transcription machinery can penetrate into the internal parts of these territories (Figure 10.9).

Each chromosome comprises a series of topologically associated domains

Some of the most powerful methods that are being used to study the internal nuclear architecture involve treating nuclei with formaldehyde, in order to induce formation of covalent bonds between the DNA and proteins in strands of chromatin that are close to one another, so those adjacent strands become linked together (Figure 10.10). The resulting network is treated with a restriction endonuclease to break it into fragments, each of which contains two linked pieces of DNA, one from each of the two regions of the genome that were adjacent in the nucleus. DNA ligase is then added to join the ends of the fragments, and the cross-links are disrupted by heating to 70°C. The end product of these manipulations will be a circle of DNA, comprising the two DNA fragments that originally were located close together in the nucleus. The method is called **chromosome conformation capture**, often abbreviated to **3C**. In its original version, 3C was used to determine whether particular interactions which were thought to occur had actually taken place. The pair of fragments present in a circle could therefore be predicted, and a

![Figure 10.8 Products of translocation between human chromosomes 9 and 22.](image)

Normal chromosomes Products of translocation

**Figure 10.8** Products of translocation between human chromosomes 9 and 22. Normal human chromosomes 9 and 22 are shown on the left, and translocation products are shown on the right. The Philadelphia chromosome is the smaller of the two translocation products. Chromosomes 9 and 22 commonly break at the positions indicated. Often the breaks are correctly repaired, but occasionally misrepair creates the hybrid products. It is thought that the relatively high frequency with which the Philadelphia chromosome arises indicates that chromosomes 9 and 22 occupy adjacent territories in the human nucleus. The chromosome 9 breakpoint lies within the ABL gene, the product of which is involved in cell signaling (Section 14.1). The translocation attaches a new coding sequence to the start of this gene, resulting in an abnormal protein that causes cell transformation and gives rise to chronic myeloid leukemia.

![Figure 10.9 Channels in chromosome territories.](image)

**Figure 10.9** Channels in chromosome territories. The view on the left shows the original model with each chromosome territory forming a block, implying that active genes are located on the surface of a territory. The view on the right shows the revised model, with channels running through the territories.
polymerase chain reaction (PCR) could be designed to detect whether that circle was present in the mixture resulting from the 3C experiment. More sophisticated versions of 3C utilize next-generation methods to sequence all the circles of linked fragments, enabling all the interactions occurring in a nucleus to be identified.

The first 3C experiments carried out with cross-linked chromatin showed that the vast majority of interactions occur between DNA sequences present in the same chromosome, with relatively few interactions between different chromosomes. These results helped establish that chromosomes occupy individual territories within the nucleus, rather than being intertwined in some way. More detailed 3C projects then revealed that a single chromosome is made up of a series of topologically associated domains (TADs), each comprising a contiguous segment of chromatin folded into coils and loops. This architecture gives rise to many intradomain interactions but few interactions between domains (Figure 10.11).

The *D. melanogaster* genome contains approximately 1000 TADs, ranging in size from 10 to 1000 kb with an average of 100 kb. In humans and mice there are 2000–3000 domains, but these are larger with an average size of about 1 Mb. TADs are present in all metazoan genomes, but they have not yet been identified in plants. The distribution of TADs along a chromosome does not change during the lifetime of a cell and is the same in all cells of a single species. There is even evidence that the domain architecture is conserved in related species, such as among different primates. This conservation reflects the synteny between genomes of related species and tells us that TADs can be looked on as functional domains: each TAD contains a set of genes that are subject to a similar expression pattern, which is set by the identities of regulatory sequences also present in the TAD. When the genes it contains are active, a TAD adopts a relatively open conformation, presumably to aid access of proteins involved in transcription. When its genes are silent, the repressed TAD takes up a more compact organization.

The preceding discussion regarding the structure of TADs, and the changes that occur when the genes they contain are switched on and off, has obvious parallels with the chromatin loops believed to be attached to the nuclear matrix. These loops are smaller than TADs, with an average length of just 16 kb in *D. melanogaster*, and have distinctive AT-rich scaffold/matrix attachment regions at their peripheries, which are not common features of the boundaries between TADs. At present, it seems likely that chromatin loops form parts of the
substructure of individual TADs, but further research is needed to elucidate the exact relationship.

**Insulators mark the boundaries of topologically associated domains**

The boundaries of topologically associated domains are marked by sequences, 1–2 kb in length, called **insulators**. Insulator sequences were first discovered in *Drosophila* and have now been identified in a range of eukaryotes. Examples are the pair of sequences called scs and scs′ (scs stands for specialized chromatin structure), which are located either side of the two hsp70 genes in the fruit fly genome (Figure 10.12).

Insulators maintain the independence of each TAD, preventing cross-talk between adjacent domains. If scs or scs′ is excised from its normal location and reinserted between a gene and the upstream regulatory modules that control expression of that gene, then the gene no longer responds to its regulatory modules: it becomes insulated from their effects (Figure 10.13A). This observation suggests that, in their normal positions, insulators prevent the genes within a domain from being influenced by the regulatory modules present in an adjacent domain (Figure 10.13B). This ability also enables insulators to overcome the **positional effect** that occurs during a gene cloning experiment with a eukaryotic host. The positional effect refers to the variability in gene expression that occurs after a new gene has been inserted into a eukaryotic chromosome. It is thought to result from the random nature of the insertion event, which could deliver the gene to a region of highly packaged chromatin, where it will be inactive, or into an area of open chromatin, where it will be expressed (Figure 10.14A). The ability of scs and scs′ to overcome the positional effect was demonstrated by placing them either side of a fruit fly gene for eye color. When flanked by the insulators, this gene was always highly expressed when it was inserted back into the *Drosophila* genome, in

![Figure 10.12](image1.png)

**Figure 10.12** Insulator sequences in the fruit fly genome. The diagram shows the region of the *Drosophila* genome containing the two hsp70 genes. The insulator sequences scs and scs′ are located on either side of the gene pair. The arrows below the two genes indicate that they lie on different strands of the double helix and so are transcribed in opposite directions.

![Figure 10.13](image2.png)

**Figure 10.13** Insulators maintain the independence of a topologically associated domain. (A) When placed between a gene and its upstream regulatory modules, an insulator sequence prevents the regulatory signals from reaching the gene. (B) In their normal positions, insulators prevent cross-talk between domains, so the regulatory modules of one gene do not influence expression of a gene in a different domain.
contrast to the variable expression that was seen when the gene was cloned without the insulators (Figure 10.14B). The deduction from this and related experiments is that insulators can bring about modifications to chromatin packaging and hence establish a TAD when inserted into a new site in the genome.

How insulators carry out their roles is not yet known, but it is presumed that the functional component is not the insulating sequence itself but the DNA-binding proteins, such as Su(Hw) in *Drosophila* and CTCF in mammals, that attach specifically to insulators. Experiments suggest that, when attached to an insulator, Su(Hw) forms an association with other proteins, such as CP190 and Mod(mdg4), with the latter acting as a molecular glue that holds pairs of insulators together, enabling the DNA between the insulator to loop out and form the TAD (Figure 10.15). In mammals, CTCF might form interactions with ring-shaped protein complexes called cohesins and condensins to achieve the same effect. However, our understanding of the roles of Su(Hw), CTCF, and other insulator binding proteins, like our knowledge of TADs, is still incomplete. There are more than 10,000 CTCF binding sites in a mammalian genome, many of which are not located in sequences that have insulator activity. The insulator binding proteins must therefore play additional roles in genome organization, and their association with insulators might be part of this general role, rather than a specific interaction that provides insulators with their special properties.

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**Figure 10.14 The positional effect.** (A) A cloned gene that is inserted into a region of highly packaged chromatin will be inactive, but one inserted into open chromatin will be expressed. (B) The results of cloning experiments without (red) and with (blue) insulator sequences. When insulators are absent, the expression level of the cloned gene is variable, depending on whether it is inserted into packaged or open chromatin. When flanked by insulators, the expression level is consistently high because the insulators establish a TAD at the insertion site.

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**Figure 10.15 The roles of Su(Hw) and Mod(mdg4) in formation of a topologically associated domain.** Su(Hw) proteins attach to the insulator sequences at either side of the region of chromatin that will become a TAD. Mod(mdg4) proteins bind to Su(Hw) and then act as a molecular glue that holds the two insulators together, establishing the TAD as a loop of chromatin.
10.2 NUCLEOSOME MODIFICATIONS AND GENOME EXPRESSION

Section 10.1 introduced the notion that chromatin structure influences genome expression: the degree of chromatin packaging displayed by a particular segment of a chromosome determines whether RNA polymerase and the other transcription proteins can gain access to the genes within that segment. But so far we have simply examined the structure of chromatin and not asked how changes in that structure are brought about. This is the next issue that we will explore.

Acetylation of histones influences many nuclear activities including genome expression

Nucleosomes appear to be the primary determinants of chromatin structure in eukaryotes, the precise chemical structure of the histone proteins contained within nucleosomes being the major factor influencing the degree of packaging displayed by a segment of chromatin. Histones can undergo various types of modification. The best-studied of these is histone acetylation, the attachment of acetyl groups to lysine amino acids in the N-terminal regions of each of the core molecules (Figure 10.16). These N-termini form tails that protrude from the nucleosome core octamer (see Figure 7.4), and their acetylation reduces the affinity of the histones for DNA and possibly also reduces the interaction between individual nucleosomes, destabilizing the 30 nm chromatin fiber. The histones in heterochromatin are generally unacetylated whereas those in active domains are acetylated, a clear indication that this type of modification is linked to DNA packaging.

The relevance of histone acetylation to genome expression was underlined in 1996 when, after several years of trying, the first examples of histone acetyltransferases (HATs), the enzymes that add acetyl groups to histones, were identified. It was realized that some proteins that had already been shown to have important influences on genome expression had HAT activity. For example, one of the first HATs to be discovered, the Tetrahymena protein called p55, was shown to be a homolog of a yeast protein, GCN5, that was known to activate assembly of the transcription initiation complex (Section 12.2). Similarly, the mammalian protein called p300/CBP, which had been ascribed a clearly defined role in activation of a variety of genes, was found to be a HAT. These observations, plus the demonstration that different types of cells display different patterns of histone acetylation, underline the prominent role that histone acetylation plays in regulating genome expression.

Individual HATs can acetylate histones in the test tube but most have negligible activity on intact nucleosomes, indicating that, in the nucleus, HATs almost certainly do not work independently but instead form multiprotein complexes, such as the SAGA and ADA complexes of yeast and the TFTC complex of humans. These complexes are typical of the large multiprotein structures that catalyze and

Figure 10.16 Positions at which acetyl groups are attached to lysines within the N-terminal regions of the four core histones. The sequences shown are those of human histones. Each sequence begins with the N-terminal amino acid.
regulate the various steps in genome expression, many examples of which we will discuss in the next few chapters. SAGA, for example, comprises at least 18 proteins with a combined molecular mass of 1.8 million. The complex forms a particle with dimensions $27 \times 17 \times 13$ nm, which means that it is larger than the nucleosome core octamer, which with its associated DNA has a diameter of 11 nm, and it is comparable in one dimension with the 30 nm chromatin fiber. As well as GCN5, the protein with HAT activity, the SAGA complex contains a set of proteins related to the TATA-binding protein (TBP), which initiates the process by which a gene is transcribed (Section 12.2), as well as five TBP-associated factors (TAFs) that help TBP fulfill its role. The complexity of SAGA and the other HAT complexes, and the presence within these complexes of proteins with distinct roles in the initiation of gene expression, indicates that the individual events that result in a gene becoming active are intimately linked, with histone acetylation being an integral part, but just one part, of the overall process.

There are at least five different families of HAT proteins. The GCN5-related acetyltransferases, or GNATs, which are components of the SAGA, ADA, and TFTC complexes, are clearly associated with activation of gene transcription but also are involved in the repair of some types of damaged DNA, in particular double-strand breaks and lesions resulting from ultraviolet irradiation (Section 16.1). A second family of HATs, called MYST after the initial letters of four of the proteins in this family, is similarly involved in transcription activation and DNA repair and has also been implicated in control of the cell cycle, though this may simply be another aspect of the DNA repair function, as the cell cycle stalls if the genome is extensively damaged (Section 15.5). Different complexes appear to acetylate different histones and some can also acetylate other proteins involved in genome expression, such as the general transcription factors TFIIE and TFIIF, which we will meet in Section 12.2. HATs are therefore emerging as versatile proteins that may have diverse functions in expression, replication, and maintenance of the genome.

**Histone deacetylation represses active regions of the genome**

Gene activation must be reversible, otherwise genes that become switched on would remain permanently active. Hence it is no surprise that there is a set of enzymes that can remove acetyl groups from histone tails, overturning the transcription-activating effects of the HATs described above. This is the role of the **histone deacetylases (HDACs)**. The link between HDAC activity and gene silencing was established in 1996, when mammalian HDAC1, the first of these enzymes to be discovered, was shown to be related to a yeast protein called Rpd3, which was known to be a repressor of transcription. The link between histone deacetylation and repression of transcription was therefore established in the same way as the link between acetylation and activation: by showing that two proteins that were initially thought to have different activities are in fact related. These are good examples of the value of homology analysis in studies of gene and protein function (Section 6.1).

HDACs, like HATs, are contained in multiprotein complexes. One of these is the mammalian Sin3 complex, which comprises at least seven proteins, including HDAC1 and HDAC2, along with others that do not have deacetylase activity but provide ancillary functions essential to the process. Examples of ancillary proteins are RBBP4 and RBBP7, which are members of the Sin3 complex and are thought to contribute the histone-binding capability. RBBP4 and RBBP7 were first recognized through their association with the retinoblastoma protein, which controls cell proliferation by inhibiting expression of various genes until their activities are required and which, when mutated, leads to cancer. This link between Sin3 and a protein implicated in cancer provides a powerful argument for the importance of histone deacetylation in gene silencing. Other deacetylation complexes include NuRD in mammals, which combines HDAC1, HDAC2, RBBP4, and RBBP7 with a different set of ancillary proteins, and yeast Sir2, which is different from other
HDACs in that it has an energy requirement. The distinctive features of Sir2 show that HDACs are more diverse than originally realized, possibly indicating that novel roles for histone deacetylation are waiting to be discovered.

Studies of HDAC complexes are beginning to reveal links between the different mechanisms for genome activation and silencing. Both Sin3 and NuRD contain proteins that bind to methylated DNA (Section 10.3), and NuRD contains proteins that are very similar to components of the nucleosome remodeling complex SWI/SNF (which will be discussed later). NuRD does in fact act as a typical nucleosome remodeling machine in vitro. Further research will almost certainly unveil additional links between what we currently look on as different types of chromatin modification systems but which in reality may simply be different facets of a single grand design.

**Acetylation is not the only type of histone modification**

Lysine acetylation/deacetylation is the best-studied form of histone modification, but it is by no means the only type. Three other kinds of covalent modification are known to occur:

- Methylation of lysine and arginine residues. Methylation was originally thought to be irreversible and hence responsible for permanent changes to chromatin structure. This view has been challenged by the discovery of enzymes that demethylate lysine and arginine residues, but it is still accepted that the effects of methylation are relatively long-term.

- Phosphorylation of serine, threonine, and tyrosine residues.

- Ubiquitination of lysine residues at the C-terminal regions of H2A and H2B. This modification involves addition of the small, common (ubiquitous) protein called ubiquitin or a related protein, rather unhelpfully called small ubiquitin-related modifier or SUMO.

- Citrullination in the N-terminal regions of H3 and H4. Citrullination is the conversion of arginine to the related amino acid called citrulline, by replacement of the terminal =NH group of the arginine side chain with an =O group.

As with acetylation, these other types of modification influence chromatin structure and have a significant impact on cellular activity. For example, phosphorylation of serine-1 of histone H2A has been associated with transcription repression, and ubiquitination of histone H2B is part of the general role that ubiquitin plays in control of the cell cycle. The effects of methylation of a pair of lysine amino acids at the fourth and ninth positions from the N-terminus of histone H3 are particularly interesting. Methylation of lysine-9 forms a binding site for the HP1 protein, which induces chromatin packaging and silences gene expression, but this event is blocked by the presence of two or three methyl groups attached to lysine-4. Methylation of lysine-4 therefore promotes an open chromatin structure and is associated with active genes. In the part of human chromosome 11 that contains the β-globin genes, and probably elsewhere, lysine-4 methylation also prevents binding of the NuRD deacetylase to histone H3, ensuring that this histone remains acetylated. Lysine-4 methylation may therefore work hand-in-hand with histone acetylation to activate regions of chromatin.

Altogether, at least 80 sites in the four core histones are known to be subject to covalent modification (Figure 10.17). Our growing awareness of the variety of histone modifications that occur, and of the way in which different modifications work together, has led to the suggestion that there is a histone code, by which the pattern of chemical modifications specifies which regions of the genome are expressed at a particular time and dictates other aspects of genome biology, such as the repair of damaged sites and coordination of genome replication with the
cell cycle (Table 10.1). This idea is still unproven, but it is clear that the pattern of specific histone modifications within the genome is linked closely to gene activity. Studies of human chromosomes 21 and 22, for example, have shown that regions within these chromosomes where lysine-4 of histone H3 is trimethylated and lysine-9 and lysine-14 are acetylated correspond to the transcription start points for active genes and that dimethylated lysine-4 is also sometimes found in these regions (Figure 10.18). As with all aspects of chromatin modification, the key question is to distinguish cause and effect: are these patterns of histone modification the reason why these particular genes are active or merely a byproduct of the processes responsible for their activation?

**Nucleosome repositioning also influences gene expression**

A second type of chromatin modification that can influence genome expression is nucleosome remodeling. This term refers to the modification or repositioning of nucleosomes within a short region of the genome, so that DNA-binding proteins can gain access to their attachment sites. This does not appear to be an essential requirement for transcription of all genes, and in at least a few cases it is possible for a protein that switches on gene expression to achieve its effect either by binding to the surfaces of nucleosomes or interacting with the linker DNA without affecting the nucleosome positions. In other examples, repositioning of nucleosomes has been clearly shown to be a prerequisite for gene activation.
This is apparent from the detection of DNase I hypersensitive sites in the vicinity of genes that are being actively transcribed. For example, the DNA just upstream of the human β-globin gene cluster contains five DNase I hypersensitive sites (Figure 10.19). Each of these is a short region of DNA that is cleaved by DNase I more easily than other parts of the gene cluster. These sites are thought to coincide with positions where nucleosomes have been modified or are absent and which are therefore accessible to binding proteins that attach to the DNA.
Similarly, activation of the hsp70 gene of D. melanogaster, which codes for a protein involved in folding other proteins (Section 13.4), is associated with the creation of a DNase I hypersensitive region upstream of the gene. Altogether, there are almost 2.9 million DNase I hypersensitive sites in the human genome. The vast majority of these are detectable only in those tissues within which the genes whose positions they mark are active.

Unlike acetylation and the other chemical modifications described previously, nucleosome remodeling does not involve covalent alterations to histone molecules. Instead, remodeling is induced by an energy-dependent process that weakens the contact between the nucleosome and the DNA with which it is associated. Three distinct types of change can occur (Figure 10.20):

- Remodeling, in the strict sense, involves a change in the structure of the nucleosome but no change in its position. The nature of the structural change is not known, but when remodeling is induced in vitro, the outcome is a doubling in size of the nucleosome and an increased DNase sensitivity of the attached DNA.

- Sliding, or cis-displacement, physically moves the nucleosome along the DNA.

- Transfer, or trans-displacement, results in the nucleosome being transferred to a second DNA molecule or to a nonadjacent part of the same molecule.

As with histone acetyltransferases, the proteins responsible for nucleosome remodeling work together in large complexes. One of these is SWI/SNF, which is made up of at least 10–12 proteins and is present in many eukaryotes. The subunit proteins include some that have DNA-binding capability and others that detect the presence of histone acetylations and methylations and which presumably direct the complex to active parts of the genome. Other subunits are able to add ubiquitin tags to lysine-120 of histone H2B (see Figure 10.17), a modification that is associated with highly transcribed regions. SWI/SNF therefore combines the two activities—nucleosome repositioning and histone modification—that are currently looked on as central to genome activation.

Figure 10.19 DNase I hypersensitive sites upstream of the human β-globin gene cluster. A series of hypersensitive sites is located in the 20 kb of DNA upstream of the start of the β-globin gene cluster. Additional hypersensitive sites are seen immediately upstream of each gene, at the position where the RNA polymerase attaches to the DNA. These hypersensitive sites are specific to different developmental stages and are seen only during the phase of development when the adjacent gene is active.

Figure 10.20 Nucleosome remodeling, sliding, and transfer.
10.3 DNA MODIFICATION AND GENOME EXPRESSION

Important alterations in genome activity can also be achieved by making chemical changes to the DNA itself. These changes are associated with the semipermanent silencing of regions of the genome, possibly entire chromosomes, and often the modified state is inherited by the progeny arising from cell division. The modifications are brought about by DNA methylation.

Genome silencing by DNA methylation

In eukaryotes, cytosine bases in chromosomal DNA molecules are sometimes changed to 5-methylcytosine by the addition of methyl groups by enzymes called DNA methyltransferases (Figure 10.21). Cytosine methylation is relatively rare in lower eukaryotes, but in vertebrates up to 10% of the total number of cytosine bases in a genome are methylated, and in plants the figure can be as high as 30%. The methylation pattern is not random, instead being limited to the cytosine in some copies of the sequences 5′-CG-3′ and, in plants, 5′-CNG-3′. Two types of methylation activity have been distinguished (Figure 10.22). The first is maintenance methylation, which, following genome replication, is responsible for adding methyl groups to the newly synthesized strand of DNA at positions opposite methylated sites on the parent strand. The maintenance activity therefore ensures that the two daughter DNA molecules retain the methylation pattern of the parent molecule, which means that the pattern can be inherited after cell division. The second activity is de novo methylation, which adds methyl groups at totally new positions and so changes the pattern of methylation in a localized region of the genome.

Both maintenance and de novo methylation result in repression of gene activity. This has been shown by experiments in which methylated or unmethylated genes have been introduced into cells by cloning and their expression levels measured: expression does not occur if the DNA sequence is methylated. The link with gene expression is also apparent when the methylation patterns in chromosomal DNAs are examined, these showing that active genes are located in unmethylated regions. For example, in humans, 40–50% of all genes are located close to CpG islands (Section 5.1), with the methylation status of the CpG island reflecting the expression pattern of the adjacent gene. Housekeeping genes, which are expressed in all tissues, have unmethylated CpG islands, whereas the CpG islands associated with tissue-specific genes are unmethylated only in those tissues in which the gene is expressed. Note that because the methylation pattern is maintained after cell division, information specifying which genes should be expressed is inherited by the daughter cells, ensuring that in a differentiated tissue the appropriate pattern of gene expression is retained even though the cells in the tissue are being replaced and/or added to by new cells.

The importance of DNA methylation is underlined by studies of human diseases. The syndrome called ICF (immunodeficiency, centromere instability, and facial anomalies), which as the name suggests has wide-ranging phenotypic effects, is associated with undermethylation of various genomic regions and is caused by a mutation in the gene for DNA methyltransferase 3b, one of the enzymes responsible for de novo methylation. The opposing situation, hypermethylation, is seen within the CpG islands of genes that exhibit altered expression patterns in certain types of cancer, although in these cases the abnormal methylation could equally well be a result rather than the cause of the disease state.

How methylation influences genome expression was a puzzle for many years. Now it is known that methyl-CpG-binding proteins (MeCPs) are components of both the Sin3 and NuRD histone deacetylase complexes. This discovery has led to a model in which methylated CpG islands are the target sites for attachment of HDAC complexes that modify the surrounding chromatin in order to silence the adjacent genes (Figure 10.23).
Methylation is involved in genomic imprinting and X inactivation

Further evidence, if it is needed, of the link between DNA methylation and genome silencing is provided by two intriguing phenomena called **genomic imprinting** and **X inactivation**.

Genomic imprinting is a relatively uncommon but important feature of mammalian genomes in which only one of a pair of genes, present on homologous chromosomes in a diploid nucleus, is expressed while the second is silenced by methylation. It also occurs in some insects (though apparently not *D. melanogaster*) and some plants. It is always the same member of a pair of genes that is imprinted and hence inactive: for some genes this is the version inherited from the mother, and for other genes it is the paternal version. Almost 200 genes in humans and mice have been shown to display imprinting, including both protein-coding genes and genes specifying noncoding RNAs. Imprinted genes are distributed around the genome but tend to occur in clusters. For example, in humans there is a 2.2 Mb segment of chromosome 15 within which there are at least ten imprinted genes, and a smaller 1 Mb region of chromosome 11 contains at least eight imprinted genes.

An example of an imprinted gene in humans is *Igf2*, which codes for a growth factor, a protein involved in signaling between cells (Section 14.1). Only the paternal gene is active (Figure 10.24), because on the chromosome inherited from the mother, various segments of DNA in the region of *Igf2* are methylated, preventing expression of this copy of the gene. A second imprinted gene, *H19*, is located some 90 kb away from *Igf2*, but the imprinting is the other way around: the maternal version of *H19* is active and the paternal version is silent. Imprinting is controlled by **imprint control elements**, DNA sequences that are found within a few kilobases of clusters of imprinted genes. These centers mediate the methylation of the imprinted regions, but the mechanism by which they do this has not yet been described in detail. There is also uncertainty regarding the function of imprinting. One possibility is that it has a role in development, because artificially created parthenogenetic mice, which have two copies of the maternal genome, fail to develop properly. In addition, several of the genetic diseases that are associated with dysfunctional imprinting, such as Prader–Willi syndrome and Angelman syndrome, are characterized by developmental abnormalities. Individual genes displaying imprinting have been implicated in physiological functions as diverse as body temperature maintenance and behaviors such as sleep and maternal care. More subtle explanations of the role of imprinting, based on the evolutionary conflicts between the males and females of a species, have also been proposed.

X inactivation is less enigmatic. This is a special form of imprinting that leads to almost total inactivation of one of the X chromosomes in a female mammalian cell. It occurs because females have two X chromosomes whereas males have only one. If both of the female X chromosomes were active, then proteins coded by genes on the X chromosome might be synthesized at twice the rate in females compared with males. To avoid this undesirable state of affairs, one of the female X chromosomes is silenced and is seen in the nucleus as a condensed structure called the **Barr body**, which is composed entirely of heterochromatin. Most of the genes on the inactivated X chromosome become silenced but, for reasons that are unknown, some 20% escape the process and remain functional.

Silencing occurs early in embryo development and is controlled by the X inactivation center (Xic), a discrete region present on each X chromosome. In a cell
undergoing X inactivation, the inactivation center on one of the X chromosomes initiates the formation of heterochromatin, which spreads out from the nucleation point until the entire chromosome is affected, with the exception of a few short segments containing those genes that remain active. The process takes several days to complete. The exact mechanism is not understood but it involves, although is not entirely dependent upon, a gene called Xist, located in the inactivation center, which is transcribed into a 17 kb noncoding RNA, copies of which coat the chromosome as heterochromatin is formed. At the same time, various histone modifications occur. Lysine-9 of histone H3 becomes methylated (recall that this modification is associated with genome inactivation; see Table 10.1), histone H4 becomes deacetylated (as usually occurs in heterochromatin), and the histone H2A molecules are replaced with a special histone, macroH2A1. Certain DNA sequences become hypermethylated by DNA methyltransferase 3a, although this appears to occur after the inactive state has been set up. X inactivation is heritable and is displayed by all cells descended from the initial one within which the inactivation took place.

In a normal diploid female, one X chromosome is inactivated and the other remains active. Remarkably, in diploid females with unusual sex chromosome constitutions, the process still results in just a single X chromosome remaining active. For example, in those rare individuals that possess just a single X chromosome, no inactivation occurs, and in those individuals with an XXX karyotype, two of the three X chromosomes are inactivated (Figure 10.25A). This means that there must be a mechanism by which the X chromosomes in the nucleus are counted and the appropriate number are inactivated. In fact, this mechanism does not simply count the X chromosomes; it also counts the autosomes and compares the two numbers. This is evident because if the cell has four X chromosomes but is otherwise diploid, then three X chromosomes are inactivated, but if it is tetraploid (that is, it has four X chromosomes and also four copies of each autosome), then two X chromosomes are inactivated (Figure 10.25B). How the cell counts its chromosomes has puzzled cytogeneticists for many years and continues to puzzle us, but the most recent research ascribes a role to a second gene in the inactivation center, called Tsix, the transcript of which is an antisense version of the Xist RNA. Overexpression of Tsix results in inactivation of an incorrect number of chromosomes, and it seems likely that this effect occurs via base pairing between the Tsix and Xist RNAs, preventing the latter from initiating the inactivation process. According to this model, the number of X chromosomes that are inactivated will depend on the amount of Tsix RNA that is made, which in turn will depend on the X chromosome copy number and might possibly be modulated by autosomal genes that also exert their effect in a copy-number-dependent manner.

**SUMMARY**

- The nuclear environment has a substantial and important impact on expression of the genome.
- A eukaryotic nucleus has a highly ordered internal architecture that includes structures associated with rRNA processing, mRNA splicing, and synthesis of small nuclear and small nucleolar RNAs.
- The most compact form of chromatin is heterochromatin, within which genes are inaccessible and cannot be expressed.
- The nucleus is thought to contain a fibrous matrix to which chromosomal DNA is attached, although the existence of this matrix has been questioned.
- Each chromosome has its own territory within the nucleus, these territories being separated from one another by nonchromatin regions within which the enzymes and other proteins involved in genome expression are located.
• A chromosome is made up of a series of topologically associated domains, each comprising a contiguous segment of chromatin folded into coils and loops.

• Each domain is delimited by a pair of insulators, which maintain the independence of the domain.

• Nucleosomes appear to be the primary determinants of genome activity in eukaryotes, not only by virtue of their positioning on a strand of DNA, but also because the precise chemical structure of the histone proteins contained within nucleosomes is the major factor determining the degree of packaging displayed by a segment of chromatin.

• Acetylation of lysine amino acids in the N-terminal regions of each of the core histones is associated with activation of a region of the genome, and deacetylation leads to genome silencing. Histones can also be modified by methylation, phosphorylation, and ubiquitination.

• There may be a histone code that specifies how particular combinations of nucleosome modifications should be interpreted by the genome.

• Nucleosome repositioning is required for the expression of some, but not all, genes.

• Regions of the genome can be also silenced by DNA methylation, the relevant enzymes possibly working in conjunction with histone deacetylases.

• Methylation is responsible for genomic imprinting, which results in one of a pair of genes on homologous chromosomes becoming silenced, and X inactivation, which leads to the almost complete inactivation of one of the X chromosomes in a female nucleus.

**SHORT ANSWER QUESTIONS**

1. Describe the methods that have been used to examine the structural organization of the nucleus.

2. Distinguish between the terms constitutive heterochromatin and facultative heterochromatin.

3. What has chromosome painting revealed about the location of chromosomes within the nucleus?

4. Translocations occur at a higher frequency between certain pairs of chromosomes. What does this tell us about the distribution of chromosomes in the nucleus?

5. How are topologically associated domains identified in a eukaryotic chromosome?

6. What is the explanation of the positional effect that sometimes occurs when a gene is cloned in a eukaryotic host?

7. What are insulator sequences and what unique properties do they possess?

8. Give examples of histone acetyltransferases and describe the role of these enzymes in nucleosome modification.

9. What is the role of histone deacetylases in the regulation of genome expression?

10. Explain what is meant by the term “histone code.”

11. Why is DNase I used to study changes in chromatin structure? What does the susceptibility of DNA to cleavage by DNase I tell us about gene expression?

12. How does DNA methylation influence genome activity?
IN-DEPTH PROBLEMS

1. To what extent can it be assumed that the picture of nuclear architecture built up by modern electron microscopy is an accurate depiction of the actual structure of the nucleus, as opposed to an artifact of the methods used to prepare cells for examination?

2. In many areas of biology it is difficult to distinguish between cause and effect. Evaluate this issue with regard to nucleosome remodeling and genome expression: does nucleosome remodeling cause changes in genome expression or is remodeling the effect of these expression changes?

3. Explore and assess the histone code hypothesis.

4. Maintenance methylation ensures that the pattern of DNA methylation displayed by two daughter DNA molecules is the same as the pattern on the parent molecule. In other words, the methylation pattern, and the information on gene expression that it conveys, is inherited. Other aspects of chromatin structure might also be inherited in a similar way. How do these phenomena affect the Mendelian view that inheritance is specified by genes?

5. What might be the means by which the numbers of X chromosomes and autosomes in a nucleus are counted so that the appropriate number of X chromosomes can be inactivated?

FURTHER READING

The internal structure of the nucleus and the matrix controversy

Chromosome territories

Chromosome domains

Covalent modification of histones


**DNA methylation, imprinting, and X inactivation**


**Nucleosome remodeling**

In Chapter 10, we learned that the activity of individual genes is influenced by the degree of packaging exhibited by the chromatin domain in which those genes are contained and also by the precise positioning of the nucleosomes in the vicinity of the genes. Although our focus was on higher-order structures such as chromatin and nucleosomes, we recognized that, at a more basic level, the accessibility of the genome is controlled by the interactions between histone proteins and the DNA to which they are attached. These interactions are an example of the central role that DNA-binding proteins play in genome expression. Histones are DNA-binding proteins, as are several of the proteins responsible for transcription of individual genes. There are also DNA-binding proteins that are involved in DNA replication, repair, and recombination, as well as a large group of related proteins that bind to RNA rather than DNA. Many DNA-binding proteins recognize specific nucleotide sequences and bind predominantly to these target sites, whereas others, such as histones, lack sequence specificity and attach at various positions in the genome.

In this chapter we will examine the special structural features of DNA-binding proteins and explore how these features enable a DNA-binding protein to attach to the genome, focusing in particular on the way in which a sequence-specific binding protein recognizes its attachment sites. Understanding the nature of and effects of these DNA-protein interactions is an important and active area of genome research, and we will therefore begin by studying the methods used to elucidate the structures of DNA-binding proteins and to identify their binding positions on a DNA molecule.

### 11.1 Methods for Studying DNA-Binding Proteins and Their Attachment Sites

The methods that are used to study the interactions between DNA-binding proteins and the genome fall into two categories:

- Various technologies, most importantly **X-ray crystallography** and **nuclear magnetic resonance (NMR) spectroscopy**, that are used to study the structures of DNA-binding proteins and in particular to identify the structural features that enable a protein to make sequence-specific attachments to a DNA molecule.

- Methods that identify, with varying degrees of accuracy, the positions on a DNA molecule at which a DNA-binding protein attaches.

**X-ray crystallography provides structural data for any protein that can be crystallized**

Once a DNA-binding protein has been purified, attempts can be made to determine its structure, either in isolation or attached to its binding site. This enables the conformation of the DNA-binding part of the protein to be studied and allows the identity and nature of contacts with the DNA molecule to be elucidated. Two techniques, X-ray crystallography and NMR spectroscopy, are central to this area of research.
X-ray crystallography, which is a long-established technique whose pedigree stretches back to the late nineteenth century, is based on X-ray diffraction. X-rays have very short wavelengths, between 0.01 and 10 nm, which is 4000 times shorter than the wavelength of visible light and is comparable with the spacings between atoms in chemical structures. When a beam of X-rays is directed onto a crystal, some of the X-rays pass straight through, but others are diffracted and emerge from the crystal at a different angle from which they entered. If the crystal is made up of many copies of the same molecule, all positioned in a regular array, then different X-rays are diffracted in similar ways, resulting in overlapping circles of diffracted waves that interfere with one another. An X-ray-sensitive photographic film or electronic detector placed across the beam reveals a series of spots, an X-ray diffraction pattern, from which the structure of the molecule in the crystal can be deduced (Figure 11.1). The relative positioning of the spots indicates the arrangement of molecules in the crystal, and the relative intensities of the spots provide information on the structure of the molecule. The more complex the molecule, the greater the number of spots and the larger the number of comparisons that must be made between them. Computational help is therefore required for all but the simplest molecules.

If successful, analysis of an X-ray diffraction pattern enables an electron density map to be constructed (Figure 11.2). For a protein, this provides a chart of the folded polypeptide from which the positioning of structural features such as α-helices and β-sheets can be determined. If the map is sufficiently detailed, the R groups of individual amino acids in the polypeptide can be identified and their orientations relative to one another can be established, allowing deductions to be made about hydrogen bonding and other chemical interactions occurring within the protein structure. In the most successful projects, a resolution of 0.1 nm is possible, which means that structures just 0.1 nm apart can be distinguished.
In proteins, most carbon–carbon bonds are 0.1–0.2 nm in length and carbon–hydrogen bonds are 0.08–0.12 nm. This means that, at 0.1 nm resolution, a very detailed three-dimensional model of the protein can be constructed. The one limitation with X-ray crystallography is that the protein must be crystallized before its structure can be studied by this method. For many proteins this is not a problem, as good quality crystals can be obtained from a supersaturated solution. Other proteins, especially membrane proteins, which have external hydrophobic regions, are more difficult or even impossible to crystallize.

**NMR spectroscopy is used to study the structures of small proteins**

Like X-ray crystallography, NMR is a long-established technique that traces its origins to the early part of the twentieth century, first being described in 1936. The principle of the technique is that rotation of an atomic nucleus generates a magnetic moment. When it is placed in an applied electromagnetic field, the spinning nucleus is oriented in one of two ways, called $\alpha$ and $\beta$ (Figure 11.3); the $\alpha$-orientation (which is aligned with the magnetic field) has slightly lower energy. In NMR spectroscopy, the magnitude of this energy separation is determined by measuring the frequency of electromagnetic radiation needed to induce the transition from $\alpha$ to $\beta$, this value being described as the resonance frequency of the nucleus being studied. The critical point is that although each type of nucleus has its own specific resonance frequency, the measured frequency is often slightly different from the standard value (typically by less than 10 parts per million) because electrons in the vicinity of the rotating nucleus shield it to a certain extent from the applied magnetic field. This chemical shift (the difference between the observed resonance energy and the standard value for the nucleus being studied) enables the chemical environment of the nucleus to be inferred and hence provides structural information. Particular types of analysis (called correlation spectroscopy, COSY, and total correlation spectroscopy, TOCSY) enable atoms linked by chemical bonds to the spinning nucleus to be identified. Other analyses (for example, nuclear Overhauser effect spectroscopy or NOESY) identify atoms that are close to the spinning nucleus in space but not directly connected to it.

To be suitable for NMR, a chemical nucleus must have an odd number of protons and/or neutrons, otherwise it will not spin when placed in an electromagnetic field. Most protein NMR projects are $^1$H studies, where the aim is to identify the chemical environments and covalent linkages of every hydrogen atom and from this information to infer the overall structure of the protein. These studies are frequently supplemented by analyses of substituted proteins in which at least some of the carbon and/or nitrogen atoms have been replaced with the rare isotopes $^{13}$C and $^{15}$N, these also giving good results with NMR.

When successful, NMR results in the same level of resolution as X-ray crystallography and so provides very detailed information on protein structure. The main advantage of NMR is that it works with molecules in solution and so avoids the problems that sometimes occur when attempting to obtain crystals of a protein for X-ray analysis. Solution studies also offer greater flexibility if the aim is to examine changes in protein structure, as occur during protein folding or in response to addition of a substrate. The disadvantage of NMR is that it is suitable only for relatively small proteins. There are several reasons for this, one being the need to identify the resonance frequencies for each, or as many as possible, of the $^1$H or other nuclei being studied. This depends on the various nuclei having different chemical shifts so that their frequencies do not overlap. The larger the protein, the greater the number of nuclei and the greater the chances that frequencies overlap and structural information is lost. Although this limits the applicability of NMR, the technique is still very valuable. There are many interesting proteins that are small enough to be studied by NMR, and important information can also be obtained by structural analysis of peptides, which, although they are not complete proteins, can act as models for aspects of protein activity such as nucleic acid binding.
Many DNA-binding proteins, such as histones, attach to DNA molecules of any nucleotide sequence, but others display sequence specificity and form stable attachments only at certain positions in the genome. A complement to structural studies of DNA-binding proteins is therefore provided by methods that enable the attachment sites for these sequence-specific proteins to be identified. Many of these methods were invented during the pre-genome era and are designed to identify protein binding sites in cloned fragments of DNA up to about 2 kb in length. The stimulus for development of these methods was the discovery that many transcription factors (proteins that control gene transcription) exert their effects by binding to sequence-specific attachment sites located immediately upstream of their target genes (Figure 11.4). This means that the sequence of a newly discovered gene, if it includes the upstream region, provides immediate access to the binding sites for at least some of the proteins that control its expression.

The various methods that are available enable the protein binding sites in a cloned DNA fragment to be located with different degrees of accuracy. The least accurate of these methods, but the easiest to carry out, makes use of the substantial difference between the electrophoresis properties of a naked DNA fragment and one that carries a bound protein. Recall that DNA fragments are separated by agarose gel electrophoresis because smaller fragments migrate through the porelike structure of the gel more quickly than larger fragments (Section 2.1). If a DNA fragment has a protein bound to it, then its mobility through the gel will be impeded, and the DNA–protein complex will form a band at a position nearer to the starting point (Figure 11.5). This is called gel retardation. In practice, the technique is carried out with a collection of restriction fragments that span the region thought to contain a protein binding site. The digest is mixed with an extract of nuclear proteins (if a eukaryote is being studied), and retarded fragments are identified by comparing the banding pattern obtained after electrophoresis with the pattern for restricted fragments that have not been mixed with proteins. A nuclear extract is used because at this stage of the project the DNA-binding protein has not usually been purified. However, if the protein is available, then the experiment can be carried out just as easily with the pure protein as with a mixed extract.

Protection assays pinpoint binding sites with greater accuracy

Gel retardation gives a general indication of the location of a protein binding site in a DNA sequence, but it does not pinpoint the site with great accuracy. Often the retarded fragment is several hundred base pairs in length, compared with the expected length of the binding site of a few tens of base pairs at most, and there is no indication of where within the retarded fragment the binding site lies. Also, if the retarded fragment is long, then it might contain separate binding sites for several proteins, or if the retarded fragment is quite small, then the binding site might include nucleotides on adjacent fragments, ones that on their own do not form a stable complex with the protein and so do not lead to gel retardation. Gel retardation studies are therefore a starting point, but other techniques are needed to provide more accurate information.

Modification protection assays can take over where gel retardation leaves off. The basis of these techniques is that if a DNA molecule carries a bound protein, then part of its nucleotide sequence will be protected from modification. Any guanines protected by the bound protein will not be methylated by a methylating nuclease, which cleaves all phosphodiester bonds except those protected by the bound protein.
Both methods utilize an experimental approach called footprinting. In nuclease footprinting, the DNA fragment being examined is labeled at one end, complexed with binding protein (as a nuclear extract or as pure protein), and treated with deoxyribonuclease I (DNase I). Normally, DNase I cleaves every phosphodiester bond, leaving only the DNA segment protected by the binding protein. This is not very useful because it can be difficult to sequence such a small fragment. It is quicker to use the more subtle approach shown in Figure 11.6. Nuclease treatment is carried out under limiting conditions, such as a low temperature and/or very little enzyme, so that on average each copy of the DNA fragment suffers a single hit, meaning that it is cleaved at just one position along its length. Although each fragment is cut just once, in the entire population of fragments all bonds are cleaved except those protected by the bound protein. The protein is now removed, the mixture electrophoresed, and the labeled fragments visualized. Each of these fragments has the label at one end and the other end cleaved by DNase I digestion. The result is a ladder of bands corresponding to fragments that differ in length by one nucleotide, with the ladder broken by a blank area in which no labeled bands occur. This blank area, or footprint, corresponds to the positions of the protected phosphodiester bonds, and hence of the bound protein, in the starting DNA.

In the second modification protection method, instead of DNase I digestion, the fragments are treated with limited amounts of dimethyl sulfate so that on average each guanine base is methylated in each fragment (Figure 11.7). Guanines that are protected by the binding protein cannot be modified. After removal of the protein, the DNA is treated with piperidine, which makes single-stranded cuts at the modified nucleotide positions. Electrophoresis is carried out in the presence of a denaturant, such as urea, so that the double-stranded molecules are separated into single strands, some of which have one labeled end and one end created by piperidine nicking. After electrophoresis, the control DNA
strands (those not incubated with the nuclear extract) give a banding pattern that indicates the positions of all guanines in the restriction fragment, and the footprint seen in the banding pattern for the test sample shows which guanines were protected.

**Modification interference identifies nucleotides central to protein binding**

Modification protection should not be confused with **modification interference**, a different technique that provides an extra dimension to the study of protein binding. Modification interference works on the basis that if a nucleotide critical for protein binding is altered, for example, by addition of a methyl group, then binding may be prevented. One of this family of techniques is illustrated in **Figure 11.8**. The DNA fragment, labeled at one end, is treated with the modification reagent, in this case dimethyl sulfate, under limiting conditions so that on average just one guanine per fragment is methylated. Now the binding protein or nuclear extract is added, and the fragments are electrophoresed. Two bands are seen: one corresponds to the DNA-protein complex, and one contains DNA without bound protein. The latter contains DNA molecules that have been prevented from attaching to the protein because the methylation treatment has modified one or more guanines that are crucial for binding. To identify which guanines are modified, the DNA fragment is purified from the gel and treated with piperidine.
As in the modification protection assay, the results of cleavage are visualized by denaturing gel electrophoresis. The length(s) of the labeled fragment(s) reveal which nucleotide(s) in the original fragment were methylated and hence identify the positions in the DNA sequence of guanines that participated in the binding reaction. Equivalent techniques can be used to identify the A, C, and T nucleotides involved in binding.

**Genomewide scans for protein attachment sites**

Gel retardation and modification assays have been used to map attachment sites for DNA-binding proteins in the regions adjacent to many genes in different species. One outcome of this work has been the identification of consensus sequences for the binding sites of several important transcription factors. One example in animals is the cyclic AMP response element (CRE), which has the consensus sequence 5′-TGACGTCA-3′, and is the recognition site for the cyclic AMP response element-binding (CREB) protein. This protein responds to elevated levels of cyclic AMP by binding to CRE sequences and activating the genes adjacent to those sequences. The cyclic AMP level in a cell is influenced by the presence of hormones that regulate physiological functions such as blood sugar level. The CREB protein is therefore the final link in a pathway that transduces the extracellular signal provided by the hormone into a change in the pattern of gene expression within the cell (Figure 11.9). The consensus sequences for most transcription factor binding sites are relatively unambiguous (compare the CRE consensus sequence with, for example, the downstream intron-exon boundary sequence in Section 5.1), and we might therefore imagine that they could be located in a genome sequence simply by scanning for these motifs. In practice this is not a regular component of a genome annotation project, because sequence scans usually result in a high proportion of false-positive identifications. These include sequences that have all the canonical features of a binding site but appear never to be used and others that have the appropriate sequence but are methylated in most cells, the additional methyl groups blocking protein binding.

Even if sequence scanning were a reliable way of identifying protein binding sites in a genome, the resulting catalog would not be particularly interesting. What we really wish to know is which sites are occupied in a particular tissue and how that occupancy pattern changes in response to external stimuli and during differentiation or development. That information enables genome annotation to move beyond the simple mapping of sequence motifs within a genome to a description of how the genome is expressed in different tissues and under different physiological conditions.

The most useful method for genomewide detection of occupied protein binding sites is chromatin immunoprecipitation sequencing or ChIP-seq. The initial step in a ChIP-seq experiment is treatment of the cells with formaldehyde to form DNA–protein cross-links, similar to the use of formaldehyde in the chromosome conformation capture or 3C method (Section 10.1). The DNA–protein complexes are extracted and sonicated to break the DNA into fragments (Figure 11.10). Because of the cross-linking, transcription factors and other DNA-binding proteins that are attached to the genome are not displaced during these extraction and fragmentation steps. The next step is to separate out of the mixture just those DNA fragments that are attached to the transcription factor of interest. This can be achieved by immunoprecipitation of the desired DNA–protein complexes with an antibody that is specific for that transcription factor. The DNA–protein cross-links
are now broken by heating to 70°C, and the DNA fragments that are released are sequenced by a next-generation method. The reads are mapped onto the genome sequence, revealing the location of those binding sites that were occupied by the transcription factor in the cells from which the extract was prepared. A related method called ChIP-on-chip or ChIP-chip achieves the same end, but instead...
of sequencing it uses hybridization to a microarray to identify the DNA fragments that are released from the purified DNA–protein complexes.

11.2 THE SPECIAL FEATURES OF DNA-BINDING PROTEINS

The structures of many DNA-binding proteins and DNA-protein complexes have been determined by methods such as X-ray crystallography and NMR. When the structures of sequence-specific DNA-binding proteins are compared, it is immediately evident that the family as a whole can be divided into a limited number of different groups according to the structure of the protein segment that interacts with the DNA molecule (Table 11.1). Each of these DNA-binding motifs is present in a range of proteins, often from very different organisms, and at least some of them probably evolved more than once. We will look at two motifs in detail—the helix-turn-helix (HTH) motif and the zinc finger—and then briefly survey the others.

The helix-turn-helix motif is present in prokaryotic and eukaryotic proteins

The helix-turn-helix (HTH) motif was the first DNA-binding structure to be identified. As the name suggests, the motif is made up of two α-helices separated by a turn (Figure 11.11). The latter is not a random conformation but a specific structure, referred to as a β-turn, made up of four amino acids, the second of which

### Table 11.1 DNA-Binding Motifs

<table>
<thead>
<tr>
<th>Motif</th>
<th>Examples of proteins with this motif</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sequence-specific DNA-binding motifs</strong></td>
<td></td>
</tr>
<tr>
<td>Helix–turn–helix family</td>
<td></td>
</tr>
<tr>
<td>Standard helix–turn–helix</td>
<td><em>Escherichia coli</em> lactose repressor</td>
</tr>
<tr>
<td>Homeodomain</td>
<td><em>Drosophila</em> Antennapedia protein</td>
</tr>
<tr>
<td>Paired homeodomain</td>
<td>Metazoan Pax transcription factors</td>
</tr>
<tr>
<td>POU domain</td>
<td>Metazoan transcription factors PIT-1, OCT-1, OCT-2</td>
</tr>
<tr>
<td>Winged helix–turn–helix</td>
<td>Mammalian E2F transcription factors</td>
</tr>
<tr>
<td><strong>Zinc-finger family</strong></td>
<td></td>
</tr>
<tr>
<td>Cys2His2 finger</td>
<td>Transcription factor TFIIA of eukaryotes</td>
</tr>
<tr>
<td>GATA zinc finger</td>
<td>GATA family of eukaryotic transcription factors</td>
</tr>
<tr>
<td>Treble clef finger</td>
<td>Metazoan nuclear receptor transcription factors</td>
</tr>
<tr>
<td><strong>Other motifs</strong></td>
<td></td>
</tr>
<tr>
<td>Basic helix–loop–helix</td>
<td>Eukaryotic MYC transcription factors</td>
</tr>
<tr>
<td>Ribbon–helix–helix</td>
<td>Bacterial MetJ, Arc, and Mnt repressors</td>
</tr>
<tr>
<td>High mobility group (HMG) box</td>
<td>Mammalian sex determination protein SRY</td>
</tr>
<tr>
<td>TBP domain</td>
<td>Eukaryotic TATA-binding protein</td>
</tr>
<tr>
<td>β-Barrel dimer</td>
<td>Papillomavirus E2 protein</td>
</tr>
<tr>
<td>Rel homology domain (RHB)</td>
<td>Mammalian transcription factor NF-κB</td>
</tr>
<tr>
<td><strong>Nonspecific DNA-binding motifs</strong></td>
<td></td>
</tr>
<tr>
<td>Histone fold</td>
<td>Eukaryotic histones</td>
</tr>
<tr>
<td>HU/HF motif*</td>
<td>Bacterial HU and IHF proteins</td>
</tr>
<tr>
<td>Polymerase cleft</td>
<td>DNA and RNA polymerases</td>
</tr>
</tbody>
</table>

*The HU/HF motif is a nonspecific DNA-binding motif in bacterial HU proteins (Section 8.1) but directs sequence-specific binding of the IHF (integration host factor) protein.

![Figure 11.11 Helix-turn-helix motif](image-url)
is usually glycine. This turn, in conjunction with the first \( \alpha \)-helix, positions the second \( \alpha \)-helix on the surface of the protein in an orientation that enables it to fit inside the major groove of a DNA molecule. This second \( \alpha \)-helix is therefore the recognition helix that makes the vital contacts which enable the DNA sequence to be read. The HTH structure is usually 20 or so amino acids in length, and so it is just a small part of the protein as a whole. Some other parts of the protein might also form attachments with the surface of the DNA molecule, primarily to aid the correct positioning of the recognition helix within the major groove.

Many prokaryotic and eukaryotic DNA-binding proteins utilize an HTH motif. In bacteria, HTH motifs are present in some of the best-studied regulatory proteins, which switch on and off the expression of individual genes. An example is the lactose repressor, which regulates expression of the lactose operon in *Escherichia coli* (Section 12.2). The various eukaryotic HTH proteins also include many whose DNA-binding properties are important in the developmental regulation of genome expression, such as the homeodomain proteins, whose roles we will examine in Section 14.3. The homeodomain is an extended HTH motif made up of 60 amino acids that form four \( \alpha \)-helices; helices 2 and 3 are separated by a \( \beta \)-turn, with helix 3 acting as the recognition helix and helix 1 stabilizing the structure (Figure 11.12).

Other versions of the HTH motif found in eukaryotes include the following:

- The POU domain is usually found in proteins that also have a homeodomain; the two motifs probably work together by binding different regions of a double helix. The name POU comes from the initial letters of the names of the first proteins found to contain this motif.

- The winged helix-turn-helix motif is another extended version of the basic HTH structure; this one has a third \( \alpha \)-helix on one side of the HTH motif and a \( \beta \)-sheet on the other side.

Many proteins, both prokaryotic and eukaryotic, possess an HTH motif, but the details of the interaction of the recognition helix with the major groove are not exactly the same in all cases. The length of the recognition helix varies (generally being longer in eukaryotic proteins), the orientation of the helix in the major groove is not always the same, and the positions within the recognition helix of the amino acids that make contacts with the nucleotides are different.

### Zinc fingers are common in eukaryotic proteins

The second type of DNA-binding motif that we will examine in detail is the zinc finger, which is rare in prokaryotic proteins but very common in eukaryotes. There appear to be more than 350 different zinc-finger proteins in the worm *Caenorhabditis elegans*, and over 10% of all mammalian genes code for zinc-finger proteins (see Table 7.5).

There are at least six different versions of the zinc finger. The first to be studied in detail was the Cys\(_2\)His\(_2\) finger, which comprises a series of 12 or so amino acids, including two cysteines and two histidines, that form a segment of \( \beta \)-sheet followed by an \( \alpha \)-helix. These two structures, which form the finger projecting from the surface of the protein, hold between them a bound zinc atom, coordinated to the two cysteines and two histidines (Figure 11.13). The \( \alpha \)-helix is the part of the motif that makes critical contacts within the major groove, and its positioning within the groove is determined by the \( \beta \)-sheet, which interacts with the sugar-phosphate backbone of the DNA, and by the zinc atom, which holds the \( \beta \)-sheet and \( \alpha \)-helix in the appropriate positions relative to one another. The \( \alpha \)-helix of a Cys\(_2\)His\(_2\) finger is therefore a recognition helix, similar to the second helix of the helix-turn-helix structure. Other versions of the zinc finger differ in the structure of the finger—some lack the \( \beta \)-sheet component and consist simply of one or more \( \alpha \)-helices—and the precise way in which the zinc atom is held in place also varies. For example, multicysteine zinc fingers lack histidines, and the zinc atom is coordinated between four cysteines. The multicysteine group includes the GATA zinc fingers, which are found in the GATA family of transcription factors.
An interesting feature of the zinc finger is that multiple copies of the finger are sometimes found in a single protein. Several proteins have two, three, or four fingers, but there are examples with many more than this: 37 in the case of one toad protein. In most cases, the individual zinc fingers are thought to make independent contacts with the DNA molecule, but sometimes the relationship between different fingers is more complex. In one particular group of transcription factors, the nuclear receptor family, two α-helices containing six cysteines combine to coordinate two zinc atoms in a single DNA-binding domain, larger than a standard zinc finger, called a treble clef finger (Figure 11.14). Within this motif, it appears that one of the α-helices enters the major groove whereas the second makes contacts with other proteins.

Other nucleic acid-binding motifs

Various other DNA-binding motifs have been discovered in different proteins:

- The basic helix-loop-helix motif, which is distinct from the HTH family, is found in a number of eukaryotic transcription factors. The first α-helix includes a region containing a high proportion of basic amino acids (such as arginine, histidine, and lysine), which binds to the major groove of the DNA (Figure 11.15). The remainder of the helix-loop-helix structure facilitates dimerization, so the active transcription factor consists of two subunits, which might have identical (homodimer) or different (heterodimer) structures.

- The ribbon-helix-helix motif is one of the few motifs that achieves sequence-specific DNA binding without making use of an α-helix as the recognition structure. Instead, the ribbon (that is, two strands of β-sheet) makes contact with the major groove (Figure 11.16). Ribbon-helix-helix motifs are found in some gene regulatory proteins in bacteria.

- The high mobility group (HMG) box domain is approximately 75 amino acids in length and comprises three α-helices that form an L shape. The high mobility group proteins are a large and diverse collection of chromatin proteins that were originally classified together because of their electrophoretic properties. They include several sequence-specific transcription factors, which typically have a single HMG box, and other DNA-binding proteins that lack sequence specificity. Members of the latter group, which are involved in processes such as DNA replication and repair, usually have more than one HMG box and recognize distorted regions of a DNA...
molecule, such as those that occur when a nucleotide has undergone chemical modification due to the action of a mutagen (Section 16.2).

RNA-binding proteins also have specific motifs that form attachments with RNA molecules, most of these acting in a sequence-independent manner. The most important are as follows:

- The **RNA recognition domain** comprises four β-strands and two α-helices in the order β-α-β-β-α-β. The two central β-strands make the critical attachments with a single-stranded RNA molecule. The RNA recognition domain is the commonest RNA-binding motif and has been found in more than 250 proteins. A similar domain is also present in a few proteins that bind to single-stranded DNA.

- The **double-stranded RNA binding domain (dsRBD)** is similar to the RNA recognition domain but has the structure α-β-β-β-α. The RNA-binding function lies between the β-strand and the α-helix at the end of the structure. As the name implies, the motif is found in proteins that bind double-stranded RNA.

- The **K homology (KH) domain** has the structure β-α-α-β-β-α, with the binding function between the pair of α-helices. There are two groups of KH domains: in one group, the three β-strands form an antiparallel β-sheet, and in the other group, two of the strands are in the parallel conformation. Proteins of the first group are found mainly in eukaryotes, and those in the second group are found mainly in prokaryotes. KH domains are also found in some single-stranded DNA-binding proteins.

Additionally, the DNA-binding homeodomain may also have RNA-binding activity in some proteins. One ribosomal protein uses a structure similar to a homeodomain to attach to rRNA, and some homeodomain proteins, such as Bicoid of Drosophila melanogaster (Section 14.3), can bind both DNA and RNA.

### 11.3 Interaction Between DNA and Its Binding Proteins

In recent years, our understanding of the interaction between the genome and DNA-binding proteins has begun to change. It has always been accepted that proteins that recognize a specific sequence as their binding site can locate this site by forming contacts with chemical groups attached to the bases that are exposed within the major and minor grooves that spiral around the double helix (see Figure 1.9). This is called direct readout and is still looked on as the predominant component of the interaction between a DNA-binding protein and its attachment site. What we are now beginning to recognize is that direct readout can be aided by the influence that the nucleotide sequence has on the precise conformation of the helix. These conformational features represent a second, less direct way in which the DNA sequence can influence protein binding.

#### Direct Readout of the Nucleotide Sequence

It was clear from the double-helix structure described by Watson and Crick (see Figure 1.9) that although the nucleotide bases are on the inside of the DNA molecule, they are not entirely buried, and some of the chemical groups attached to the purine and pyrimidine bases are accessible from outside the helix. Direct readout of the nucleotide sequence should therefore be possible without breaking the base pairs and opening up the molecule.

In order to form chemical bonds with groups attached to the nucleotide bases, a binding protein must make contacts within one or both of the grooves on the surface of the helix. With the B-form of DNA, the identity and orientation of the exposed parts of the bases within the major groove is such that most
sequences can be read unambiguously, whereas within the minor groove the identification is less precise (Figure 11.17). Direct readout of the B-form therefore predominantly involves contacts in the major groove. With other DNA types there is much less information on the contacts formed with binding proteins, but the picture is likely to be quite different. In the A-form, for example, the major groove is deep and narrow and less easily penetrated by any part of a protein molecule (see Figure 1.11). The shallower minor groove is therefore likely to play a more substantial role in direct readout. With Z-DNA, the major groove is virtually nonexistent and direct readout is possible to a certain extent without moving beyond the surface of the helix.

**The nucleotide sequence has a number of indirect effects on helix structure**

Originally it was thought that cellular DNA molecules have fairly uniform structures, made up mainly of the B-form of the double helix. Some short segments might be in the A-form, and there might be some Z-DNA tracts, especially near the ends of a molecule, but the majority of the length of a double helix would be unvarying B-DNA. We now recognize that DNA is much more polymorphic and that it is possible for the A-, B-, and Z-DNA configurations, and intermediates between them, to coexist within a single DNA molecule, with different parts of the molecule having different structures. There may also be regions that are identified as B-DNA but in which the major and/or minor grooves have atypical dimensions. These conformational variations are sequence-dependent, in part as a result of the base-stacking interactions that occur between adjacent base pairs. As well as being responsible, along with base pairing, for the stability of the helix, base stacking also influences the amount of rotation that occurs around the covalent bonds within individual nucleotides and hence determines the conformation of the helix at a particular position. The rotational possibilities in one base pair are influenced, via base-stacking interactions, by the identities of the neighboring base pairs. This means that the nucleotide sequence indirectly affects the overall conformation of the helix, possibly providing structural information that a binding protein can use to help it locate its appropriate attachment site on a DNA molecule. At present this is just a theoretical possibility, as no protein that specifically recognizes a non-canonical B-form of the helix has been identified, but many researchers believe that helix conformation is likely to play some role in the interaction between DNA and proteins.

A second type of conformational change is **DNA bending**. This does not refer to the natural flexibility of DNA, which allows it to form circles and supercoils, but instead to localized positions where the nucleotide sequence causes the DNA to bend. Like other conformational variations, DNA bending is sequence-dependent. In particular, a DNA molecule in which one polynucleotide contains two or more groups of repeated adenines, where each group comprises 3–5 adenines with the individual groups separated by 10 or 11 nucleotides, will bend at the 3′-end of the adenine-rich region. As with helix conformation, it is not yet known to what extent DNA bending influences protein binding, although several examples are known of DNA-binding proteins that induce bends or kinks in the double helix after they have attached.

**Contacts between DNA and proteins**

The contacts formed between DNA and its binding proteins are noncovalent. Within the major groove, hydrogen bonds form between the nucleotide bases and the R groups of amino acids in the recognition structure of the protein, whereas in the minor groove, hydrophobic interactions are more important. On the surface of the DNA helix, the major interactions are electrostatic, between the negative charges on the phosphate component of each nucleotide and the positive charges on the R groups of amino acids such as lysine and arginine, although some hydrogen bonding also occurs. In some cases, hydrogen bonding on the surface of the
helix or in the major groove occurs directly between DNA and protein; in others, it is mediated by water molecules. Few generalizations can be made: at this level of DNA-protein interaction, each example has its own unique features, and the details of the bonding have to be worked out by structural studies rather than by comparisons with other proteins.

Most proteins that recognize specific nucleotide sequences are also able to bind nonspecifically to other parts of a DNA molecule. In fact, it has been suggested that the amount of DNA in a cell is so large, and the numbers of each DNA-binding protein are so small, that the proteins spend most of their time attached nonspecifically to DNA. The distinction between nonspecific and specific forms of binding is that the latter is more favorable in thermodynamic terms. As a result, a protein is able to bind to its specific site even though there are literally millions of other sites to which it could attach nonspecifically. To achieve this thermodynamic favorability, the specific binding process must involve the greatest possible number of DNA-protein contacts, which explains in part why the recognition structures of many DNA-binding motifs have evolved to fit snugly into the major groove of the helix, where the opportunity for DNA-protein contacts is greatest. It also explains why some DNA-protein interactions result in conformational changes to one partner or the other, increasing still further the complementarity of the interacting surfaces and allowing additional bonding to occur.

The need to maximize contacts in order to ensure specificity is also one of the reasons why many DNA-binding proteins are dimers, consisting of two proteins attached to one another. This is the case for most HTH proteins and many of the zinc-finger type. Dimerization occurs in such a way that the DNA-binding motifs of the two proteins are both able to access the helix, possibly with some degree of cooperativity between them, so that the resulting number of contacts is greater than twice the number achievable by a monomer. As well as their DNA-binding motifs, many proteins contain additional characteristic domains that participate in the protein–protein contacts that result in dimer formation. One of these is the leucine zipper, in which an α-helix that coils more tightly than normal presents a series of leucines on one of its faces. These can form contacts with the leucines on the helix of a second polypeptide, forming the dimeric protein (Figure 11.18). As mentioned above, the basic helix–loop–helix motif is also able to form protein–protein contacts that result in homo- or heterodimer formation.

An intriguing question is whether the specificity of DNA binding can be understood in sufficient detail for the sequence of a protein’s target site to be predicted from examination of the structure of the recognition helix of a DNA-binding motif. To date, this objective has largely eluded us, but it has been possible to deduce some rules for the interaction involving certain types of zinc finger. In these proteins, four amino acids—three in the recognition helix and one immediately adjacent to it—form critical attachments with the nucleotide bases of the target site. By comparing the sequences of amino acids in the recognition helices of different zinc fingers with the sequences of nucleotides at the binding sites, it has been possible to identify a set of rules governing the interaction. These enable the nucleotide sequence specificity of a new zinc-finger protein to be predicted, admittedly with the possibility of some ambiguity, once the amino acid composition of its recognition helix is known.

**SUMMARY**

- The central players in genome expression and other aspects of genome activity are DNA-binding proteins that attach to the genome in order to perform their biochemical functions.
- The structures of these proteins have been studied by X-ray crystallography and nuclear magnetic resonance spectroscopy.
- The binding positions of these proteins on DNA molecules can be identified by gel retardation analysis and delineated in greater detail by modification
assays. Genomewide scans for the binding sites of a protein can be carried out by chromatin immunoprecipitation sequencing.

- DNA-binding proteins are able to attach to specific DNA sequences by virtue of special domains that form interactions with the double helix.
- The helix–turn–helix motif is a common domain in prokaryotic and eukaryotic DNA-binding proteins.
- Zinc fingers are common DNA-binding domains in eukaryotic proteins.
- There are also RNA-binding proteins with domains specific for attachment to RNA polynucleotides.
- Some proteins recognize their binding sites by direct readout of the DNA sequence, which is possible by the protein making contacts within the major groove of the double helix. In the major groove the identity of nucleotides can be determined from the positions of the chemical groups attached to the purine and pyrimidine nucleotides.
- Direct readout can be influenced by various indirect effects that the nucleotide sequence has on the conformation of the helix, including the formation of bends in A-rich sequences.
- Many DNA-binding proteins act as dimers, contacting the helix at two positions simultaneously. Special structures on the protein surface, such as leucine zippers, aid dimerization.

**SHORT ANSWER QUESTIONS**

1. Describe how X-ray crystallography is used to study protein structures.
2. Outline the strengths and weaknesses of nuclear magnetic resonance spectroscopy as a means of studying protein structure.
3. Explain how gel retardation analysis is used to identify protein-binding sites in a DNA molecule.
4. How are modification assays used to delineate the positions at which proteins bind to DNA molecules?
5. Describe the techniques that are available for scanning a genome to locate the positions of the attachment sites of a particular DNA-binding protein.
6. Describe the different types of helix–turn–helix motif that are known in prokaryotic and/or eukaryotic DNA-binding proteins.
7. What are the general properties of the Cys$_2$His$_2$ zinc-finger motif and how does this motif bind to DNA?
8. Describe three examples of RNA-binding domains.
9. Explain how direct readout of a nucleotide sequence is possible without detaching the two polynucleotides of a double helix.
10. What indirect effects does nucleotide sequence have on the structure of the double helix, and how might these effects influence the attachment of DNA-binding proteins?
11. Describe the features of a DNA region at which bending occurs.
12. What is a leucine zipper?
IN-DEPTH PROBLEMS

1. DNA does not form crystals but X-ray diffraction analysis was very important in the work that led to discovery of the double-helix structure. Explain how X-ray diffraction analysis can be used with DNA.

2. The resolution achievable by NMR is directly related to the field strength of the magnet that is used. Explore how this relationship has affected development of NMR over the last 20 years, and speculate about the future potential of the procedure.

3. Describe how a DNA chip or microarray might be used in a genomewide scan for protein binding sites in a DNA molecule.

4. Why are there so many different types of DNA-binding motif?

5. To what extent is it possible to use the amino acid sequence of a zinc finger in order to deduce the nucleotide sequence of the binding site for a protein containing one or more copies of that finger?

FURTHER READING

**X-ray crystallography and NMR spectroscopy**


**Methodology for identifying protein-binding sites**


**DNA- and RNA-binding motifs**


**Interactions between DNA and DNA-binding proteins**


The transcriptome is the collection of RNA molecules present in a cell. These RNA molecules are the transcripts of the genes that are active in that particular cell or whose recent activity gave rise to transcripts that have not yet been degraded. In the past, the transcriptome has been defined as simply the mRNA content of the cell and hence a reflection of the capacity of the cell to make proteins. In recent years, the term has been expanded to encompass all of a cell’s RNA, acknowledging the greater perception that we now have of the importance of noncoding RNA in diverse cellular activities.

We begin this chapter by examining the components of prokaryotic and eukaryotic transcriptomes and surveying the methods used to catalog the RNAs present in individual transcriptomes. We are already familiar with those methods, as they are the same ones that are used to map RNAs onto genomes during annotation projects (Section 5.3), so in this chapter we simply need to consider the technical aspects that are relevant to transcriptome cataloging. We will then study the events that result in synthesis, degradation, and processing of the individual components of a transcriptome and how those events are controlled. This will enable us to understand how the composition of a transcriptome is maintained by the cell and altered in response to changing environmental or physiological conditions. Finally, we will explore how transcriptome studies are contributing to research in areas such as cancer biology and the response of plants to environmental stress.

12.1 Components of the Transcriptome

In most prokaryotes and eukaryotes, the coding components of the transcriptome, messenger RNAs (mRNAs), make up less than 5% of the total RNA content of the cell. The remainder of the transcriptome consists of noncoding RNA molecules that are not translated into protein but nonetheless have important functional roles in the cell (Section 1.2). Two types of noncoding RNA have been known since the 1950s. These are ribosomal RNAs (rRNAs), which are components of ribosomes, and transfer RNAs (tRNAs), which carry amino acids to the ribosome and ensure that these amino acids are linked together in the order specified by the nucleotide sequence of the mRNA that is being translated (Section 13.3).

In addition to rRNA and tRNA, the cells of most organisms also contain a variety of other noncoding RNAs, several of which have been discovered very recently and whose functions are still not clear. In eukaryotes, these RNAs are divided into two groups on the basis of size. Those shorter than 200 nucleotides are called short noncoding RNAs (sncRNAs), and the longer ones, unsurprisingly, are called long noncoding RNAs (lncRNAs). This division is not generally used with prokaryotic noncoding RNAs, because apart from rRNAs, these are all shorter than 200 nucleotides.

The mRNA fraction of a transcriptome is small but complex

Although only a small part of a transcriptome, rarely more than 5% of the total RNA, the mRNA fraction is complex, containing copies of many different genes. Studies of human transcriptomes, for example, have shown that between 10,000
and 15,000 genes are expressed in a single tissue, with the cerebellum and testes being most complex in this regard, while skeletal muscle and liver are least complex. These gene numbers do not, however, reveal the true complexity of the mRNA content of a transcriptome. As we will begin to appreciate as we progress through this chapter, alternative splicing (Section 7.3), as well as the presence of multiple start- and endpoints for some transcripts, means that an individual gene can give rise to many different mRNAs (Figure 12.1). Because of these alternative synthesis and processing pathways, the 10,000–15,000 active genes in a cell can give rise to a transcriptome that contains over 100,000 different mRNAs.

Although the identities of different mRNAs in a transcriptome can be cataloged with some degree of accuracy, measuring the copy numbers of individual mRNAs is much more difficult. The relative abundances of different mRNAs can be assessed from the intensity of hybridization occurring at each position in a tiling array or from the numbers of reads that map to each gene in an RNA-seq database (Section 5.3), but converting these data into absolute numbers for individual transcripts is problematic. It would require controls containing known mRNA copy numbers from which a calibration curve could be constructed, and this is rarely attempted in a microarray or RNA-seq experiment. Quantitative PCR (Section 2.2), with suitable controls, can be used to estimate copy numbers of individual mRNAs, and estimates can also be obtained by fluorescent in situ hybridization (Section 3.5), which can be carried out in a format that enables the RNA contents of individual cells to be examined. These methods can assay the copy numbers of the transcripts of individual genes, but neither is capable of a rigorous distinction between all possible splice variants. Taking account of these limitations, the available evidence suggests that a typical mammalian cell contains approximately 200,000 mRNA molecules in total, implying a mean copy number of about 15 for the mRNAs derived from a single gene. Undoubtedly, there is considerable variation around this mean. For example, in cerebral cells the CREB transcription factor that binds to cAMP response element (CRE) sites (Section 11.1) has been estimated to have an mRNA copy number of less than 25, but a repressor protein that also binds to CRE sites has an mRNA copy number of up to 240. There are also some tissues that have highly specialized biochemistries, which are reflected by transcriptomes in which one or a few mRNAs predominate. Wheat seeds are an example. The cells in the endosperm of wheat seeds synthesize large amounts of gliadin proteins, which accumulate in the dormant grain and provide a source of amino acids for the germinating seedling. Within the developing endosperm, gliadin mRNAs can make up as much as 30% of the total mRNA content of certain cells.
Short noncoding RNAs have diverse functions

Although tRNAs were discovered in 1958, the existence of other types of snRNA was not suspected until the 1960s when gel electrophoresis was first applied to RNA extracts. The gels revealed the presence in eukaryotes of small RNAs, with relatively high uridine content, located primarily in the nucleus. To reflect the uridine content, these molecules are sometimes called U-RNAs, but the more commonly used term is small nuclear RNA (snRNA). The individual types are called U1-snRNA, U2-snRNA, etc., and their typical sizes are in the range 105–190 nucleotides, though longer examples are known. Most snRNAs (the Sm subgroup) are transported to the cytoplasm, where they attach to proteins to form complexes called small nuclear ribonucleoproteins (snRNPs) that then return to the nucleus, but a minor set (the Lsm subgroup) are converted into snRNPs in the nucleus, where they remain throughout their functional life. Most snRNPs are found within the nuclear regions called speckles (see Figure 10.2B), where they associate together to form spliceosomes, the structures that splice, or remove introns from, pre-mRNA transcripts (Section 12.4). Splicing has always been looked on as the major function of snRNAs, but some other roles are also known, in particular for U7-snRNA, which is not present in spliceosomes and instead is involved specifically in the processing of histone pre-mRNAs.

Eukaryotic nuclei also contain a second major group of small noncoding RNAs, called small nucleolar RNAs (snoRNAs). As the name indicates, these molecules are located in the nucleoli (Section 10.1), where they participate in the chemical modification of rRNAs (Section 1.2). As with snRNAs, snoRNAs are divided into two subgroups, in this case depending on whether they aid in methylation or pseudouridylation reactions (see Figure 1.18). Other snoRNAs are responsible for chemical modification of snRNAs, which takes place in the Cajal bodies (see Figure 10.2A). This group of snoRNAs is therefore sometimes called small Cajal body-specific RNAs (scaRNAs).

Outside the nucleus there are additional groups of small noncoding RNAs that have a variety of roles including regulation of genome expression. These RNAs were first discovered in the late 1990s, when processes responsible for RNA silencing in Caenorhabditis elegans and various species of plants were first described at the molecular level (Section 12.3). Since 2000, we have begun to realize that there are several types of these small regulatory RNAs and that they play central roles in controlling expression of individual genes. The main groups are as follows:

- **Short interfering RNAs** (siRNAs) are 20–25 nucleotides in length and are involved in the RNA interference pathway, which results in silencing of particular mRNAs. Silencing occurs because an siRNA base-pairs to a region in the target mRNA, forming a double-stranded structure that is cut by a nuclease. We will examine the details of the process in Section 12.3.

- **MicroRNAs** (miRNAs) are similar to siRNAs and also participate in RNA interference. The difference is that miRNAs are synthesized by cleavage of precursor molecules that form stem–loop structures, whereas the precursors of siRNAs are linear molecules (Figure 12.2).

- **Piwi-interacting RNAs** (piRNAs) are 25–30 nucleotides and hence slightly longer than siRNAs or miRNAs. They associate with piwi proteins, which were first discovered in Drosophila melanogaster but are now known to be present in many metazoans. Although piRNAs are the largest group of small regulatory RNAs in animal cells, their roles are not fully understood. They participate in RNA interference but are also involved in other processes that result in repression of gene activity and, in particular, prevent expression of the genes present in retrotransposons (Section 9.2), both by mRNA silencing and also by methylation of the DNA copies of these elements.

Additional types of small noncoding RNAs have more specialist roles. **Vault RNAs** are present in protein–RNA complexes called vaults, which are found in
most eukaryotic cells but whose functions are not known. Vaults tend to be associated with nuclear pore complexes and so may aid transport of molecules into and out of the nucleus. Other protein–RNA structures include the eukaryotic signal recognition particle, which is involved in movement of newly translated polypeptides into the endoplasmic reticulum. This particle is a complex of six proteins and one noncoding RNA, called the 7SL RNA. Another eukaryotic noncoding RNA, the 7SK RNA, is a component of a protein–RNA complex that controls the activity of a second protein complex called P-TEFb (positive transcriptional elongation factor b). P-TEFb regulates the rate at which protein-coding genes are transcribed into mRNA. Various other noncoding RNAs are components of enzymes such as ribonuclease P, which is involved in tRNA processing, and telomerase, the enzyme that prevents a chromosomal DNA molecule from decreasing in length every time the chromosome replicates (Section 15.4).

The types of snRNA that we have discussed so far appear to be unique to eukaryotes. Archaea and bacteria possess proteins similar to those found in eukaryotic snRNPs, and the prokaryotic versions might associate with RNAs to form snRNPs of their own, but what role these structures might have in RNA processing or other cellular activities is not known. Bacterial transcriptomes contain noncoding RNAs from repetitive extragenic palindromic (REP) sequences and clustered regularly interspaced short palindromic repeats (CRISPRs), which we encountered in Section 8.2 when we studied prokaryotic genomes, as well as transfer-messenger RNA, which forms part of a recovery system that enables damaged mRNAs to be translated.

**Long noncoding RNAs are enigmatic transcripts**

One of the more remarkable discoveries arising from genome annotation projects has been the realization that eukaryotic transcriptomes contain many long noncoding RNAs that collectively cover a large proportion of the intergenic space within a genome. The human genome, for example, is thought to specify over 50,000 IncRNA transcripts, none of which contains an open reading frame of greater than 100 codons and none of which is thought to be translated into a functional protein. The majority of these IncRNAs are synthesized only in certain tissues or at certain developmental stages. Their individual copy numbers are, on average, much lower than mRNA copy numbers, and in many cases there may be as few as two or three copies of a particular IncRNA per cell. As well as IncRNAs that are located entirely within an intergenic region, sometimes called long intergenic noncoding RNAs (lincRNAs), some map within the introns of mRNA transcription units, and others overlap with protein-coding exons, though often they are transcribed from the other strand of the double helix and hence give rise to an antisense version of the mRNA. Additional complexity arises from the presence of introns in some IncRNAs (Figure 12.3). In some parts of a eukaryotic genome, multiple IncRNAs form overlapping arrays of sense and antisense transcripts covering tens of kilobases of sequence.
Although the huge numbers of lncRNAs argue that they must have some purpose, assigning functions to these RNAs has proven difficult. The group includes the Xist and Tsix RNAs, which mediate the X inactivation process in female mammals, as well as other RNAs that play similar roles during genomic imprinting (Section 10.3). Other lncRNAs are transcripts of pseudogenes, and although a few of these lncRNAs are translated there is no convincing evidence that any of the resulting proteins are functional. When we discussed the issue of pseudogene functionality in Section 7.3, we emphasized the need to distinguish between a protein that participates in a biochemical process, and therefore seems to play a role in a cell, and proteins that display true functionality in an evolutionary sense, which are therefore subject to positive selection pressure. The same issues arise when we try to assess whether an lncRNA is functional. For example, some lncRNAs contain binding sites for transcription factors and so can act as decoys for these regulatory proteins. The presence of the decoy lncRNA in a transcriptome can therefore result in down-regulation of transcription of one or more protein-coding genes (Figure 12.4). This could be a genuine regulatory function, but an alternative explanation is that the lncRNA simply interferes with the transcription of these genes.

If many lncRNAs lack a genuine function, then why are they synthesized? It has been suggested that at least some lncRNAs represent transcriptional noise. Within the vast intergenic regions of a eukaryotic genome, there will inevitably be sequences that arise by chance mutation and which resemble the promoters and other sequences that direct transcription of a functional gene. The cell will be unable to prevent these sequences from acting as transcription signals, especially if they lie within a functional domain and hence are influenced by the regulatory signals that direct transcription of the genuine genes in that domain. According to this hypothesis, many lncRNAs are junk that the cell is unable to get rid of.

(A) PANDA absent, genes expressed

(B) PANDA present, genes not expressed

NF-YA transcription factor

CCNB1

FAS

BBC3

PMAIP1

PANDA IncRNA

Figure 12.3 Examples of possible map positions for long noncoding RNAs (lncRNAs) and long intergenic noncoding RNAs (lincRNAs).

Figure 12.4 PANDA, an example of a decoy lncRNA in the human transcriptome. PANDA, although it is an RNA molecule, is able to bind the transcription factor NF-YA. (A) When PANDA is absent, NF-YA attaches upstream of various target genes, including ones that induce apoptosis, or programmed cell death. (B) When PANDA lncRNAs are present, NF-YA binds to the RNA sites, reducing occupancy of the sites upstream of the target genes. These genes are therefore switched off and apoptosis does not occur. It has been proposed that PANDA mediates the balance between cell cycle arrest and apoptosis in cells that have damaged genomes. DNA damage activates a second transcription factor, p53, which switches on the PANDA gene. Synthesis of the lncRNA then prevents or delays activation of the apoptosis genes, thereby providing the cell with an opportunity to repair the damage and continue its active life rather than immediately dying.
Despite doubts about the functions of lncRNAs, there is a growing list of human diseases and developmental abnormalities that are associated with transcription of one or more of these RNAs. For example, over 500 lncRNAs have been associated with one or more types of human cancer. These lncRNAs include prostate cancer-associated transcript 6 (PCAT6), which is a 1.2 kb lncRNA transcribed from human chromosome 1. Copy numbers of PCAT6 are higher in the transcriptomes of cancerous prostate tissues compared with normal prostates, and the copy numbers have a significant correlation with the potential of the cancer to undergo metastasis. Similar results have been reported for the association of PCAT6 with lung cancer, where there is evidence that the transcript plays an active role in cancer progression. The many associations between lncRNAs and disease that are being discovered every year will ensure that research remains focused on these transcripts.

**Microarray analysis and RNA sequencing are used to study the contents of transcriptomes**

We are already familiar with the methods used to study transcriptomes, as these are the same methods that are used for genomewide mapping of RNA transcripts during genome annotation (Section 5.3). Hybridization of RNA or complementary DNA preparations to DNA chips—tiling arrays or microarrays, the latter usually representing the coding sequences in a genome—has been used extensively in the past to identify the RNAs present in a transcriptome. More recently, with the growing ease of next-generation sequencing, RNA-seq methods have become increasingly important, despite the computational challenges.

One difference between the use of microarray analysis and RNA-seq in transcript mapping and in transcriptome analysis is that the latter often requires quantitative data. With RNA mapping, the only important information is that a transcript is present in the transcriptome. To understand the content of the transcriptome in detail, we also wish to know the relative abundances of individual RNAs and to be able to compare these abundances in different transcriptomes. Often the key distinction between two tissues, or between the normal and diseased versions of a single tissue, is not the presence or absence of particular transcripts but their relative amounts, which reflect the patterns of up- or down-regulation of gene expression that are occurring in those tissues.

If a microarray is being used, then the differences between hybridization intensities for the same gene with two different RNA samples must represent genuine differences in RNA amount and not be due to experimental factors such as the amount of target DNA on different copies of the array, the efficiency with which the probe has been labeled, or the effectiveness of the hybridization process. Even in a single laboratory, these factors can rarely be controlled with absolute precision, and exact reproducibility between different laboratories is more or less impossible. This means that the data analysis must include normalization procedures that enable results from different array experiments to be accurately compared. The arrays must therefore include negative controls so that the background can be determined in each experiment, as well as positive controls that should always give identical signals. For vertebrate transcriptomes, the actin gene is often used as a positive control, as its expression level tends to be fairly constant in a particular tissue, regardless of the developmental stage or disease state. A more satisfactory alternative is to design the experiment so that the two transcriptomes can be directly compared, in a single analysis using a single array. This is done by labeling the cDNA preparations with different fluorescent probes and then scanning the array at the appropriate wavelengths to determine the relative intensities of the two fluorescent signals at each position, and hence to determine the differences between the RNA contents of the two transcriptomes (Figure 12.5).

If RNA-seq is being used, then the relative amounts of transcripts in different transcriptomes are assessed from the numbers of reads that map to the gene of interest. The rationale is, of course, that if an RNA is relatively abundant in transcriptome A but less abundant in transcriptome B, then there will be more reads
derived from the RNA in the RNA-seq data set for transcriptome A compared with that for transcriptome B. As with microarray analysis, genes whose expression levels are expected to be the same in the two tissues are used as controls to enable the RNA-seq data to be normalized.

If it is presumed that accurate comparisons can be made between two or more transcriptomes, then quite complex differences in gene expression patterns can be distinguished. Genes that display similar expression patterns are likely to be ones with related functions, so as well as cataloging transcriptome components, it is also possible to address questions relating to the functional annotation of a genome. To do this, a rigorous method is needed for comparing the expression profiles of different genes. The standard method, which can be applied to both microarray and RNA-seq data, is called hierarchical clustering. This involves comparing the expression levels of every pair of genes in the transcriptomes that have been analyzed and assigning a value that indicates the degree of relatedness between those expression levels. These data can then be expressed as a **dendrogram**, in which genes with related expression profiles are clustered together (*Figure 12.6*). The dendrogram therefore gives a clear visual indication of the functional relationships between genes.

**12.2 SYNTHESIS OF THE COMPONENTS OF THE TRANSCRIPTOME**

The composition of a transcriptome is determined by the balance between synthesis and degradation of the individual RNAs that it contains. For any RNA, a steady state is reached when its rate of synthesis (that is, the number of new molecules being added per unit time) is equal to the rate of degradation (that is, the number of molecules being destroyed per unit time). This steady state is achieved when the expression levels of different genes can be compared and standardized. The expression levels of genes are typically represented by a **dendrogram**, which is used to visualize the relationships between genes. **Figure 12.5** shows a comparison of two transcriptomes in a single experiment, while **Figure 12.6** illustrates how hierarchical clustering can be used to compare the expression profiles of five genes in seven transcriptomes. The dendrogram in **Figure 12.6** shows the degree of relatedness between the expression profiles of the five genes.
Chapter 12: Transcriptomes

of copies made per unit time) equals its rate of degradation (Figure 12.7). To increase the amount of an RNA in a transcriptome, the rate of synthesis must increase or the rate of degradation must decrease. To reduce the amount of an RNA, the rate of synthesis must decrease or the rate of degradation must increase. Understanding how RNAs are synthesized and turned over is therefore central to understanding how the composition of a transcriptome responds to external stimuli, such as the presence of hormones or an alteration in the environment, and also provides us with an appreciation of how gene expression patterns change during differentiation, development, and disease. We will begin to address these issues by examining how RNAs, especially the mRNAs that code for proteins, are synthesized.

RNA polymerases are molecular machines for making RNA

In Section 1.2 we learned that the enzymes responsible for transcription of DNA into RNA are called DNA-dependent RNA polymerases. These enzymes synthesize RNA in the 5′→3′ direction, base pairing between the RNA and the DNA template ensuring that the sequence of nucleotides in the transcript is complementary to the sequence of the DNA molecule that is being transcribed (see Figure 1.13).

Transcription of eukaryotic nuclear genes requires three different RNA polymerases: RNA polymerase I, RNA polymerase II, and RNA polymerase III. Each is a multisubunit protein (8–12 subunits) with a molecular mass in excess of 500 kDa. Structurally these polymerases are quite similar to one another, but functionally they are quite distinct. Each works on a different set of genes, with no interchangeability (Table 12.1). Most research attention has been directed at RNA polymerase II, as this enzyme transcribes genes that code for proteins. It also synthesizes the Sm subgroup of snRNAs, as well as some snoRNAs, siRNAs, miRNAs, piRNAs, and most lncRNAs. RNA polymerase III transcribes a variety of snRNA genes, including those for tRNAs. RNA polymerase I transcribes the multicycopy repeat units containing the 28S, 5.8S, and 18S rRNA genes. Plants have two additional RNA polymerases, IV and V, which are related to RNA polymerase II and transcribe particular classes of sRNA genes.

Archaea possess a single RNA polymerase that is structurally very similar to the eukaryotic enzymes. But this is not typical of prokaryotes in general because the bacterial RNA polymerase is very different, consisting of just six subunits, with its composition described as $\alpha_2\beta\beta'$ωσ (two $\alpha$ subunits, one each of $\beta$ and the related
12.2 SYNTHESIS OF THE COMPONENTS OF THE TRANSCRIPTOME

### Table 12.1 Functions of the Three Eukaryotic Nuclear RNA Polymerases

<table>
<thead>
<tr>
<th>Polymerase</th>
<th>Types of RNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>RNA polymerase I</td>
<td>28S, 5.8S, and 18S rRNAs</td>
</tr>
<tr>
<td>RNA polymerase II</td>
<td>mRNA, 5s family of snRNAs, some snoRNAs, siRNA*, miRNA, piRNA, lncRNA</td>
</tr>
<tr>
<td>RNA polymerase III</td>
<td>tRNAs, 5S rRNA, Lsm family of snRNAs, some snoRNAs, 7SL RNA, 7SK RNA</td>
</tr>
</tbody>
</table>

*In plants, some siRNAs are transcribed by RNA polymerase IV or V.

β', one ω, and one σ). The α, β, β’, and ω subunits are structural homologs of subunits present in the eukaryotic RNA polymerases, but the σ subunit has its own special properties, in terms of both its structure and, as we will see later, its function.

The chloroplast genome encodes an RNA polymerase that is very similar to the bacterial enzyme, reflecting the bacterial origins of these organelles (Section 8.3). In addition, the chloroplasts of monocotyledonous and dicotyledonous plants import from the cytoplasm one or two, respectively, single-subunit RNA polymerases that are coded by nuclear genes. Many genes can be transcribed by both the chloroplast and nuclear-encoded RNA polymerases, though the chloroplast version appears to be used more frequently in photosynthetic tissue. A few genes can be transcribed only by the nuclear-encoded polymerases; interestingly, these include the rpoB gene, which specifies the β subunit of the chloroplast RNA polymerase. Mitochondria also have their own RNA polymerases, in this case a single-subunit enzyme coded by a nuclear gene. There is no endogenous mitochondrial RNA polymerase.

The bacterial RNA polymerase can synthesize RNA at a rate of several hundred nucleotides per minute. The average *Escherichia coli* gene, which is just a few thousand nucleotides in length, can therefore be transcribed in a few minutes. The eukaryotic RNA polymerase II has a more rapid synthesis rate than the bacterial polymerase, up to 2000 nucleotides per minute, but can take hours to synthesize a single RNA because many eukaryotic genes are much longer than bacterial ones. For example, the 2400 kb transcript of the human dystrophin gene takes about 20 hours to synthesize. Most RNA polymerases have an accuracy of one error per $10^4$–$10^5$ nucleotide additions, this level of inaccuracy being tolerated because RNAs are usually multicopy, and transcripts that contain errors form only a small proportion of the overall pool. The error rate is minimized by backtracking, which is stimulated by the bulge in a DNA–RNA duplex that occurs at a position that is not base-paired (Figure 12.8). The polymerase undergoes a slight structural rearrangement, which enables it to slide back along the template and cut the RNA upstream of the error. The polymerase then reverts to its synthesis structure and moves forward again to continue transcription.

![Figure 12.8](image-url) Backtracking enables an RNA polymerase to correct a transcription error.
Transcription start points are indicated by promoter sequences

It is essential for transcription to initiate at the correct position on a DNA molecule, immediately upstream of the individual gene that must be copied into RNA. This position is marked by a target sequence, loosely called a promoter, that is recognized either by the RNA polymerase itself or by a DNA-binding protein that, once attached to the DNA, forms a platform onto which the RNA polymerase binds (Figure 12.9).

The bacterial RNA polymerase recognizes a bipartite promoter sequence, whose two components are located approximately 10 bp and 35 bp upstream of the position where transcription should begin. In E. coli, the promoter consensus sequences are as follows:

-35 box: 5’-TTGACA-3’
-10 box: 5’-TATAAT-3’

The spacing between the two boxes is important because it places the two motifs on the same face of the double helix, facilitating their interaction with the σ subunit, which is the DNA-binding component of the bacterial RNA polymerase. The consensus sequences given above refer to the standard version of the bacterial polymerase, which contains the 70 subunit, so-called because its molecular mass is approximately 70 kDa. E. coli and other bacteria can also make a variety of other σ subunits, each one specific for a different -35 sequence. An example is the 32 subunit, which is synthesized when the bacterium is exposed to a heat shock. This subunit recognizes a -35 sequence that is found upstream of genes coding for proteins that help the bacterium withstand high temperatures (Figure 12.10). These proteins include chaperones that protect other proteins from heat degradation and enzymes that repair heat-induced DNA damage. Other σ subunits are used during nutrient starvation and nitrogen limitation. Through use of these alternative σ subunits, E. coli is able to remodel its transcriptome in response to changing environmental and nutritional demands. Alternative σ subunits are also used by other bacteria: for example, Klebsiella pneumoniae uses a σ54 subunit to switch on genes involved in nitrogen fixation, and Bacillus species use a whole range of different σ subunits to switch on and off groups of genes during the changeover from normal growth to formation of spores (Section 14.3).

In eukaryotes, the term promoter is used to describe all the sequences that are important in initiating transcription of a gene. For some genes these sequences can be numerous and diverse in their functions, including not only the core promoter, sometimes called the basal promoter, which is the site at which the RNA polymerase attaches, but also one or more upstream promoter elements which, as their name implies, lie upstream of the core promoter. Transcription initiation can usually occur in the absence of the upstream elements, but only in an inefficient way. This indicates that the proteins that bind to the upstream elements include at least some that are activators of transcription and therefore promote gene expression. Inclusion of these sequences in the promoter is therefore justified.

Each of the three eukaryotic RNA polymerases recognizes a different type of promoter sequence; indeed, the difference between the promoters defines which genes are transcribed by which polymerases. The details for vertebrates are as follows (Figure 12.11):

- RNA polymerase I promoters consist of a core promoter, spanning the transcription startpoint, between nucleotides -45 and +20, and an upstream control element (UCE) about 100 bp further upstream.
RNA polymerase II promoters are variable and can stretch for several kilobases upstream of the transcription start site. The core promoter consists of two main segments: the −25 or TATA box (consensus 5′-TATAAAAR-3′, where W is A or T, and R is A or G) and the initiator (Inr) sequence (mammalian consensus 5′-YANWYY-3′, where Y is C or T, and N is any nucleotide) located around nucleotide +1. Some genes transcribed by RNA polymerase II have only one of these two components of the core promoter and some, surprisingly, have neither. The latter are called null genes. They are still transcribed, although the start position for transcription is more variable than for a gene with a TATA box and/or Inr sequence. Some genes also have additional sequences that can be looked on as part of the core promoter. Examples include the following:

- The downstream promoter element (DPE; located at positions +28 to +32) has a variable sequence but has been identified through its ability to bind TFIID, a protein complex that plays a central role in the initiation of transcription.
- A 7 bp GC-rich motif called the B response element (BRE) is located immediately upstream or downstream of the TATA box. This motif is recognized by TFIIB, another component of the initiation complex.
- The proximal sequence element (PSE) is located between positions −45 and −60 upstream of those snRNA genes that are transcribed by RNA polymerase II.

As well as the components of the core promoter, genes transcribed by RNA polymerase II have various upstream promoter elements, usually located within 2 kb of the transcription start site, whose functions are described later in this chapter.

RNA polymerase III promoters fall into at least three categories. Two of these categories are unusual in that the important sequences are located within the genes whose transcription they promote. Usually these sequences span 50–100 bp and comprise two conserved boxes separated by a variable region. The third category of RNA polymerase III promoter is similar to those for RNA polymerase II, having a TATA box and a range of additional promoter elements (sometimes including the PSE mentioned above) located upstream of the target gene. Interestingly, this arrangement is seen with the Lsm class of snRNA genes, which are transcribed by RNA polymerase III, which means that these genes have similar promoter sequences to the Sm group that are transcribed by RNA polymerase II.

An additional level of complexity is seen with some eukaryotic genes that have alternative promoters that give rise to different versions of the transcript specified by the gene. An example is provided by the human dystrophin gene, which has been extensively studied because defects in this gene result in the genetic disease called Duchenne muscular dystrophy. The dystrophin gene is one of the largest known in the human genome, stretching over 2.4 Mb and containing 78
intron. It has at least seven alternative promoters, which are used in different tissues and which direct synthesis of mRNAs of different lengths (Figure 12.12). Alternative promoters are also used to generate related versions of some proteins at different stages in development and to enable a single cell to synthesize similar proteins with slightly different biochemical properties. The last point indicates that although they are usually referred to as alternative promoters, these are more correctly called multiple promoters, as more than one may be active at a single time. Indeed, this may be the normal situation for many genes. For example, a genomewide survey has revealed that 10,500 promoters are active in human fibroblast cells but that these promoters are driving expression of fewer than 8000 genes, indicating that a substantial number of genes in these cells are being expressed from two or more promoters simultaneously.

**Synthesis of bacterial RNA is regulated by repressor and activator proteins**

The consensus sequence for a bacterial promoter is quite variable, with a range of different motifs being permissible at both the −35 and −10 boxes (Table 12.2). These variations, together with less well-defined sequence features around the transcription start site and in the first 50 or so nucleotides of the transcription unit, affect the efficiency of the promoter. Efficiency is defined as the number of productive initiations that are promoted per second, where a productive initiation is one that results in RNA polymerase clearing the promoter and beginning synthesis of a full-length transcript. Different promoters vary 1000-fold in their efficiencies, the most efficient promoters (called strong promoters) directing 1000 times as many productive initiations as the weakest promoters. We refer to these as differences in the basal rate of transcription initiation.

Promoter structure determines the basal rate of transcription initiation but does not provide any general means by which transcription of a gene can respond to changes in the environment or to the biochemical requirements of the cell. Instead, these transient changes are brought about by regulatory proteins that have a negative (repressing) or positive (activating) effect on the basal transcription rate for a particular gene.

The foundation of our understanding of regulatory control over transcription initiation in bacteria was laid in the early 1960s by François Jacob, Jacques Monod, and other geneticists who studied the lactose operon, the group of three genes whose protein products convert lactose to glucose and galactose (see

<table>
<thead>
<tr>
<th>Genes</th>
<th>Protein product</th>
<th>−35 box</th>
<th>−10 box</th>
</tr>
</thead>
<tbody>
<tr>
<td>Consensus</td>
<td></td>
<td>5′-TTGACA-3′</td>
<td>5′-TATAAT-3′</td>
</tr>
<tr>
<td>argF</td>
<td>Ornithine transcarbamoylase</td>
<td>5′-TTGTGA-3′</td>
<td>5′-AATAAT-3′</td>
</tr>
<tr>
<td>can</td>
<td>Carbonic anhydrase</td>
<td>5′-TTTAAA-3′</td>
<td>5′-TATATT-3′</td>
</tr>
<tr>
<td>dnaB</td>
<td>DnaB helicase</td>
<td>5′-TCGTCA-3′</td>
<td>5′-TAAAGT-3′</td>
</tr>
<tr>
<td>gcd</td>
<td>Glucose dehydrogenase</td>
<td>5′-ATGACG-3′</td>
<td>5′-TATAAT-3′</td>
</tr>
<tr>
<td>gldA</td>
<td>Citrate synthase</td>
<td>5′-TTGACA-3′</td>
<td>5′-TACAAA-3′</td>
</tr>
<tr>
<td>ligB</td>
<td>DNA ligase</td>
<td>5′-GTCACA-3′</td>
<td>5′-TAAAAG-3′</td>
</tr>
</tbody>
</table>
Most of the original scheme for regulation of the lactose operon has been confirmed by DNA sequencing of the control region and by structural studies of the repressor bound to its operator. One complication has been the discovery that the repressor has three potential binding sites, centered on nucleotide positions −82, +11, and +412. The operator defined by genetic studies is the sequence located at +11 (see Figure 12.13A), and this is the only one of the three sites whose occupancy by the repressor would be expected to prevent access of the RNA polymerase to the promoter. But the other two sites also play some role in repression as their removal, individually or together, significantly impairs the ability of the repressor to switch off transcription. The repressor is a tetramer of four identical subunits, and the genetic evidence suggests that the repressor has an inherent flexibility that allows it to adopt the alternative conformations needed for repression and transcription.

Figure 12.13 Regulation of the *E. coli* lactose operon. (A) The operator sequence lies immediately downstream of the promoter for the lactose operon. Note that this sequence has regions of inverted symmetry: when each strand is read in the 5′ → 3′ direction, the sequence is the same. This enables two subunits of the tetrameric repressor protein to make contact with a single operator sequence. (B) In the original model for lactose regulation, the lactose repressor is looked on as a simple blocking device that binds to the operator and prevents RNA polymerase from gaining access to the promoter. The three genes in the operon are therefore switched off. This is the situation in the absence of lactose, although transcription is not completely blocked because the repressor occasionally detaches, allowing a few transcripts to be made. Because of this basal level of transcription, the bacterium always possesses a few copies of each of the three enzymes coded by the operon, probably fewer than five of each. This means that when the bacterium encounters a source of lactose it is able to transport a few molecules into the cell and split these into glucose and galactose. An intermediate in this reaction is allolactose, an isomer of lactose, which induces expression of the lactose operon by binding to the repressor, causing a change in the repressor conformation so it is no longer able to attach to the operator. This allows the RNA polymerase to bind to the promoter and transcribe the three genes. When fully induced, approximately 5000 copies of each protein product are present in the cell. When the lactose supply is used up and allolactose is no longer present, the repressor reattaches to the operator and the operon is switched off. The transcripts of the operon (which have a half-life of less than 3 min) decay, and the enzymes are no longer made.
subunits that work in pairs to attach to a single operator, so the repressor has the capacity to bind to two of the three operator sites at once. It is likely that binding of one pair of subunits to the +11 site is enhanced or stabilized by attachment of the other pair of subunits to the −82 or +412 site. It is also possible that the repressor can bind to a pair of operator sequences in such a way that it does not block attachment of the polymerase to the promoter but prevents the polymerase from leaving the promoter region.

As well as down-regulation by repressors, transcription of many bacterial operons and individual genes can be up-regulated by activating proteins. An example is provided by the **catabolite activator protein** (also called the CRP activator). This protein binds to a recognition sequence at various sites in the bacterial genome and activates transcription initiation at downstream promoters by interacting with the α subunit of the RNA polymerase. Inherent in this activation is the creation of a sharp 90° bend in the double helix in the region of the binding site when the catabolite activator protein is attached. Productive initiation of transcription at these promoters is dependent on the presence of bound catabolite activator protein; if the protein is absent, then the genes controlled by the promoter are not transcribed.

Attachment of the catabolite activator protein to its binding site occurs only when the bacterium has an inadequate supply of glucose. The availability of glucose is monitored by a protein called IIA\(^{Glc}\), which is a component of a multiprotein complex that transports sugars into the bacterium. When glucose is being transported into the cell, IIA\(^{Glc}\) becomes dephosphorylated (that is, additional phosphate groups previously added to the protein by post-translational modification are removed). The dephosphorylated version of IIA\(^{Glc}\) inhibits adenylate cyclase, the enzyme that converts ATP into cyclic AMP (cAMP) (**Figure 12.14**). This means that if glucose levels are high, the cAMP content of the cell is low. The catabolite activator protein can bind to its target sites only in the presence of cAMP, so when glucose is present, the protein remains detached and the operons it controls are switched off. One of these target sites is adjacent to the promoter of the lactose operon. If the bacterium has supplies of both glucose and lactose, the binding site for the catabolite activator protein is unoccupied, and in the absence of activation the lactose operon is not transcribed, even though the lactose repressor is not bound. If the bacterium uses up all the glucose, then the cAMP level rises and the catabolite activator protein binds to its target sites, including the site upstream of the lactose operon. Transcription of the lactose genes is

![Figure 12.14 Role of the catabolite activator protein](image)

(A) Transport of glucose into the bacterium results in dephosphorylation of IIA\(^{Glc}\). Dephosphorylated IIA\(^{Glc}\) inhibits adenylate cyclase, reducing the conversion of ATP into cAMP. (B) Catabolite activator protein (CAP) can attach to its DNA binding sites only in the presence of CAMP. If glucose is present, the cAMP level is low, so CAP does not bind to the DNA and does not activate RNA polymerase. Once the glucose has been used up, the cAMP level rises, allowing CAP to bind to the DNA and activate transcription of the target genes.
therefore activated, enabling the bacterium to start using lactose as its primary energy source. The catabolite activator protein therefore remodels the \textit{E. coli} transcriptome so the bacterium makes the most efficient use of the available sugars. This ability of \textit{E. coli} and other bacteria to metabolize one sugar in preference to another, and then switch to that second sugar when the preferred one is used up, was first discovered in 1941 by Jacques Monod, who used the French word \textit{diauxie} to describe it (Figure 12.15).

\textbf{Synthesis of bacterial RNA is also regulated by control over transcription termination}

Although initiation of transcription is looked on as the main control point in the process leading to synthesis of a bacterial RNA, it is not the only step that can be influenced in order to adjust the composition of a transcriptome.

Current thinking views transcription as a discontinuous process, with the polymerase pausing regularly and making a choice between continuing elongation, by adding more nucleotides to the transcript, or terminating, by dissociating from the template. Which choice is selected depends on which alternative is more favorable in thermodynamic terms. This model emphasizes that, in order for termination to occur, the polymerase has to reach a position on the template where dissociation is more favorable than continued RNA synthesis. Consistent with this model, many of the positions in a bacterial genome where transcription terminates are marked by an inverted palindrome, a sequence that can be folded into a stem–loop structure after transcription into RNA (Figure 12.16). Formation of this stem–loop is thought to weaken the overall interaction between the polymerase and the template, favoring dissociation over continued transcription. At intrinsic terminator sequences, attachment of the polymerase is further weakened by a run of As in the template, which leads to the transcript being held to the DNA by a series of A–U base pairs, which have only two hydrogen bonds each, compared with three for each G–C pair (Figure 12.17A). Alternatively, at Rho-dependent
Figure 12.17 Alternative modes for termination of transcription in bacteria. (A) At an intrinsic terminator, attachment of the polymerase to the DNA is weakened by a run of As in the template. (B) Termination at a Rho-dependent terminator. Rho is a helicase that follows the RNA polymerase along the transcript. When the polymerase stalls at a stem–loop structure, Rho catches up and breaks the RNA–DNA base pairs, releasing the transcript.

terminators, a helicase enzyme breaks the base pairs between the DNA and transcript when the polymerase pauses at the stem–loop structure (Figure 12.17B).

The ability of stem–loop structures to terminate bacterial transcription is exploited by a regulatory process called attenuation. The tryptophan operon of E. coli (see Figure 8.9B) illustrates how it works. In this operon, two stem–loops can form in the region between the start of the transcript and the beginning of trpE, the first gene in the operon. The smaller of these loops acts as a Rho-dependent terminator, but the larger one does not. The larger loop overlaps with the termination hairpin, so only one of the two hairpins can form at any given time. Which loop is formed depends on the relative positioning between the RNA polymerase and a ribosome that attaches to the 5′-end of the transcript as soon as it is synthesized, in order to translate the coding parts of the RNA into protein (Figure 12.18). If the ribosome stalls so that it does not keep up with the polymerase, then the larger stem–loop forms and transcription continues. However, if the ribosome keeps pace with the RNA polymerase, then it disrupts the larger stem–loop, which means the terminator structure is able to form, and transcription stops. Ribosome stalling can occur because, upstream of the termination signal, there is a short open reading frame coding for a 14-amino-acid peptide that includes two tryptophans. If the amount of free tryptophan is limiting, then the ribosome stalls as it attempts to synthesize this peptide, enabling the polymerase to continue making its transcript. Because the transcript contains copies of the genes coding for the biosynthesis of tryptophan, its continued elongation addresses the requirement that the cell has for this amino acid. When the amount of tryptophan in the cell reaches a satisfactory level, the attenuation system prevents further transcription of the tryptophan operon, because now the ribosome does not stall while making the short peptide and instead keeps pace with the polymerase, allowing the termination signal to form.

The E. coli tryptophan operon is controlled not only by attenuation but also by a repressor protein. Exactly how attenuation and repression work together to regulate expression of the operon is not known, but it is thought that repression provides the basic on–off switch and attenuation modulates the precise level of gene expression that occurs. Attenuation is used to regulate transcription of several operons involved in amino acid biosynthesis in E. coli and other bacteria, and it is also used to remodel transcriptomes in response to other types of nutrient limitation and to environmental changes such as temperature fluctuations. A common theme among these different attenuation systems is the presence of alternative stem–loop structures in the transcript, but the mechanisms used to influence formation of the terminator stem–loop are more variable. Some attenuators use ribosome stalling in the same way as the E. coli tryptophan operon, but others, called riboswitches, are regulated directly by the attachment of small molecules, such as amino acids, which disrupt the larger of the two stem–loops so termination occurs, and a third class is regulated by attachment of an RNA-binding protein.

Synthesis of eukaryotic RNA is regulated primarily by activator proteins

The key lesson that we have learned from our examination of transcriptional control in bacteria is that transcription initiation can be influenced by DNA-binding
12.2 SYNTHESIS OF THE COMPONENTS OF THE TRANSCRIPTOME

proteins that recognize specific sequences located near the attachment site for RNA polymerase. This is also the basis of transcriptional control in eukaryotes but with one difference. The bacterial RNA polymerase has a strong affinity for its promoter and the basal rate of transcription initiation is relatively high for all but the weakest promoters. With most eukaryotic genes, the reverse is true. RNA polymerase II and III initiation complexes do not assemble efficiently and the basal rate of transcription initiation is therefore very low, regardless of how strong the promoter is. In order to achieve effective initiation, formation of the complex must be activated by additional proteins. This means that, compared with bacteria, eukaryotes use different strategies to control transcription initiation, with activators playing a much more prominent role than repressor proteins.

Unlike the bacterial RNA polymerase, eukaryotic polymerases do not bind directly to their core promoter sequences. For genes transcribed by RNA polymerase II, initial contact is made by the general transcription factor (GTF) TFIIID, which is a complex made up of the TATA-binding protein (TBP) and at least 12 TBP-associated factors, or TAFs. TBP is a sequence-specific protein with an unusual DNA-binding domain that makes contact with the minor groove in the region of the TATA box. The TAFs assist in attachment of TBP to the TATA box and, in conjunction with other proteins called TAF- and initiator-dependent cofactors (TICs), possibly also participate in recognition of the Inr sequence, especially at those promoters that lack a TATA box.

X-ray crystallographic studies of TBP show that it has a saddlelike shape that wraps partially around the double helix, forming a platform onto which the RNA polymerase is subsequently positioned (Figure 12.19). Recruitment of RNA polymerase is aided by three more transcription factors, TFIIA, TFIIB, and TFIIIE, with the addition of TFIIH completing what is called the preinitiation
complex. In older models, construction of the preinitiation complex was a stepwise process, involving the sequential addition of TFIIA, TFIIB, RNA polymerase II, TFIIF, TFIIE, and finally TFIIH. It now seems more likely that at least some of the transcription factors form an assembly with RNA polymerase prior to attachment of this larger complex to the platform provided by TBP.

Whichever model is correct, completion of the preinitiation complex is followed by addition of phosphate groups to the C-terminal domain (CTD) of the largest subunit of RNA polymerase. In mammals, this domain consists of 52 repeats of the seven-amino-acid sequence Tyr-Ser-Pro-Thr-Ser-Pro-Ser. Two of the three serines in each repeat unit can be modified by addition of a phosphate group, catalyzed by a kinase activity provided by two of the subunits of TFIIH. Phosphorylation causes a substantial change in the ionic properties of the polymerase, enabling it to leave the initiation complex and begin synthesizing RNA.

The above account describes how the transcription of a eukaryotic protein-coding gene is initiated but does not explain how the rate of transcription is set at the appropriate level between zero and maximal. This is the role of DNA-binding proteins called transcription factors, which are separate and distinct from the general transcription factors, such as TFIID, that are contained within the preinitiation complex. An example of a regulatory transcription factor is the CREB protein, which activates its target genes by binding to CRE sequences in response to elevated cAMP levels (Section 11.1). Other transcription factors transduce signals from sources as diverse as steroid hormones and heat shock. The binding sites for transcription factors are located either close to or more distant from the gene whose expression they control (Figure 12.20):

- Proximal binding sites are upstream promoter elements, most of which are located within 2 kb of the transcription start site of the target gene. Attachment of a transcription factor to one of these sites influences transcription only of the gene within whose promoter the element is located.

- More distant binding sites are located within enhancers, which can be positioned anywhere within the functional domain that contains their target gene. A single enhancer can influence transcription of multiple genes within its domain, but it is prevented by insulator sequences from affecting genes in neighboring domains (Section 10.1).

How does the attachment of a transcription factor to its binding site activate or repress transcription initiation? The answer is that a bound transcription factor makes physical contact with a multisubunit protein called the mediator, which in turn makes contacts with different components of the preinitiation complex (Figure 12.21). The mediator is thought to influence several events involved in assembly of the complex, including positioning of the polymerase on TBP and activation of the polymerase by CTD phosphorylation. The mediator, therefore, as its name implies, mediates the signal from the transcription factor to the preinitiation complex.
Because of the low level of basal initiation occurring at an RNA polymerase II promoter, most transcription factors for this polymerase are activators. Only a few proteins that repress transcription initiation are known, these proteins binding to upstream promoter elements or to more distant sites in silencers. Some influence genome expression in a general way, through histone deacetylation (Section 10.2) or DNA methylation (Section 10.3), but others have more specific effects at individual promoters. Other proteins activate or repress transcription depending on the circumstances. Pit-1, which is the first of the three proteins after which the POU domain is named (Section 11.2), activates some genes and represses others, depending on the sequence of the DNA binding site. The presence in this site of two additional nucleotides induces a change in the conformation of Pit-1, enabling it to interact with a second protein called N-CoR and repress transcription of the target gene (Figure 12.22).

### 12.3 DEGRADATION OF THE COMPONENTS OF THE TRANSCRIPTOME

Now that we understand how the components of a transcriptome are synthesized, we must turn our attention to how they are degraded. As we have already discussed (see Figure 12.7), the processes responsible for synthesis and degradation of individual transcripts act in unison to determine the composition of a transcriptome and to enable that composition to change in response to the prevailing conditions.

**Several processes are known for nonspecific RNA turnover**

The rate of degradation of an RNA can be estimated by determining its **half-life** in a transcriptome. This is the period of time required for the amount of an individual type of RNA to fall to half its initial value, if it is assumed that there is no new synthesis of the molecule. Half-lives can be measured by **pulse labeling**. The cells being studied are briefly provided with a labeled substrate for RNA synthesis, such as radioactive 4-thiouracil, in which one of the oxygen atoms attached to carbon 4 of the normal uracil molecule has been replaced with a $^{35}$S atom. RNAs that are synthesized during the period of pulse labeling will contain the labeled nucleotide, but those made before or after will not. The degradation rates of the labeled molecules are then followed by measuring the amounts of label present in...
RNA extracts prepared at intervals after the period of pulse labeling. Experiments of this type have shown that there is considerable variation between and within organisms. Bacterial mRNAs are generally turned over very rapidly, with half-lives rarely longer than a few minutes, this being a reflection of the rapid changes in protein synthesis patterns that can occur in an actively growing bacterium with a generation time of 20 minutes or so. Eukaryotic mRNAs are longer-lived, with half-lives of, on average, 10–20 minutes for yeast and several hours for mammals. There have been fewer studies of noncoding RNA half-lives, but those that have been reported suggest that, in eukaryotes, tRNAs and rRNAs are both turned over more slowly than mRNAs, with tRNA half-lives between 9 hours and several days and rRNA half-lives up to 8 days. The figures for long noncoding RNAs are also variable, from less than 2 hours to over 16 hours in mouse neuroblastoma cells.

The half-lives measured by pulse labeling indicate that most types of RNA are subject to continual turnover. This means that, in a general sense, the composition of the transcriptome is responsive to changes in the rate of synthesis of individual transcripts. Much of this turnover is probably nonspecific, acting on all RNAs of a particular type and not discriminating between the transcripts of individual genes. In bacteria, nonspecific mRNA degradation is carried out by the degradosome, a multiprotein structure whose components include polynucleotide phosphorylase (PNPase), which removes nucleotides sequentially from the 3’-end of an mRNA; RNA helicase B, which opens up stem–loop structures such as terminator sequences and hence aids the progression of PNPase along the mRNA; and an endonuclease called RNase E, which makes internal cuts in RNA molecules. The eukaryotic equivalent of the degradosome is the exosome, which comprises a ring of six proteins, each of which has ribonuclease activity, with three RNA-binding proteins attached to the top of the ring. Other ribonucleases associate with the exosome in a transient manner. It is thought that RNAs to be degraded are initially captured by the binding proteins and then threaded through the channel in the middle of the ring, where they are exposed to the ribonuclease activities of the ring proteins (Figure 12.23).

Exosomes are present in both the cytoplasm and the nucleus of a eukaryotic cell. The main role of the nuclear exosomes appears to be the rapid turnover of aberrant RNAs that have not been transcribed or processed correctly and which are therefore not released into the cytoplasm. Aberrant mRNAs are detected by a surveillance mechanism, which identifies ones that lack a termination codon, which might occur if the DNA has been copied incorrectly, or have a termination codon at an unexpected position, indicating that the exons have been joined together incorrectly during RNA splicing. Surveillance involves a protein complex that scans mRNAs for these errors and directs aberrant transcripts to the exosome or some other degradation pathway. Other surveillance systems look for errors in the chemical modification of tRNAs. Some modification errors can be tolerated but others appear to be more critical, and tRNAs containing them are rapidly degraded, either by the exosome or by a second turnover pathway specific for tRNAs.

RNA silencing was first identified as a means of destroying invading viral RNA

The nonspecific turnover of RNAs ensures that a change in the transcription rate for a particular gene will result in a change in the steady-state concentration of the transcript of that gene in the transcriptome. One of the major areas of development in our understanding of genomes over the last 20 years has been the growing realization that most types of organisms also have mechanisms that enable individual transcripts, in particular mRNAs, to be degraded, possibly resulting in their rapid and complete removal from a transcriptome. For many years it has been known that eukaryotes possess RNA degradation mechanisms that protect their cells from attack by foreign RNAs, such as the genomes of viruses. Originally called RNA silencing, this process is already familiar to us under its alternative
name of RNA interference, as its underlying mechanism has been utilized by genome researchers as a means of inactivating selected genes in order to study their function (Section 6.2).

The target for RNA silencing must be double-stranded, which excludes cellular mRNAs but encompasses viral genomes, many of which are either double-stranded RNA in their native state or replicate via a double-stranded RNA intermediate (Section 9.1). The double-stranded RNA is recognized by binding proteins that form an attachment site for a ribonuclease called Dicer, which cuts the molecule into short interfering RNAs (siRNAs) of 20–25 nucleotides in length (Figure 12.24). This inactivates the virus genome, but what if the virus genes have already been transcribed? If this has occurred, then the harmful effects of the virus will already have been initiated and RNA silencing would appear to have failed in its attempt to protect the cell from damage. One of the more remarkable discoveries of recent years has revealed a second stage of the interference process that is directed specifically at viral mRNAs. The siRNAs produced by cleavage of the viral genome are separated into individual strands, and one strand of each siRNA subsequently base-pairs with any viral mRNAs that are present in the cell. The double-stranded regions that are formed are target sites for assembly of the RNA-induced silencing complex (RISC), which includes an endonuclease of the Argonaute family that cleaves and hence silences the mRNA.

The work that resulted in the initial description of the molecular process underlying RNA interference was carried out in the late 1990s with C. elegans. Since then, RNA interference has been shown to occur in most eukaryotes, the few exceptions including Saccharomyces cerevisiae, and interference has been linked to various events that involve RNA degradation but were previously thought to be unrelated. For example, the movement of some types of transposable element involves a double-stranded RNA intermediate that can be degraded by a process now known to be RNA interference. This is one way in which eukaryotes prevent the wholesale proliferation of transposons within their genomes. Genetic engineers had also been puzzled by the ability of some organisms, especially plants, to silence new genes that had been inserted into their genomes by cloning techniques. We now know that this type of silencing can occur if the transgene is inserted, by chance, upstream of a promoter that directs synthesis of an antisense RNA copy of all or part of the gene. This RNA can then base-pair with the sense mRNA produced from the transgene’s own promoter to form a double-stranded RNA that triggers the RNA interference pathway (Figure 12.25). Other phenomena in diverse organisms, variously known as quelling, cosuppression, and post-transcriptional gene silencing, are all now known to be different guises of RNA interference.

![Figure 12.24](image-url) The RNA silencing pathway. The double-stranded viral RNA is cut by Dicer, giving double-stranded siRNAs. Single-stranded versions of the siRNAs then base-pair with viral mRNAs, inducing assembly of an RNA-induced silencing complex (RISC), which contains an Argonaute endonuclease that cleaves and hence silences the mRNA.

![Figure 12.25](image-url) RNA interference explains why transgenes are sometimes inactive. For clarity, the mRNA and antisense RNA are shown being transcribed from different copies of the inserted transgene. They could also come from a single transgene that is transcribed from both its own promoter and an endogenous promoter.
MicroRNAs regulate genome expression by causing specific target mRNAs to be degraded

A link between silencing of viral RNAs and specific degradation of endogenous mRNAs was made when it was discovered that many eukaryotes have more than one type of Dicer protein. The fruit fly D. melanogaster, for example, has two Dicer enzymes. It turns out that the second type of Dicer in Drosophila works not with viral RNAs but with endogenous molecules called foldback RNAs, which are coded by the fruit fly DNA and synthesized by RNA polymerase II. The name foldback is given to these RNAs because they can form intranstrand base pairs, giving rise to a stem–loop structure (Figure 12.26). The stem can be cut by Dicer, releasing short double-stranded molecules, each about 21 bp, called microRNAs (miRNAs). One strand of each double-stranded molecule is degraded, giving functional miRNAs. A few miRNAs are obtained by a slightly different method, not by transcription of a gene for a precursor miRNA but from an intron cut out of the mRNA of a protein-coding gene. Part of the intron RNA folds up to form the stem–loop structure, which is then processed by Dicer as described above.

Each miRNA is complementary to part of a cellular mRNA and hence basepairs with this target, stimulating assembly of a RISC. Often the miRNA annealing site is present in the 3′-untranslated region of the target mRNA, sometimes in multiple copies (Figure 12.27). Cleavage by Argonaute therefore does not disrupt the coding region of the mRNA but will lead to detachment of the poly(A) tail. Loss of the poly(A) tail might interfere with the initiation of translation, which is inefficient if this part of the mRNA is absent. Alternatively, removal of the poly(A) tail might target the mRNA for degradation by the exosome or one of the other non-specific pathways for mRNA turnover. Whatever the precise mechanism, cleavage by Argonaute leads to the mRNA being silenced.

The first miRNA silencing system to be characterized involved the C. elegans genes called lin-4 and let-7, both of which code for foldback RNAs that generate miRNAs after cleavage by Dicer. A mutation in either of these two genes causes defects in the worm’s development pathway, indicating that this type of RNA degradation is not simply a means of getting rid of aberrant or potentially harmful miRNAs but instead plays a fundamental role in regulating composition of the transcriptome. Further support for this notion was provided by other studies of C. elegans miRNAs, which revealed that these molecules are involved in biological events as diverse as cell death, specification of neuron cell types, and control of fat storage. Genome analysis shows that most humans have the capacity to synthesize about 1000 miRNAs, but together these can target mRNAs from over 10,000 genes, possibly because miRNAs from different genes share the same miRNA binding sequence or possibly because a precise match between miRNA and mRNA is not needed in order for the mRNA to be captured by a RISC. Some miRNA genes are located close to the protein-coding genes whose mRNAs are targeted by the miRNA. In these cases, it is possible that the same regulatory proteins control both mRNA and miRNA synthesis. This would allow synthesis of the miRNA to be directly coordinated with repression of the protein-coding gene. The mRNA would therefore be degraded immediately after its synthesis is switched off. But in many other cases the miRNA and protein genes are not co-located, and the way in which mRNA synthesis and degradation are coordinated is not clear.

12.4 Influence of RNA processing on the composition of a transcriptome

As well as synthesis and degradation, most RNAs are processed by cutting and, in some cases, joining reactions that convert the initial transcripts into functional molecules. For many types of RNA, these processing events are part of their routine synthesis pathway and do not include any significant post-transcriptional control points that influence the composition of the transcriptome. We believe that this is true for the processing of pre-rRNAs and pre-tRNAs (Section 1.2) and
for the splicing pathways that result in removal of introns from the precursors of some tRNAs, rRNAs, and organellar transcripts.

One aspect of RNA processing that is of definite importance in influencing transcriptome composition is the splicing of eukaryotic pre-mRNA. This is because alternative splicing pathways can generate a range of mRNAs from a single pre-mRNA. We must therefore study how the splicing of pre-mRNAs is regulated in order to complete our understanding of the events that determine the composition of a eukaryotic transcriptome.

The splicing pathway for eukaryotic pre-mRNA introns

With the vast majority of pre-mRNA introns, the first two nucleotides of the intron sequence are 5'-GU-3' and the last two are 5'-AG-3'. They are therefore called GU-AG introns. A small minority do not fall into this class and, based on their boundary sequences, were originally called AU-AC introns, although now that more examples have been characterized it has become clear that these sequences are variable, and not all introns of this type have the AU and/or AC motifs. The splicing pathway is similar for both types of intron, with differences only in the details, and we will therefore focus our attention on the GU-AG group.

The conserved GU and AG motifs were recognized soon after introns were discovered, and it was immediately assumed that they must be important in the splicing process. As intron sequences started to accumulate in the databases, it was realized that these motifs are merely parts of the longer consensus sequences, spanning the 5'- and 3'-splice sites, that we met in Section 5.1 when we considered the methods used to identify intron boundaries when a eukaryotic genome sequence is annotated. Other conserved sequences are present in some but not all eukaryotes. Introns in higher eukaryotes usually have a polypyrimidine tract, which is a pyrimidine-rich region located just upstream of the 3'-end of the intron sequence (Figure 12.28). This tract is less frequently seen in yeast introns, but these have an invariant 5'-UACUAAC-3' sequence, located between 18 and 140 nucleotides upstream of the 3'-splice site, which is not present in higher eukaryotes.

The conserved sequence motifs indicate important regions of GU-AG introns, regions that we would anticipate either acting as recognition sequences for RNA-binding proteins involved in splicing or playing some other central role in the process. Early attempts to understand splicing were hindered by technical problems (in particular, difficulties in developing a cell-free splicing system with which the process could be probed in detail), but during the 1990s there was an explosion of information. This work showed that the splicing pathway can be divided into two steps (Figure 12.29):

- Cleavage of the 5'-splice site, also called the donor site, occurs by a trans-esterification reaction promoted by the hydroxyl group attached to the 2'-carbon of an adenosine nucleotide located within the intron sequence. In yeast, this branch site is the last adenosine in the conserved UACUAAC sequence. The result of the hydroxyl attack is cleavage of the phosphodiester bond at the donor site, accompanied by formation of a new 5'-2' phosphodiester bond linking the first nucleotide of the intron (the G of the GU motif) with the internal adenosine. This means that the intron loops back on itself to create a lariat structure.
• Cleavage of the 3′-splice site (the acceptor site) and joining of the exons result from a second transesterification reaction, this one promoted by the 3′-OH group attached to the end of the upstream exon. This group attacks the phosphodiester bond at the acceptor site, cleaving it and releasing the intron as the lariat structure, which is subsequently converted back to a linear RNA and degraded. At the same time, the 3′-end of the upstream exon joins to the newly formed 5′-end of the downstream exon, completing the splicing process.

In a chemical sense, intron splicing is not a great challenge for the cell. The difficulty lies with the topological problem caused by the substantial distance that might lie between splice sites, possibly a few tens of kilobases, representing 100 nm or more if the mRNA is in the form of a linear chain. A means is therefore needed of bringing the splice sites into proximity. This is the role of small nuclear ribonucleoproteins (snRNPs) (Section 12.1), which, together with other accessory proteins, attach to specific positions in the pre-mRNA and form a series of complexes, the most important of which is the spliceosome, the structure within which the actual splicing reactions occur.

Splicing initiates with formation of complex E (Figure 12.30). This complex comprises U1-snRNP, which binds to the donor site, partly by RNA-RNA base pairing, and the protein factors SF1, U2AF1, and U2AF2, which make protein-RNA contacts with the branch site, the polypyrimidine tract, and possibly the acceptor site. The next step is conversion of complex E to complex A, also called the prespliceosome complex, which occurs when U2-snRNP attaches to the branch site, probably not by base pairing but instead by an interaction between the branch site and one of the proteins associated with U2-snRNP. U1- and U2-snRNPs have an affinity for each other, and this draws the donor site toward the branch point. Complex B (the precatalytic spliceosome) is then formed when U4-, U5-, and U6-snRNPs attach to the intron. Their arrival results in additional interactions that bring the acceptor site close to the donor site and the branch point. U1- and U4-snRNPs then leave the complex, giving rise to the spliceosome. All three key positions in the intron are now in proximity and the cutting and joining reactions can take place, catalyzed by U2- and U6-snRNPs. The initial product of the splicing reaction is a post-spool lysis complex, which dissociates into the spliced mRNA and the intron lariat, the latter still attached to U2-, U5-, and U6-snRNPs.

The splicing process must have a high degree of precision

Precision is essential in splicing. If a cut is made just one nucleotide away from the actual exon–intron boundary position, then the open reading frame of the transcript will be disrupted and the mRNA that is made will be nonfunctional. At a different level, precision is needed in the selection of splice sites. All donor sites have a similar sequence, as do all acceptor sites, so if a pre-mRNA contains two or more
introns, then there is the possibility that the wrong splice sites could be joined, resulting in exon skipping, which is the loss of an exon from the mature mRNA (Figure 12.31A). Equally unfortunate would be selection of a cryptic splice site, a site within an intron or exon that has sequence similarity with the consensus motifs of real splice sites (Figure 12.31B). Cryptic sites are present in most pre-mRNAs and must be ignored by the splicing apparatus.

Key players in the control of splice site selection are short sequences called exonic splicing enhancers (ESE), exonic splicing silencers (ESS), intronic splicing enhancers (ISE), and intronic splicing silencers (ISS). These sequences act as binding sites for proteins that, when attached to an enhancer or silencer sequence, activate or repress selection of the adjacent splice site. Most of the proteins that bind to enhancers are SR proteins, so-called because they contain a domain that is rich in serine (abbreviated S) and arginine (abbreviated R). SR proteins attach to the CTD of the largest subunit of RNA polymerase II soon after the CTD becomes phosphorylated during the initiation stage of transcription. They ride with the polymerase as it synthesizes the transcript and are deposited at their splicing enhancer sequences as soon as these are transcribed. Electron
microscopic studies have shown that transcription and splicing occur together, and the discovery of splicing factors that have an affinity for RNA polymerase II provides a biochemical basis for this observation. When attached to a splicing enhancer, SR proteins appear to participate in several steps in the splicing pathway, including the establishment of a connection between bound U1-snRNP and the bound U2AF proteins in complex E. This is perhaps the clue to their role in splice site selection, formation of complex E being the critical stage in the splicing process, as it identifies which sites will be linked.

Less is known about the mode of action of exonic and intronic splicing silencers. These sequences act as binding sites for heterogeneous nuclear ribonucleoproteins (hnRNPs). However, as their name implies, these are a broad group of RNA–protein complexes, which play several roles in the nucleus, most of which involve binding to RNAs. The particular types called hnRNP1 and hnRNPL appear to bind at specific positions in introns and exons and are associated with splicing regulation. Their mode of action might simply be to block access to one or more of the splice sites, branch site, and/or polypyrimidine site, interfering with the assembly of complex E and therefore preventing splicing.

Enhancer and silencer elements specify alternative splicing pathways

Alternative splicing is now looked on as a major component of the processes that determine the composition of a transcriptome. As we discussed in Section 7.3, 75% of all human protein-coding genes, representing 95% of those with two or more introns, undergo alternative splicing, giving rise to an average of four different spliced mRNAs, or isoforms, per gene. These alternative splicing events can be placed in four categories:

- **Exon skipping**, which results in one or more exons being left out of the final mRNA (see Figure 12.31A).
- **Alternative site selection**, in which the usual donor or acceptor site is ignored and a second site is used in its place. This is equivalent to cryptic splice site selection (see Figure 12.31B).
- **Alternative exons**, where the mRNA contains one or the other of a pair of exons but not both at the same time.
- **Intron retention**, which occurs when an intron that is usually spliced out of the pre-mRNA is retained in the final mRNA. The retained intron might be in the upstream or downstream untranslated region of the mRNA, but some examples are known where the intron contains an open reading frame and so contributes to the amino acid sequence of the protein coded by the mRNA.

Two examples will illustrate the importance and complexity of alternative splicing. The first of these concerns sex determination, a fundamental aspect of the biology of any organism, which in Drosophila is determined by an alternative splicing cascade. The first gene in this cascade is sxl, whose transcript contains an optional exon that, when spliced to the one preceding it, results in an inactive version of protein SXL. In females the splicing pathway is such that this exon is
skipped so that functional SXL is made (Figure 12.32). SXL promotes selection of an alternative splice site in a second transcript, tra, by directing U2AF2 away from its normal acceptor site to a second site further downstream. The resulting female-specific TRA protein is again involved in alternative splicing, this time by interacting with SR proteins to form a multifactor complex that attaches to a silencing element within an exon of a third pre-mRNA, dsx, promoting selection of a secondary, female-specific splice site in this transcript. The male and female versions of the DSX proteins are the primary determinants of Drosophila sex.

Figure 12.32 Regulation of splicing during expression of genes involved in sex determination in Drosophila.

(A) The cascade begins with sex-specific alternative splicing of the sxl pre-mRNA. In males, all exons are present in the mRNA, but this means that a truncated protein is produced because exon 3 contains a termination codon. In females, exon 3 is skipped, leading to a full-length, functional SXL protein. (B) In females, SXL blocks the acceptor site in the first intron of the tra pre-mRNA. U2AF2 is unable to locate this site and instead directs splicing to a cryptic site in exon 2. This results in an mRNA that codes for a functional TRA protein. In males, there is no SXL, so the acceptor site is not blocked and a dysfunctional mRNA is produced. (C) In males, exon 4 of the dsx pre-mRNA is skipped. The resulting mRNA codes for a male-specific DSX protein. In females, TRA stabilizes the attachment of SR proteins to an exonic splicing enhancer located within exon 4, so this exon is not skipped, resulting in the mRNA that codes for the female-specific DSX protein. The two versions of DSX are the primary determinants of male and female physiologies. The female dsx mRNA ends with exon 4 because the intron between exons 4 and 5 has no donor site, meaning that exon 5 cannot be ligated to the end of exon 4. Instead, a polyadenylation site at the end of exon 4 is recognized in females.
Chapter 12: Transcriptomes

The second example of alternative splicing illustrates the multiplicity of mRNAs synthesized from some primary transcripts. The human slo gene codes for a membrane protein that regulates the entry and exit of potassium ions into and out of cells. The gene has 35 exons, eight of which are involved in alternative splicing events (Figure 12.33). The alternative splicing pathways involve different combinations of the eight optional exons, leading to over 500 distinct mRNAs, each specifying a membrane protein with slightly different functional properties. The human slo genes are active in the inner ear and determine the auditory properties of hair cells on the basilar membrane of the cochlea. Different hair cells respond to different sound frequencies between 20 and 20,000 Hz, their individual capabilities determined in part by the properties of their Slo proteins. Alternative splicing of slo genes in cochlear hair cells therefore determines the auditory range of humans.

At present we can describe a substantial number of alternative splicing pathways, and for some steps in these pathways we know how the choice between alternative outcomes is regulated. Our expectation is that, underlying the apparent complexity, there is a splicing code of some description, which explains the outcomes of the various interactions that can occur between enhancers, silencers, and their binding proteins. At present, our knowledge of that code is rudimentary, and as such there are still important gaps in our understanding of how alternative splicing is controlled.

12.5 TRANSCRIPTOMES IN RESEARCH

We have explored the coding and noncoding contents of a transcriptome, learned how the composition of a transcriptome is cataloged, and studied the biological events involved in synthesis, degradation, and processing of the RNAs present in a transcriptome. To conclude this chapter, we now examine the importance of transcriptome studies in biological research.

Transcriptome analysis as an aid to genome annotation

In Section 5.3 we looked at the various ways in which RNA mapping is used during a genome annotation project, in order to identify the segments of a genome sequence that are transcribed. Those methods included the analysis of microarray data and RNA-seq reads, which are the two principal techniques used to analyze transcriptome compositions. Indeed, transcriptome analysis and RNA mapping are essentially the same procedure, the one difference being that the former is carried out in order to understand the RNA content of a cell and the latter to identify genes in a DNA sequence. The complementarity of the two types of analysis means that in many projects no distinction is made between them, and analysis of a transcriptome goes hand in hand with annotation of a genome. To illustrate the importance of transcriptome analysis in genome annotation, we will examine how the cataloging of D. melanogaster transcriptomes has added unprecedented detail to the annotation of the fruit fly genome.

Because gene expression patterns, and hence transcriptomes, vary in different tissues of a multicellular organism and change in response to environmental factors, a number of different RNA libraries must be prepared and sequenced in order to use transcriptome analysis to obtain a comprehensive genome annotation. In one particular project, Drosophila larvae, pupae, and adults were dissected and polyadenylated RNA, the bulk of which comprises transcripts of protein-coding genes (Section 5.3), was purified from a variety of tissues including different parts of the nervous, digestive, reproductive, and endocrine systems as well as from the epidermis and muscles. Additional libraries were prepared from adult flies that had been exposed to heat shock and cold shock and from adults and larvae.
Chapter 12: Transcriptomes

Exposed to environmental contaminants such as heavy metals, herbicides, and pesticides. Altogether 12.4 billion sequence reads were obtained, which could be assembled into 304,788 transcripts.

The number of transcripts identified in this project greatly exceeds the approximately 14,000 protein-coding genes that are recognized in the D. melanogaster genome annotation (see Table 7.4). This discrepancy can, of course, be explained by the use of alternative promoters and alternative splicing pathways, which enable a single gene to specify more than one transcript. Some transcripts also have alternative polyadenylation sites, meaning that they can be cut at two or more alternative positions prior to attachment of the poly(A) tail that marks the 3′-end of an RNA synthesized by RNA polymerase II (see Figure 12.1). Overall, the number of genes shown to give rise to more than one transcript was not unexpected: 42% of the total compared to 95% for mammals. The importance of alternative splicing in specifying the male and female characteristics of fruit flies was also emphasized, with 575 alternative splicing events occurring in a sex-specific manner, most of these in transcripts synthesized in the male testes or female ovaries. Many other examples of a single gene giving rise to a series of alternatively spliced mRNA isoforms, with different tissue specificities, were identified.

What was surprising about the results of this analysis of the Drosophila transcriptome was the extreme complexity of the transcription patterns for a small number of genes. Forty-seven genes were identified that each gave rise to over 1000 different transcripts, these isoforms making up 50% of the transcriptome catalog as a whole (Figure 12.34). Many of the transcripts of these 47 genes were tissue-specific, with 56% of the RNAs present only in the embryo, and 27% present only in neuronal tissue. A second surprise was that many of these multiple alternative transcripts specified the same protein, the differences between the RNA isoforms not affecting the open reading frame. An example is provided by the pUf68 gene, also called half pint (Drosophila geneticists have a penchant for assigning interesting names to fruit fly genes), which codes for a protein that regulates alternative splicing of a series of genes in adult ovarian tissues. Mutations in pUf68 result in defects in the development of eggs, which are smaller than normal, giving rise to the half pint name. The primary transcript of pUf68 can follow a variety of alternative splicing pathways, giving rise to over 100 isoforms, but there is only one half pint protein. This is because the alternative splicing events involve only exons that are present in the upstream untranslated region of the pre-mRNA and so do not affect the open reading frame. How do we interpret results such as these? As with long noncoding RNAs, there is a temptation to dismiss the multiplicity of transcripts, all coding for the same protein, as transcriptional noise. According to this hypothesis, the route taken to synthesis of the final mRNA is immaterial, so long as that mRNA is made in the correct tissue at the correct time. Whatever the explanation, the discovery in fruit flies and other eukaryotes of

![Figure 12.34 Alternative splicing in Drosophila transcriptomes.](A) Genes are placed into classes depending on how many transcripts each gives rise to. The majority of genes, 79%, give rise to less than five transcripts. At the other end of the scale, 47 genes (0.2% of the total) each specify over 1000 RNAs. (B) Contribution of the transcripts of each class of gene to the transcriptome as a whole. Half of all RNAs derive from the 47 genes that give rise to over 1000 transcripts each. (Data from Brown JB, Boley N, Eisman R, et al. [2014] Nature 512:393–399.)
individual genes that give rise to a multitude of RNA isoforms adds complexity to transcriptome studies and emphasizes the importance of detailed analyses of transcriptomes as an adjunct to genome annotation.

**Cancer transcriptomes**

Transcriptome restructuring as a result of cancer was first described in 1997, when it was shown that 289 mRNAs are present in significantly different amounts in the transcriptomes of normal colon epithelial cells compared with cancerous colon cells; about half of these mRNAs also display an altered abundance in pancreatic cancer cells. These were important observations, as understanding the differences between the transcriptomes of normal and cancerous cells points to biochemical differences and hence to new ways of treating the cancers. Transcriptome studies also have applications in cancer diagnosis. The initial breakthrough in this area came in 1999, when it was shown that the transcriptome of acute lymphoblastic leukemia cells is different from that of acute myeloid leukemia cells. In this work, 27 lymphoblastic and 11 myeloid cancers were studied, and although all the transcriptomes were slightly different, the distinctions between the two types of cancer were sufficient for unambiguous identifications to be made. The significance of this work lay with the improved remission rates that are achievable if a cancer is identified accurately at an early stage, before clear morphological indicators are seen. This is not relevant with these two types of leukemia because these can be distinguished by nongenetic means, but it is important with other cancers such as non-Hodgkin’s lymphoma. The commonest version of this disease is called diffuse large B-cell lymphoma, and for many years it was thought that all tumors of this type were the same. Transcriptome studies changed this view and showed that B-cell lymphoma can be divided into two distinct subtypes. The distinctions between the transcriptomes of the two subtypes enable each one to be related to a different class of B cells, stimulating and directing the search for specific treatments that are tailored to each lymphoma.

Transcriptome analysis is also providing new information on breast cancer. In the early 2000s, this approach enabled breast cancer to be divided into five subtypes, called luminal A, luminal B, HER2-positive, basal-like, and normal-like. Each of these subtypes has different characteristics and, in effect, they are distinct diseases that affect the same tissue. Detailed examination of breast cancer transcriptomes has also identified groups of genes whose expression profiles enable the likely progress of breast cancer to be predicted, in particular the risk of metastasis, which results in cells from one cancer spreading to other places in the body and initiating new tumors. The ability to use transcriptome profiling to predict the risk of metastasis at an early stage of the disease was an unexpected discovery, as it had been thought that the cellular switches that initiated metastasis were not activated until a much later step in progression of the cancer. Transcriptome studies have also revealed unexpected aspects of the genetic basis to metastasis. In particular, the risk of metastasis appears to be associated with the presence in tumor transcriptomes of mRNAs usually found in tissues that are recovering from trauma such as wounding, not just in the case of breast cancer but also with cancers of other epithelial tissues. This observation has contributed to the hypothesis that tumors are wounds that do not heal, which has stimulated new avenues of research into possible therapies.

Transcriptomes are therefore having an important impact on our understanding of breast and other cancers. The challenge now lies not so much in adding greater detail to the transcriptome catalogs for different cancers and different stages of cancer, but with the need to translate the information that we now have about cancer transcriptomes into the clinical setting. Microarray and RNA-seq analyses are expensive to carry out and are not yet routinely used in predicting the prognosis of individual patients. Considerable work is also needed in order to assess the actual importance in cancer of a gene whose expression is up- or down-regulated during progression of that disease. Many changes in expression
pattern are likely to result from the presence of the cancer, rather than playing an active role in the development of a tumor. Transcriptome analysis, by providing a detailed description of gene expression changes, is a starting point to identify genes central to cancer progression, but many other types of genetic and biochemical studies are needed to exploit these initial findings.

**Transcriptomes and the responses of plants to stress**

To many nonbiologists, the notion that plants lead stressful lives is rather surprising and at odds with the perception of a plant spending its life basking in the warm sunlight. In fact, the ability of plants to respond to stresses such as high or low temperature, drought, or the presence of insect pests is extremely important. Being immobile, plants cannot mitigate these stresses by running away or by migrating to a new area where a particular stress is less severe in its impact. Plants therefore have to find biochemical and physiological solutions to the challenges provided by environmental stresses. As well as their intrinsic interest, these solutions have applied importance because engineering them to be more effective is looked on as a potential means of improving the yields of crop plants and maintaining those yields as the climate changes.

Plant stress responses are so important that they have been studied for many decades by biologists, using a variety of different research approaches. This work has led to a good understanding of the basis to several individual types of stress response. For example, most plants synthesize proteins called dehydrins in response to environmental stresses that result in cell dehydration: these stresses including drought, high temperature, or high saline conditions. Dehydrins contain a large proportion of hydrophilic amino acids and have open coil structures, so many of these amino acids are exposed and can form hydrogen bonds with water molecules. As a result, the dehydrin proteins help to maintain the cellular water content.

Although individual stress responses have been characterized, the response of a plant to a mixture of stresses is less well understood. A combination of stresses is not unusual in the natural environment where, for example, high humidity might be associated with an increase in bacterial pathogen numbers. When exposed to two or more stresses, does the plant have to operate each individual response process independently, or do these processes share components so that the plant is able to mount a coordinated response that deals with the stresses collectively? These questions have been addressed by studies of transcriptomes in *Arabidopsis thaliana* exposed to different combinations of temperature stress, light intensity, salinity, and presence of bacterial pathogens, the last of these stresses elicited by applying peptides from the bacterial flagellin protein to the leaves. A total of 210 microarray experiments were performed to analyze the responses of 10 ecotypes of *A. thaliana* to different combinations of stresses. An ecotype is a population of plants from a single geographical location, adapted to the environmental conditions of that region. For each combination of stresses, RNAs that fell into one of five categories were identified (Figure 12.35):

- **Independent-mode RNAs** respond to either stress A or stress B by increasing or decreasing in amount and respond in the same way to the combined stress A + B. Clearly, these RNAs code for proteins needed for the response to one stress but not the other. Examples are chloroplast proteins that play a specialized role in enabling the plant to withstand high light intensity.

- **Similar-mode RNAs** show the same change in abundance in response to stress A, stress B, and stresses A + B. These RNAs code for proteins that form part of a common response to A and B.

- **Combinatorial-mode RNAs** show the same response to A or B but a different response to A + B. In some cases a combinatorial RNA responds to the
individual stresses but not to the combined stresses, and in other cases a response is only seen when the stresses are combined.

- Canceled-mode RNAs respond differently to the two individual stresses and do not respond to the combined stress.

- Prioritized-mode RNAs respond to stresses A and B differently, but in this case the response to A + B is the same as one of the individual responses.

Considering these five categories of response, we can see that the three latter types—combinatorial, canceled, and prioritized modes—are the most interesting, as the responses of these RNAs to a combination of stresses cannot be predicted from their responses to the individual stresses. In other words, these categories indicate components of the individual stress pathways that function differently when two stresses are combined. Canceled-mode RNAs, for example, specify proteins that are needed to combat one stress but are not needed when that stress occurs in combination with a second stress. In contrast, the combinatorial mode includes RNAs whose products are needed only when two stresses are experienced at the same time.

For most combinations of stresses, the majority of transcripts that displayed a response fell into one of the three unpredictable modes, these three modes in total accounting for, on average, 85% of the transcripts. This observation indicates that there is substantial interconnectivity between stress response pathways and that the biochemical events that enable the plant to respond to one stress are often different from the events that address two or more stresses. Importantly, the canceled-mode transcripts indicate cases where the response to one stress might actually interfere with the response to a second stress. This appears to be the case when plants are exposed to high temperature at the same time as high salinity. Detailed analysis of the transcriptomes for this pair of stresses suggested that the response to heat stress dominated the response to salinity, implying that plants experiencing such a combination in their natural environment are unlikely to thrive. Transcriptome analysis as a whole therefore resets the agenda for more broadly based studies of plant stress responses.

Figure 12.35 Transcriptional modes for RNAs involved in Arabidopsis stress responses. Each panel shows the possible responses of a transcript to stress A, stress B, and the combined stress A + B. Dashed lines indicate the amount of transcript in the absence of the stress, with lines above and below this level indicating an increase or decrease, respectively, in the amount of transcript. (Data from Rasmussen S, Barah P, Suarez-Rodriguez MC et al. [2013] *Plant Physiol.* 161:1783–1794.)
SUMMARY

- The transcriptome is the collection of RNA molecules present in a cell.
- The mRNA fraction makes up a relatively small part of the transcriptome of most cells, but comprises many different transcripts.
- Transcriptomes contain a variety of short noncoding RNAs with diverse functions.
- Transcriptomes also contain long noncoding RNAs whose functions, if any, are not well understood.
- The content of a transcriptome can be cataloged by microarray studies and by RNA sequencing.
- The components of a transcriptome are synthesized by RNA polymerases. In eukaryotes, there are three different RNA polymerases, each responsible for transcribing a different set of nuclear genes.
- The positions at which transcription must begin are marked by promoter sequences. Some genes have alternative promoters that give rise to different versions of the transcript specified by the gene.
- Initiation of transcription in bacteria is regulated by the combined action of activator and repressor proteins. For some genes, additional control is also exerted by the premature termination of transcripts that are not needed.
- Initiation of transcription in eukaryotes is regulated mainly by activator proteins called transcription factors, which interact with the initiation complex via the mediator protein.
- RNA can be turned over by nonspecific processes and by RNA silencing, the latter enabling specific mRNAs to be targeted for degradation.
- Most RNAs are processed by cutting and, in some cases, joining reactions that convert the initial transcripts into the functional molecules.
- Some RNAs are also processed by splicing—the removal of introns. Splicing of pre-mRNAs is particularly important in eukaryotes as the transcripts of some genes are able to follow alternative splicing pathways, giving rise to different mRNAs and hence different protein products.
- During alternative splicing, splice site selection is controlled by short enhancer and silencer sequences, which are the binding sites for regulatory proteins.
- Transcriptome analysis can aid genome annotation by identifying those parts of a sequence that are transcribed. The transcriptome restructuring that occurs during cancer enables different subtypes of cancer to be distinguished and also reveals changes in gene expression patterns associated with the disease. The response of plants to environmental stress is also being studied by transcriptome analysis.

SHORT ANSWER QUESTIONS

1. Describe the various types of small noncoding RNAs that are found in eukaryotic transcriptomes.

2. What evidence is there to suggest that at least some long noncoding RNAs are functional?
3. Outline how a transcriptome is characterized by (A) microarray analysis and (B) RNA sequencing.

4. Compare the structures of the three eukaryotic RNA polymerases with that of the bacterial enzyme. What are the key features of the RNA polymerases that transcribe organelle genes?

5. Distinguish between the structures of the promoters for (A) the bacterial RNA polymerase, (B) RNA polymerase I, (C) RNA polymerase II, and (D) RNA polymerase III.

6. How is the synthesis of bacterial mRNAs regulated?

7. Describe the role of the mediator protein during initiation of eukaryotic transcription.

8. Outline the processes for nonspecific degradation of RNA in bacteria and eukaryotes.

9. Explain how specific RNAs are targeted for degradation by siRNAs and miRNAs.

10. Outline the splicing pathway for a eukaryotic pre-mRNA.

11. What is alternative splicing and how is it regulated?

12. In what ways are transcriptome studies being used in cancer research?

IN-DEPTH PROBLEMS

1. Construct a hypothesis to explain why eukaryotes have three RNA polymerases. Can your hypothesis be tested?

2. A model for control of transcription of the lactose operon in *Escherichia coli* was first proposed by François Jacob and Jacques Monod in 1961 (Jacob, F. and Monod, J. [1961] Genetic regulatory mechanisms in the synthesis of proteins. *J. Mol. Biol.* 3:318–356). Explain the extent to which their work, which was based almost entirely on genetic analysis, provided an accurate description of the molecular events that are now known to occur.

3. To what extent is *E. coli* a good model for the regulation of transcription initiation in eukaryotes? Justify your opinion by providing specific examples of how extrapolations from *E. coli* have been helpful and/or unhelpful in the development of our understanding of equivalent events in eukaryotes.

4. Discontinuous genes are common in higher organisms but virtually absent in bacteria. Discuss the possible reasons for this.

5. To what extent has the study of AU-AC introns provided insights into the details of GU-AG intron splicing?

FURTHER READING

**Components of transcriptomes**


Methods for studying transcriptomes


RNA polymerases, promoters, and RNA synthesis


Control of RNA synthesis


Degradation of RNA


Splicing


**Case studies**


The proteome is the collection of protein molecules present in a cell. The proteome is therefore the final link between the genome and the biochemical capability of the cell, and characterization of the proteomes of different cells is one of the keys to understanding how the genome operates and how dysfunctional genome activity can lead to diseases. Transcriptome studies can address these issues only in part. Examination of the transcriptome gives an accurate indication of which genes are active in a particular cell but gives a less accurate indication of the proteins that are present. This is because the factors that influence protein content include not only the amount of mRNA that is available but also the rate at which the mRNAs are translated into protein and the rate at which the proteins are degraded. Additionally, the protein that is the initial product of translation may not be active, as some proteins must undergo physical and/or chemical modification before becoming functional. Determining the amount of the active form of a protein is therefore critical to understanding the biochemistry of a cell or tissue.

The issues that we must examine regarding the proteome are very similar to the issues that interested us in Chapter 12 when we studied transcriptomes. First, we will explore the various methods that are used to catalog the components of a proteome and to understand how a proteome functions within a cell. Then we must study the events involved in synthesis, degradation, and processing of the components of a proteome. Finally, we will examine more closely the link between the proteome and the biochemistry of the cell.

13.1 STUDYING THE COMPOSITION OF A PROTEOME

The methodology used to study proteomes is called proteomics. Strictly speaking, proteomics is a collection of diverse techniques that are related only in their ability to provide information on a proteome. That information encompasses not only the identities of the constituent proteins that are present but also factors such as the functions of individual proteins and their localization within the cell. The particular technique that is used to study the composition of a proteome is called protein profiling or expression proteomics.

Protein profiling is usually carried out in two stages:

- In the first stage, the individual proteins in a proteome are separated from one another.
- In the second stage, the proteins are identified, usually by mass spectrometry.

This basic format encompasses two different approaches called top-down and bottom-up proteomics (Figure 13.1). The difference is that in top-down
proteomics, individual proteins are directly examined by mass spectrometry, whereas in bottom-up proteomics, the proteins are broken into peptides by treatment with a sequence-specific protease, such as trypsin, prior to mass spectrometry.

**The separation stage of a protein profiling project**

In order to characterize a proteome, it is first necessary to prepare pure samples of its constituent proteins. How difficult this is depends on the complexity of the proteome. Separation of the 10,000–20,000 proteins in some mammalian proteomes requires more sophisticated methods than are needed for the less complex proteomes of bacteria or mammalian cell fractions (for example, mitochondria), which might contain fewer than 1000 proteins. The choice of separation technique is therefore dictated in part by the complexity of the proteome that is being studied.

Polyacrylamide gel electrophoresis (PAGE) is the standard method for separating the proteins in a complex mixture. Depending on the composition of the gel and the conditions under which the electrophoresis is carried out, different chemical and physical properties of proteins can be used as the basis for their separation. One technique makes use of the detergent called sodium dodecyl sulfate (SDS), which denatures proteins and confers a negative charge that is roughly equivalent to the length of the unfolded polypeptide. Under these conditions, the proteins separate according to their molecular masses, the smallest proteins migrating more quickly toward the positive electrode. Alternatively, proteins can be separated by isoelectric focusing in a gel containing chemicals that establish a pH gradient when the electrical charge is applied. In this type of gel, a protein migrates to its isoelectric point, the position in the gradient where its net charge is zero.

When a complex proteome is being studied, the two versions of PAGE are often combined in **two-dimensional gel electrophoresis**. In the first dimension, the proteins are separated by isoelectric focusing. The gel is then soaked in sodium dodecyl sulfate and rotated by 90°, and a second electrophoresis, separating the proteins according to their sizes, is carried out at a right angle to the first (Figure 13.2). This approach can separate several thousand proteins in a single gel, the proteins revealed as a complex pattern of spots when the gel is stained (Figure 13.3). Individual spots can therefore be cut out of the gel, and the proteins they contain can be purified. Cutting out 20,000 spots would clearly be a laborious process, and in practice two-dimensional PAGE is not used if the aim is to catalog every protein in a proteome. Instead, it is used to identify interesting proteins, such as those that have different abundances in two or more related proteomes: the healthy and diseased versions of a tissue, for example.

An alternative approach to protein separation by PAGE is provided by **column chromatography**. This method involves passing the protein mixture through a column packed with a solid matrix. The proteins in the mixture move through the matrix at different rates and so become separated into bands. The solution emerging from the column can then be collected as a series of fractions, with each individual protein present in a different fraction (Figure 13.4). The identity of the solid phase (the matrix or resin) and the composition of the mobile phase (the liquid used to move the proteins through the column) specify which of the variable
physicochemical properties of proteins are used to achieve separation. In protein profiling, the two most commonly used types of column chromatography are as follows:

- **In reverse-phase liquid chromatography (RPLC),** the solid phase is a matrix of silica particles whose surfaces are covered with nonpolar chemical groups such as hydrocarbons. The mobile phase is a mixture of water and an organic solvent such as methanol or acetonitrile. Most proteins have hydrophobic areas on their surfaces, which bind to the nonpolar matrix, but the stability of this attachment decreases as the organic content of the liquid phase increases. Gradually changing the ratio of the aqueous and organic components of the mobile phase therefore results in the elution of proteins according to their degree of surface hydrophobicity.

- **Ion-exchange chromatography** separates proteins according to their net electric charges. The matrix consists of polystyrene beads that carry either positive or negative charges. If the beads are positively charged, then proteins with a net negative charge will bind to them, and vice versa. The proteins can be eluted with a salt gradient, set up by gradually increasing the salt concentration of the buffer being passed through the column. The charged salt ions compete with the proteins for the binding sites on the resin, so proteins with low charges are eluted at low salt concentration, and those with higher charges are eluted at higher salt concentrations. The salt gradient therefore separates proteins according to their net charges. Alternatively, a pH gradient can be used. The net charge of a protein depends on the pH and, as described above, is zero at the pH corresponding to that protein’s isoelectric point. Gradually changing the pH of the mobile phase will result in the elution of proteins with different isoelectric points, again achieving their separation.

Compared with two-dimensional PAGE, column chromatography is less laborious to carry out and has the advantage that individual proteins can be collected as they elute from the column, avoiding the postseparation purification step needed to obtain a protein from its spot on a polyacrylamide gel. Column chromatography is usually carried out in a capillary tube with an internal diameter of
less than 1 mm, with the liquid phase being pumped at high pressure. This procedure, called **high-performance liquid chromatography (HPLC)**, has a high resolving power and enables proteins with very similar chromatographic properties to be separated (Figure 13.5).

To increase the resolving power, different types of chromatography column can be linked together, with each consecutive fraction from one column being fed into a second column, in which a further round of separation by a different procedure is carried out (Figure 13.6A). In this way, quite complex mixtures of proteins can be fully separated. Alternatively, proteins can initially be separated by one-dimensional PAGE, using either the SDS version or isoelectric focusing, and the resulting gel cut into segments (Figure 13.6B). The set of proteins present in each segment is then entered sequentially into the column chromatography system. If the top-down approach is being used, then rather than collecting protein fractions as they emerge from the column—the offline mode—the column can be directly attached to the mass spectrometer. Each protein is therefore analyzed by the spectrometer as it elutes from the column (Figure 13.6C). This online mode cannot be used with the standard version of bottom-up proteomics, because each protein must be treated with a protease to cut it into fragments prior to injection into the mass spectrometer. The online mode is, however, possible with the modification of the bottom-up approach called **shotgun proteomics**. With this method, the proteins are treated with the protease before the column.
chromatography step. These proteins could be the entire proteome, if it is not overly complex, or the mixture obtained from a segment of a one-dimensional PAGE gel. In either case, the column now separates peptides rather than intact proteins, and the online mode is used to inject the eluted molecules directly into the mass spectrometer.

The identification stage of a protein profiling project

Separation of the components of a proteome is followed by identification of the individual proteins, either directly, in top-down proteomics, or from the peptides resulting from proteolytic cleavage, if a bottom-up method is being used. Identification of proteins and peptides used to be a difficult proposition but advances in mass spectrometry, driven in part by the requirements of proteomics, have largely solved this problem.

Mass spectrometry is a means of identifying a compound from the mass-to-charge ratio (designated m/z) of the ions that are produced when molecules of the compound are exposed to a high-energy field. The first type of mass spectrometry to be used widely in protein profiling was matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF). This technique forms the basis for peptide mass fingerprinting, the bottom-up procedure that identifies individual peptides in the mixture obtained by protease cleavage of a protein purified by two-dimensional PAGE or column chromatography. The first step in MALDI-TOF is ionization of the peptides. This is achieved by absorbing the mixture into an organic crystalline matrix, usually made of a phenylpropanoid compound called sinapinic acid, which is excited with a UV laser. The excitation initially ionizes the matrix, with protons then donated to or removed from the peptide molecules to give the molecular ions [M + H]⁺ and [M − H]⁻, respectively, where M is the peptide. Ionization also results in vaporization of the peptides, which are then accelerated along the tube of the mass spectrometer by an electric field. The flight path can be direct from the ionization source to a detector, but often the ions are initially directed at a reflectron, which reflects the ion beam toward the detector (Figure 13.7). As well as enabling a longer flight path to be built into a machine of a defined size, the reflectron acts as a focusing device, ensuring that all ions with the same m/z value travel through the mass spectrometer at the same speed. This is critical because a time-of-flight spectrometer uses the time that an ion takes to reach the detector in order to calculate the mass-to-charge ratio for that ion. As the charge is always +1 or −1, the time of flight can easily be converted to a molecular mass, which in turn allows the amino acid composition of the peptide to be deduced. If a number of peptides from a single protein spot in a two-dimensional gel are analyzed, then the resulting compositional information can be related to the genome sequence in order to identify the gene

Figure 13.7 Use of MALDI-TOF in protein profiling. (A). In the matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF) mass spectrometer, the peptides are ionized by a pulse of energy from a laser and then accelerated down the column to the reflectron and onto the detector. The time of flight of each peptide depends on its mass-to-charge ratio. (B) The data are visualized as a spectrum indicating the m/z values of the peptides. The computer converts the m/z values into molecular masses and compares these masses with the predicted masses of all the peptides that would be obtained by protease treatment of all the proteins encoded by the genome of the organism under study. The protein that gave rise to the detected peptides can therefore be identified.
that specifies that protein. The amino acid compositions of the peptides derived from a single protein can also be used to check that the gene sequence is correct and, in particular, to ensure that exon-intron boundaries have been correctly located. This not only helps to ensure that the genome annotation is accurate but also allows alternative splicing pathways to be identified in cases where two or more proteins are derived from the same gene.

MALDI-TOF is usually able to identify a protein that has been purified from a gel or by column chromatography. It is less suitable for shotgun proteomics, because the larger number of peptides that are produced when a mixture of proteins is treated with a protease increases the possibility that two peptides will have similar m/z values and hence be indistinguishable when examined by MALDI-TOF. Two innovations have been introduced in recent years to improve the resolution of peptide mass spectrometry and hence provide support for the shotgun methods. The first of these is the use of electrospray ionization, which can be performed online between HPLC and mass spectrometry. A high voltage is applied to the solution emerging from the HPLC, generating an aerosol of charged droplets that evaporate, transferring their charges to the peptides dissolved within them. The advantage of this ionization method is that, as well as the [M + H]$^+$ and [M – H]$^-$ molecular ions, each with a single ionized group, multiple ionized molecules such as [M + nH]$^{n+}$ are also obtained. Generation of multiple ions with different m/z values from individual peptides increases the amount of information that can be used to infer the composition of that peptide.

The second innovation is to break peptides down into smaller fragments within the mass spectrometer. Fragmentation can be induced during the ionization step by use of a hard ionization method, one that injects greater quantities of energy into the molecules being ionized, causing bonds within those molecules to break. However, in peptide mass spectrometry, fragmentation is usually delayed until a later stage, by inducing collisions between the peptide molecular ions and inert atoms such as helium. These collisions cause peptide bonds to break, resulting in a variety of fragment ions whose m/z values reveal the composition of the original peptide. If sufficient fragment ions are obtained, then it might be possible to use the data to work out the sequence of the peptide. Knowing the sequence of the peptide enables a much more precise identification of the parent protein than is possible simply from compositional information. Collision-induced fragmentation is also utilized in top-down proteomics, because analysis of molecular ions derived from intact proteins is usually insufficient to distinguish all of the different proteins in a proteome. Fragment ions must therefore be obtained before a protein can be identified unambiguously.

Innovations involving ionization methods and the use of fragmentation have been accompanied by a diversification in the types of mass spectrometer employed in proteomics research. As well as time-of-flight configurations, other types of mass analyzer used with peptides and proteins include the following (Figure 13.8):

- The **quadrupole** mass analyzer has four magnetic rods placed parallel to one another, surrounding a central channel through which the ions must pass. Oscillating electrical fields are applied to the rods, deflecting the ions in a complex way so that their trajectories wiggle as they pass through the quadrupole. Gradually changing the field strengths enables ions with different m/z values to pass through the quadrupole without colliding with the rods.

- The **Fourier transform ion cyclotron resonance (FT-ICR)** mass analyzer includes an ion trap that captures individual ions and further excites them within a cyclotron, so they accelerate along an outward spiral; the vector of this spiral revealing the m/z ratio.

Mass analyzers can also be linked in series, further increasing the amount of information that can be obtained regarding a single peptide or protein. This is called **tandem mass spectrometry**. A typical configuration involves analysis of
the molecular ions in the first mass analyzer, followed by fragmentation and analysis of the fragment ions in the second mass analyzer.

**Comparing the compositions of two proteomes**

Often the aim of a protein profiling project is not to identify every protein in a single proteome but to understand the differences between the protein compositions of two different proteomes. If the differences are relatively large, then they will be apparent simply by looking at the stained gels after two-dimensional electrophoresis. However, important changes in the biochemical properties of a proteome can result from relatively minor changes in the amounts of individual proteins, and methods for detecting small-scale changes are therefore essential.

One possibility is to label the constituents of two proteomes with different fluorescent markers, and then run them together in a single two-dimensional gel. This is the same strategy as is used for comparing pairs of transcriptomes (see Figure 12.5). Visualization of the two-dimensional gel at different wavelengths enables the intensities of equivalent spots to be judged more accurately than is possible when two separate gels are obtained.

A more accurate alternative in bottom-up proteomics is to label peptides with an **isotope-coded affinity tag (ICAT)**. These are chemical groups that can be attached to a peptide. In one system, the tags are short hydrocarbon chains that contain either the common $^{12}\text{C}$ isotope of carbon or the less common $^{13}\text{C}$ isotope (Figure 13.9). The proteins in the proteomes are separated in the normal manner, and equivalent proteins from each proteome are recovered and treated with protease. One set of peptides is then labeled with $^{12}\text{C}$ tags and the other with $^{13}\text{C}$ tags. Because the $^{12}\text{C}$ and $^{13}\text{C}$ tags have different masses, the $m/z$ ratio of the molecular ion obtained from a peptide labeled with a $^{12}\text{C}$ tag will be different from that of an identical peptide labeled with a $^{13}\text{C}$ tag. The peptides from the two proteomes are therefore run through the mass spectrometer together. A pair of identical peptides (one from each proteome) will occupy slightly different positions on the resulting mass spectrum, because of their distinctive $m/z$ ratios (Figure 13.10). Comparison of the peak heights allows the relative abundance of each peptide to be estimated.

The main disadvantage of ICAT labeling is that peptides with $^{12}\text{C}$ and $^{13}\text{C}$ tags can have slightly different chromatographic properties, especially during RPLC, so they might emerge from the column at slightly different times. The different masses are also a hindrance in tandem mass spectrometry, as the $^{12}\text{C}$- and $^{13}\text{C}$-labeled peptides pass through the first mass analyzer at different rates, so their fragment ions will be collected by the second mass analyzer at different times. These problems are avoided by **isobaric labeling**. In this method, each tag consists of three parts: a reactive region that forms the attachment with the peptide; a balance region, which is labeled with $^{12}\text{C}$ and $^{13}\text{C}$; and a reporter region, which is also labeled with $^{12}\text{C}$ and $^{13}\text{C}$ (Figure 13.11). The labeling is designed in such a way that each tag has the same mass: in other words they are isobaric. Pairs of tagged peptides from two proteomes therefore give molecular ions that have the same $m/z$ values and so behave in the same way in the first mass analyzer. However, the labels are distributed differently within the balance and reporter regions, so cleavage of a tag in the second mass analyzer releases a fragment ion.

**Figure 13.9 Typical isotope-coded affinity tag for proteome studies.**

The iodoacetyl group reacts with cysteine and hence forms an attachment to the peptide. The linker region contains either $^{12}\text{C}$ or $^{13}\text{C}$ atoms and so provides the isotope coding function. The terminal biotin group enables tagged peptides to be separated from untagged ones by affinity chromatography on a column matrix carrying avidin groups. Untagged peptides (ones that lack a cysteine group) can therefore be discarded prior to mass spectrometry.
whose mass is characteristic of that tag. The relative amounts of the reporter fragment ions, as detected in the second mass analyzer, gives the relative amounts of the peptides in the two proteomes.

A final strategy that can sometimes be used with microorganisms and eukaryotic cell lines is metabolic labeling. If a microbial or cell culture is supplied with nutrients that contain $^{13}$C rather than $^{12}$C atoms, then all the proteins in the proteome will become labeled with the heavy isotope. If one of the proteomes that is being studied is obtained in this way, then there is no need to add tags to individual peptides, as all the proteins will be prelabeled. This approach therefore enables a rapid, high-throughput means of comparing the relative amounts of all the proteins in a proteome, albeit with the possible drawbacks described above regarding differential mobility during column chromatography and the first stage of tandem mass spectrometry.

### Analytical protein arrays offer an alternative approach to protein profiling

Gel and/or column separation followed by mass spectrometry offers an effective but laborious and expensive means of profiling the contents of a proteome. These approaches are necessary during the initial characterization of a proteome, but in many research projects the objective is not to catalog the entire content of a proteome but to understand changes that occur within a proteome, for example, in response to extracellular stimuli and during the transition from a healthy tissue to a diseased one. For these applications, a more rapid method of assessing the relative abundance of different proteins is desirable.

**Protein arrays** provide the main alternative to the top-down and bottom-up approaches to protein profiling. A protein array is similar to a DNA array (Section 12.1), the difference being that the immobilized molecules are proteins rather than oligonucleotides. There are several types of protein array, including a version used to detect protein–protein interactions, which we will study in Section 13.2. The particular type of protein array used in protein profiling is called an analytical protein array or an antibody array, the second name indicating that the array carries a series of antibodies, each one specific for a different protein in the proteome for which the microarray is designed. When a sample of the proteome is applied to the array, individual proteins bind to their antibodies and become captured on the array. The amount of binding at each position is dependent on the abundance of that protein in the proteome. The captured proteins are usually detected with a second, polyclonal antibody that binds to all the proteins in the
proteome. This antibody is fluorescently labeled and so gives signals for those positions on the array where a protein has been captured (Figure 13.12). As with a DNA microarray, the intensities of the resulting fluorescent signals can be used to assay the amount of each protein in the proteome.

The main difficulty in designing an analytical protein array is ensuring that each antibody is specific for its target protein and does not cross-react with any other proteins. Cross-reaction will occur if the epitope recognized by an antibody is a common surface feature shared by two or more distinct proteins. However, once a non-cross-reacting array has been designed, then multiple copies can be fabricated and its actual usage is relatively straightforward. Although hundreds of thousands of antibodies can be accommodated on a single chip, most antibody arrays are designed for the assay of particular components of a proteome and hence carry fewer than 1000 antibodies. Typical applications would be screening for the presence or absence and relative abundance of cytokines in different human tissues, for which commercial arrays targeting 640 proteins are available.

13.2 IDENTIFYING PROTEINS THAT INTERACT WITH ONE ANOTHER

Important data pertaining to the function of a proteome can be obtained by identifying pairs and groups of proteins that interact with one another. At a detailed level, this information is often valuable when attempts are made to assign a function to a newly discovered gene or protein (Chapter 6) because an interaction with a second, well-characterized protein can often indicate the role of an unknown protein. For example, an interaction with a protein that is located on the cell surface might indicate that the unknown protein is involved in cell–cell signaling (Section 14.1).

Identifying pairs of interacting proteins

There are several methods for studying protein–protein interactions, the two most useful being phage display and the yeast two-hybrid system. In phage display a special type of cloning vector is used, based on λ bacteriophage or one of the filamentous bacteriophages such as M13. The vector is designed so that when a new gene is inserted, it is expressed in such a way that its protein product becomes fused with one of the phage coat proteins (Figure 13.13A). The phage protein therefore carries the foreign protein into the phage coat, where it is displayed
Figure 13.13 Phage display. (A) The cloning vector used for phage display is a bacteriophage genome with a unique restriction site located within a gene for a coat protein. The technique was originally carried out with the gene III coat protein of the filamentous phage called f1, but it has now been extended to other phages including λ. To create a display phage, the DNA sequence coding for the test protein is ligated into the restriction site so that a fused reading frame is produced: this is one in which the series of codons continues unbroken from the test gene into the coat protein gene. After transformation of E. coli, this recombinant molecule directs synthesis of a hybrid protein made up of the test protein fused to the coat protein. Phage particles produced by these transformed bacteria therefore display the test protein in their coats. (B) Using a phage display library. The test protein is immobilized within a well of a microtiter tray, and the phage display library is added. After one or more washes, the phages that are retained in the well are those displaying a protein that interacts with the test protein.

in a form that enables it to interact with other proteins that the phage encounters. There are several ways in which phage display can be used to study protein interactions. In one method, the test protein is displayed and interactions are sought with a series of purified proteins or protein fragments of known function. This approach is limited because it takes time to carry out each test, so it is feasible only if some prior information has been obtained about likely interactions. A more powerful strategy is to prepare a phage display library, a collection of clones displaying a range of proteins, and identify which members of the library interact with the test protein (Figure 13.13B).

The yeast two-hybrid system detects protein interactions in a more complex way. In Section 12.2 we saw that transcription factors are responsible for controlling the expression of genes in eukaryotes. To carry out this function, a transcription factor must bind to a DNA sequence upstream of a gene and stimulate the mediator protein that regulates the initiation of transcription (see Figure 12.21). These two abilities, DNA binding and mediator activation, are specified by different parts of the transcription factor. Some transcription factors can be cleaved into two segments, where one segment contains the DNA-binding domain and the other contains the activation domain. In the cell, the two segments interact to form the functional transcription factor.

The two-hybrid system makes use of a Saccharomyces cerevisiae strain that lacks a transcription factor for a reporter gene. This gene is therefore switched off. An artificial gene that codes for the DNA-binding domain of the transcription factor is ligated to the gene for the protein whose interactions we wish to study. This protein can come from any organism, not just yeast: in the example shown in Figure 13.14A, it is a human protein. After introduction into yeast, this construct specifies synthesis of a fusion protein made up of the DNA-binding domain of the transcription factor attached to the human protein. The recombinant yeast strain is still unable to express the reporter gene because the modified transcription factor only binds to DNA; it cannot influence the mediator protein. Activation only occurs after the yeast strain has been co-transformed with a second construct,
one comprising the coding sequence for the activation domain fused to a DNA fragment that specifies a protein able to interact with the human protein that is being tested (Figure 13.14B). As with phage display, if there is some prior knowledge about possible interactions, then individual DNA fragments can be tested one by one in the two-hybrid system. Usually, however, the gene for the activation domain is ligated with a mixture of DNA fragments so that many different constructs are made. After transformation, cells are plated out and those that express the reporter gene are identified. These are cells that have taken up a copy of the gene for the activation domain fused to a DNA fragment that encodes a protein able to interact with the test protein.

It is also possible to use a protein array to study protein–protein interactions. Unlike the arrays used in protein profiling, the immobilized proteins are not antibodies but instead are the actual proteins whose possible interactions we wish to assay. A fluorescently labeled version of the test protein is applied to the array, the positions of the resulting signals indicating proteins with which the test protein interacts (Figure 13.15). Although this approach enables interactions between the test protein and a broad range of other proteins to be checked in a single experiment, it is not usually the first choice for this type of work, and phage display and the two-hybrid system remain the standard methods for studying protein–protein interactions. Protein arrays are more popular for testing interactions with DNA fragments—for example, to identify proteins that bind to a particular DNA sequence—and with small molecules such as some drugs.

**Figure 13.14 The yeast two-hybrid system.** (A) On the left, a gene for a human protein has been ligated to the gene for the DNA-binding domain of a yeast transcription factor. After transformation of yeast, this construct specifies a fusion protein, part human protein and part yeast transcription factor. On the right, various human DNA fragments have been ligated to the gene for the activation domain of the transcription factor; these constructs specify a variety of fusion proteins. (B) The two sets of constructs are mixed and co-transformed into yeast. A colony in which the reporter gene is expressed contains fusion proteins whose human segments interact, thereby bringing the DNA-binding and activation domains into proximity and stimulating the mediator protein.

**Figure 13.15 Using a protein array to test protein–protein interactions.** The array carries a series of different proteins. Detection of the fluorescent signal indicates which proteins bind to the test protein.
Identifying the components of multiprotein complexes

Phage display and the yeast two-hybrid system are effective methods for identifying pairs of proteins that interact with one another, but identifying such links reveals only the basic level of protein–protein interactions. Many cellular activities are carried out by multiprotein complexes, such as the mediator protein (Section 12.2) or the spliceosome that is responsible for the removal of introns from pre-mRNA (Section 12.4). Complexes such as these typically comprise a set of core proteins, which are present at all times, along with a variety of ancillary proteins that associate with the complex under particular circumstances. Identifying the core and ancillary proteins is a critical step toward understanding how these complexes carry out their functions. These proteins might be identified pair-by-pair by a long series of phage display or two-hybrid experiments, but a more direct route to determining the composition of multiprotein complexes is clearly needed.

In principle, a phage display library can be used to identify the members of a multiprotein complex, as in this procedure all proteins that interact with the test protein are identified in a single experiment (see Figure 13.13B). The problem is that large proteins are displayed inefficiently because they disrupt the phage replication cycle. To circumvent this problem, it is generally necessary to display a short peptide, representing part of a cellular protein, rather than the entire protein. The displayed peptide may therefore be unable to interact with all members of the complex within which the intact protein is located, because the peptide lacks some of the protein–protein attachment sites present in the intact form (Figure 13.16). A method that avoids this problem, because it works with intact proteins, is affinity chromatography. In affinity chromatography, the test protein is attached to a chromatographic matrix and placed in a column. The cell extract is passed through the column in a low-salt buffer, which allows formation of the hydrogen bonds that hold proteins together in a complex (Figure 13.17A). The proteins that interact with the bound test protein are therefore retained in the column, while all the others are washed away. The interacting proteins are then eluted with a high-salt buffer. A disadvantage of this procedure is the need to purify the test protein, which is time-consuming and hence difficult to use as the basis for a large screening program. In the more sophisticated method called tandem
affinity purification (TAP), which was developed as a means of studying protein complexes in *S. cerevisiae*, the gene for the test protein is modified so that the test protein, when synthesized, has a C-terminal extension that can bind to a second protein called calmodulin. The cell extract is prepared under gentle conditions so that multiprotein complexes do not break down, and then passed through an affinity chromatography column packed with a resin containing attached calmodulin molecules. This results in immobilization of both the test protein and others with which it is associated (Figure 13.17B). In both techniques, the identities of the purified proteins are determined by mass spectrometry. When used in a large-scale screen of 1739 yeast genes, TAP identified 232 multiprotein complexes, providing new insights into the functions of 344 genes, many of which had not previously been characterized by experimental means.

A second disadvantage of affinity chromatography methods is that a single member of a multiprotein complex is used as the bait for isolation of other proteins from that complex. In practice, if a member of a complex does not interact directly with the bait, then it may not be isolated (Figure 13.18). These methods therefore identify groups of proteins that are present in a complex but do not necessarily provide the total protein complement of the complex. Developing ways of purifying intact complexes is therefore a major goal of current research. In co-immunoprecipitation, a cell extract is prepared under gentle conditions so that complexes remain intact. An antibody specific for the test protein is then added, which results in precipitation of this protein and all other members of the complex within which it is present. Treatment of the collection of proteins with a protease, followed by bottom-up proteomics, then enables the members of the complex to be identified. This version of shotgun proteomics is called the multidimensional protein identification technique (MudPIT). The method was first used to study the large subunit of the yeast ribosome and resulted in identification of 11 proteins that had not previously been known to be associated with this complex.

### Identifying proteins with functional interactions

Proteins do not need to form physical associations with one another in order to have a functional interaction. For example, in bacteria such as *Escherichia coli*, the enzymes lactose permease and β-galactosidase have a functional interaction in that they are both involved in utilization of lactose as a carbon source. But there is no physical interaction between these two proteins; the permease is located in the cell membrane and transports lactose into the cell, while β-galactosidase, which splits lactose into glucose and galactose, is present in the cell cytoplasm (see Figure 8.9A). Many enzymes that work together in the same biochemical pathway never form physical interactions with one another, and if studies were to be based solely on detection of physical associations between proteins, then many functional interactions would be overlooked.

Several methods can be used to identify proteins that have functional interactions. Most of these do not involve direct study of the proteins themselves and hence, strictly speaking, do not come under the general heading of proteomics. Nonetheless, it is convenient to consider them here because the information they yield is often considered along with the results of proteomics studies. These methods include the following:

- **Comparative genomics** can be used in various ways to identify groups of proteins that have functional relationships. One approach is based on the observation that pairs of proteins that are separate molecules in some organisms are fused into a single polypeptide in others. An example is provided by the yeast gene *HIS2*, which codes for an enzyme involved in histidine biosynthesis. In *E. coli*, two genes are homologous to *HIS2*. One of these, itself called *his2*, has sequence similarity with the 5′-region of the yeast gene, and the second, *his10*, is similar to the 3′-region (Figure 13.19). The implication is that the proteins coded by *his2* and *his10* interact within

![Figure 13.18](image-url) A disadvantage of affinity chromatography. If the bait protein (labeled with a B) does not interact directly with one or more proteins in the complex, then those proteins might not be isolated.

![Figure 13.19](image-url) Using homology analysis to deduce protein–protein interactions. The 5′-region of the yeast *HIS2* gene is homologous to *E. coli his2*, and the 3′-region is homologous to *E. coli his10*. 
the \textit{E. coli} proteome to provide part of the histidine biosynthesis activity. Analysis of the sequence databases reveals many examples of this type, where two proteins in one organism have become fused into a single protein in another organism. A similar approach is based on examination of bacterial operons. An operon consists of two or more genes that are transcribed together and usually have a functional relationship (Section 8.2). For example, the genes for lactose permease and \(\beta\)-galactosidase of \textit{E. coli} are present in the same operon, along with the gene for a third protein involved in lactose utilization (see Figure 8.9A). The identities of genes in bacterial operons can therefore be used to infer functional interactions between the proteins coded by homologous genes in a eukaryotic genome.

- Transcriptome studies can identify functional interactions between proteins, as the mRNAs for functionally related proteins often display similar expression profiles under different conditions.
- Gene inactivation studies can be informative. If a change in phenotype is observed only when two or more genes are inactivated together, then it can be inferred that those genes function together in generation of the phenotype.

**Protein interaction maps display the interactions within a proteome**

The information from phage display, two-hybrid analyses, and other methods for identifying pairs and groups of proteins that associate with one another in a cell enables a \textbf{protein interaction map} to be constructed. In this type of map each protein is depicted by a dot, or \textbf{node}, with pairs of interacting proteins linked by lines, or \textbf{edges}. The resulting network displays all the interactions that occur between the components of a proteome. The first of these maps were constructed in 2001 for relatively simple proteomes, almost entirely from two-hybrid experiments. These included maps for the bacterium \textit{Helicobacter pylori}, comprising over 1200 interactions involving almost half the proteins in the proteome, and for 2240 interactions between 1870 proteins from the \textit{S. cerevisiae} proteome (Figure 13.20A). More recently, the application of additional techniques has led to more detailed versions of the \textit{S. cerevisiae} map, as well as maps for humans and other eukaryotes (Figure 13.20B). Each protein interaction map forms a part of the broader \textbf{interactome} for the species. The interactome comprises all of the molecular interactions that occur, including those involving small molecules that regulate protein activity and between DNA-binding proteins and the genes whose expression they control.

What interesting features have emerged from these protein interaction maps? The most intriguing discovery is that each network is built up around a small number of proteins that have many interactions and form \textbf{hubs} in the network, along with a much larger number of proteins with few individual connections.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{protein_interaction_maps.png}
\caption{Protein interaction maps. (A) Initial version of the \textit{S. cerevisiae} map, published in 2001. Each dot represents a protein, with connecting lines indicating interactions between pairs of proteins. Red dots are essential proteins: an inactivating mutation in the gene for one of these proteins is lethal. Mutations in the genes for proteins indicated by green dots are nonlethal, and mutations in genes for proteins shown in orange lead to slow growth. The effects of mutations in genes for proteins shown as yellow dots were not known when the map was constructed. (A, from Jeong H, Mason SP, Barabási AL & Oltvai ZN [2001] \textit{Nature} 411:41–42. With permission from Macmillan Publishers Limited. B, from Stelzl U, Worm U, Lalowski M et al. [2005] 122:957–968. With permission from Elsevier.)}
\end{figure}
This architecture is thought to minimize the impact on the proteome of the disruptive effects of mutations that might inactivate individual proteins. Only if a mutation affects one of the proteins at a highly interconnected node will the network as a whole be damaged. This hypothesis is consistent with the discovery, from gene inactivation studies (Section 6.2), that a substantial number of yeast proteins are apparently redundant, meaning that if the protein activity is destroyed, the proteome as a whole continues to function normally, with no discernible impact on the phenotype of the cell. Examination of the expression profiles of the hub proteins and their direct partners enables these hubs to be divided into two groups. The first group includes those hub proteins that interact with all their partners simultaneously. These have been called party hubs, and their removal has little effect on the overall structure of the network (Figure 13.21B). In contrast, removal of the second group, the date hubs, which interact with different partners at different times, breaks the network into a series of small subnetworks (Figure 13.21C). The implication is that the party hubs work within individual biological processes and do not contribute greatly to the overall organization of the proteome. The date hubs, on the other hand, are the key players that provide organization to the proteome by linking biological processes to one another.

Most of the protein interaction maps that have been constructed so far are incomplete, simply because not all of the interactions occurring in the proteome under study have been identified. In fact, it is debatable whether it will ever be possible to construct a fully comprehensive interaction map for any proteome, bearing in mind the limitations in scope and sensitivity of the methods used to study protein–protein interactions. The accuracy of these methods also needs to be considered to ensure that the resulting networks do not contain spurious links. Both problems are illustrated by the current status of the human protein interaction map. When all reported interactions are taken into account, this network comprises almost 30,000 proteins with 350,000 interactions, but the numbers drop to 16,000 proteins and 116,000 interactions if only those interactions that have been checked by two different methods are included. Clearly these networks are far from complete: they account for only a fraction of 70,000 proteins thought to be present in the human proteome, and many of the interactions that have been identified are unconfirmed. Despite these limitations, protein interaction maps are proving valuable as a means of probing the link between the proteome and the biochemistry of the cell. In particular, different genes that, when defective, give rise to the same disease often specify proteins that occupy a distinct disease module within a network (Figure 13.22). The implication is that loss or perturbation of the biochemical function performed by those interlinked components of the proteome gives rise to the symptoms of the disease. Even if the map is incomplete, identification of proteins within a module that were not previously associated with the disease enables the biochemical basis of the defect to be understood in greater detail. The discovery that there are sometimes overlaps between the modules for distinct diseases, such as multiple sclerosis and rheumatoid arthritis.
13.3 Synthesis and degradation of the components of the proteome

The composition of a proteome is determined by the balance between the synthesis and degradation of the individual proteins that it contains. Except for the words proteome and proteins, this is the same sentence that we used to describe the composition of the transcriptome (Section 12.2). The principles are exactly the same, and the dynamics of synthesis and degradation illustrated in Figure 12.7 for RNA could equally well refer to proteins. To understand how a proteome is maintained and how a proteome changes in response to external stimuli and during differentiation, development, and disease, we must therefore study the same topics as in Chapter 12—synthesis, degradation, and processing—but this time with reference to protein rather than RNA.

Ribosomes are molecular machines for making proteins

Proteins are synthesized by the large RNA-protein complexes called **ribosomes**. An *E. coli* cell contains approximately 20,000 ribosomes, distributed throughout its cytoplasm. The average human cell contains more than a million ribosomes, some free in the cytoplasm and some attached to the outer surface of the endoplasmic reticulum, the membranous network of tubes and vesicles that permeates the cell.

Originally, ribosomes were looked on as passive partners in protein synthesis, merely the structures on which mRNAs were translated into polypeptides. This view has changed over the years, and ribosomes are now considered to play two active roles:

- Ribosomes **coordinate** protein synthesis by placing the mRNA, tRNAs, and associated protein factors in their correct positions relative to one another.
- Components of ribosomes, including the rRNAs, **catalyze** at least some of the chemical reactions that occur during protein synthesis, including the central reaction that results in synthesis of the peptide bond that links two amino acids together.

(see Figure 13.22), also provides important insights into **comorbidity**, which is the tendency for patients suffering from one disease to display symptoms associated with other diseases.
When the involvement of ribosomes in protein synthesis became clear in the 1950s, biologists quickly realized that a detailed knowledge of ribosome structure would be necessary in order to understand how mRNAs are translated into polypeptides. Originally called microsomes, ribosomes were first observed in the early decades of the twentieth century as tiny particles almost beyond the resolving power of light microscopy. In the 1940s and 1950s, the first electron micrographs showed that bacterial ribosomes are oval-shaped, with dimensions of $29 \times 21$ nm, rather smaller than eukaryotic ribosomes, which vary a little in size depending on species but average about $32 \times 22$ nm. Compositional studies then revealed that a ribosome comprises two subunits, referred to as large and small, each subunit made up of one or more rRNAs and a collection of ribosomal proteins (Figure 13.23). We now know that ribosomes dissociate into their subunits when they are not actively participating in protein synthesis, and the subunits remain in the cytoplasm until they are used for a new round of translation.

Once the basic composition of eukaryotic and bacterial ribosomes had been worked out, attention became focused on the way in which the various rRNAs and proteins fit together. Important information was provided by the first rRNA sequences. Comparisons between these sequences identified conserved regions that can base-pair to form complex two-dimensional structures (Figure 13.24). This suggested that the rRNAs provide a scaffolding within the ribosome, to which the proteins are attached, an interpretation that underemphasizes the active role that rRNAs play in protein synthesis but which nonetheless was a useful foundation on which to base subsequent research.

Much of that subsequent research has concentrated on the bacterial ribosome, which is smaller than the eukaryotic version and available in large amounts from extracts of cells grown to high density in liquid cultures. A number of technical approaches have been used to study the bacterial ribosome:

- **Nuclease protection experiments** (Section 7.1) enabled contacts between rRNAs and proteins to be identified.
- **Protein–protein cross-linking** identified pairs or groups of proteins that are located close to one another in the ribosome.
- **Electron microscopy** gradually became more sophisticated, enabling the overall structure of the ribosome to be resolved in greater detail. For
example, innovations such as immunoelectron microscopy, in which ribosomes are labeled with antibodies specific for individual ribosomal proteins before examination, have been used to locate the positions of these proteins on the surface of the ribosome.

- **Site-directed hydroxyl radical probing**, which makes use of the ability of Fe(II) ions to generate hydroxyl radicals that cleave RNA phosphodiester bonds located within 1 nm of the site of radical production, has been used to determine the exact positioning of ribosomal proteins in the *E. coli* ribosome. For example, to determine the position of S5, different amino acids within this protein were labeled with Fe(II) and hydroxyl radicals were induced in reconstituted ribosomes. The positions at which the 16S rRNA was cleaved were then used to infer the topology of the rRNA in the vicinity of S5 protein (Figure 13.25).

In recent years these techniques have been increasingly supplemented by X-ray crystallography (Section 11.1), which has been responsible for the most exciting insights into ribosome structure. Analyzing the massive amount of X-ray diffraction data produced by crystals of an object as large as a ribosome is a huge task, particularly at the level needed to obtain a structure that is detailed enough to be informative about the way in which the ribosome works. This challenge has been met, and detailed structures are now known for entire ribosomes, including ones attached to mRNA and tRNAs (Figure 13.26A). These studies have shown that, in bacteria, attachment of the two ribosome subunits to one another results in the formation of two sites at which an aminoacyl-tRNA (a tRNA with an attached amino acid) can bind. These are called the **P** or peptidyl site and the **A** or aminoacyl site. The P site is occupied by the aminoacyl-tRNA whose amino acid has just been attached to the end of the growing polypeptide, and the A site is entered by the aminoacyl-tRNA carrying the next amino acid that will be used. There is also a third site, the **E** or exit site, through which the tRNA departs after its amino acid has been attached to the polypeptide (Figure 13.26B). The structures revealed by X-ray diffraction analysis show that these sites are located in a cavity between the large and small subunits of the ribosome, the mRNA threading through a channel formed mainly by the small subunit. After each amino acid addition, the ribosome adopts a less compact structure, with the two subunits rotating slightly in opposite directions. This opens up the space between the subunits and enables the ribosome to slide along the mRNA in order to read the next codon in the open reading frame.

![Figure 13.25](image1.png) Positions within *E. coli* 16S rRNA that form contacts with ribosomal protein S5. The distribution of the contact positions (shown in pink) for this single ribosomal protein emphasizes the extent to which the base-paired secondary structure of the rRNA is further folded within the three-dimensional structure of the ribosome.

![Figure 13.26](image2.png) Detailed structure of a bacterial ribosome. (A) Structure of a ribosome in the process of translating an mRNA. The tRNAs positioned in the A, P, and E sites are indicated in pink, green, and yellow, respectively. (B) Diagram showing relative positions of the A, P, and E sites and the channel through which the mRNA is translocated. (A, From Schmeing TM & Ramakrishnan V [2009] Nature 461:1234–1242. With permission from Macmillan Publishers Limited)
During stress, bacteria inactivate their ribosomes in order to downsize the proteome

As well as revealing the structures of active ribosomes, X-ray crystallography has also helped to elucidate the events that enable a bacterium to bring about a general reduction in the size of its proteome during periods of stress, such as nutrient limitation. The latter is signaled by the presence in the A sites of ribosomes of tRNAs that do not have attached amino acids. These tRNAs lack amino acids because the amino acid pool in the cytoplasm has become depleted due to the starvation conditions. In E. coli, the presence of deacylated tRNA in the A site is detected by the L11 ribosomal protein, initiating the stringent response. L11 activates a ribosome-associated protein called RelA, which converts guanosine 5'-triphosphate (GTP) to guanosine 5'-triphosphate 3'-diphosphate (pppGpp) by transferring a diphosphate from adenosine 5'-triphosphate (ATP) to a GTP molecule. Guanine pentaphosphatase then converts pppGpp to ppGpp (Figure 13.27), which is an alarmone, a stress response molecule that modifies a broad range of cellular activities in order to help the cell deal with the stress. In starving bacteria, one of these responses is a general decrease in transcription but an increase in transcription of genes involved in amino acid biosynthesis. These changes are brought about by ppGpp binding to the β and β' subunits of the bacterial RNA polymerase and altering the affinity of the polymerase for different types of promoters.

By switching on amino acid biosynthesis, the bacterium is able to carry out essential maintenance of its proteome while its rides out the stress conditions and waits for the external nutrient supply to increase. Because its overall metabolic activity has declined, the bacterium also decreases the size of its proteome by globally reducing the rate of protein synthesis. This is achieved, at least in part, by ppGpp binding to the translation initiation factor IF-2. This factor is required during the first stage of protein synthesis, when the initiation complex is assembled at the ribosome binding site, which is located close to the initiation codon of the open reading frame present in the mRNA. The initiation complex comprises the two subunits of the ribosome, the aminoacylated tRNA that recognizes the initiation codon, and three protein initiation factors. The specific roles of IF-2 include hydrolysis of GTP to provide the energy needed to complete assembly of the initiation complex. However, IF-2 has a higher affinity for ppGpp compared with GTP and so binds the former when it is present. In the absence of GTP hydrolysis, translation stalls at the initiation phase, resulting in a global decrease in protein synthesis and hence a reduction in the size of the proteome.

The downshift in protein synthesis that occurs during starvation means that the bacterium now has an excess of ribosomes. The surplus ribosomes are not broken down but instead stored in inactive form. The A and P sites become blocked by a protein called the ribosome modulation factor, and pairs of ribosomes interact with a second protein, the hibernation promotion factor, forming ribosome dimers. The dimers are inactive but can be reconverted to

![Figure 13.27](structure_of_the_alarmone_guanosine_5'-diphosphate_3'-diphosphate_(ppGpp).svg)
If you can be up-regulated as soon as the environmental conditions improve.

**Initiation factors mediate large-scale remodeling of eukaryotic proteomes**

Initiation factors are also the mediators of large-scale changes in proteome composition in eukaryotes. Although ribosomal architecture is similar in bacteria and eukaryotes, there are distinctions in the way in which protein synthesis is carried out in the two types of organism, in particular during the initiation phase. Rather than being assembled at the initiation codon, the eukaryotic initiation complex is constructed at the 5′-end of the mRNA and then **scans** along the mRNA until it locates the initiation codon. The first component of the initiation complex, called the [preinitiation complex](#), comprises the small subunit of the ribosome, a ternary complex made up of initiation factor eIF-2 bound to the [initiator tRNA](#) and a molecule of GTP, and three additional initiation factors, eIF-1, eIF-1A, and eIF-3 ([Figure 13.28](#)). After assembly, the preinitiation complex associates with the cap structure at the 5′-end of the mRNA (see [Figure 1.17A](#)), and additional initiation factors are recruited prior to commencement of the scanning process.

The initiation process can be repressed by phosphorylation of eIF-2, which prevents this initiation factor from binding the molecule of GTP that it needs before it can transport the initiator tRNA to the small subunit of the ribosome. Phosphorylation of eIF-2 occurs during stresses, such as heat shock, and results in a decrease in the overall level of protein synthesis. This is not, however, a means of reducing the size of the proteome, as occurs during the bacterial stringent response. Instead, eIF-2 phosphorylation results in proteome remodeling, with an increase in the amounts of various proteins involved in protection of the cell against stress. This occurs because of a second route to initiation of translation in eukaryotes, which avoids the scanning process and hence circumvents the inactivation of eIF-2. This alternative pathway involves assembly of the preinitiation complex at an [internal ribosome entry site (IRES)](#), which is similar in function to the ribosome binding site of bacteria, although the positions of IRESs relative to the initiation codon are more variable than the bacterial versions. The ability of eukaryotic translation to initiate in this way was first recognized with the picornaviruses, a group of viruses with RNA genomes, including human poliovirus and rhinovirus, the latter responsible for the common cold. Transcripts from these viruses are not capped but instead have an IRES, meaning that picornaviruses can block protein synthesis in the host cell without affecting translation of their own transcripts. Remarkably, no viral proteins are required for recognition of an IRES by a host ribosome. In other words, the normal eukaryotic cell possesses proteins and/or other factors that enable it to initiate translation by the IRES method. This discovery prompted a search for IRES sequences in eukaryotic mRNAs, with some 150 so far discovered. The presence of an IRES enables these mRNAs to undergo preferential translation at times of stress when eIF-2 and the scanning process are inactivated.

Although eIF-2 is the best-studied example of the global regulation of eukaryotic protein synthesis, recent research has shown that there are several other mechanisms for repressing cap-dependent initiation and switching translation to IRES-mRNAs. One of these occurs during mitosis when a second initiation factor, eIF-4E, becomes phosphorylated. This factor is a part of the [cap binding complex](#), which aids attachment of the preinitiation complex to the 5′-end of the mRNA. Phosphorylation of eIF-4E reduces the affinity of the cap binding complex for the mRNA, repressing cap-dependent translation and promoting translation of IRES-mRNAs coding for proteins needed specifically during mitosis. These proteins are unlikely to be identical to those required during stress conditions, indicating that repression of cap-dependent initiation does not simply result in an increase in translation of IRES-mRNAs. Instead, there must be mechanisms for modulating the IRES process so that specific groups of transcripts are translated according to the needs of the cell. This point is further emphasized by a third example of cap-dependent translation repression, which occurs when a second component
of the cap binding complex, eIF-4G, is cleaved. This is associated with apoptosis, the IRES-mRNAs that are subsequently translated specifying proteins that aid in this process. Clearly the cell needs to distinguish its IRES-mediated response to mitosis from its pathway to apoptosis. How the translation of different sets of IRES-mRNAs is regulated is currently not understood, but the discovery of IRES trans-acting factors (ITAFs), which appear to control the affinity of the ribosome for different IRESs, points a way toward a possible explanation.

The translation of individual mRNAs can also be regulated

As well as the large-scale changes to proteome size or composition that can be brought about by global regulation of translation initiation, the translation rates of individual mRNAs can also be regulated. In bacteria, several of these transcript-specific regulation processes involve attachment of an RNA-binding protein to the target mRNA, the attached protein blocking access to the ribosome binding site and hence inhibiting formation of the translation initiation complex. The most frequently cited example involves the operons for the ribosomal protein genes of *E. coli* (**Figure 13.29A**). The leader region of the mRNA transcribed from each operon contains a sequence that acts as a binding site for one of the proteins coded by the operon. When this protein is synthesized it can either attach to its position on the ribosomal RNA or bind to the leader region of the mRNA. The rRNA attachment is favored and occurs if there are free rRNAs in the cell. Once all the free rRNAs have been assembled into ribosomes, the ribosomal protein binds to its mRNA, preventing translation initiation and hence switching off further synthesis of the ribosomal proteins coded by that particular mRNA. Similar events involving other mRNAs ensure that synthesis of each ribosomal protein is coordinated with the amount of free rRNA in the cell.

A second example of transcript-specific regulation by RNA-binding proteins, which occurs in mammals, involves the mRNA for ferritin, an iron-storage protein (**Figure 13.29B**). In the absence of iron, ferritin synthesis is inhibited by proteins that bind to sequences called iron-response elements located in the leader region of the ferritin mRNA. The bound proteins block the ribosome as it attempts to scan along the mRNA in search of the initiation codon. When iron is present, the binding proteins detach and the mRNA is translated. Interestingly, the mRNA for a related protein, the transferrin receptor involved in the uptake of iron, also has iron-response elements, but in this case detachment of the binding proteins in the presence of iron results not in translation of the mRNA but in its degradation. This is logical because when iron is present in the cell, there is less requirement for transferrin receptor activity because there is less need to import iron from outside.

Initiation of translation of some bacterial mRNAs can also be regulated by short RNAs that attach to recognition sequences within the mRNAs, but this does not always result in translation being prevented, as some short RNAs can also activate translation of one or more of their target mRNAs. An example is the *E. coli* RNA called OxyS, which is 109 nucleotides in length and regulates translation of 40 or so mRNAs. Synthesis of OxyS is activated by hydrogen peroxide and other reactive oxygen compounds that can cause oxidative damage to the cell. Once synthesized, OxyS switches off translation of mRNAs whose products would be deleterious under these circumstances and switches on translation of other mRNAs whose products help protect the bacterium from oxidative damage. Prevention of translation occurs in a similar way to the examples we have discussed above involving RNA-binding proteins: attachment of the regulatory RNA simply blocks access to the ribosome binding site. Activation of translation involves a more subtle mechanism. In these cases, the target mRNA is able to form a stem–loop structure, with the ribosome binding site present in the stem region and hence inaccessible to the components of the initiation complex. Attachment of the regulatory RNA disrupts the stem–loop, exposing the ribosome binding site so that translation initiation can now occur.

Stem–loops and other secondary structures can also form in the 5′-untranslated regions of many eukaryotic mRNAs. During scanning of the initiation
complex along the mRNA, these secondary structures are disrupted by the initiation factor eIF-4A, which is a helicase and so is able to break the intramolecular base pairs. However, the presence of secondary structure delays progress of the initiation complex and hence reduces the translational efficiency, which is the rate at which proteins are synthesized from the mRNA. Because of differences in the translational efficiencies of different mRNAs, the proteome is not simply a reflection of the transcriptome, with the relative abundances of each protein matching the relative abundances of their mRNAs. Instead, those mRNAs with high translational efficiency give rise to greater amounts of protein per transcript. Current research is exploring the possibility that transcript-specific regulation of eukaryotic protein synthesis can be exerted by regulatory proteins or RNAs that influence the formation of secondary structures in mRNAs and hence increase or decrease their translational efficiency.

Degradation of the components of the proteome

The ability of a proteome to change over time requires the removal of proteins whose functions are no longer required. This removal must be highly selective so that only the correct proteins are degraded, and must also be rapid, in order to account for the abrupt changes that occur under certain conditions, for example, during key transitions in the cell cycle.

For many years, protein degradation was an unfashionable subject, and it was not until the 1990s that real progress was made in understanding how specific proteolytic events are linked with processes such as the cell cycle and differentiation. Even now, our knowledge centers largely on descriptions of general protein breakdown pathways and less on regulation of the pathways and the mechanisms used to target specific proteins. There appear to be a number of different types of breakdown pathway whose interconnectivities have not yet been traced. This is particularly true in bacteria, which seem to have a range of proteases that work together in controlled degradation of proteins. In eukaryotes, most breakdown occurs via a single system, involving ubiquitin and the proteasome.

A link between ubiquitin and protein degradation was first established in 1975 when it was shown that this abundant 76-amino-acid protein is involved in energy-dependent proteolysis reactions in rabbit cells. Subsequent research identified enzymes that attach ubiquitin molecules, singly or in chains, to lysine amino acids in proteins that are targeted for breakdown (Figure 13.30). There are also ubiquitin-like proteins, such as SUMO, that act in the same way as ubiquitin. Ubiquitination is a three-step process in which a ubiquitin molecule is initially attached to an activator protein, then transferred to a conjugating enzyme, and finally transferred to the target protein by a ubiquitin ligase enzyme. How does this process recognize the correct proteins, those that must be degraded? The answer appears to lie with the specificity of the conjugating and ligase enzymes. Most species possess multiple versions of the conjugating enzymes and many types of ubiquitin ligase. In humans, for example, there are 35 conjugating enzymes and several hundred ligases. It is thought that different pairs of conjugating enzyme and ligase have specificity for different proteins. Activation of different enzyme pairs, in response to intra- or extracellular signals, is probably the key to specific degradation of particular proteins and groups of proteins.

The second component of the ubiquitin-dependent degradation pathway is the proteasome, the structure within which ubiquitinated proteins are broken down. In eukaryotes, the proteasome is a large, multisubunit structure comprising a hollow cylinder with a cap at either end (Figure 13.31). The cylinder has four rings, each made up of seven proteins. The proteins present in the two inner rings are proteases, whose active sites are located on the inner ring surface. Archaea also have proteasomes of about the same size but they are less complex, being composed of multiple copies of just two proteins. Ubiquitinated proteins might interact directly with the cap structure of the proteasome, or the interaction might be via a ubiquitin-receptor protein. Before entry into the proteasome, the protein to be degraded must be at least partially unfolded and its ubiquitin

![Figure 13.30](image-url) Ubiquitin attachment targets proteins for degradation. To act as a label for degradation, chains of linked ubiquitin molecules are linked to lysine amino acids in the target protein.
labels must be removed. These steps require energy, obtained from hydrolysis of ATP, catalyzed by proteins present in the cap. The protein can then enter the proteasome, within which it is cleaved into short peptides 4–10 amino acids in length. These are released back into the cytoplasm, where they are broken down into individual amino acids that can be reutilized in protein synthesis.

13.4 INFLUENCE OF PROTEIN PROCESSING ON THE COMPOSITION OF THE PROTEOME

Translation is not the end of the genome expression pathway. The polypeptide that emerges from the ribosome is inactive, and before taking on its functional role in the cell it must undergo post-translational processing. The most fundamental of these processing events is protein folding, as the vast majority of polypeptides are inactive until they have adopted their correct tertiary structure. Some polypeptides are also cleaved in various ways to give the functional proteins, and many undergo chemical modification. The proteome therefore contains not only functional proteins but also a large variety of preprocessed forms, some of which can be retained in their inactive state for substantial periods before processing is completed and the functional protein is produced. We must therefore study protein processing in order to gain a complete understanding of the components of a proteome.

The amino acid sequence contains instructions for protein folding

One of the central principles of molecular biology is that all of the information that a polypeptide needs in order to fold into its correct three-dimensional structure is contained within its amino acid sequence. This link between sequence and structure was first established in the 1950s by experiments with ribonuclease that had been purified from bovine pancreas and resuspended in buffer. Ribonuclease is a small protein, just 124 amino acids in length and containing four disulfide bridges, with a tertiary structure that is made up predominantly of β-sheet, with very little α-helix. Addition of urea, a compound that disrupts hydrogen bonding, resulted in a decrease in the activity of the enzyme (measured by testing its ability to cut RNA) and an increase in the viscosity of the solution (Figure 13.32), indicating that the protein was being denatured by unfolding into an unstructured
polypeptide chain. The critical observation was that when the urea was removed by dialysis, the viscosity decreased and the enzyme activity reappeared. The conclusion is that the protein refolds spontaneously when the denaturant (in this case, urea) is removed. In these initial experiments, the four disulfide bonds remained intact because they were not disrupted by urea, but the same result occurred when the urea treatment was combined with addition of a reducing agent to break the disulfide bonds: the activity was still regained upon renaturation. This shows that the disulfide bonds are not critical to the protein’s ability to refold; they merely stabilize the tertiary structure once it has been adopted.

It was quickly realized that the folding process cannot be random, as it would take too long for a protein to explore all the possible conformations before finally finding the correct one. It was estimated that a polypeptide of 100 amino acids would take about $10^{37}$ seconds to check all conformations, which is much longer than the age of the universe. The implication is that a protein follows a folding pathway, with each step in the process directing the protein toward the following step, so the protein is led toward its correct tertiary structure (Figure 13.33). Researchers currently favor the molten globule model, in which the initial step is the rapid collapse of the polypeptide into a compact structure, with slightly larger dimensions than the final protein, driven by the desire of the hydrophobic amino acid side chains to avoid water. Collapse into this molten globule might automatically fold some of the polypeptide into its $\alpha$-helices and $\beta$-sheets. Because the globule is molten, it can change conformation rapidly, identifying additional folds so that the correct tertiary structure gradually emerges. For larger proteins, this step might involve construction of correctly folded subdomains, which are then brought together to make the final tertiary structure. The whole process can take just a few seconds. In thermodynamic terms, the polypeptide is looked on as passing down a funnel, gradually taking up less random conformations (Figure 13.34). The funnel narrows because at each stage in the folding pathway there are fewer available options for the next steps toward the final structure. There are, however, side funnels into which the protein can be diverted, leading to an incorrect structure. If an incorrect structure is sufficiently unstable, then partial or complete unfolding may occur, allowing the protein to return to the main funnel and pursue a productive route toward its correct conformation.

Experiments conducted in vitro have been useful in establishing the basic principles of protein folding, but they may not be a good model for the folding of proteins in the cell. In particular, a cellular protein might begin to fold before it has been fully synthesized, a scenario that is very difficult to replicate in a test tube experiment. These considerations have prompted extensive research into protein folding in living cells, with particular focus on the molecular chaperones, which are proteins that help other proteins to fold. The molecular chaperones can be divided into various groups, the most important of which are the following:

- **The Hsp70 chaperones** include the *E. coli* Hsp70 protein coded by the dnaK gene and sometimes called DnaK protein.

- **The chaperonins**, the main versions of which are the GroEL/GroES complex present in bacteria and eukaryotic organelles, and TRiC and related structures which are found in eukaryotic cytoplasm and in archaea.
Molecular chaperones do not specify the tertiary structure of a protein, they merely help the protein find that correct structure. The two types of chaperone do this in different ways. Members of the Hsp70 family bind to hydrophobic regions of unfolded proteins, including proteins that are still being translated (Figure 13.35). They hold the protein in an open conformation to aid folding, presumably by modulating the association between those parts of the polypeptide that form interactions in the folded protein. Exactly how this is achieved is not understood, but it involves repeated binding and detachment of the Hsp70 protein, each cycle of which requires energy provided by hydrolysis of ATP. In addition to protein folding, the Hsp70 chaperones are also involved in other processes that require shielding of hydrophobic regions in proteins, such as transport through membranes, association of proteins into multisubunit complexes, and disaggregation of proteins that have been damaged by heat stress.

The chaperonins work in a quite different way. GroEL and GroES form a multisubunit structure that looks like a hollowed-out bullet with a central cavity (Figure 13.36). A single unfolded protein enters the cavity and emerges folded. The mechanism for this is not known, but it is postulated that the GroEL/GroES complex acts as a cage, preventing the unfolded protein from aggregating with other proteins, and that the inside surface of the cavity changes from hydrophobic to hydrophilic in such a way as to promote the burial of hydrophobic amino acids...
within the protein. This is not the only hypothesis: other researchers hold that the cavity unfolds proteins that have folded incorrectly, passing these unfolded proteins back to the cytoplasm so they can have a second attempt at adopting their correct tertiary structure.

Although both the Hsp70 family of chaperones and the TRiC chaperonins are present in eukaryotes, it seems that in these organisms protein folding depends mainly on the action of the Hsp70 proteins, with only 10% of the proteome, mainly components of the cytoskeleton and proteins involved in the cell cycle, being folded by TRiC.

Some proteins are activated by proteolytic cleavage

Processing by proteolytic cleavage is common in eukaryotes but less frequent in bacteria. Cleavage is often used as a means of activating a secreted polypeptide whose biochemical activities might be deleterious to the cell producing the protein. These proteins are synthesized in an inactive form and then activated after secretion, so the cell is not harmed. The cellular proteome therefore contains the intact but inactive version of the polypeptide, and the secreted proteome contains the cleaved, active version. An example is provided by melittin, the most abundant protein in bee venom and the one responsible for causing cell lysis after injection of the bee sting into the person or animal being stung. Melittin lyses cells in bees as well as animals and so must initially be synthesized as an inactive precursor. This precursor, promelittin, has 22 additional amino acids at its N-terminus. The presequence is removed by an extracellular protease that cuts at 11 positions, releasing the active venom protein. The protease does not cleave within the active sequence because its mode of action is to release dipeptides with the sequence X-Y, where X is alanine, aspartic acid, or glutamic acid and Y is alanine or proline; these motifs do not occur in the active sequence (Figure 13.37).

A similar type of processing occurs with insulin, the protein made in the islets of Langerhans in the vertebrate pancreas and responsible for controlling blood sugar levels. Insulin is synthesized as preproinsulin, which is 105 amino acids in length (Figure 13.38). The processing pathway involves the removal of the first 24 amino acids to give proinsulin, followed by two additional cuts that excise a central segment, leaving two active parts of the protein, the A and B chains, which are linked together by formation of two disulfide bonds to form mature insulin. The first segment to be removed, the 24 amino acids from the N-terminus, is a signal peptide, a highly hydrophobic stretch of amino acids that aids transfer of the newly synthesized protein into the endoplasmic reticulum. This is the first stage of the transport pathway that eventually results in secretion of insulin from the cell.

Other proteins, called polyproteins, are cut into segments, giving rise to a range of active products, often with different functions. For example, the polyprotein called proopiomelanocortin, made in the vertebrate pituitary gland, contains at least 11 different peptide hormones. Proopiomelanocortin is initially made as a precursor of 267 amino acids, 26 of which form a signal peptide that is removed when the protein is transferred from the cytoplasm to secretory vesicles inside the cell. Proopiomelanocortin has a variety of internal cleavage sites recognized by proteolytic enzymes, but not all of these sites are cleaved in every tissue: the combinations, and hence the products formed, depend on the identities of the proteolytic enzymes that are present (Figure 13.39). For example, in the corticotropic cells of the anterior pituitary gland, adrenocorticotropic hormone and lipotropins are produced. In the melanotropic cells of the intermediate lobe of the pituitary, a different set of cleavage sites is used, generating melanotropins. Altogether, 11 different peptides can be obtained by alternative patterns of proteolytic cleavage.

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**Figure 13.37** Processing of promelittin, the bee-sting venom. Arrows indicate the cut sites.

<table>
<thead>
<tr>
<th>Cut sites</th>
</tr>
</thead>
<tbody>
<tr>
<td>A P E P E P A P E P E A D A E A D P E A G I G A V K V L T T G L P A I S W I K R K R Q Q G</td>
</tr>
</tbody>
</table>
of proopiomelanocortin. The contribution made by proopiomelanocortin to the proteome is therefore tissue-specific.

Several types of virus that infect eukaryotic cells use polyproteins as a way of reducing the sizes of their genomes, as a single polyprotein gene with one promoter and one termination sequence takes up less space inside the virus capsid than a series of individual genes. The proteomes of cells infected with one of these viruses might therefore contain a complex mixture of precursor proteins and mature, processed products. The human immunodeficiency virus (HIV-1) is an example. During its replication cycle, HIV-1 synthesizes the Gag polyprotein, which is cleaved into four proteins and two short spacer peptides. Three of these four proteins form structural components of the HIV capsid and the fourth, called p6, is involved in the process by which virus particles are released from the cell. HIV-1 also synthesizes an extended version of Gag, called Gag-Pol, where the additional segment is processed to give two enzymes involved in replication of the HIV genome, as well as the protease that makes the cuts in the Gag and Gag-Pol polyproteins. A few molecules of this protease are stored in each new virus particle that is produced, and hence are available to cut up the polyproteins synthesized during the next round of virus replication.
Important changes in protein activity can be brought about by chemical modification

Genomes have the capacity to code for 22 different amino acids: the 20 specified by the standard genetic code, selenocysteine, and (at least in archaea) pyrrolysine. The latter two are inserted into polypeptides by context-dependent reassignment of 5′-UGA-3′ and 5′-UAG-3′ codons, respectively (Section 1.3). This repertoire is increased dramatically by post-translational chemical modification of proteins, which results in a vast array of different amino acid types. The simpler types of modification occur in all organisms; the more complex ones, especially glycosylation, are rare in bacteria.

The simplest types of chemical modification involve addition of a small chemical group (for example, an acetyl, methyl, or phosphate group; Table 13.1) to an amino acid side chain or to the amino or carboxyl groups of the terminal amino acids in a polypeptide. Over 150 different modified amino acids have been documented in different proteins, with each modification carried out in a highly specific manner, the same amino acids being modified in the same way in every copy of the protein. A more complex type of modification is glycosylation, which involves the attachment of large carbohydrate side chains, called glycans, to polypeptides. There are two general types of glycosylation (Figure 13.40):

- **O-linked glycosylation** is the attachment of a glycan to the hydroxyl group of a serine or threonine.

- **N-linked glycosylation** involves attachment of a glycan to the amino group on the side chain of asparagine.

Glycosylation can result in attachment to the protein of structures comprising branched networks of 12 or more sugar units of various types. These glycans help to target proteins to particular sites in cells and increase the stability of proteins.

**Table 13.1 Examples of Post-Translational Chemical Modifications**

<table>
<thead>
<tr>
<th>Modification</th>
<th>Amino acid(s) modified</th>
<th>Examples of proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Addition of small chemical groups</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acetylation</td>
<td>Lys</td>
<td>Histones</td>
</tr>
<tr>
<td>Methylation</td>
<td>Lys</td>
<td>Histones</td>
</tr>
<tr>
<td>Phosphorylation</td>
<td>Ser, Thr, Tyr</td>
<td>Some proteins involved in signal transduction</td>
</tr>
<tr>
<td>Hydroxylation</td>
<td>Pro, Lys</td>
<td>Collagen</td>
</tr>
<tr>
<td>Carbamoylation</td>
<td>Lys</td>
<td>Ribulose-bisphosphate carboxylase</td>
</tr>
<tr>
<td>N-formylation</td>
<td>N-terminal Gly</td>
<td>Melittin</td>
</tr>
<tr>
<td><strong>Addition of sugar side chains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>O-linked glycosylation</td>
<td>Ser, Thr</td>
<td>Many membrane proteins and secreted proteins</td>
</tr>
<tr>
<td>N-linked glycosylation</td>
<td>Asn</td>
<td>Many membrane proteins and secreted proteins</td>
</tr>
<tr>
<td><strong>Addition of lipid side chains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acylation</td>
<td>Ser, Thr, Cys</td>
<td>Many membrane proteins</td>
</tr>
<tr>
<td>N-myristoylation</td>
<td>N-terminal Gly</td>
<td>Some protein kinases involved in signal transduction</td>
</tr>
<tr>
<td><strong>Addition of biotin</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Biotinylolation</td>
<td>Lys</td>
<td>Various carboxylase enzymes</td>
</tr>
</tbody>
</table>
circulating in the bloodstream. Another type of large-scale modification involves attachment of long-chain lipids, often to serine or cysteine. This process is called acylation and occurs with many proteins that become associated with membranes. A less common modification is biotinylation, in which a molecule of biotin is attached to a small number of enzymes that catalyze the carboxylation of organic acids such as acetate and propionate.

Identifying the chemical modifications carried by individual proteins is an essential part of protein profiling. This is because chemical modification often plays a central role in determining the precise biochemical activity of the target protein. An example is provided by the enzyme called ribulose-bisphosphate carboxylase (Rubisco), which is the major component of the proteome of plant leaves and is looked on as the most abundant protein in the biosphere. Rubisco catalyzes a key step in the photosynthesis pathway, where carbon, in the form of carbon dioxide, is fixed by reaction with ribulose 1,5-bisphosphate. The enzyme’s active site (more precisely, sites, as there are eight identical ones per protein) includes a lysine that has been modified by addition of a carboxyl group to give the carbamoyl derivative of this amino acid (Figure 13.41). In low-light conditions, carbamoylation of the lysine does not occur, so Rubisco activity is reduced. The chemical modification is therefore a means of controlling the enzyme so that the rate of carbon fixation is coordinated with the amount of energy that the plant is absorbing from sunlight. Many other enzymes are converted from active to inactive forms by chemical modification. Phosphorylation is particularly important in this regard, especially in signal transduction pathways, where it is used to activate proteins that carry signals from cell-surface receptors to transcription factors and other regulatory proteins (Section 14.1).

During protein profiling, a chemical modification can be detected because it will result in a characteristic change in the m/z value of the peptide or protein that carries the new chemical group. Sometimes, however, the modified form of the protein is present in relatively low amounts, in which case an enrichment procedure might be needed before the modified protein can be detected. Phosphorylated proteins fall into this category, as signal transduction can be effective even if only a few copies of the relevant proteins have been activated. Phosphorylated peptides can be partially purified from a proteome by immobilized metal ion affinity chromatography, which makes use of the interaction that occurs between a phosphate group and metal ions such as Fe$^{3+}$, Ca$^{2+}$, and Zr$^{4+}$. Purification is not complete, because nonphosphorylated peptides can also be retained in the affinity column, but the recovered sample is sufficiently enriched in phosphopeptides to enable their detection. Affinity chromatography can also be used to enrich proteomes for other types of modified protein. Glycosylated proteins, for example, can be captured by lectins, which are plant or animal proteins with specific sugar-binding properties. An example is concanavalin A from jack bean (Canavalia ensiformis), which binds to terminal glucose and mannose units in O-linked glycans.

![Figure 13.40 Glycosylation.](image)

(A) O-linked glycosylation

![Figure 13.41 Formation of the carbamoyl derivative of lysine.](image)
13.5 BEYOND THE PROTEOME

Protein synthesis is traditionally looked on as the final stage of genome expression, but this view obscures the true role of the proteome as part of the final link that connects the genome with the biochemistry of the cell. Exploring the nature of this link is proving to be one of the most exciting and productive areas of biological research and has led to new concepts surrounding the metabolome and systems biology.

The metabolome is the complete set of metabolites present in a cell

Often in biology, the most important steps forward do not result from some groundbreaking experiment but instead arise because biologists devise a new way of thinking about a problem. The introduction of the concept of the metabolome is an example. The metabolome is defined as the complete collection of metabolites present in a cell or tissue under a particular set of conditions. In other words, a metabolome is a biochemical blueprint, and its study, which is called metabolomics or biochemical profiling, gives a precise description of the biochemistry underlying different physiological states, including disease states, that can be adopted by a cell or tissue. By converting the biochemistry of a cell into an itemized set of metabolites, metabolomics provides a data set that can be directly linked to the equivalent, itemized information which emerges from proteomics and other studies of genome expression.

A metabolome can be characterized by chemical techniques such as infrared spectroscopy, mass spectrometry, and nuclear magnetic resonance spectroscopy which, individually and in combination, can identify and quantify the various small molecules that make up the metabolites in a cell. When these data are combined with knowledge about the reaction rates for various steps in well-characterized biochemical pathways such as glycolysis and the tricarboxylic acid cycle, it is possible to model the metabolic flux, the rate of flow of metabolites through the network of pathways that make up the cellular biochemistry (Figure 13.42). Changes

Figure 13.42 Metabolic flux studies of sugar utilization by E. coli. This example of metabolic flux analysis shows the flow of metabolites through the central energy-generating pathways during aerobic growth of E. coli on glucose (left) and galactose (right). The flux analysis shows that galactose utilization involves greater use of the glyoxylate shunt and modified use of the tricarboxylic acid (TCA) cycle compared with glucose utilization. Abbreviations: G6P, glucose 6-phosphate; 6PG, 6-phosphogluconate; F6P, fructose 6-phosphate; PSP, pentose 5-phosphate; FBP, fructose bisphosphate; T3P, 3-phosphoglycerate; PEP, phosphoenolpyruvate; PYR, pyruvate; ACE-CoA, acetyl-CoA; ACE, acetate; MAL, malate; OAA, oxaloacetate; FUM, fumarate; GOX, glyoxylate; CIT, citrate; SUC, succinate; SUC-CoA, succinyl-CoA; ICT, isocitrate; 2-OG, 2-oxoglutarate. (Adapted from Haverkorn van Rijsewijk BRB, Nanchen A, Nallet S et al. [2011] Mol Syst Biol 7:477. With permission from EMBO Press.)
in the metabolome can then be defined in terms of perturbations in the flux of metabolites through one or more parts of the network, providing a very sophisticated description of the biochemical basis for changes in the physiological state. This leads to the possibility of **metabolic engineering**, in which changes are made to the genome by mutation or recombinant DNA techniques in order to influence the cellular biochemistry in a predetermined way, for example to increase the synthesis of an antibiotic by a microorganism.

At present, metabolomics is most advanced with organisms such as bacteria and yeast, whose biochemistries are relatively simple. Considerable research is currently being directed toward the human metabolome, with the objective of describing the metabolic profiles of healthy tissues, of disease states, and of tissues in patients undergoing drug treatments. It is hoped that when these studies reach maturity, it will be possible to use the metabolic information to design drugs that reverse or mitigate the particular flux abnormalities that occur in the disease state. Biochemical profiling could also indicate any unwanted side effects of drug treatment, enabling modifications to be made to the chemical structure of the drug or to its mode of use, so that these side effects are minimized.

**Systems biology provides an integrated description of cellular activity**

The emphasis that is now placed on concepts such as metabolic flux illustrates the importance of understanding not simply the molecules—RNAs, proteins, and metabolites—whose synthesis the genome directs but also the biological systems that result from the coordinated activity of those molecules. This new emphasis on systems is a direct result of the leap that has been made in recent years from genes to genomes. One of the underlying principles of the pre-genome era of molecular biology was the one gene–one enzyme hypothesis, first put forward by George Beadle and Edward Tatum in the 1940s. By “one gene–one enzyme”, Beadle and Tatum were emphasizing that a single gene codes for a single protein, which, if the protein is an enzyme, directs a single biochemical reaction. The **trpC** gene of *E. coli*, for example, codes for the enzyme indole-3-glycerol-phosphate synthase, which converts 1-(2-carboxyphenylamino)-1’-deoxyribulose 5’-phosphate into indole-3-glycerol phosphate. However, this enzyme does not work in isolation: its activity forms part of the biochemical pathway that results in synthesis of tryptophan. The other enzymes in this pathway are specified by the genes **trpA**, **trpB**, **trpD**, and **trpE**, which together with **trpC** form the tryptophan operon of *E. coli* (see Figure 8.9B). The tryptophan biosynthesis pathway is therefore a simple biological system, and the tryptophan operon is the set of genes that specify that pathway. But simply transcribing and translating the genes in the operon will not lead to the synthesis of tryptophan. Successful operation of the system requires that the biosynthetic enzymes be present at the appropriate place in the cell, in the appropriate relative amounts, at the appropriate time. The system is therefore dependent on factors such as the rate of synthesis of the proteins coded by the genes, correct folding of these proteins into functional enzymes, the rate of degradation of the enzyme molecules, their localization in the cell, and the presence of the necessary amounts of the metabolites that act as substrates and cofactors for tryptophan synthesis. This simple biological system is starting to assume quite considerable complexity. And yet, with this system we are considering only 5 of the 4315 genes in the *E. coli* genome.

An early illustration of the power of the systems approach to biology was provided by work done during the mid-2000s on synthesis of the *E. coli* flagellum. Pre-genome studies had shown that flagellar synthesis requires 51 genes organized into 12 operons, which are activated in three groups (Figure 13.43). The first group to be activated comprises a single operon containing two genes that code for a protein that acts as a master regulator, switching on expression of the second group of seven operons, whose genes together specify components of the basal structure of the flagellum. One of these genes codes for a second regulatory protein that switches on the remaining four operons, which direct synthesis of
Chapter 13: Proteomes

The system responsible for flagellum biosynthesis in E. coli.

the flagellum filament and the biochemical system that enables the bacterium to respond to chemical stimuli by rotating its flagellum in order to swim toward an attractant. Careful use of reporter genes attached to individual operons has revealed the precise order in which the operons in each group are activated and has enabled activation coefficients, which are measures of the relative rates of expression, to be assigned to each operon. The resulting information is sufficient for the system to be modeled on a computer, enabling the detailed roles of the two regulatory proteins to be determined. From the computer models, the effects of subtle changes to the system (such as a change in the properties of one of the regulators) can be predicted and then tested by further experiments with the biological system.

Following the early successes with bacterial systems, the modeling approach has been applied to more complex biological processes in eukaryotes, including humans. An interesting example of the use of systems biology to study human disease was prompted by the discovery that the chaperone protein called GRP78 is associated with resistance of some types of breast cancer to drugs such as tamoxifen or ICI 182,780, which interfere with the ability of estrogen to stimulate the proliferation of cancerous cells. When synthesis of GRP78 is reduced, the cancerous cells undergo apoptosis and so do not proliferate. In contrast, when GRP78 is overexpressed, the cancer becomes drug-resistant and there is an increase in autophagy, the process in which cancerous cells respond to the stress caused by drug treatment by degrading and recycling damaged cell components. An analysis of the interplay between protein folding (the normal function of GRP78), apoptosis, and autophagy might therefore indicate how breast cancer cells develop drug resistance, which in turn could aid the design of better anti-estrogen drugs. The key question is whether our current understanding of the molecular basis of protein folding, apoptosis, and autophagy is sufficiently comprehensive for analyses of these processes to give reliable indications of the possible basis to drug resistance. To answer this question, the available data were used to construct separate models of the molecular events underlying protein folding, apoptosis, and autophagy in healthy tissues. These three models were then integrated by adding information on the cross-talk between the processes in breast cancer tissues (Figure 13.44). Cross-talk refers to the way in which events occurring during one process influence events in a second process.
Mathematical approaches were then used to test how well the integrated model explains the observed results of GRP78 repression and overexpression in breast cancer cells in the presence or absence of anti-estrogen drugs. The outcomes predicted by the mathematical analysis turned out to be in close agreement with experimental data, indicating that our current knowledge of protein folding, apoptosis, and autophagy is robust and that models of the type shown in Figure 13.44 can be used in interpretation and application of experimental data on the molecular basis for drug resistance in breast cancer.
SUMMARY

- The proteome is the collection of protein molecules present in a cell.
- The composition of a proteome can be examined by top-down and bottom-up processes. In top-down proteomics individual proteins are directly examined by mass spectrometry, whereas in bottom-up proteomics the proteins are broken into peptides by treatment with a sequence-specific protease, such as trypsin, prior to mass spectrometry.
- The initial stage in both proteomic approaches is separation of the proteins in the proteome. This can be achieved by two-dimensional polyacrylamide gel electrophoresis or by column chromatography.
- Two proteomes can be compared by labeling the components with different fluorescent markers or with isotope-coded affinity tags.
- Protein arrays can also be used to study proteomes.
- Pairs and groups of proteins that interact with one another can be identified by phage display, the yeast two-hybrid system, tandem affinity purification, and the multidimensional protein identification technique.
- Protein interaction maps display the detailed interactions between the components of a proteome.
- Proteins are synthesized by ribosomes, which coordinate the events occurring during protein synthesis and also catalyze at least some of the chemical reactions. Bacteria are able to inactivate their ribosomes during periods of stress.
- In eukaryotes, initiation factors mediate the large-scale remodeling of proteome structure. For example, phosphorylation of eIF-2 occurs during stresses such as heat shock and results in a decrease in the overall level of protein synthesis. The translation of individual mRNAs can also be regulated.
- Proteins must be turned over so that proteomes can respond to changing conditions. The degradation pathway involves tagging the targeted protein with ubiquitin followed by breakdown in a proteasome.
- Most polypeptides are inactive until they have been folded into their three-dimensional structures. The amino acid sequence contains the information for folding, but folding of large proteins is a complex process that can result in diversion of the partially folded protein down an incorrect side pathway. Molecular chaperones help other proteins to fold, reducing the number of folding errors that occur.
- Some proteins are processed by internal cleavage, possibly as a means of converting an inactive protein into the active form. Polyproteins are cleaved to give multiple protein products.
- Chemical modification by glycosylation, acylation, or addition of small chemical groups can bring about changes in protein activity.
- The metabolome is defined as the complete collection of metabolites present in a cell or tissue under a particular set of conditions.
- Systems biology attempts to provide an integrated description of cellular activity.

SHORT ANSWER QUESTIONS

1. Distinguish between the methods used in top-down and bottom-up proteomics.
2. Describe how the components of a proteome are separated prior to the mass spectrometry stage of protein profiling.
3. Outline the different mass spectrometric methods that are used in protein profiling.

4. How can differences between the compositions of two proteomes be identified?

5. What methods are available for identification of proteins that interact with one another?

6. Compare the compositions of the bacterial and eukaryotic ribosomes.

7. Explain how bacteria downsize their proteomes during periods of stress.

8. Describe how large and small changes in eukaryotic proteomes can be brought about by initiation factor phosphorylation and by transcript-specific regulation.

9. How are proteins degraded?

10. Outline the key features of the protein folding pathway and describe the role of molecular chaperones in protein folding.

11. Summarize the roles in protein processing of (A) proteolytic cleavage and (B) chemical modification.

12. Describe the objectives of metabolomics and systems biology.

**IN-DEPTH PROBLEMS**

1. Under what circumstances might a pair of proteins have a functional relationship but no physical interaction? Are there possible scenarios where the reverse might be true—a pair of proteins display a physical interaction but have no functional relationship?

2. Discuss the role of the hubs in a protein interaction map.

3. There appears to be no biological reason why a DNA polynucleotide could not be directly translated into protein, without the intermediate role played by mRNA. What advantages do eukaryotic cells gain from the existence of mRNA?

4. To what extent have studies of ribosome structure been of value in understanding the detailed process by which proteins are synthesized?

5. Explain why systems biology is receiving so much attention at the present time.

**FURTHER READING**

- **Methods for studying the composition of a proteome**


**Identifying protein interactions**


**Protein synthesis**


**Factors influencing the composition of a proteome**


**Protein degradation**


**Protein folding and other protein processing events**


**Metabolomics and systems biology**


We began Part III of *Genomes* by looking inside the nucleus and asking how the packaging of chromosomal DNA into nucleosomes and chromatin fibers influences the accessibility of individual genes and how that accessibility sets the overall pattern of genome expression. We then remained in the nucleus in Chapters 11 and 12, examining how the interaction between DNA-binding proteins and the genome results in synthesis of the transcriptome, before transferring to the cytoplasm in Chapter 13 to explore the synthesis and maintenance of the proteome. Throughout those four chapters, our attention was firmly on the genome and the RNAs and proteins whose synthesis the genome specifies. However, a genome does not exist purely for its own benefit. The genome resides in a cell that might itself be an organism or which forms part of a larger multicellular organism. Through the transcriptome and proteome, the genome specifies the biochemical activities occurring in the cell in which it resides, enabling the cell to generate energy and to grow and divide. The genome must therefore be responsive to the extracellular environment, so that the biochemical activities of the cell are continually in tune with the available nutrient supply and the prevailing physical and chemical conditions. Cells in multicellular organisms must also respond to hormones and growth factors, which may signal changes in the environment to which the organism must adapt but which also, importantly, coordinate the biochemical activities of different cells so that the multicellular organism functions as a unified entity. The response of the genome to these extracellular signals is usually transient: the expression pattern reverts to its original form when the signals cease or takes on a new pattern if one set of signals is replaced by a second set.

Other changes in genome activity are permanent, or at least semipermanent, and result in the cell’s biochemical signature becoming altered in a way that is not readily reversible. These changes lead to **differentiation**, the adoption by the cell of a specialized physiological role. Differentiation pathways are known in many unicellular organisms, but we more frequently associate differentiation with multicellular organisms, in which a variety of specialized cell types (over 400 types in humans) are organized into tissues and organs. The assembly and modification of these complex multicellular structures over time constitutes the **development** pathway for the organism, the successful completion of which requires temporal control over genome activity, so that the developmental events occur in the correct order and at the appropriate times.

In this chapter, we will explore how the genome acts within the context of cell and organism by responding to extracellular signals and by driving the biochemical changes that underlie differentiation and development. Underlying all these processes is the regulation of genome expression, which has been a constant theme in our discussion of the transcriptome and proteome during the previous two chapters. This means that we have already examined many of the ways in which genome expression can be regulated, and our objective now is not simply to reiterate this material but instead, as the title of the chapter indicates, to understand how these regulatory systems operate within the contexts of the cell and organism.
14.1 THE RESPONSE OF THE GENOME TO EXTERNAL SIGNALS

For unicellular organisms, the most important external signals relate to nutrient availability, as these cells live in variable environments in which the identities and relative amounts of the nutrients change over time. The genomes of unicellular organisms therefore include genes for uptake and utilization of a range of nutrients, and changes in nutrient availability are shadowed by changes in genome activity, so that at any one time only those genes needed to utilize the available nutrients are expressed. The response of the lactose operon of *Escherichia coli* to the available glucose and lactose supply (Section 12.2) provides a typical example of the way in which expression of a bacterial genome can be influenced by the nutrient status of the external environment. Most cells in multicellular organisms live in less variable environments, but the maintenance of these environments requires coordination between the activities of different cells. For these cells, the major external stimuli are therefore hormones, growth factors, and related compounds that convey signals within the organism and stimulate coordinated changes in genome activity.

To exert an effect on genome activity, the nutrient, hormone, growth factor, or other extracellular compound that represents the external signal must influence events within the cell. There are two ways in which it can do this (Figure 14.1):

- directly, by acting as a signaling compound that is transported across the cell membrane and into the cell
- indirectly, by binding to a cell surface receptor that transmits a signal into the cell.

Signal transmission, by direct or indirect means, is one of the major research areas in cell biology, with attention focused in particular on its relevance to the abnormal biochemical activities that underlie cancer. Many examples of signal transmission have been discovered, some found in a variety of organisms and others restricted to just a few species. In this section we will survey the most important of these signaling pathways.

**Signal transmission by import of the extracellular signaling compound**

First, we will consider those signaling compounds that are able to pass through the cell membrane and into the cell. The simplest way in which an imported signaling compound could influence genome expression is by acting in the same way as one of the various regulatory proteins that we have met in previous chapters, for example by activating or repressing assembly of the transcription initiation complex (Section 12.2) or by interacting with a splicing enhancer or silencer (Section 12.4). This might appear to be an attractively straightforward way of regulating genome activity, but it is not a common mechanism. The reason for this is not clear but probably relates, at least partly, to the difficulty in designing a protein that combines the hydrophobic properties needed for effective transport across a membrane with the hydrophilic properties needed for migration through the aqueous cytoplasm to the protein’s site of action in the nucleus or elsewhere in the cell.

The one clear example of a signaling compound that can function in this way is provided by lactoferrin, a mammalian protein found mainly in milk and, to a lesser extent, in the bloodstream. The specific function of lactoferrin has been difficult to pin down, but it seems to play a role in the body’s defenses against microbial attack. As its name suggests, lactoferrin is able to bind iron, and it is thought that at least part of its protective role arises from its ability to reduce free iron levels in milk, thereby starving invading microbes of this essential cofactor. It might therefore appear unlikely that lactoferrin would have a role in genome expression, but it has been known since the early 1980s that the protein is multitalented and, among other things, can bind to DNA. This property was linked to a second function of lactoferrin—stimulation of the blood cells involved in the immune response—when in 1992 it was shown that the protein is taken up by immune
cells, enters their nuclei, and attaches to the genome. Subsequently, it was shown that the DNA binding is sequence-specific and that bound lactoferrin activates transcription of the interleukin 1β gene, at least in cultured cells. These results might appear to suggest that lactoferrin is a transcription factor, but we should be cautious in reaching this conclusion. The lactoferrin gene has two alternative promoters (Section 12.2). Transcription from one of these promoters gives the full-length protein that is secreted and then taken up by other cells. Transcription from the second promoter results in a shortened form, called δ-lactoferrin, which lacks the signal peptide at the N-terminus of the full-length protein. δ-Lactoferrin, therefore, is not secreted but has been shown to act as a transcription factor in the cells in which it is synthesized. It is therefore possible that the transcriptional activity of secreted lactoferrin is an artifact, resulting from this protein attaching to what are, in fact, δ-lactoferrin binding sites. This spurious binding may be observable in experimental systems such as cultured cells but may not have any relevance in intact tissues.

Although few, if any, imported signaling compounds are able themselves to act as activators or repressors of genome expression, many have the ability to influence the activity of regulatory proteins that are already present in the cell. We encountered one example of this type of regulation in Section 12.2 when we studied the lactose operon of *E. coli*. This operon responds to extracellular levels of lactose, where lactose acts as a signaling molecule that enters the cell and, after conversion to its isomer allolactose, influences the DNA-binding properties of the lactose repressor, hence determining whether or not the lactose operon is transcribed (see Figure 12.13). Many other bacterial operons coding for genes involved in sugar utilization are controlled in a similar way. Interaction between a signaling compound and a transcription factor is also a common means of regulating genome activity in eukaryotes. A good example is provided by the control system that maintains the intracellular metal ion content at an appropriate level. Cells need metal ions such as copper and zinc as cofactors in biochemical reactions, but these metals are toxic if they accumulate in the cell above a certain level. Their uptake therefore has to be carefully controlled so that the cell contains sufficient metal ions when the environment is lacking in metal compounds but does not overaccumulate metal ions when the environmental concentrations are high. The strategies used are illustrated by the copper control system of *Saccharomyces cerevisiae*. This yeast has two copper-dependent transcription factors, Mac1p and Ace1p. Both of these proteins bind copper ions, and the binding induces conformational changes that enable the factors to stimulate expression of their target genes (Figure 14.2). For Mac1p these target genes code for copper-uptake proteins, whereas for Ace1p they are genes coding for proteins such as superoxide dismutase that are involved in copper detoxification. The balance between the activities of Mac1p and Ace1p, controlled by the metal ions acting as signaling molecules, ensures that the copper content of the cell remains within acceptable levels.

Transcription activators are also the targets for steroid hormones, which are signaling compounds that coordinate a range of physiological activities in the cells of higher eukaryotes. They include the sex hormones (estrogens for female sex development and androgens for male sex development) and the glucocorticoid and mineralocorticoid hormones. Steroids are hydrophobic and so easily penetrate the cell membrane. Once inside the cell, each hormone binds to a specific steroid receptor protein, which is usually located in the cytoplasm. After binding, the activated receptor migrates into the nucleus, where it attaches to a hormone response element upstream of a target gene. Once bound, the receptor acts as a transcription activator. Response elements for each receptor are located upstream of 50–100 genes, often within enhancers, so a single steroid hormone can induce a large-scale change in the biochemical properties of the cell. All steroid receptors are structurally similar, not only with regard to their DNA-binding domains but also in other parts of their protein structures (Figure 14.3). Recognition of these similarities has led to the identification of a number of putative or orphan steroid receptors whose hormonal partners and cellular functions are not yet known. The structural similarities have also shown that a second set of receptor proteins, such as copper and zinc as cofactors in biochemical reactions, but these metals are toxic if they accumulate in the cell above a certain level. Their uptake therefore has to be carefully controlled so that the cell contains sufficient metal ions when the environment is lacking in metal compounds but does not overaccumulate metal ions when the environmental concentrations are high. The strategies used are illustrated by the copper control system of *Saccharomyces cerevisiae*. This yeast has two copper-dependent transcription factors, Mac1p and Ace1p. Both of these proteins bind copper ions, and the binding induces conformational changes that enable the factors to stimulate expression of their target genes (Figure 14.2). For Mac1p these target genes code for copper-uptake proteins, whereas for Ace1p they are genes coding for proteins such as superoxide dismutase that are involved in copper detoxification. The balance between the activities of Mac1p and Ace1p, controlled by the metal ions acting as signaling molecules, ensures that the copper content of the cell remains within acceptable levels.
the **nuclear receptor superfamily**, belongs to the same general class as steroid receptors, although the hormones that they work with are not steroids. As their name suggests, these receptors are located in the nucleus rather than the cytoplasm. They include the receptors for vitamin D$_3$, whose roles include control of bone development, and thyroxine, which stimulates the tadpole-to-frog metamorphosis.

Steroid and nuclear receptors are dimers, each subunit of which possesses one of the treble clef zinc fingers that are characteristic of this group of proteins (see **Figure 11.14**). Each of these zinc fingers recognizes and binds to a 6 bp sequence in its hormone response element. For most steroid receptors, the pair of 6 bp sequences is arranged as a direct or inverted repeat, separated by a 0–4 bp spacer (**Figure 14.4**). The response elements for nuclear receptors are similar except that the recognition sequences are more frequently direct repeats. The spacer serves simply to ensure that the distance between the recognition sequences is appropriate for the orientation of the zinc fingers in the receptor protein. This means that different receptor proteins can possess the same pair of zinc fingers but recognize different response elements, specificity being maintained by the orientation of the fingers and the spacing between the recognition sequences.

### Receptor proteins transmit signals across cell membranes

Many extracellular signaling compounds are unable to enter the cell because they are too hydrophilic to penetrate the lipid membrane and the cell lacks a specific transport mechanism for their uptake. In order to influence genome activity, these signaling compounds must bind to cell surface receptors that transfer the signal across the cell membrane and into the cell (see **Figure 14.1**). A cell surface receptor is a protein that spans the membrane, with a site for binding the signaling compound on the outer surface. Binding of the signaling compound results in a conformational change in the receptor, inducing a biochemical event within the cell, that event forming the first step in the intracellular stage of the **signal transduction** pathway.

There are several types of cell surface receptor, but most of those mediating changes in genome expression are **kinase** or **kinase-associated receptors**. As the names indicate, these receptors induce the intracellular stage of the signal transduction pathway by adding a phosphate group to a cytoplasmic protein. The most important examples are the **tyrosine kinase receptors**, which add phosphates to tyrosine amino acids in their target proteins. Most tyrosine kinase receptors are dimers of identical subunits, each subunit comprising an extracellular binding domain and an intracellular kinase activity separated by a hydrophobic transmembrane region of some 25–35 amino acids (**Figure 14.5**). Tyrosine kinase receptors recognize a variety of extracellular signaling compounds, including **cytokines** and other growth factors, as well as hormones such as insulin. In the absence of the signaling compound, the two subunits of a receptor are disassociated. Attachment of the signal results in the subunits coming together to form the dimer, which activates the internal kinase activity, initiating the intracellular signal transduction pathway.

Tyrosine is not the only amino acid that can be phosphorylated in order to activate an intracellular protein. The **serine-threonine kinase receptors** act in a similar way to tyrosine kinase receptors, but they phosphorylate serines and/or threonines on their intracellular target proteins. The signaling compounds recognized by this type of receptor include transforming growth factor β (TGFβ) and members of the bone morphogenetic protein family of cytokines.

Both the tyrosine and serine-threonine groups of receptors possess kinase activity and so directly phosphorylate their internal target proteins. The **tyrosine**
14.1 The Response of the Genome to External Signals

**Kinase-Associated Receptors** are slightly different, as they do not themselves have kinase activity. Instead, they act indirectly by influencing the activity of intracellular tyrosine kinases. We will study an example of this type of receptor in the next section.

**Some Signal Transduction Pathways Have Few Steps Between Receptor and Genome**

With some signal transduction systems, stimulation of the cell surface receptor by attachment of the extracellular signaling compound has a direct effect on the activity of a transcription factor. This is the simplest system by which an extracellular signal can be transduced into a genomic response.

The direct system is used by the cell surface receptors for many cytokines, such as interleukins and interferons, which are extracellular polypeptides that control cell growth and division and influence genome expression via the JAK/STAT pathway. This pathway is found in all vertebrates, and related pathways are present in many invertebrates, including *Drosophila melanogaster* and *Caenorhabditis elegans*. In vertebrates, the receptors are members of the tyrosine kinase-associated family, each receptor protein being associated with an internal Janus kinase (JAK). Cytokine binding induces dimerization of the receptor, moving its pair of JAKs close enough together to phosphorylate one another (Figure 14.6). Phosphorylation activates the JAKs, which now phosphorylate transcription factors called STATs (signal transducers and activators of transcription). Phosphorylation causes pairs of STATs to form dimers that move to the nucleus, where they activate expression of a variety of genes.

Seven STATs have so far been identified in mammals, called STAT1, STAT2, STAT3, STAT4, STAT5A, STAT5B, and STAT6. Each of these except STAT2 can form homodimers, and heterodimers can form between STAT1 and STAT2, STAT1 and STAT3, and STAT5A and STAT5B. The composition of the dimers formed depends on the identity of the extracellular compound and also its concentration. With just nine possible dimers, the JAK/STAT system might appear to have limited flexibility, but the expectation that a particular dimer will always activate the same set of genes turns out to be incorrect. Interleukin 6 and interleukin 10, for example, have opposite effects on genome expression in human myeloid cells: interleukin 6 induces an inflammatory response and interleukin 10 suppresses inflammation, even though both exert their response by attaching to receptors that construct STAT3 dimers. How the same dimer can mediate different genome responses depending on the nature of the original signal is not yet understood.

![Figure 14.5 A tyrosine kinase receptor](image)

This is the epidermal growth factor (EGF) receptor, which is a typical example of a tyrosine kinase receptor. Attachment of an EGF molecule to each of the receptor subunits induces dimerization, which activates the kinase specified by the intracellular domains of the two monomers.

![Figure 14.6 The JAK/STAT pathway](image)
The DNA-binding domain of a STAT protein is made up of three loops emerging from a barrel-shaped β-sheet. This structure, called the immunoglobulin fold, is found in many proteins but is not usually associated with DNA binding. Other proteins that use the immunoglobulin fold for this purpose include the transcription factors NK-kB and Rel. The similarities between these proteins refer only to the tertiary structures of the DNA-binding domains because STATS, NK-kB, and Rel, as a whole, have very little amino acid sequence identity. The consensus sequence of the DNA binding sites for most STAT dimers was originally defined as 5′-TTN5,6AA-3′, largely from studies in which purified STATs were tested for their ability to bind to oligonucleotides of known sequence. Variations in the surrounding nucleotides and in the internal series of Ns are thought to specify the actual binding sites for different types of dimers. This particular type of binding site is called the interferon γ-stimulated gene response (GAS) element. Some heterodimers attach to additional proteins, such as p48, and then recognize a different binding site, called the interferon-stimulated response element (ISRE), whose consensus sequence is 5′-AGTTTNNTTCC-3′. Chromatin immunoprecipitation sequencing (ChiP-seq; see Section 11.1) has revealed thousands of binding sites for each STAT dimer in the human genome, at positions adjacent to target genes and also in enhancers. However, many of these binding sites lack a GAS or ISRE sequence, suggesting that the association of STATs with the genome is more complicated than originally thought. This added complexity is underlined by studies showing that unphosphorylated STATs also influence genome expression under some conditions and that the phosphorylated versions can form polymeric oligomers made up of more than two individual STATs. These observations may go some way toward explaining how the same STAT dimer can have different effects depending on the identity of the signaling compound, as described above for interleukins 6 and 10.

Some signal transduction pathways have many steps between receptor and genome

The relative simplicity of the JAK/STAT pathway contrasts with the more prevalent forms of signal transduction, in which the receptor represents just the first in a series of steps that lead eventually to the activation of one or more transcription factors. A number of these cascade pathways have been delineated in different organisms, the most important being the MAP (mitogen-activated protein) kinase or MAPK/ERK pathway.

The MAP kinase pathway responds to many extracellular signals, including mitogens, which are compounds that have similar effects to cytokines but more specifically stimulate cell division. The initial steps in the pathway are centered around the Ras proteins, three of which are known in mammalian cells (H-, K-, and N-Ras). Ras is a G-protein, a type of small protein that binds a molecule of either guanosine 5′-diphosphate (GDP) or guanosine 5′-triphosphate (GTP). When GDP is bound, Ras is inactive, but if the GDP is replaced with GTP, Ras becomes activated. Which nucleotide is bound to Ras depends on the balance between the activities of guanine nucleotide exchange factors (GEFs), which activate Ras by replacing the GDP with GTP, and GTPase-activating proteins (GAPs), which inactivate Ras by stimulating Ras to convert its bound GTP to GDP (Figure 14.7).

How exactly does the presence of the extracellular signal result in Ras activation or inactivation? Ras is attached to the inner surface of the cell membrane in the vicinity of the mitogen receptor. Binding of the signaling compound results in dimerization of the receptor and mutual phosphorylation of the internal parts of its two subunits (Figure 14.8). GEFs such as the SOS protein bind to the phosphorylated receptor and activate Ras by replacing its GDP with GTP. The phosphorylated receptor might also recruit GAPs such as RasGAP, which have the opposite effect on Ras, deactivating it by stimulating the conversion of GTP to GDP. The balance between GEF and GAP activity therefore determines the nucleotide status of Ras and modulates further transmission of the signal along the transduction pathway.
Ras, when bound to GTP, is able to activate Raf, the next protein in the MAP kinase pathway. Raf is itself a kinase, but it is usually inactive because the protein is folded in such a way that the N-terminal region of its polypeptide blocks access to the catalytic site. The Ras–GTP complex is able to bind to Raf, causing a conformational change in the latter that unblocks the active site. Raf then initiates a cascade of phosphorylation reactions (Figure 14.9). It phosphorylates Mek, activating this protein so that it, in turn, phosphorylates a MAP kinase. The activated MAP kinase now moves into the nucleus where it activates, again by phosphorylation, a series of transcription factors. The MAP kinase also phosphorylates another protein kinase, this one called Rsk, which phosphorylates and activates a second set of factors. Additional flexibility is provided by the possibility of replacing one or more of the proteins in the MAP kinase pathway with related proteins, ones with slightly different specificities, and so switching on another suite of transcription activators.

The MAP kinase pathway is used by vertebrate cells, and equivalent pathways, using intermediates similar to those identified in mammals, are known in other organisms. Each step in these cascade pathways involves a physical interaction between two proteins, often resulting in the downstream member of the pair becoming phosphorylated. Phosphorylation activates the downstream protein, enabling it to form a connection with the next protein in the cascade. These interactions involve special protein–protein binding domains, such as the ones called SH2 and SH3, which bind to receptor domains on their partner proteins. The receptor domains contain one or more tyrosines that must be phosphorylated in order for docking to take place. Hence the upstream protein contains the receptor domain, whose phosphorylation status determines whether the protein can bind its downstream partner and thereby propagate the signal (Figure 14.10).

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**Figure 14.8** Activation of Ras at the start of the MAP kinase pathway. Binding of the mitogen to the receptor subunit results in receptor dimerization and autophosphorylation. A guanine nucleotide exchange factor (GEF), such as the SOS protein, associates with the receptor and activates Ras by replacing its GDP molecule with a GTP.

**Figure 14.9** The second part of the MAP kinase pathway. MK is the MAP kinase. Elk-1, c-Myc, and SRF (serum response factor) are examples of transcription factors activated at the end of the pathway.

**Figure 14.10** A scheme for interaction of proteins in a signaling cascade. The upstream protein is phosphorylated and hence able to bind its downstream partner. Binding leads to phosphorylation of the receptor domain in the downstream protein, propagating the signal.
Some signal transduction pathways operate via second messengers

Some signal transduction cascades do not involve direct transfer of the external signal to the genome but instead utilize an indirect means of influencing transcription. These pathways make use of second messengers, which are less specific internal signaling compounds that transduce the signal from a cell surface receptor in several directions so that a variety of cellular activities, not just transcription, respond to the one signal.

In Section 12.2 we saw how glucose modulates the bacterial catabolite activator protein by influencing the cellular cyclic adenosine monophosphate (cAMP) level. Cyclic nucleotides are also important second messengers in eukaryotic cells. Some cell surface receptors have guanylate cyclase activity, and so convert GTP to cyclic guanosine monophosphate (cGMP), but most receptors in this family work indirectly by influencing the activity of cytoplasmic cyclases and decyclases. These cyclases and decyclases determine the cellular levels of cGMP and cAMP, which in turn control the activities of various target enzymes. An example of a target enzyme is protein kinase A, which is stimulated by cAMP. One of the functions of protein kinase A is to phosphorylate, and hence activate, the transcription factor called CREB. As well as activating its specific target genes, CREB stimulates the activity of a variety of other genes by interacting with a second protein, p300, to form a complex called p300/CBP, which is able to modify histone proteins and so affect chromatin structure and nucleosome positioning (Section 10.2).

As well as being activated indirectly by cAMP, p300/CBP responds to another second messenger, calcium. The calcium ion concentration within the lumen of the endoplasmic reticulum is higher than that in the rest of the cell, so proteins that open calcium channels in the endoplasmic reticulum allow calcium ions to flow into the cytoplasm. This can be induced by extracellular signals that activate tyrosine kinase receptors, which in turn activate phospholipases that cleave phosphatidylinositol 4,5-bisphosphate [PtdIns(4,5)P₂], a lipid component of the inner cell membrane, into inositol 1,4,5-trisphosphate [Ins(1,4,5)P₃] and 1,2-diacylglycerol (DAG). Ins(1,4,5)P₃ opens the calcium channels in the endoplasmic reticulum (Figure 14.11). Ins(1,4,5)P₃ and DAG are themselves second messengers that can initiate other signal transduction cascades. Both calcium- and lipid-induced cascades target transcription factors, but only indirectly: the primary targets are other proteins. Calcium, for example, binds to and activates the protein called calmodulin, which regulates a variety of enzymes, including protein kinases, ATPases, phosphatases, and nucleotide cyclases.

14.2 Changes in genome activity resulting in cellular differentiation

Transient changes in genome activity are, by definition, readily reversible: the genome expression pattern reverts to its original state when the external stimulus is removed. In contrast, the permanent and semipermanent changes in genome activity that underlie cellular differentiation must persist for long periods and ideally should be maintained even when the stimulus that originally induced them has disappeared. We therefore anticipate that the regulatory mechanisms bringing about these longer-term changes will involve systems in addition to the modulation of transcription factors. This expectation is correct, the mechanisms that result in differentiation including changes in chromatin structure and physical rearrangement of the genome.

Some differentiation processes involve changes to chromatin structure

We studied some of the effects that chromatin structure can have on genome expression in Section 10.2. These effects range from the modulation of transcription initiation at an individual promoter by nucleosome repositioning through to the silencing of large segments of DNA locked up in higher-order chromatin structures. The latter is an important means of bringing about long-term changes in genome activity and is implicated in a number of differentiation
events. One example involves the Polycomb group (PcG) proteins, which were first discovered in D. melanogaster but are now known to have homologs in other organisms including mammals and plants. Polycomb proteins recognize DNA sequences called Polycomb response elements, their attachment inducing localized formation of heterochromatin, the condensed form of chromatin that prevents transcription of the genes that it contains (Figure 14.12). Each response element is approximately 10 kb in length and contains multiple copies of the DNA binding sites for PcG proteins such as Pleiohomeotic (PHO) and Pleiohomeotic-like (PHOL), which form a PhoRC complex that probably acts as the primary DNA recognition component of the Polycomb system (Figure 14.13). Once PhoRC is bound to the DNA, it recruits a second group of PcG proteins, called Polycomb repressive complex 2 (PRC2). One of the members of this complex, EZH2, is a histone methyltransferase that trimethylates lysine-27 of histone H3 and possibly also methylates lysine-9 (Section 10.2). These are repressive histone modifications (see Table 10.1) and hence induce heterochromatin formation. The methylations are then recognized by Polycomb repressive complex 1 (PRC1), which contains a second histone-modifying enzyme. This enzyme adds a ubiquitin group to lysine-119 of histone H2A, leading to further compaction of the heterochromatin, which propagates along the DNA for tens of kilobases in either direction. The EZH2 protein also attracts DNA methyltransferases into the bound PcG complex, so the genome region becomes further silenced by DNA methylation (Section 10.3).

In Drosophila, the regions that become silenced contain homeotic selector genes that, as we will see in Section 14.3, specify the development of individual body parts of the fly. As only one body part must be specified at a particular position in the fruit fly, it is important that a cell expresses only the correct homeotic gene. This is ensured by the action of the PcG proteins, which permanently silence the homeotic genes that must be switched off. PcG proteins do not, however, determine which genes will be silenced, as expression of these genes has already been repressed before the proteins bind to their response elements. The role of the PcG proteins is therefore to maintain rather than initiate gene silencing. An important point is that the heterochromatin that is formed is heritable: after division, the two new cells retain the heterochromatin established in the parent cell. This type of regulation of genome activity is therefore permanent not only in a single cell but also in a cell lineage.

The trithorax group (trxG) proteins act in a similar manner to the PcG proteins but have the opposite effect, maintaining an open chromatin state in the regions of active genes. Their targets include the same homeotic genes that are silenced, in different body parts, by PcG proteins. How the opposing actions of PcG and trxG proteins are controlled is not yet known. The notion that there is a straightforward toggle switch, with PcG switching genes off and trxG switching them on again, is probably overly simplistic. This would imply that PcG proteins would be associated with the target regions of the genome when those regions are repressed, and trxG proteins would be present when the regions are active. In fact, both groups of protein appear to be present all the time. One hypothesis is that the heterochromatin induced by the PcG proteins is the default state, and the role of trxG is to modify PcG action in the region of those genes whose activities are needed in a particular cell.
Yeast mating types are determined by gene conversion events

A permanent change in the pattern of genome expression can also be brought about by changing the physical structure of the genome. Here we are not referring to changes in chromatin structure as just discussed, whereby the sequence of the genome is unchanged but certain regions become inaccessible because they are compacted into heterochromatin. By physical structure we mean the nucleotide sequence, and the changes we refer to include gene conversion, in which one part of the genome is deleted and replaced with a copy of a segment from a different part of the genome, and rearrangements that result in parts of the genome that were previously separate becoming linked to one another (Figure 14.14). These events constitute an effective, if drastic, way of bringing about the permanent change in genome expression needed to maintain a differentiated state.

Yeast mating types provide an example of the use of gene conversion to bring about cellular differentiation. Mating type is the equivalent of sex in yeasts and other eukaryotic microorganisms. Because these organisms reproduce mainly by vegetative cell division, there is the possibility that a population, being derived from just one or a few ancestral cells, will be largely or completely composed of a single mating type and so will not be able to reproduce sexually. In *S. cerevisiae* and some other species, this problem is avoided by the process called mating-type switching.

The two *S. cerevisiae* mating types are called **a** and **α**. Each mating type secretes a short polypeptide pheromone (12 amino acids for **a** and 13 for **α**) that binds to receptors on the surfaces of cells of the opposite mating type. Binding of the pheromone initiates a MAP kinase signal transduction pathway that alters the genome expression profile within the cell, leading to subtle morphological and physiological changes that convert the cell into a gamete able to participate in sexual reproduction. Mixing two haploid strains of opposite mating type therefore stimulates formation of gametes that fuse to produce a diploid zygote. Meiosis occurs within the zygote, giving rise to a tetrad of four haploid ascospores, contained in a structure called an ascus. The ascus bursts open, releasing the ascospores, which then divide by mitosis to produce new haploid vegetative cells (Figure 14.15).

The mating type is specified by the MAT gene, located on chromosome III. This gene has two alleles, *MATa* and *MATα*: a haploid yeast cell displays the mating type corresponding to whichever allele it possesses. Elsewhere on chromosome III are two additional MAT-like genes, called *HMRa* and *HMLα*. These have the same sequences as *MATa* and *MATα*, respectively, but neither gene is expressed because upstream of each one is a silencer that represses transcription initiation. These two genes are called silent mating-type cassettes. Their silencing involves the Sir proteins, several of which have histone deacetylase activity (Section 10.2), indicating that silencing involves changes in the chromatin structure in the region of *HMRa* and *HMLα*.

Mating-type switching is initiated by the HO endonuclease, which makes a double-strand cut at a 24 bp sequence located within the MAT gene (Figure 14.16). This enables the gene conversion event to take place. We will examine the details of gene conversion in Section 17.1; all that concerns us at the moment is that one of the free 3’-ends produced by the endonuclease can be extended by DNA synthesis, using one of the two silent cassettes as the template. The newly synthesized DNA subsequently replaces the DNA currently at the MAT locus. The silent cassette chosen as the template is usually the one that is different to the allele originally at

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**Figure 14.14** Gene conversion and DNA rearrangement. In this simple example, gene conversion is illustrated by the orange gene being replaced with a second copy of the blue gene. DNA rearrangement is illustrated by the relative positions of the two genes changing, so that they become linked to one another.

**Figure 14.15** Life cycle of the yeast *Saccharomyces cerevisiae*. 
Genome rearrangements are responsible for immunoglobulin and T-cell receptor diversity

In vertebrates, there are two striking examples of the use of DNA rearrangements to achieve permanent changes in genome activity. These two examples, which are very similar, are responsible for the generation of immunoglobulin and T-cell receptor diversity.

Immunoglobulins and T-cell receptors are related proteins that are synthesized by B and T lymphocytes, respectively. Both types of protein become attached to the outer surfaces of their cells, and immunoglobulins are also released into the bloodstream. The proteins help to protect the body against invasion by bacteria, viruses, and other unwanted substances by binding to these antigens, as they are called. During its lifetime, an organism could be exposed to a vast range of antigens, which means that the immune system must be able to synthesize an equally vast range of immunoglobulin and T-cell receptor proteins. In fact, humans can make approximately $10^8$ different immunoglobulin and T-cell receptor proteins. But there are only 20,441 protein-coding genes in the human genome, so where do all these immunoglobulin and T-cell receptor proteins come from?

To understand the answer, we will look at the structure of a typical immunoglobulin protein. Each immunoglobulin is a tetramer of four polypeptides linked by disulfide bonds (Figure 14.17). There are two longer heavy chains and two shorter light chains. When the sequences of different heavy chains are compared, it becomes clear that the variability between them lies mainly in the N-terminal regions of these polypeptides, while the C-terminal parts are very similar, or constant, in all heavy chains. The same is true for the light chains, except that two families, $\kappa$ and $\lambda$, can be distinguished that differ in the sequences of their constant regions.

In vertebrate genomes, there are no complete genes for the immunoglobulin heavy and light polypeptides. Instead, these proteins are specified by gene segments. The heavy-chain segments are located within a 1 Mb region of chromosome 14 and comprise up to 11 constant-region ($C_H$) gene segments, preceded by 123–129 $V_H$ gene segments, 27 $D_H$ gene segments, and 9 $J_H$ gene segments, these last three types coding for different versions of the $V$ (variable), $D$ (diverse), and $J$ (joining) components of the variable part of the heavy chain (Figure 14.18). A

MAT, so replacement with the newly synthesized strand converts the MAT gene from MATa to MATα, or vice versa. This results in mating-type switching.

The MAT genes code for regulatory proteins (one in the case of MATa and two for MATα) that interact with a transcription activator, MCM1. The MATα and MATa gene products have different effects on MCM1 and so specify different allele-specific genome expression patterns. These expression patterns are maintained in a semipermanent fashion until another MAT gene conversion occurs.

**KEY**

- Variable region
- Heavy chain
- Constant region
- Light chain
- Disulfide bond

**Figure 14.16** Mating-type switching in yeast. In this example, the cell begins as mating type a. The HO endonuclease cuts the MATa locus, initiating gene conversion by the HMLα locus. The result is that the mating type switches to α.

**Figure 14.17** Immunoglobulin structure. Each immunoglobulin protein is made up of two heavy and two light chains, linked by disulfide bonds. Each heavy chain is 446 amino acids in length and consists of a variable region (shown in pink) spanning amino acids 1–108 followed by a constant region. Each light chain is 214 amino acids in length, again with an N-terminal variable region of 108 amino acids. Additional disulfide bonds form between different parts of individual chains; these and other interactions fold the protein into a more complex three-dimensional structure.
similar arrangement is seen with the light-chain loci on chromosomes 2 (κ locus) and 22 (λ locus); the only difference is that the light chains do not have D segments.

During the early stage of B-lymphocyte development, the immunoglobulin loci in its genome undergo rearrangements. Within the heavy-chain locus, these rearrangements first link one of the V<sub>H</sub> gene segments with one of the D<sub>H</sub> gene segments and then link this V-D combination with a J<sub>H</sub> gene segment (Figure 14.19). These rearrangements occur by an unusual type of recombination, catalyzed by a pair of proteins called RAG1 and RAG2, with the positions at which breakage and reunion reactions must occur in order to link the gene segments marked by a series of 8 and 9 bp consensus sequences. The end result is an exon that contains the complete open reading frame specifying the V<sub>H</sub>, D<sub>H</sub>, and J<sub>H</sub> segments of the immunoglobulin protein. This exon becomes linked to a C<sub>H</sub> segment exon by splicing during the transcription process, creating a complete heavy-chain mRNA that can be translated into an immunoglobulin protein that is specific for just that one lymphocyte. A similar series of DNA rearrangements results in the lymphocyte’s light-chain V-J exon being constructed at either the κ or λ locus; once again, splicing attaches a light-chain C segment exon when the mRNA is synthesized.

Despite its name, the constant region is not identical in every immunoglobulin protein. The small variations that occur result in five different classes of immunoglobulin—IgA, IgD, IgE, IgG, and IgM—each with its own specialized role in the immune system. Initially, each B lymphocyte synthesizes an IgM molecule, the C<sub>H</sub> segment of which is specified by the C<sub>μ</sub> sequence that lies at the 5′-end of the C<sub>H</sub> segment cluster. As shown in Figure 14.19, later in its development the immature cell might also synthesize some IgD proteins, utilizing the second C<sub>H</sub> sequence in the cluster (C<sub>δ</sub>), the exon for this sequence becoming attached to the V-D-J segment by alternative splicing. Later in their lifetimes, when they have reached maturity, some B lymphocytes undergo a second type of class switching, which results in a complete change in the type of immunoglobulin that the lymphocyte synthesizes. This second class switching requires a further recombination event that deletes the C<sub>μ</sub> and C<sub>δ</sub> sequences along with the part of the
chromosome between this region and the C_H segment specifying the class of immunoglobulin that the lymphocyte will now synthesize. For example, for the lymphocyte to switch to synthesis of IgG, the most prevalent type of immunoglobulin made by mature lymphocytes, the deletion will place one of the Cγ segments, which specify the IgG heavy chain, at the 5’-end of the cluster (Figure 14.20). Class switching is therefore a second example of genome rearrangement occurring during B-lymphocyte development. The mechanism is distinct from V-D-J joining, and the recombination event does not involve the RAG proteins.

Diversity of T-cell receptors is based on similar rearrangements that link V, D, J, and C gene segments in different combinations to produce cell-specific genes. Each receptor comprises a pair of β molecules, which are similar to the immunoglobulin heavy chain, and two α molecules, which resemble the immunoglobulin κ light chains. The T-cell receptors become embedded in the cell membrane and enable the lymphocyte to recognize and respond to extracellular antigens.

14.3 CHANGES IN GENOME ACTIVITY UNDERLYING DEVELOPMENT

The developmental pathway of a multicellular eukaryote begins with a fertilized egg cell and ends with an adult form of the organism. In between lies a complex series of genetic, cellular, and physiological events that must occur in the correct order, in the correct cells, and at the appropriate times if the pathway is to reach a successful culmination. With humans, this developmental pathway results in an adult containing 10^{13} cells differentiated into approximately 400 specialized types, the activity of each individual cell coordinated with that of every other cell. Developmental processes of such complexity might appear intractable, even to the powerful investigative tools of modern molecular biology, but remarkably good progress toward understanding them has been made in recent years. The research that has underpinned this progress has been designed around three guiding principles:

- It should be possible to describe and comprehend the genetic and biochemical events that underlie differentiation of individual cell types. This in turn means that an understanding of how specialized tissues, and even complex body parts, are constructed should be within reach.

- The signaling processes that coordinate events in different cells should be amenable to study. We saw in Section 14.1 that progress has been made in describing these systems at the molecular level.

- There should be similarities and parallels between developmental processes in different organisms, reflecting common evolutionary origins. This means that information relevant to human development can be obtained from studies of model organisms chosen for the relative simplicity of their developmental pathways.

Developmental biology encompasses areas of genetics, molecular biology, cell biology, physiology, biochemistry, and systems biology. We are concerned only with the role of the genome in development and so will not attempt a wide-ranging overview of developmental research in all its guises. Instead, we will concentrate on four model systems of increasing complexity in order to investigate the types of change in genome activity that occur during development.
**Bacteriophage λ**: a genetic switch enables a choice to be made between alternative developmental pathways

A bacteriophage that infects *E. coli* might seem an odd place to begin a study of the role of the genome in development. But this is exactly where molecular biologists began the lengthy program of research that today is revealing the underlying genomic basis to development in humans and other vertebrates. We will therefore follow this same progression from the relatively simple to the relatively complex.

In Section 9.1 we learned that lysogenic bacteriophages such as λ can follow two alternative replication pathways after infection of a host cell. As well as the lytic pathway, during which new phages are assembled and released from the cell soon after the initial infection (after 45 minutes for λ), these phages can also pursue a lysogenic cycle characterized by insertion of the phage DNA into the host chromosome. The integrated prophage remains quiescent for many bacterial generations until a chemical or physical stimulus linked to DNA damage induces excision of the λ genome, rapid assembly of phages, and lysis of the host cell (see Figures 9.3 and 9.4). How does the phage decide whether to follow the lytic or lysogenic cycle? To answer this question, we first need to understand the series of genetic events that underlies lysogeny.

The λ genome contains two promoters, *P*₁ and *P*₂, which are recognized by the *E. coli* RNA polymerase as soon as λ injects its DNA into the cell. The polymerase therefore transcribes the two immediate-early λ genes, called *N* and *cro* (Figure 14.21). The product of gene *N* is an antiterminator protein, which attaches to the DNA and causes the host RNA polymerase to ignore the termination signals that it encounters immediately downstream of the *N* and *cro* coding sequences. The polymerase therefore transcribes the delayed-early genes. These genes include *clII* and *clIII*, which together activate transcription of *cl*. This is an important gene as it codes for the λ repressor protein, the key master switch that shuts down the lytic cycle and maintains lysogeny. The repressor does this by binding to the operators *O₁* and *O₂*, which are adjacent to *P*₁ and *P*₂, respectively (Figure 14.22). As a result almost the entire λ genome is silenced, because *P*₁ and *P*₂ direct transcription not only of the immediate-early and delayed-early genes but also of the late genes, which code for the proteins needed for assembly of new phages and host cell lysis. One of the few genes to remain active is *int*, which is transcribed from its own promoter. The integrase protein coded by this gene catalyzes the site-specific recombination by which λ DNA is inserted into the host genome. Lysogeny is maintained for numerous cell divisions because the cl gene is continuously expressed, albeit at a low level, so that the amount of cl repressor present in the cell is always enough to keep *P*₁ and *P*₂ switched off. This continued expression of cl occurs because the cl repressor, when bound to *O₂*, not only blocks transcription from *P*₂ but also stimulates transcription from its own promoter, *P*₃RM. The dual role of the cl repressor is therefore the key to lysogeny.

![Diagram](image-url)

*Figure 14.21 Transcription of the immediate-early and delayed-early λ genes.* (A) Transcription from promoters *P*₁ and *P*₂ initially results in the synthesis of two immediate-early mRNAs, terminating at positions *t*₁ and *t*₂. (B) The mRNA transcribed from *P*₂ to *t*₁ codes for the N protein, which attaches at the antitermination sites *nutL* and *nutR*. Now the RNA polymerase continues transcription downstream of *t*₁ and *t*₂. Transcription from *P*₃ also ignores terminator *t*₃ and continues until *t*₄ is reached.
Once \( cI \) is expressed, the repressor protein prevents entry into the lytic cycle and ensures that lysogeny is set up and maintained. But \( \lambda \) does not always enter the lysogenic cycle: on some occasions, an infection proceeds immediately to host lysis. This is because of the activity of the second immediate-early gene, \( cro \), which also codes for a repressor, but in this case one that prevents transcription of \( cI \) (Figure 14.23). The decision between lysis and lysogeny is therefore determined by the outcome of a race between \( cI \) and \( cro \). If the \( cI \) repressor is synthesized more quickly than the Cro repressor, then genome expression is blocked and lysogeny follows. However, if \( cro \) wins the race, then the Cro repressor blocks \( cI \) expression before enough \( cI \) repressor has been synthesized to silence the genome. As a result, the phage enters the lytic infection cycle. The decision appears to be random, depending on chance events that lead to either \( cI \) or Cro repressor accumulating more quickly in the cell, although environmental conditions can have an influence. Growth on a rich medium, for example, shifts the balance toward the lytic cycle, presumably because it is beneficial to produce new phages when the host cells are proliferating. This shift is brought about by activation of proteases that degrade the \( cII \) protein, reducing the ability of the \( cII-cIII \) combination to switch on transcription of the \( cI \) gene.

If the bacteriophage enters the lysogenic cycle, then this state is maintained as long as the \( cI \) repressor is bound to the operators \( O_L \) and \( O_R \). The prophage will therefore be induced if the level of active \( cI \) repressor declines below a certain point. This may happen by chance, leading to spontaneous induction, or may occur in response to physical or chemical stimuli. These stimuli activate a general protective mechanism in \( E. \coli \), the \textbf{SOS response}. Part of the SOS response is expression of an \( E. \coli \) gene, \textit{recA}, whose product inactivates the \( cI \) repressor by cleaving it in half. This switches on expression of the early genes, enabling the phage to enter the lytic cycle. Inactivation of the \( cI \) repressor also means that transcription of \( cI \) is no longer stimulated, avoiding the possibility of lysogeny being reestablished through the synthesis of more \( cI \) repressor. Inactivation of the \( cI \) repressor therefore leads to induction of the prophage.

What do we learn from this model system?

- A simple genetic switch can determine which of two developmental pathways is followed by a cell.
- Genetic switches can involve a combination of activation and repression of different promoters.
- It is possible to reprogram a developmental pathway, and transfer to an alternative pathway, in response to appropriate stimuli.

\textbf{Bacillus sporulation: coordination of activities in two distinct cell types}

The second system that we will examine is formation of spores by the bacterium \textit{Bacillus subtilis}. As with \( \lambda \) lysogeny, this is not, strictly speaking, a developmental pathway, merely a type of cellular differentiation, but the process illustrates two
of the fundamental issues that have to be addressed when genuine development in multicellular organisms is studied. These issues are how a series of changes in genome activity over time is controlled and how signaling establishes coordination between events occurring in different cells. The advantages of Bacillus as a model system are that it is easy to grow in the laboratory and is amenable to study by genetic and molecular biological techniques such as analysis of mutants and sequencing of genes.

Bacillus is one of several genera of bacteria that produce endospores in response to unfavorable environmental conditions. These spores are highly resistant to physical and chemical abuse and can survive for decades or even centuries. Resistance is due to the specialized nature of the spore coat, which is impermeable to many chemicals, and to biochemical changes within the spore that retard the decay of DNA and other polymers and enable the spore to survive a prolonged period of dormancy.

In the laboratory, sporulation is usually induced by nutrient starvation. This causes the bacteria to abandon their normal vegetative mode of cell division, which involves synthesis of a septum (or cross-wall) in the center of the cell. Instead the cells construct an unusual septum, one that is thinner than normal, at one end of the cell (Figure 14.24). This produces two cellular compartments, the smaller of which is called the prespore and the larger the mother cell. As sporulation proceeds, the prespore becomes entirely engulfed by the mother cell. By now the two cells are committed to different but coordinated differentiation pathways, the prespore undergoing the biochemical changes that enable it to become

Figure 14.24 Sporulation in Bacillus subtilis. The top part of the diagram shows the normal vegetative mode of cell division, involving formation of a septum across the center of the bacterium and resulting in two identical daughter cells. The lower part of the diagram shows sporulation, in which the septum forms near one end of the cell, leading to a mother cell and a prespore of different sizes. Eventually the mother cell completely engulfs the prespore. At the end of the process, the mature resistant spore is released.
dormant, and the mother cell constructing the resistant coat around the spore and eventually dying.

Changes in genome activity during sporulation are controlled largely by the synthesis of special σ subunits that change the promoter specificity of the Bacillus RNA polymerase (Section 12.2). We have seen how this simple control system is used by E. coli in response to heat stress (see Figure 12.10). It is also the key to changes in genome activity that occur during sporulation. The standard B. subtilis subunits are called σ^A and σ^H. These subunits are synthesized in vegetative cells and enable the RNA polymerase to recognize promoters for all the genes it needs to transcribe in order to maintain normal growth and cell division. In the prespore and mother cell these subunits are replaced by σ^E and σ^F, respectively, which recognize different promoter sequences and so give rise to large-scale changes in genome expression patterns. The master switch from vegetative growth to spore formation is provided by a protein called SpoOA, which is present in vegetative cells but in an inactive form. This protein is activated by phosphorylation, via a cascade of protein kinases that respond to various extracellular signals that indicate the presence of an environmental stress such as lack of nutrients. The initial response is provided by two kinases, called KinA and KinB, which phosphorylate themselves and then pass the phosphate via SpoOF and SpoOB to SpoOA (Figure 14.25). Activated SpoOA is a transcription factor that modulates the expression of various genes transcribed by the vegetative RNA polymerase and hence recognized by the regular σ^A and σ^H subunits. The genes that are switched on include those for σ^E and σ^F, resulting in the switch to prespore and mother cell differentiation (Figure 14.26).

Initially, both σ^A and σ^E are present in each of the two differentiating cells. This is not exactly what is wanted because σ^F is the prespore-specific subunit and so should be active only in this cell, and σ^E is mother-cell-specific. A means is therefore needed to activate or inactivate the appropriate subunit in the correct cell. This is thought to be achieved as follows (Figure 14.27):

- σ^F is activated by release from a complex with a second protein, SpoIIB. This is controlled by a third protein, SpoIIA, which, when unphosphorylated, can also attach to SpoIIB and prevent the latter from binding to σ^F. If SpoIIA is unphosphorylated, then σ^F is released and is active; if SpoIIA is phosphorylated, then σ^F remains bound to SpoIIB and so is inactive. In the mother cell, SpoIIB phosphorylates SpoIIA and so keeps σ^F in its bound, inactive state. However, in the prespore, SpoIIB’s attempts to phosphorylate SpoIIA are antagonized by yet another protein, SpoIIE, and so σ^F is released and becomes active. SpoIIE’s ability to antagonize SpoIIB in the prespore but not the mother cell derives from the fact that SpoIIE molecules are bound to the membrane on the surface of the septum. Because the prespore is much smaller than the mother cell but the septum surface area is similar in both, the concentration of SpoIIE is greater in the prespore, and this enables it to antagonize SpoIIB.

- σ^E is activated by proteolytic cleavage of a precursor protein. The protease that carries out this cleavage is the SpoIIGA protein, which spans the septum between the prespore and mother cell, with the protease domain on the mother-cell side. Activation of SpoIIGA requires the presence of σ^F in the prespore, but the precise nature of this link is not clear. One possibility is that SpoIIGA is activated by binding of SpoIIA to a receptor domain on the prespore side. The gene for SpoIIR is one of those whose promoter is recognized specifically by σ^F, so activation of the protease and conversion of pre-σ^E to active σ^E occurs once σ^F-directed transcription is underway in the prespore. If this model is correct, then it would constitute a typical receptor-mediated signal transduction system (Section 14.1).

Activation of σ^F and σ^E is just the beginning of the story. In the prespore, about 1 hour after its activation, σ^F responds to an unknown signal (possibly from the mother cell), which results in a slight change in genome activity in the spore. This
This protease then activates a proteolytic cascade with SpoIIAB, which is indirectly influenced by the concentration of membrane-bound SpoIIE. In the mother cell, σF is activated by proteolytic cleavage by SpoIIAG, possibly in response to the presence in the prespore of the σE-dependent protein SpoIIA. Abbreviations: AA, SpoIIA; AB, SpoIIAB; E, σE; F, σF; GA, SpoIIAG; R, SpoIIIR.

To summarize, the key features of Bacillus sporulation are as follows:

- The master protein, SpoOA, responds to external stimuli via a cascade of phosphorylation events to determine if and when the switch to sporulation should occur.
- A succession of σ subunits in prespore and mother cell brings about time-dependent changes in genome activity in the two cells.
- Cell–cell signaling ensures that the events occurring in prespore and mother cell are coordinated.

**Caenorhabditis elegans:** the genetic basis of positional information and the determination of cell fate

Research with the microscopic nematode worm *C. elegans* (Figure 14.29) was initiated by Sydney Brenner in the 1960s with the aim of utilizing it as a simple model for multicellular eukaryotic development. *C. elegans* is easy to grow in the laboratory and has a short generation time, taking just 3.5 days for the fertilized egg to develop into a mature adult. The worm is transparent at all stages of its life cycle, so internal examination is possible without killing the animal. This is an important point because it has enabled researchers to follow the entire development of the worm at the cellular level. Every cell division in the pathway from fertilized egg to adult worm has been charted, and every point at which a cell adopts a specialized role has been identified. This pathway is more or less invariant: the pattern of cell division and differentiation is virtually the same in all cultures.
14.3 Changes in Genome Activity Underlying Development

Chapter 14: Genome Expression in the Context of Cell and Organism Development

This appears to be due in large part to cell–cell signaling, which induces each cell to follow its appropriate differentiation pathway. To illustrate this we will look at development of the *C. elegans* vulva.

Most *C. elegans* worms are hermaphrodites, meaning that they have both male and female sex organs. The vulva is part of the female sex apparatus, being the tube through which sperm enter and fertilized eggs are laid. The adult vulva comprises 22 cells, which are the progeny of three ancestral cells originally located in a row on the undersurface of the developing worm (*Figure 14.30*). Each of these ancestral cells becomes committed to the differentiation pathway that leads to production of vulva cells. The central cell, called P6.p, adopts the primary vulva cell fate and divides to produce eight new cells. The other two cells, P5.p and P7.p, take on the secondary vulva cell fate and divide into seven cells each. These 22 cells then reorganize their positions to construct the vulva.

A critical aspect of vulva development is that it must occur in the correct position relative to the gonad, the structure containing the egg cells. If the vulva develops in the wrong place, then the gonad will not receive sperm and the egg cells will never be fertilized. The positional information needed by the vulva progenitor cells is provided by a cell within the gonad called the anchor cell (*Figure 14.31*). The importance of the anchor cell has been demonstrated by experiments in which it is artificially destroyed in the embryonic worm: in the absence of the anchor cell, a vulva does not develop. The implication is that the anchor cell secretes an extracellular signaling compound that induces P6.p, the cell closest to the anchor cell, to the primary vulva cell fate. P5.p and P7.p (shown in light orange) are further away from the anchor cell and so are exposed to a lower concentration of LIN-3 and become secondary vulva cells. As described in the text, there is evidence that commitment of the secondary cells to their fates is also influenced by signals from the primary vulva cell.

*Figure 14.30* Cell divisions resulting in production of the vulva cells of *C. elegans*. Three ancestral cells divide in a programmed manner to produce 22 progeny cells, which reorganize their positions relative to one another to construct the vulva.

*Figure 14.31* Postulated role of the anchor cell in determining cell fate during vulva development in *C. elegans*. It is thought that release of the signaling compound LIN-3 by the anchor cell commits P6.p (shown in pink), the cell closest to the anchor cell, to the primary vulva cell fate. P5.p and P7.p (shown in light orange) are further away from the anchor cell and so are exposed to a lower concentration of LIN-3 and become secondary vulva cells. As described in the text, there is evidence that commitment of the secondary cells to their fates is also influenced by signals from the primary vulva cell.
of a MAP kinase-like protein, which in turn switches on a variety of transcription factors. The identity of this MAP kinase, and hence of the transcription factors that are switched on, depends on the number of LET-23 receptors that are activated, which in turn depends on the extracellular LIN-3 concentration, explaining how the primary and secondary fates can be specified by the distance of the recipient cells from the anchor cell.

In the second signaling system, P6.p, having adopted the primary cell fate, synthesizes DSL proteins, some of which become embedded in the cell membrane while some are secreted. Both the embedded and secreted versions are able to interact with LIN-12 receptor proteins on P5.p and P7.p, inducing a second intracellular signal transduction pathway that further contributes to adoption of the secondary cell fate. The importance of this pathway is supported by the abnormal features displayed by certain mutants in which more than three cells become committed to vulva development. With these mutants there is more than one primary cell, but each one is invariably surrounded by two secondary cells, suggesting that in the living worm, adoption of the secondary cell fate is dependent on the presence of an adjacent primary cell.

In summary, the general concepts to emerge from the study of vulva development in C. elegans are as follows:

- In a multicellular organism, positional information is important: the correct structure must develop at the appropriate place.
- The commitment to differentiation of a small number of progenitor cells can lead to construction of a multicellular structure.
- Cell–cell signaling can utilize a concentration gradient to induce different responses in cells at different positions relative to the signaling cell.

**Fruit flies: conversion of positional information into a segmented body plan**

The last organism whose development we will study is D. melanogaster. The experimental history of the fruit fly dates back to 1910 when Thomas Hunt Morgan first used this organism as a model system in genetic research. For Morgan, the advantages of Drosophila were its small size (enabling large numbers to be studied in a single experiment), its minimal nutritional requirements (the flies like bananas), and the presence in natural populations of occasional variants with easily recognized genetic characteristics such as unusual eye colors. The body plan of the adult fly, as well as that of the larva, is built from a series of segments, each with a different structural role. This is clearest in the thorax, which has three segments, each carrying one pair of legs, and the abdomen, which is made up of eight segments, but it is also true for the head, even though in the head the segmented structure is less visible (Figure 14.32). The early embryo, on the other hand, is a single syncytium comprising a mass of cytoplasm and multiple nuclei (Figure 14.33). The major contribution that Drosophila has made to our understanding of development has been through the insights it has provided into how this undifferentiated embryo acquires positional information that eventually results in the construction of complex body parts at the correct places in the adult organism.

Initially, the positional information that the embryo needs is a definition of which end is the front (anterior) and which is the back (posterior), as well as similar information relating to up (dorsal) and down (ventral). This information is provided by concentration gradients of proteins that become established in the syncytium. The majority of these proteins are not synthesized from genes in the embryo but are specified by maternal-effect genes whose mRNAs are injected into the embryo by the mother. The bicoid gene, for example, is transcribed in the maternal nurse cells, which are in contact with the egg cells, and the mRNA is injected into the anterior end of the unfertilized egg. This position is defined by
the orientation of the egg cell in the egg chamber. The bicoid mRNA remains in the anterior region of the egg cell, attached by its 3'-untranslated region to the cell's cytoskeleton, and the Bicoid protein diffuses through the syncytium, setting up a concentration gradient from highest at the anterior end to lowest at the posterior end (Figure 14.34). Additional maternal-effect proteins, such as Hunchback, Nanos, Caudal, and Torso, contribute in a similar way to the anterior–posterior axis, while Dorsal and others set the dorsal–ventral axis. As a result, each point in the syncytium acquires its own unique chemical signature defined by the relative amounts of the various maternal-effect proteins.

This basic positional information is made more precise by expression of the gap genes. Three of the anterior–posterior gradient proteins—Bicoid, Hunchback, and Caudal—are transcription factors that target the gap genes in the nuclei that now line the inside of the embryo (see Figure 14.33). The identities of the gap genes expressed in a particular nucleus depend on the relative concentrations of the gradient proteins and hence on the position of the nucleus along the anterior–posterior axis. Some gap genes are activated directly by Bicoid, Hunchback, and Caudal; examples are buttonhead, empty spiracles, and orthodenticle, which are activated by Bicoid. Other gap genes are switched on indirectly, as is the case with huckebein and tailless, which respond to transcription factors that are activated by Torso. This complex interplay results in the positional information in the embryo, now carried by the relative concentrations of the gap gene products, becoming more detailed (Figure 14.35).

The next set of genes to be activated, the pair-rule genes, establish the basic segmentation pattern. Transcription of these genes responds to the relative concentrations of the gap gene products and occurs in nuclei that have become enclosed in cells. The pair-rule gene products therefore do not diffuse through the syncytium but remain localized within the cells that express them. The result is that the embryo can now be looked upon as comprising a series of stripes, each stripe consisting of a set of cells expressing a particular pair-rule gene. In a further round of gene activation, the segment polarity genes become switched on, providing greater definition to the stripes by setting the sizes and precise locations...
Figure 14.35 Role of gap gene products in conferring positional information during embryo development in *D. melanogaster*. The concentration gradient of each gap gene product is denoted by the colored bars. The parts of the embryo that give rise to the head, thorax, and abdomen regions of the adult fly are indicated.

Figure 14.36 Antennapedia and Bithorax gene complexes of *D. melanogaster*. Both complexes are located on fruit fly chromosome 3, with ANT-C upstream of BX-C. The genes are usually drawn in the order shown, although this means that they are transcribed from right to left. The diagram does not reflect the actual lengths of the genes. The full gene names are as follows: *lab*, *labial palps*; *pb*, *proboscipedia*; *Dfd*, *Deformed*; *Scr*, *Sex combs reduced*; *Antp*, *Antennapedia*; *Ubx*, *Ultrabithorax*; *abdA*, *abdominal A*; *AbdB*, *Abdominal B*. In ANT-C, the non-selector genes *zerknüllt* and *bicoid* occur between *pb* and *Dfd*, and *fushi tarazu* lies between *Scr* and *Antp*.

The correct selector gene is expressed in each segment because the activation of each one is responsive to the positional information represented by the distributions of gap gene and pair-rule gene products. The selector gene products are themselves transcription factors, each containing a homeodomain version of the helix-turn-helix DNA-binding structure (Section 11.2). Each selector gene product switches on the set of genes needed to initiate development of the specified segment. Maintenance of the differentiated state is ensured partly by the repressive effect of each selector gene product on expression of the other selector genes and partly by the work of the Polycomb group proteins, which, as we saw in Section 14.2, construct inactive chromatin over the selector genes that are not expressed in a particular cell.

The homeodomains of the various *Drosophila* selector genes are strikingly similar. This observation led researchers in the 1980s to search for other homeotic
genes by using the homeodomain as a probe in hybridization experiments. First, the *Drosophila* genome was searched, resulting in isolation of several previously unknown homeodomain-containing genes. These have not turned out to be selector genes but other genes coding for transcription factors involved in development. Examples include the pair-rule genes *even-skipped* and *fushi tarazu* and the segment polarity gene *engrailed*. Researchers then probed the genomes of other organisms and discovered that homeodomains are present in genes in a wide variety of animals, including humans. Examination of these genes showed that some are homeotic selectors, organized into clusters similar to ANT-C and BX-C, and that these genes have equivalent functions to the *Drosophila* versions, specifying construction of the body plan. For example, mutations in the HoxC8 gene of mouse results in an animal that has an extra pair of ribs, due to conversion of a lumbar vertebra (normally in the lower back) into a thoracic vertebra (from which the ribs emerge). Other Hox mutations in animals lead to limb deformations, such as absence of the lower arm or extra digits on the hands or feet.

We now look on the ANT-C and BX-C clusters of selector genes in *Drosophila* as two parts of a single complex, usually referred to as the homeotic gene complex or HOM-C. In vertebrates there are four homeotic gene clusters, called HoxA–HoxD. When these four clusters are aligned with one another and with HOM-C (Figure 14.37), similarities are seen between genes at equivalent positions, such that the evolutionary history of the homeotic selector gene clusters can be traced from insects through to humans (see Section 18.2). As in *Drosophila*, the order of genes in the vertebrate clusters reflects the order of the structures specified by the genes in the adult body plan. This is clearly seen with the mouse HoxB cluster, which controls development of the nervous system (Figure 14.38). The remarkable conclusion is that, at this fundamental level, developmental processes in fruit flies and other simple eukaryotes are similar to the processes occurring in humans and other complex organisms. The discovery that studies of fruit flies are directly relevant to human development has opened up vast vistas of research possibilities.
Homeotic genes also underlie plant development

The power of *Drosophila* as a model system for development extends even beyond vertebrates. Developmental processes in plants are, in most respects, very different from those of fruit flies and other animals, but at the genetic level there are certain similarities, sufficient for the knowledge gained about *Drosophila* development to be of value in interpreting similar research carried out with plants. In particular, the recognition that a limited number of homeotic selector genes control the *Drosophila* body plan has led to a model for plant development which postulates that the structure of the flower is determined by a small number of homeotic genes.

All flowers are constructed along similar lines, made up of four concentric whorls, each comprising a different floral organ (Figure 14.39). The outer whorl, number 1, contains sepals, which are modified leaves that envelop and protect the bud during its early development. The next whorl, number 2, contains the distinctive petals, and within these are whorls 3 (stamens, the male reproductive organs) and 4 (carpels, the female reproductive organs).

Most of the research on plant development has been carried out with *Antirrhinum* (the snapdragon) and *Arabidopsis thaliana*, a small vetch that has been adopted as a model species, partly because it has a genome of only 135 Mb (see Table 7.2), one of the smallest known among flowering plants. Although these plants do not appear to contain homeodomain proteins, they do have genes that, when mutated, lead to homeotic changes in the floral architecture, such as replacement of sepals by carpels. Analysis of these mutants has led to the ABC model, which states that there are three types of homeotic genes—A, B, and C—which control flower development as follows:

- Whorl 1 is specified by A-type genes: examples in *Arabidopsis* are apetala1 and apetala2.
- Whorl 2 is specified by A genes acting in concert with B genes; examples of B genes include apetala3 and pistillata.
- Whorl 3 is specified by the B genes plus the C gene, agamous.
- Whorl 4 is specified by the C gene acting on its own.

As anticipated from the work with *Drosophila*, the A, B, and C homeotic gene products are transcription factors. All except the APETALA2 protein contain the same DNA-binding domain, the MADS box, which is also found in other proteins involved in plant development, including SEPALLATA1, -2, and -3, which work with the A, B, and C proteins in defining the detailed structure of the flower. Other components of the flower development system include at least one master gene, called floricaula in *Antirrhinum* and leafy in *Arabidopsis*, which controls the switch from vegetative to reproductive growth, initiating flower development, and also has a role in establishing the pattern of homeotic gene expression. Plants also have Polycomb group proteins (Section 14.2), with CURLY LEAF and SWINGER of *Arabidopsis* thought to be the key histone methyltransferases responsible for silencing regions of chromatin containing those homeotic genes that are inactive in a particular whorl.

**SUMMARY**

- Transient alterations in genome expression patterns enable a cell to respond to changes in the external environment, these changes including the presence or absence of signaling compounds that coordinate biochemical activities in different cells.
- More permanent changes in genome expression underlie differentiation and development.
Transient changes in genome expression patterns occur predominantly in response to external stimuli that influence the transcription of individual genes.

Some extracellular signaling compounds are imported into the cell and directly influence transcription, an example being lactoferrin in mammals.

Steroid hormones also enter the cell but influence genome expression via receptor proteins that act as transcription activators.

Effects of other signaling compounds are mediated by cell surface receptors, many of which dimerize in response to the extracellular signal, initiating a signal transduction pathway that leads to the genome.

With some signal transduction systems, stimulation of the cell surface receptor by attachment of the extracellular signaling compound has a direct effect on the activity of a transcription factor. An example is the JAK/STAT pathway.

Other signal transduction pathways, including the MAP kinase pathway, have several steps between receptor and genome. Some of these pathways make use of second messengers such as cyclic nucleotides and calcium ions, which influence a number of cellular activities including genome expression.

Differentiation processes involve semipermanent changes in genome expression, which can be brought about by changes to chromatin structure, gene conversion events, or genome rearrangements.

The lysogenic infection cycle of bacteriophage λ demonstrates ways in which simple genetic switches can determine which of two developmental pathways are followed.

Studies of sporulation in Bacillus subtilis have illustrated how time-dependent changes in genome expression can be brought about and how cell–cell signaling can regulate a developmental pathway.

Mechanisms for the determination of cell fate have been revealed by studies of vulva development in C. elegans.

The most informative pathway for developmental genetics has been embryogenesis in the fruit fly, which has shown how a complex body plan can be specified by controlled patterns of genome expression.

Work with Drosophila has also revealed the existence of homeotic selector genes, which control developmental processes not only in flies but also in vertebrates and in plants.

### SHORT ANSWER QUESTIONS

1. Outline the differences between differentiation and development, and describe the basis for these differences.

2. Describe the evidence that suggests that lactoferrin can act as a transcription factor.

3. How do steroid hormones influence genome expression?

4. Compare the various types of cell surface receptor proteins that are known.

5. Describe the (A) JAK/STAT and (B) MAP kinase signal transduction pathways.

6. Outline the roles of second messengers in the control of genome expression.

7. Describe how the Polycomb group proteins influence genome expression.

8. How can cells of the vertebrate immune system produce so many different immunoglobulins from a small set of genes?
9. Outline the process by which the choice between the lytic and lysogenic pathways is regulated in bacteriophage λ.

10. During sporulation in Bacillus, $\sigma^F$ and $\sigma^E$ are present in both the prespore and the mother cell. How is $\sigma^F$ activated in the prespore?

11. How does the anchor cell of C. elegans induce the vulva progenitor cells to differentiate into vulva cells? Why do the vulva progenitor cells follow different pathways upon receiving the signal from the anchor cell?

12. Describe how studies of embryogenesis in Drosophila led to discovery of homeotic selector genes in vertebrates.

**IN-DEPTH PROBLEMS**

1. What methods might be used to identify those parts of a cell surface receptor protein that are exposed on the outer surface of the plasma membrane of an animal cell?

2. Describe how studies of signal transduction have improved our understanding of the abnormal biochemical activities that underlie cancer.

3. Are Caenorhabditis elegans and Drosophila melanogaster good model organisms for development in higher eukaryotes?

4. What would be the key features of an ideal model organism for development in higher eukaryotes?

5. What can be inferred about genome evolution from the discovery that Drosophila has a single homeotic gene complex whereas vertebrates have four? Would you expect any group of organisms to have more than four homeotic gene clusters?

**FURTHER READING**

**Imported extracellular signaling compounds**


**Cell surface receptor proteins and signal transduction pathways**


**Changes in genome activity during differentiation**


**Simple development pathways**


**Development in Caenorhabditis elegans**


**Embryogenesis in fruit flies and homeotic selector genes in vertebrates**


**Flower development in plants**


The primary function of a genome is to specify the biochemical signature of the cell in which it resides. We have seen that the genome achieves this objective by synthesis and maintenance of a transcriptome and proteome whose individual RNA and protein components carry out and regulate the cell’s biochemical activities. In order to continue performing this function, the genome must replicate every time that the cell divides. This means that the entire DNA content of the cell must be copied at the appropriate period in the cell cycle, and the resulting DNA molecules must be distributed to the daughter cells so that each one receives a complete copy of the genome.

When studying genome replication, it is easy to become absorbed in the molecular details and to lose sight of the broader implications of the process. For example, at the molecular level, we look on the accuracy of genome replication as vital in ensuring that daughter cells acquire precise copies of the genome, so that those cells can function in the same way as their parent or can adopt new functions in accordance with the genetic programming contained in the nucleotide sequence of the genome. However, at a higher level, absolute and inviolate identity between parent and daughter genomes would make evolution impossible, because the latter depends on the generation of genome variants that give rise to organisms with modified characteristics and different degrees of fitness to their environment. This variation is responsible not only for the differences between species but also for the differences between members of a single species.

In Part IV of Genomes we will explore the link between DNA replication and genome evolution. We will begin in this chapter by examining the elaborate process by which the genome is replicated, a process that spans the interface between molecular biology, biochemistry, and cell biology. In Chapters 16 and 17 we will see how variation in genome sequences is introduced by mutation and recombination, and in Chapter 18 we will study the ways in which these processes are thought to have shaped the structures and genetic contents of genomes over evolutionary time.

15.1 THE TOPOLOGY OF GENOME REPLICATION

Genome replication has been studied since Watson and Crick first discovered the double-helical structure of DNA back in 1953. The primary concern in the years
from 1953 to 1958 was the **topological problem**. This problem arises from the need to unwind the double helix in order to make copies of its two polynucleotides. The issue assumed center stage in the mid-1950s because it was the main stumbling block to acceptance of the double helix as the correct structure for DNA. Before we study the molecular events occurring during genome replication, we must first understand how the cell solves this topological problem.

### The double-helical structure complicates the replication process

In their paper in *Nature* announcing the discovery of the double-helical structure of DNA, Watson and Crick made one of the most famous statements in molecular biology:

> “It has not escaped our notice that the specific pairing we have postulated immediately suggests a possible copying mechanism for the genetic material.”

The pairing process to which they refer is one where each strand of the double helix acts as a template for synthesis of a second complementary strand, the end result being that both daughter double helices are identical to the parent molecule (Figure 15.1). The scheme is almost implicit in the double-helical structure, but it presents problems, as admitted by Watson and Crick in a second paper published in *Nature* just a month after the report of the structure. This paper describes the postulated replication process in more detail but points out the difficulties that arise from the need to unwind the double helix. The most trivial of these difficulties is the possibility of the daughter molecules getting tangled up. More critical is the rotation that would accompany the unwinding; with one turn occurring for every 10 bp of the double helix, complete replication of the DNA molecule in human chromosome 1, which is 250 Mb in length, would require 25 million rotations of the chromosomal DNA. It is difficult to imagine how this could occur within the constrained volume of the nucleus, but the unwinding of a linear chromosomal DNA molecule is not physically impossible. In contrast, a circular double-stranded molecule, such as a bacterial or bacteriophage genome, having no free ends, would not be able to rotate in the required manner and so, apparently, could not be replicated by the Watson–Crick scheme. Finding an answer to this dilemma was a major preoccupation of molecular biology during the 1950s.

The topological problem was considered so serious by some molecular biologists, notably Max Delbrück, that there was initially some resistance to accepting the double helix as the correct structure of DNA. The difficulty relates to the plectonemic nature of the double helix, which is the topological arrangement that prevents the two strands of a coil from being separated without unwinding. The problem would therefore be resolved if the double helix were in fact paranemic, because this would mean that the two strands could be separated simply by moving each one sideways without unwinding the molecule. It was suggested that the double helix could be converted into a paranemic structure by supercoiling in the direction opposite to the turn of the helix itself, or that within a DNA molecule the right-handed helix proposed by Watson and Crick might be balanced by equal lengths of a left-handed helical structure. The possibility that double-stranded DNA was not a helix at all, but instead a side-by-side ribbon structure, was also briefly considered, and this idea surprisingly was revived in the late 1970s. Each of these proposed solutions to the topological problem was individually rejected for one reason or another, most of them because they required alterations to the double-helical structure that were not compatible with the X-ray diffraction results and other experimental data pertaining to DNA structure.

The first real progress toward a solution to the topological problem came in 1954, when Delbrück proposed a breakage-and-reunion model for separating the strands of the double helix. In this model, the strands are separated not by unwinding the helix with accompanying rotation of the molecule but by breaking one of the strands, passing the second strand through the gap, and rejoining the first strand. This scheme is very close to the correct solution to the topological problem, as it is one of the ways in which DNA topoisomerases work...
(see Figure 15.4A), but unfortunately Delbrück overcomplicated the issue by attempting to combine breakage and reunion with the DNA synthesis that occurs during the actual replication process. This led him to propose a model for DNA replication that results in each polynucleotide in the daughter molecule being made up partly of parental DNA and partly of newly synthesized DNA. This dispersive mode of replication contrasts with the semiconservative system proposed by Watson and Crick (Figure 15.2). A third possibility is that replication is fully conservative, where one of the daughter double helices is made entirely of newly synthesized DNA and the other comprises the two parental strands. Models for conservative replication are difficult to devise, but one can imagine that this type of replication might be accomplished without unwinding the parent helix.

The Meselson–Stahl experiment proved that replication is semiconservative

Delbrück’s breakage-and-reunion model was important because it stimulated experiments designed to test between the three modes of DNA replication illustrated in Figure 15.2. Radioactive isotopes had recently been introduced into molecular biology, so attempts were made to use DNA labeling to distinguish newly synthesized DNA from the parental polynucleotides. Each mode of replication predicts a different distribution of newly synthesized DNA, and hence of radioactive label, in the double helices resulting after two or more rounds of replication. Analysis of the radioactive contents of these molecules should therefore determine which replication scheme operates in living cells. Unfortunately, it proved impossible to obtain a clear-cut result, largely because of the difficulty in measuring the precise amount of radioactivity in the DNA molecules. The analysis was also complicated by the rapid decay of the $^{32}$P isotope that was used as the label.

The breakthrough was eventually made by Matthew Meselson and Franklin Stahl who, in 1958, carried out the required experiment not with a radioactive label but with $^{15}$N, the nonradioactive heavy isotope of nitrogen. Now it was possible to analyze the replicated double helices by density gradient centrifugation, because a DNA molecule labeled with $^{15}$N has a higher buoyant density than an unlabeled molecule. Meselson and Stahl started with a culture of Escherichia coli cells that had been grown with $^{15}$NH$_4$Cl and whose DNA molecules therefore contained heavy nitrogen. The cells were transferred to normal medium, and samples
taken after 20 and 40 min, corresponding to one and two cell divisions, respectively. DNA was extracted from each sample and the molecules examined by density gradient centrifugation (Figure 15.3A). After one round of DNA replication, the daughter molecules synthesized in the presence of normal nitrogen formed a single band in the density gradient, indicating that each double helix was made up of equal amounts of newly synthesized and parental DNA. This result immediately enabled the conservative mode of replication to be discounted, as it predicts that there would be two bands after one round of replication (Figure 15.3B), but it

Figure 15.3 The Meselson–Stahl experiment. (A) The experiment carried out by Meselson and Stahl involved growing a culture of *Escherichia coli* in a medium containing $^{15}$NH$_4$Cl (ammonium chloride labeled with the heavy isotope of nitrogen). Cells were then transferred to normal medium (containing $^{14}$NH$_4$Cl), and samples taken after 20 min (one cell division) and 40 min (two cell divisions). DNA was extracted from each sample and the molecules analyzed by density gradient centrifugation. After 20 min, all the DNA contained similar amounts of $^{14}$N and $^{15}$N, but after 40 min, two bands were seen, one corresponding to hybrid $^{14}$N/$^{15}$N-DNA and the other to DNA molecules made entirely from $^{14}$N. (B) The predicted outcome of the experiment for each of the three possible modes of DNA replication. The banding pattern seen after 20 min enables conservative replication to be discounted because this scheme predicts that after one round of replication there will be two different types of double helix, one containing just $^{15}$N and the other containing just $^{14}$N. The single $^{14}$N/$^{15}$N-DNA band that was actually seen after 20 min is compatible with both dispersive and semiconservative replication, but the two bands seen after 40 min are consistent only with semiconservative replication. Dispersive replication continues to give hybrid $^{14}$N/$^{15}$N molecules after two rounds of replication, whereas the granddaughter molecules produced at this stage by semiconservative replication include two that are made entirely of $^{14}$N-DNA.
did not provide a distinction between Delbrück’s dispersive model and the semi-conservative process favored by Watson and Crick. The distinction was possible, however, when the DNA molecules resulting after two rounds of replication were examined. Now the density gradient revealed two bands of DNA, the first corresponding to a hybrid composed of equal parts of newly synthesized and old DNA and the second corresponding to molecules made up entirely of new DNA. This result agrees with the semi-conservative scheme but is incompatible with dispersive replication, which predicts that after two rounds of replication all molecules would still be hybrids.

**DNA topoisomerases provide a solution to the topological problem**

The Meselson–Stahl experiment proved that DNA replication in living cells follows the semi-conservative scheme proposed by Watson and Crick and hence indicated that the cell must have a solution to the topological problem. This solution was not understood by molecular biologists until some 25 years later, when the activities of the enzymes called DNA topoisomerases were characterized.

DNA topoisomerases solve the topological problem by counteracting the overwinding that otherwise would be introduced into the molecule by progression of the replication fork. They do this by carrying out breakage-and-reunion reactions similar but not identical to that envisaged by Delbrück. We recognize two broad classes of DNA topoisomerases, called types I and II, the distinction being that a type I enzyme breaks just one strand of the double helix, whereas type II enzymes break both strands. Type I enzymes are further subdivided into types IA and IB, depending on what they do after making the single-strand break (Figure 15.4):

- A type IA topoisomerase uses the single-strand break as a gate through which it passes the uncut polynucleotide. The two ends of the broken strand are then religated. This mode of action results in the linking number (the number of times one strand crosses the other in a circular molecule) being changed by one.

![Figure 15.4](image-url)

Figure 15.4 Mode of action of type I topoisomerases. (A) A type IA topoisomerase makes a cut in one strand and passes the second strand through this gap. (B) A type IB topoisomerase cuts one strand and then rotates the cut strand around the uncut one.
A type IB topoisomerase acts as a molecular rotor, relaxing the torsional stress in an overwound helix by swiveling the cut strand around the uncut one. The linking number is therefore reduced by multiples of one.

Type II topoisomerases are also divided into A and B subgroups, but both IIA and IIB enzymes act in the same way: the double-strand break creates a gate through which a second segment of the helix is passed prior to religation of the cut ends (Figure 15.5). This process changes the linking number by two.

Cutting one or both DNA strands might appear to be a drastic solution to the topological problem, leading to the possibility that the topoisomerase might occasionally fail to rejoin a strand, generating a break that would interfere with the replication process. This possibility is reduced by the mode of action of these enzymes. One end of each cut polynucleotide becomes covalently attached to a tyrosine amino acid at the active site of the enzyme, ensuring that this end of the polynucleotide is held tightly in place while the free end(s) is being manipulated. With type IA and II enzymes, the polynucleotide–tyrosine linkage involves a phosphate group attached to the free 5'-end of the cut polynucleotide, and with type IB enzymes, the linkage is via a 3'-phosphate group.

The four groups of topoisomerases—IA, IB, IIA, and IIB—have distinct structures and each probably evolved separately. Type IA topoisomerases were first discovered in *E. coli* and for some time were thought to be specific to prokaryotes, but a type IA enzyme, called Top3, is now known to be present in most eukaryotes.

![Figure 15.5 Mode of action of type II topoisomerase.](image-url)
However, the main eukaryotic topoisomerases are type IB enzymes. To date no definite type IB topoisomerase has been identified in prokaryotes, although genome annotations indicate that some archaea possess genes that code for type IB-like topoisomerases. Type IIA enzymes are known in all species, and type IIB enzymes are found in archaea and plants.

Replication is not the only activity that is complicated by the topology of the double helix, and it is becoming increasingly clear that DNA topoisomerases have equally important roles during transcription, recombination, and other processes that can result in over- or underwinding of DNA. In eukaryotes, type II topoisomerases are also responsible for separating DNA molecules that become intertwined during chromosome division. Most topoisomerases are only able to relax overwound DNA, but some prokaryotic enzymes, such as the bacterial DNA gyrase (a type IIA topoisomerase) and the archaeal reverse gyrase (a type IA enzyme), can carry out the reverse reaction and introduce supercoils into DNA molecules (see Figure 8.1).

Variations on the semiconservative theme

No exceptions to the semiconservative mode of DNA replication are known, but there are several variations on this basic theme. DNA copying via a replication fork, as shown in Figure 15.1, is the predominant system, used by chromosomal DNA molecules in eukaryotes and by the circular genomes of prokaryotes. Some smaller circular molecules use a slightly different process called displacement replication. In these molecules, the point at which replication begins is marked by a D-loop, a region of approximately 500 bp where the double helix is disrupted by the presence of an RNA molecule base-paired to one of the DNA strands (Figure 15.6). This RNA molecule acts as the starting point for synthesis of one of the daughter polynucleotides. This polynucleotide is synthesized by continuous copying of one strand of the helix, while the second strand is displaced and subsequently copied after synthesis of the first daughter genome has been completed. Displacement replication has been studied extensively because it is thought to be the primary mode used to replicate the mitochondrial genomes of humans and other vertebrates.

![Figure 15.6 Displacement replication.](image-url)
The advantage of displacement replication compared to the regular form of semiconservative replication is not clear. In contrast, the special type of displacement process called rolling-circle replication is an efficient mechanism for rapid synthesis of multiple copies of a circular genome. Rolling-circle replication, which is used by λ and various other bacteriophages, initiates at a nick made in one of the parent polynucleotides. The free 3′-end that results is extended, displacing the 5′-end of the polynucleotide. Continued DNA synthesis rolls off a complete copy of the genome, and further synthesis eventually results in a series of genomes linked head to tail (Figure 15.7). These genomes are single-stranded and linear, but they can easily be converted to double-stranded circular molecules by complementary strand synthesis, followed by cleavage at the junction points between genomes and circularization of the resulting segments.

15.2 THE INITIATION PHASE OF GENOME REPLICATION

In order to initiate a round of genome replication, the double helix must be opened up at a particular point and the replication machinery must be assembled at the two nascent replication forks that are created. Initiation of replication is not a random process and always begins at the same or similar positions on a DNA molecule, these points being called the origins of replication. Once initiated, the replication forks emerge from the origin and progress in opposite directions along the DNA: replication is therefore bidirectional with most genomes (Figure 15.8). A circular bacterial genome has a single origin of replication, meaning that several thousand kilobases of DNA are copied by each replication fork. This situation differs from that seen with eukaryotic chromosomes, which have multiple origins and whose replication forks progress for shorter distances. The yeast Saccharomyces cerevisiae, for example, has about 400 origins, corresponding to 1 per 15.25 kb of DNA, and humans have 30,000–50,000 origins, or 1 for every 65–110 kb of DNA.

Initiation at the E. coli origin of replication

We know substantially more about initiation of replication in bacteria than in eukaryotes. The E. coli origin of replication is referred to as oriC. By transferring segments of DNA from the oriC region into plasmids that lack their own origins, it has been estimated that the E. coli origin of replication spans approximately 245 bp of DNA. Compared to other bacterial species, the E. coli origin is relatively short: lengths in general range from 100 to 1000 bp. Despite these variations, most bacterial origins have a very similar organization, comprising an AT-rich DNA
unwinding element (DUE) and 5–12 binding sites displaying differing degrees of affinity for a protein called DnaA (Figure 15.9A). The high-affinity sites, of which there are three in the *E. coli* origin, are permanently occupied by DnaA proteins, while the other sites are filled immediately before replication commences. With 5–12 sites, it might be imagined that 5–12 copies of DnaA attach, but in fact bound DnaA proteins cooperate with unbound molecules until some 10–20 copies are associated with the origin. Attachment occurs only when the DNA is negatively supercoiled, as is the normal situation for a bacterial chromosome (Section 8.1).

The result of DnaA binding is that the double helix opens up (melts) within the AT-rich DUE (Figure 15.9B). The exact mechanism is unknown, but DnaA does not appear to possess the enzymatic activity needed to break base pairs, and it is therefore assumed that the helix is melted by torsional stresses introduced by attachment of the DnaA proteins. An attractive model imagines that the DnaA proteins form a barrel-like structure around which the helix is wound. Melting of the helix is promoted by the HU proteins, the most abundant of the bacterial DNA packaging proteins (Section 8.1).

Melting of the helix initiates a series of events that construct a nascent replication fork at either end of the open region. The first step is the attachment of a prepriming complex at each of these two positions. Each prepriming complex initially comprises 12 proteins, six copies of DnaB and six copies of DnaC, but DnaC has a transitory role and is released from the complex soon after it is formed: its function is probably just to aid the attachment of DnaB. DnaB is a helicase, an enzyme that can break base pairs. DnaB begins to increase the single-stranded region within the origin, enabling the enzymes involved in the elongation phase of genome replication to attach. This represents the end of the initiation phase of replication in *E. coli*, as the replication forks now start to progress away from the origin and DNA copying begins.

**Origins of replication have been clearly defined in yeast**

The technique used to delineate the *E. coli* oriC sequence, involving transfer of DNA segments into a nonreplicating plasmid, has also proved valuable in identifying origins of replication in the yeast *S. cerevisiae*. Origins identified in this way are called autonomously replicating sequences (ARS). A typical yeast origin is shorter than *E. coli* oriC, usually less than 200 bp in length. Like the *E. coli* origin, the yeast sequence contains discrete regions, called subdomains, each with a different functional role (Figure 15.10A). The most important is subdomain A, also called the autonomous consensus sequence (ACS). This 11 bp sequence is found at over 12,000 positions in the *S. cerevisiae* genome, but only 400 of these act as replication origins under most circumstances. The ACS, along with the adjacent subdomain B1, makes up the origin recognition sequence, a stretch of some 40 bp in total that is the binding site for the origin recognition complex (ORC), a set of six proteins that attach to the origin (Figure 15.10B). ORCs are
attached to the yeast replication origins at all times and are involved in the regulation of genome replication, acting as mediators between replication origins and the regulatory signals that coordinate the initiation of DNA replication with the cell cycle (Section 15.5).

There is no evidence that the ORC is directly responsible for melting the helix. We must therefore look elsewhere in yeast origins for sequences that provide this function. This leads us to the two other conserved sequences in a typical yeast origin, subdomains B2 and B3 (see Figure 15.10A). Our current understanding suggests that melting is induced by attachment of ARS binding factor 1 (ABF1) to subdomain B3 (see Figure 15.10B) and that subdomain B2, which is AT-rich and also acts as the binding site for the MCM helicase, is the position at which the two strands of the helix are first separated. As in E. coli, melting of the helix within a yeast replication origin is followed by attachment of the replication enzymes, completing the initiation process and enabling the replication forks to begin their progress along the DNA.

**Origins in higher eukaryotes have been less easy to identify**

The identification of replication origins in humans and other higher eukaryotes has proved to be less straightforward. **Initiation regions** (parts of the chromosomal DNA where replication initiates) were originally delineated by various biochemical methods: for example, by allowing replication to initiate in the presence of labeled nucleotides and then arresting the process, purifying the newly synthesized DNA, and determining the positions of the labeled, nascent strands in the genome. These experiments suggested that there are specific regions in mammalian chromosomes where replication begins, but it quickly became clear that these regions do not contain replication origins equivalent to those in yeast. Further doubts about the specificity of mammalian replication origins were raised by experiments showing that various fragments of mammalian DNA greater than 2 kb in length can confer replicative ability on replication-deficient plasmids introduced into human cells, and that segments of bacterial DNA are only slightly less efficient in this regard. It was also shown that the mammalian ORC, although made of proteins that are homologous to their yeast counterparts, does not have sequence-specific DNA binding ability and appears to attach to the genome at multiple positions, many of which are not used as replication origins during a particular cell cycle.

The most recent attempts to characterize replication origins in higher eukaryotes have utilized genomewide screens to identify the positions at which DNA synthesis initiates during genome replication. Various methods have been adopted, the most successful being the following (Figure 15.11):

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**Figure 15.11** Methods for identifying the positions of replication origins. (Left) Short nascent strand (SNS) sequencing, in which the short strands representing newly synthesized DNA are sequenced. (Right) Bubble-seq, which involves treating the DNA with a restriction endonuclease and sequencing fragments that contain replication bubbles. With both methods, the sequence reads are mapped onto the genome annotation in order to identify the positions of replication origins.
In short nascent strand (SNS) sequencing, DNA is extracted immediately after initiation of a round of genome replication, and the short fragments representing newly synthesized DNA are purified and sequenced. The sequences are then mapped onto the reference genome in order to identify the positions where replication was initiated. A problem with this method is that it can be difficult to distinguish short nascent fragments from fragments resulting from the random DNA breakage that occurs during DNA extraction.

Bubble-seq again involves extracting DNA shortly after genome replication has begun, but the DNA is then cut with a restriction endonuclease and the replication bubbles, which form when the two replication forks proceed away from an origin in opposite directions, are purified. Sequencing of these bubbles indicates the replication initiation points.

These approaches have enabled thousands of initiation sites to be identified in various cell types, but there have been concerns about the lack of agreement between the results obtained when different methods are used. In some cases there is as little as 33% identity between the initiation regions located by SNS sequencing compared with those found by bubble-seq. Results with the same method can also be discordant: there was only 14% overlap between the initiation regions identified in human HeLa cells when two modifications of SNS sequencing were used. These anomalies might be explained in part by a single cell type using different origins for different rounds of replication, but it is also likely that only a fraction of the active initiation origins are being identified in any single experiment.

Examination of the many initiation regions that have been identified by these high-throughput methods has still not revealed any obvious characteristic features among replication origins in higher eukaryotes. It is now accepted that the origins in these species are not clearly defined and that the signals that determine where replication initiates are provided largely by the chromatin structure rather than the DNA sequence. In agreement with this model, it is known that ORCs attach preferentially at regions where the DNA displays negative supercoiling, which is induced by removal of nucleosomes. There may also be histone modifications that mark those origins that will be active in a particular round of genome replication.

15.3 EVENTS AT THE REPLICATION FORK

The central players in genome replication are the DNA polymerases that synthesize the daughter strands of DNA. DNA polymerases are a diverse group of enzymes that can be divided into at least seven groups on the basis of their structural and catalytic properties (Table 15.1). These enzymes include not just the DNA-dependent polymerases that are involved in genome replication and in repair of damaged DNA (Section 16.2) but also reverse transcriptase, which is an RNA-dependent DNA polymerase, and the template-independent DNA polymerase called terminal deoxynucleotidyl transferase (Section 2.1). To understand the events occurring at the replication fork, we must first understand the properties of the DNA polymerases that carry out genome replication.

DNA polymerases are molecular machines for making (and degrading) DNA

The principal chemical reaction catalyzed by a DNA polymerase is the 5’ → 3’ synthesis of a DNA polynucleotide. We learned in Section 2.1 that some DNA polymerases combine this function with at least one exonuclease activity, which means that these enzymes can degrade polynucleotides as well as synthesize them (see Figure 2.7):

- A 3’ → 5’ exonuclease activity is possessed by many bacterial and eukaryotic template-dependent DNA polymerases, enabling the enzyme to remove nucleotides from the 3’-end of the strand that it has just synthesized. This
15.3 Even though a proofreading activity whose function is to correct the occasional base-pairing error that might occur during strand synthesis.

- A 5′ → 3′ exonuclease activity is less common but is possessed by some polymerases whose function in replication requires that they must be able to remove at least part of a polynucleotide that is already attached to the template strand that the polymerase is copying.

The search for DNA polymerases began in the mid-1950s, as soon as it was realized that DNA synthesis was the key to replication of genes. It was thought that bacteria would probably have just a single DNA polymerase, and when the enzyme now called DNA polymerase I was isolated by Arthur Kornberg in 1957, there was a widespread assumption that this was the main replicating enzyme. The discovery that inactivation of the *E. coli* polA gene, which codes for DNA polymerase I, was not lethal (cells were still able to replicate their genomes) therefore came as something of a surprise, especially when a similar result was obtained with inactivation of polB, coding for a second enzyme, DNA polymerase II. It was not until 1972 that the main replicating polymerase of *E. coli*, DNA polymerase III, was eventually isolated. Both DNA polymerases I and III are involved in genome replication, as we will see in the next section. DNA polymerase II, as well as other bacterial DNA polymerases such as IV and V, is mainly involved in the repair of damaged DNA (Section 16.2).

DNA polymerases I and II are single polypeptides, but DNA polymerase III, befitting its role as the main replicating enzyme, is a multisubunit protein. The subunit called α is responsible for synthesizing the new polynucleotide, with the other subunits playing ancillary roles in the replication process. For example, the ε subunit specifies a 3′ → 5′ exonuclease activity, and the β subunit acts as a sliding clamp, holding the polymerase complex tightly to the template strand but at the same time allowing it to move along that strand as it makes the new polynucleotide.

Eukaryotes have at least 15 DNA polymerases, which in mammals are distinguished by Greek suffixes (α, β, γ, δ, etc.), an unfortunate choice of nomenclature as it tempts confusion with the identically named subunits of *E. coli* DNA polymerase III. The main replicating enzymes are DNA polymerase δ and DNA polymerase ε, which work in conjunction with an accessory protein called the

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<th>Table 15.1 DNA Polymerases</th>
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proliferating cell nuclear antigen (PCNA). PCNA is a homotrimer that encircles the DNA and provides the attachment point for DNA polymerases and other proteins involved in replication. DNA polymerase α also has an important function in genome replication, and DNA polymerase γ, although coded by a nuclear gene, is responsible for replicating the mitochondrial genome. As with the prokaryotic enzymes, most of the other eukaryotic DNA polymerases are involved in the repair of damaged DNA.

DNA polymerases have limitations that complicate genome replication

DNA polymerases have two limitations that complicate genome replication. The first of these is their ability to synthesize DNA only in the 5’ → 3’ direction. This means that one strand of the parent double helix, called the leading strand, can be copied in a continuous manner, but replication of the lagging strand has to be carried out in a discontinuous fashion, resulting in a series of short segments that must be ligated together to produce the intact daughter strand (Figure 15.12). The discontinuous nature of lagging-strand replication was confirmed in 1969 when Okazaki fragments, as these segments are now called, were first isolated from *E. coli*. In bacteria, Okazaki fragments are 1000–2000 nucleotides in length, but in eukaryotes the equivalent fragments appear to be much shorter, perhaps less than 200 nucleotides in length. The latter is an interesting observation that might indicate that each round of discontinuous synthesis replicates the DNA associated with a single nucleosome, comprising between 140 and 150 bp wound around the core particle plus 50–70 bp of linker DNA (Section 7.1).

The second limitation of DNA synthesis by DNA polymerases, which is also illustrated in Figure 15.12, is the need for a primer to initiate synthesis of each new polynucleotide. It is not known for certain why DNA polymerases cannot begin synthesis on an entirely single-stranded template, but this may relate to the proofreading activity of these enzymes, which is essential for the accuracy of replication. The logic is as follows. If the nucleotide that has just been added to the 3’-end of the growing polynucleotide is not base-paired to the template, then an error has been made, and to correct this error, the 3’ → 5’ exonuclease of the polymerase must operate instead of continued 5’ → 3’ polymerization. In other words, the 5’ → 3’ polymerase function is active only if the 3’-terminal nucleotide is base-paired to the template. A template that is entirely single-stranded has no base-paired 3’ nucleotide and so requires a primer, to provide that nucleotide, in order for the polymerase to be activated.

Whatever the reason, priming is a necessity in DNA replication but does not present too much of a problem. Although DNA polymerases cannot deal with an entirely single-stranded template, RNA polymerases have no difficulty in this respect, so the primers for DNA replication are made of RNA. In bacteria, primers are synthesized by primase, a special RNA polymerase unrelated to the transcribing enzyme, with each primer being 10–12 nucleotides in length. Once the primer has been completed, strand synthesis is continued by DNA polymerase III (Figure 15.13A). In eukaryotes, the situation is slightly more complex because the primase is tightly bound to DNA polymerase α and cooperates with this enzyme in synthesis of the first few nucleotides of a new polynucleotide. This primase synthesizes an RNA primer of 7–12 nucleotides and then hands over to DNA polymerase α, which extends the RNA primer by adding about 20 nucleotides of DNA. This DNA stretch often has a few ribonucleotides mixed in, but it is not clear if these are incorporated by DNA polymerase α or by intermittent activity of the primase. After completion of the RNA–DNA primer, DNA synthesis is continued by DNA polymerase ε on the leading strand and by DNA polymerase δ on the lagging strand (Figure 15.13B).

Priming needs to occur just once on the leading strand, within the replication origin, because once primed, the leading-strand copy is synthesized continuously until replication is completed. On the lagging strand, priming is a repeated process that must occur every time a new Okazaki fragment is initiated. In *E. coli*, which makes Okazaki fragments of 1000–2000 nucleotides in length,
approximately 4000 priming events are needed every time the genome is replicated. In eukaryotes, the Okazaki fragments are much shorter and priming is a highly repetitive event.

**Okazaki fragments must be joined together to complete lagging-strand replication**

Many of the events at the replication fork are similar in both bacteria and eukaryotes. Progress of the replication fork is maintained by helicase activity, with the torsional stress that results from overwinding of the helix ahead of the fork relieved by DNA topoisomerases. Single-stranded DNA is naturally sticky and the two separated polynucleotides produced by helicase action would immediately re-form base pairs after passage of the enzyme, if allowed. The single strands are also highly susceptible to nuclease attack and are likely to be degraded if they are not protected in some way. To avoid these unwanted outcomes, **single-strand binding proteins** (SSBs) attach to the polynucleotides and prevent them from reassociating or being degraded (**Figure 15.14A**). Detachment of the SSBs, which must occur when the replication complex arrives to copy the single strands, is brought about by a second set of proteins called **replication mediator proteins** (RMPs). DNA synthesis in *E. coli* is carried out by two linked copies of DNA polymerase III, one for leading-strand replication and one for the lagging strand (**Figure 15.15**), aided by a single γ complex (sometimes called the clamp loader). The main role of the γ complex is to interact with the β subunit (the sliding clamp) of each polymerase and hence control the attachment and removal of the enzyme from the template, a function that is required primarily during lagging-strand replication when the enzyme has to attach and detach repeatedly at
the start and end of each Okazaki fragment. The combination of these two DNA polymerases, along with the primase needed for repetitive initiation of Okazaki fragments, is called the replisome. In eukaryotes, the DNA polymerase ε and δ enzymes that copy the leading and lagging strands, respectively, do not associate into a dimeric complex but instead remain separate. The function performed by the γ complex of the *E. coli* polymerase—controlling attachment and detachment of the DNA polymerase from the lagging strand—is carried out by a multisubunit accessory protein called replication factor C (RFC).

Just one issue must be resolved in order to complete the DNA synthesis phase of genome replication. The initial copy of the lagging strand comprises a series of Okazaki fragments. The RNA primers of these fragments must be replaced with DNA, and adjacent fragments must be joined, in order to generate the final version of the lagging-strand copy. This is one aspect of genome replication where the events occurring in bacteria and eukaryotes are significantly different.

Removal of the RNA primer of an Okazaki fragment can be achieved if the DNA polymerase that is making the next fragment in the series has a 5′ → 3′ exonuclease activity. This polymerase could continue making its DNA copy of the lagging strand into the region initially occupied by the 5′-end of the adjacent Okazaki fragment, by using the exonuclease to remove the RNA nucleotides and then recopying that segment into DNA. Unfortunately, DNA polymerase III, which copies the lagging strand in bacteria, lacks the required 5′ → 3′ exonuclease activity (Table 15.2). DNA polymerase III therefore releases the lagging strand when it reaches the 5′-end of the adjacent Okazaki fragment. Its place is taken by DNA polymerase I, which does have a 5′ → 3′ exonuclease activity and so removes the primer, and usually the start of the DNA component of the Okazaki fragment as well, extending the 3′-end of the fragment it is synthesizing into the region of the template that is exposed (Figure 15.16A). The two Okazaki fragments now abut, with the terminal regions of both composed entirely of DNA. All that remains is for the missing phosphodiester bond to be put in place by a DNA ligase, linking the two fragments and completing replication of this region of the lagging strand.

In eukaryotes, the problem is more extreme as there appears to be no eukaryotic DNA polymerase with the 5′ → 3′ exonuclease activity needed to degrade the RNA primers of the Okazaki fragments. The solution to this problem is therefore

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**Figure 15.15** A model for parallel synthesis of the leading- and lagging-strand copies by a dimer of DNA polymerase III enzymes. It is thought that the lagging strand loops through its copy of the DNA polymerase III enzyme, in the manner shown, so that both the leading and lagging strands can be copied as the dimer moves along the molecule being replicated. The two components of the DNA polymerase III dimer are not identical because they share a single γ complex, which is made up of subunit γ in association with subunits δ, δ′, χ, and ψ.

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**Table 15.2 Exonuclease Activities of DNA Polymerases Involved in Replication of Bacterial and Eukaryotic Genomes**

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Bacterial DNA polymerases</th>
<th>Eukaryotic DNA polymerases</th>
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<tbody>
<tr>
<td></td>
<td>3′ → 5′</td>
<td>5′ → 3′</td>
</tr>
<tr>
<td>DNA polymerase I</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>DNA polymerase III</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>DNA polymerase α</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>DNA polymerase δ</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>DNA polymerase ε</td>
<td>Yes</td>
<td>No</td>
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Very different from that described for bacteria. The central player is the flap endonuclease (FEN1), which has an unusual endonuclease activity that enables it to cut the phosphodiester bond at the branch point at the base of the flap that is formed when the 5' end of a polynucleotide is displaced from its template. FEN1 therefore associates with DNA polymerase δ as it approaches the RNA primer of the adjacent Okazaki fragment. The base pairs holding the primer to the lagging strand are broken by a helicase enzyme, enabling the primer to be pushed aside by the polymerase as it extends the adjacent Okazaki fragment into the exposed region, resulting in the flap structure that can be cut off by FEN1. A DNA ligase can then form the final phosphodiester bond to link the Okazaki fragments. This scheme raises the possibility that both the RNA primer and all of the DNA originally synthesized by the priming enzyme, DNA polymerase α, are removed. This is an attractive option because DNA polymerase α has no 3' --> 5' proofreading activity (see Table 15.2) and therefore synthesizes DNA in a relatively error-prone manner. Removal of this region as part of the flap cleaved by FEN1, followed by resynthesis by DNA polymerase δ (which does have a proofreading activity and so makes a highly accurate copy of the template), would prevent any errors made by DNA polymerase α from becoming permanent features of the daughter double helix.

15.4 TERMINATION OF GENOME REPLICATION

Replication forks proceed along linear genomes, or around circular ones, generally unimpeded except when a region that is being transcribed is encountered. DNA synthesis occurs at approximately five times the rate of RNA synthesis, so the replication complex can easily overtake an RNA polymerase, but this probably does not happen: instead, it is thought that the replication fork pauses behind the RNA polymerase, proceeding only when the transcript has been completed. Eventually the replication fork reaches the end of the molecule or meets a second replication fork moving in the opposite direction. What happens next is one of the least understood aspects of genome replication.

**Figure 15.16** The series of events involved in joining up adjacent Okazaki fragments during DNA replication.

(A) DNA polymerase III of *E. coli* lacks a 5' --> 3' exonuclease activity and so stops making DNA when it reaches the RNA primer of the adjacent Okazaki fragment. At this point, DNA synthesis is continued by DNA polymerase I, which does have a 5' --> 3' exonuclease activity and so can remove the RNA primer and replace it with DNA. DNA polymerase I usually also replaces some of the DNA from the Okazaki fragment before detaching from the template. This leaves a single missing phosphodiester bond, which is synthesized by DNA ligase, completing this step in the replication process. (B) The eukaryotic DNA polymerase δ also lacks a 5' --> 3' exonuclease activity, but it continues making DNA into the region occupied by the primer of the adjacent Okazaki fragment, the 5' end of this fragment being displaced by a helicase. The resulting flap structure is then cut off by FEN1.
Replication of the *E. coli* genome terminates within a defined region

Bacterial genomes are replicated bidirectionally from a single point (see Figure 15.8A), which means that the two replication forks should meet at a position diametrically opposite the origin of replication on the genome map. However, if one fork is delayed, possibly because it has to replicate extensive regions where transcription is occurring, then it might be possible for the other fork to overshoot the halfway point and continue replication on the other side of the genome (Figure 15.17). It is not immediately apparent why this should be undesirable, as the daughter molecules presumably would be unaffected, but it is not allowed to happen because of the presence of terminator sequences. Ten of these have been identified in the *E. coli* genome (Figure 15.18A), each one acting as the recognition site for a sequence-specific DNA-binding protein called the terminator utilization substance (Tus).

The mode of action of Tus is quite unusual. When it is bound to a terminator sequence, a Tus protein allows a replication fork to pass if the fork is moving in one direction but blocks progress if the fork is moving in the opposite direction around the genome. The directionality is set by the orientation of the Tus protein on the double helix (Figure 15.18B). Exactly how Tus exerts its effect is not yet understood, but it has been assumed that the Tus protein interacts directly with the DnaB helicase, which is responsible for progression of the replication fork. This assumption has been challenged by recent research supporting the alternative possibility that the key interactions are between Tus and the replication fork itself.

The progress of individual replication forks has been studied in an experimental system in which replisome proteins are absent. Without these proteins, a fork will not move naturally along a DNA double helix. To bring about fork progression in the experimental system, a magnetic bead was attached to the end of one of the polynucleotides, and the end of the second polynucleotide was immobilized by attachment to a solid support. The two polynucleotides were then pulled apart by manipulating the magnetic bead with a magnetic tweezer, a device comprising a set of magnets whose positions and field strengths can be varied in such a way that the magnetic bead, and its attached polynucleotide, can be moved about in a controlled manner (Figure 15.19). Upon moving the magnetic bead away from the solid support, the helix is opened up, producing a fork that can be moved along the helix simply by pulling the ends of the two polynucleotides further apart.

Figure 15.17 A situation that is not allowed to occur during replication of the circular *E. coli* genome. One of the replication forks has proceeded some distance past the halfway point. This does not happen during DNA replication in *E. coli* because of the action of the Tus proteins.
These experiments have shown that if the orientation of the terminator sequence is such that the fork approaches a Tus protein from the permissive direction, then movement of the fork is not impeded, but when the fork approaches from the nonpermissive direction, its progress is prevented by Tus. The implication is that interactions between Tus and the DNA being replicated are at least partly responsible for the ability of Tus to block progression of the replication fork. The results do not exclude a role for a protein–protein interaction between Tus and DnaB, but they do show that such an interaction is unlikely to be the full explanation of the mode of action of Tus.

Whatever the mechanism, the orientation of the terminator sequences, and hence of the bound Tus proteins, in the *E. coli* genome is such that both replication forks become trapped within a relatively short region on the opposite side of the genome to the origin (see Figure 15.18A). This ensures that termination always occurs at or near the same position. Each leading strand becomes ligated to the lagging strand of the other fork and the replisomes are dissembled. The result is two interlinked daughter molecules, which are separated by topoisomerase IV, a type II enzyme.

**Little is known about termination of replication in eukaryotes**

Much less is known about the termination of DNA replication in eukaryotes. Fork pausing elements, similar in function to bacterial terminator sequences, have been identified in yeast chromosomes, associated with the type II DNA topoisomerase Top2 that separates interlinked daughter molecules. The importance of these sequences is unclear, as some replication forks terminate in regions not marked by these sequences, and forks can pass through these sequences with the aid of helicase enzymes. In higher eukaryotes, there is little evidence for pause or capture sites, and each replication fork is thought to continue its progression until it meets a fork coming in the other direction. The leading strands then pass one another and each becomes ligated to the lagging strand from the other fork.

Inside eukaryotic nuclei, DNA synthesis occurs within replication factories and replication foci, which are discrete, immobile regions that contain all the relevant proteins and through which the DNA threads as it is being replicated. Up to 10 replication origins are thought to be associated with a single factory,
the replicated DNA between these origins looping out of the factory as it is synthesized. This model reduces the need for discrete termination sequences in the DNA as it implies that the progression of adjacent forks could be coordinated within a single factory, so the forks meet at an appropriate position. The factory model also helps to solve the difficult question of how the daughter DNA molecules produced in a eukaryotic nucleus are prevented from becoming impossibly tangled up. Although type II DNA topoisomerases have the ability to untangle DNA molecules, it is generally assumed that tangling is kept to a minimum so that extensive breakage-and-reunion reactions, as catalyzed by topoisomerases, can be avoided. It is envisaged that the replication of individual segments of a chromosome in a single factory enables the daughter molecules to be maintained in a specific arrangement that at least partially avoids their entanglement.

Cohesins also play an important role. These are multisubunit proteins that form ring structures that are attached to the daughter molecules immediately after passage of the replication fork. The cohesins then maintain the alignment of the sister chromatids until the anaphase stage of nuclear division, when they are cleaved by cutting proteins, enabling the daughter chromosomes to be separated (Figure 15.20).

**Telomerase completes replication of chromosomal DNA molecules, at least in some cells**

There is one final problem that we must consider before leaving the replication process. This concerns the steps that have to be taken to prevent the ends of a linear, double-stranded molecule from gradually getting shorter during successive rounds of chromosomal DNA replication. There are two ways in which this shortening might occur:

- **The extreme 3'-end of the lagging strand might not be copied because the final Okazaki fragment cannot be primed, as the natural position for the priming site is beyond the end of the template (Figure 15.21A).** The absence of this Okazaki fragment means that the lagging-strand copy is shorter than it should be. If the copy remains this length, then when it acts as a parent polynucleotide in the next round of replication, the resulting daughter molecule will be shorter than its grandparent.

- **If the primer for the last Okazaki fragment is placed at the extreme 3'-end of the lagging strand, then shortening will still occur, although to a lesser extent, because this terminal RNA primer cannot be converted into DNA by the standard processes for primer removal (Figure 15.21B).** This is because the methods for primer removal require extension of the 3'-end of an adjacent Okazaki fragment, which cannot occur at the very end of the molecule.

Once this problem had been recognized, attention was directed at the telomeres, the unusual DNA sequences at the ends of eukaryotic chromosomes. We noted in Section 7.1 that telomeric DNA is made up of a type of minisatellite sequence, composed of multiple copies of a short repeat motif, 5'-TTAGGG-3' in most higher eukaryotes, a few hundred copies of this sequence occurring in tandem repeats at each end of every chromosome. The solution to the shortening problem lies with the way in which this telomeric DNA is made. Most of the telomeric DNA is copied in the normal fashion during DNA replication, but this is not the only way in which it can be synthesized. To compensate for the limitations of the replication process, telomeres can be extended by an independent mechanism catalyzed by the enzyme **telomerase.** This is an unusual enzyme in that it consists of both protein and RNA. In the human enzyme, the RNA component is 450 nucleotides in length and contains near its 5'-end the sequence 5'-CUAACCCCUAAC-3', whose central region is the reverse complement of the human telomere repeat sequence 5'-TTAGGG-3'. This complementarity enables telomerase to extend the telomeric DNA at the 3'-end of a polynucleotide by the copying mechanism shown in Figure 15.22, in which the telomerase RNA is used...
as a template for each extension step and DNA synthesis is carried out by the protein component of the enzyme, which is a reverse transcriptase. The correctness of this model is indicated by comparisons between telomere repeat sequences and the telomerase RNAs of other species (Table 15.3): in all organisms that have been examined, the telomerase RNA contains a sequence that enables it to make copies of the repeat motif present at the organism’s telomeres. An interesting feature is that, in all organisms, the strand synthesized by telomerase has a preponderance of G nucleotides and is therefore referred to as the G-rich strand.
15.4 Termination of Genome Replication

Telomerase can only synthesize this G-rich strand. It is not clear how the other polynucleotide (the C-rich strand) is extended, but it is presumed that when the G-rich strand is long enough, the primase–DNA polymerase α complex attaches at its end and initiates synthesis of complementary DNA in the normal way (Figure 15.23). This requires the use of a new RNA primer, so the C-rich strand will still be shorter than the G-rich one, but the important point is that the overall length of the chromosomal DNA has not been reduced. In mammalian cells, after extension, the end of the telomere may form a t-loop, in which the free 3\' -end loops back, invades the double helix, and forms base pairs with its complementary sequence on the C-rich strand (Figure 15.24). This reaction is promoted by the telomere-binding protein TRF2 and may provide additional stabilization of a chromosome end that does not require further extension.

The activity of telomerase must clearly be controlled very carefully to ensure that the appropriate length extension is made at each chromosome end. One part of this regulatory mechanism is provided by the TRF1 proteins, which bind to the telomere repeat sequences and inhibit the activity of telomerase (Section 7.1). As the telomere shortens, the number of bound TRF1 proteins decreases, enabling

**Table 15.3 Sequences of Telomere Repeats and Telomerase RNAs in Various Organisms**

<table>
<thead>
<tr>
<th>Species</th>
<th>Telomere repeat sequence</th>
<th>Telomerase RNA template sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human</td>
<td>5'-TTAGGG-3'</td>
<td>5'-CUAACCUAACC-3'</td>
</tr>
<tr>
<td>Oxytricha</td>
<td>5'-TTTTGGGG-3'</td>
<td>5'-CAAAACCCCAAACC-3'</td>
</tr>
<tr>
<td>Tetrahymena</td>
<td>5'-TTGGGG-3'</td>
<td>5'-CAACCCCAA-3'</td>
</tr>
</tbody>
</table>

**Figure 15.22** Extension of the end of a human chromosome by telomerase.
The 3\'-end of a human chromosomal DNA molecule is shown. The sequence comprises repeats of the human telomere motif 5'-TTAGGG-3'. The telomerase RNA base-pairs to the end of the DNA molecule, which is extended a short distance. The telomerase RNA then translocates to a new base-pairing position slightly further along the DNA polynucleotide and the molecule is extended by a few more nucleotides. The process can be repeated until the chromosome end has been extended by a sufficient amount.

**Figure 15.23** Completion of the extension process at the end of a chromosome. It is believed that after telomerase has extended the 3\'-end by a sufficient amount, a new Okazaki fragment is primed and synthesized, converting the 3\'-extension into a completely double-stranded end.
Stem cells are progenitor cells that divide continually. With some.

Telomere length is implicated in cell senescence and cancer

Perhaps surprisingly, telomerase is not active in all mammalian cells. The enzyme is functional in the early embryo, but after birth it is active only in the reproductive cells and stem cells. Stem cells are progenitor cells that divide continually throughout the lifetime of an organism, producing new cells to maintain organs and tissues in a functioning state. Somatic cells, which lack telomerase activity, undergo chromosome shortening every time they divide. Eventually, after many cell divisions, the chromosome ends could become so truncated that essential genes are lost, but this is unlikely to be a major cause of the defects that can occur in cells lacking telomerase activity. Instead, the critical factor is the need to maintain a protein cap on each chromosome end, to protect these ends from the effects of the DNA repair enzymes that join together uncapped ends that are produced by accidental breakage of a chromosome (Section 16.2). The proteins that form this protective cap, such as TRF2 in humans, recognize the telomere repeats as their binding sequences and so have no attachment points after the telomeres have been deleted. If these proteins are absent, then the repair enzymes can make inappropriate linkages between the ends of intact, although shortened, chromosomes; this is probably the underlying cause of the disruption to the cell cycle that results from telomere shortening.

Telomere shortening will therefore lead to the termination of a cell lineage. For several years, biologists have attempted to link this process with cell senescence, a phenomenon originally observed in cell cultures. All normal cell cultures have a limited lifetime: after a certain number of divisions, the cells enter a senescent state in which they remain alive but cannot divide (Figure 15.25). With some mammalian cell lines, notably cultures of fibroblasts (connective tissue cells), senescence can be delayed by engineering the cells so that they synthesize active telomerase. These experiments suggest a clear relationship between telomere shortening and senescence, but the exactness of the link has been questioned, and any extrapolation from cell senescence to aging of the organism is fraught with difficulties.

Not all cell lines display senescence. Cancerous cells are able to divide continuously in culture, their immortality being looked upon as analogous to tumor growth in an intact organism. With several types of cancer, this absence of senescence is associated with activation of telomerase, sometimes to the extent that telomere length is maintained through multiple cell divisions, but often in such a way that the telomeres become longer than normal because the telomerase is overactive. These observations suggest that drugs that inhibit telomerase activity might be useful in cancer treatment. Attempts to inactivate telomerase in cancer cells have focused on both the protein and RNA components of the enzyme. The protein has been used to prepare vaccines that contain antibodies that should bind to and inactivate any telomerase proteins that they encounter. In clinical trials, these vaccines have been able to reduce the number of cancerous cells circulating in the bloodstream of patients, reducing the chances of their cancer spreading to other parts of their body. Whether the vaccines can reduce the growth of existing tumors, or prevent a cancer from becoming established in the first place, is not yet clear. A second approach makes use of a short oligonucleotide that is complementary to part of the RNA component of telomerase. The idea is that this oligonucleotide will inhibit a telomerase enzyme by binding to its RNA molecule.
15.4 TERMINATION OF GENOME REPLICATION

**Figure 15.26** Use of an oligonucleotide to inhibit telomerase activity. The oligonucleotide is complementary to part of the RNA component of telomerase and hence competes with the chromosomal DNA for attachment to the telomerase RNA. Telomere extension is therefore inhibited.

(Figure 15.26). Vaccines and oligonucleotides targeted at telomerase are among the most promising of the various strategies being used to combat cancer, but a stumbling block at the moment is the ability of some cells to maintain their telomeres, and hence avoid senescence, by an alternative process that does not require telomerase. This involves the transfer of telomere repeats from DNA molecules that have not yet reached a critically short length to other DNA molecules that are close to the danger point. This process is switched on in some cancer cells, counteracting the effects of telomerase inactivation.

*Drosophila* has a unique solution to the end-shortening problem

When the amino acid sequences of the protein subunits of telomerase enzymes are compared with those of other reverse transcriptases, the closest similarities are seen with the reverse transcriptases coded by the non-long terminal repeat (non-LTR) retroelements called retroposons (Section 9.2). This is a fascinating observation when taken in conjunction with the unusual structure of the telomeres of *Drosophila*. These telomeres are not made up of the short repeated sequences seen in most other organisms but instead consist of tandem arrays of much longer repeats, 6 or 10 kb in length. These repeats are full-length and truncated copies of three *Drosophila* retroposons, called *HeT-A*, *TART*, and *TAHRE*. They are maintained by a process analogous to that carried out by telomerase (Figure 15.27), with a template RNA obtained by transcription of the telomeric retroposons being copied by the reverse transcriptase coded by the *TART* and *TAHRE* sequences (*HeT-A* does not have a reverse transcriptase gene).

The unusual structure of the *Drosophila* telomere could simply be a quirk of nature, but the attractive possibility that the telomes of other organisms are degraded retroposons, as suggested by the similarities between telomerase and retroposon reverse transcriptases, cannot be discounted.

**Figure 15.27** A model for maintenance of *Drosophila* telomeres. In this example, a *TART* sequence is located at the extreme end of the telomere. DNA replication has left the typical 3' overhang, which is extended by reverse transcription of a RNA copy of a *TART* sequence located within the intact region of the telomere. A new Okazaki fragment can then be primed in order to complete the extension process.
15.5 REGULATION OF EUKARYOTIC GENOME REPLICATION

It is essential that genome replication is regulated so that the process is completed by the time the cell actually divides. Eukaryotes possess a sophisticated set of controls that coordinate genome replication with the cell cycle and are able to arrest the replication process under certain circumstances, for example if the DNA is damaged and must be repaired before copying can be completed. We will end this chapter by looking at these regulatory mechanisms.

Genome replication must be synchronized with the cell cycle

The concept of a cell cycle emerged from light microscopic studies carried out by the early cell biologists. Their observations showed that dividing cells pass through repeated cycles of mitosis, which is the period when nuclear and cell division occurs (see Figure 3.16), and interphase, a less dramatic period when few dynamic changes can be detected with the light microscope. It was understood that chromosomes divide during interphase, so when DNA was identified as the genetic material, interphase took on a new importance as the period when genome replication takes place. This led to a reinterpretation of the cell cycle as a four-stage process (Figure 15.28), comprising the following:

- **Mitosis** or M phase, the period when the nucleus and cell divide
- **Gap 1** or G1 phase, an interval when transcription, translation, and other general cellular activities occur
- **Synthesis** or S phase, when the genome is replicated
- **Gap 2** or G2 phase, a second interval period

It is clearly important that the S and M phases are coordinated so that the genome is completely replicated, but replicated only once, before mitosis occurs. To ensure this happens, there is a series of cell cycle checkpoints that act as key transition stages, progression past a checkpoint committing the cell to the next phase of the cycle. The most important of these with regard to genome replication is the G1–S checkpoint, because only when this point has been passed is the cell able to replicate its DNA. If the cell suffers trauma or the genome has been damaged in some way, for example, by mutation of key genes, then the cell cycle can be arrested at this checkpoint while the damage is repaired. Later in the cell cycle, the G2–M checkpoint ensures that the cell is ready to enter mitosis, the central requirement being that the genome has been replicated correctly, with every part replicated once and no parts replicated multiple times.

Origin licensing is the prerequisite for passing the G1–S checkpoint

Studies, primarily with *S. cerevisiae*, have led to a model for the regulation of genome replication that defines origin licensing as the basis to preparation of the cell for passage through the G1–S checkpoint. Origin licensing involves construction on a replication origin of a set of proteins, called the pre-replication complex (pre-RC). Each origin is already marked by the six proteins that make up the origin recognition complex (ORC), which is assembled onto multiple origins soon after completion of the previous cell division. To become licensed, an origin must initially recruit two additional proteins, Cdc9 and Cdt1. Cdc9 is an ATPase that releases energy from a molecule of ATP that is present in the ORC (Figure 15.29). The energy is used to drive attachment of the MCM2–7 complex, which comprises six proteins that form the core components of the helicase that breaks base pairs at the eukaryotic replication fork. Cdc9 and Cdt1 depart as the MCM2–7 complex arrives, attachment of the MCM2–7 complex to the ORC completing construction of the pre-RC. These events appear to be similar in most eukaryotes.

Origin licensing enables the cell to pass through the G1–S checkpoint, but the pre-RCs are themselves inactive and unable to initiate genome replication.
To become active, each pre-RC must be converted into a preinitiation complex (pre-IC). In *S. cerevisiae*, this conversion is initiated soon after the beginning of S phase by a cyclin-dependent kinase (CDK) and by Dbf4-dependent kinase (DDK), which together phosphorylate target proteins, including the MCM2–7 components of the inactive helicase, in order to initiate a cascade of reactions that result in addition of Cdc45 and GINS to the pre-RC, converting it into a pre-IC (Figure 15.30). GINS, which is itself a complex of four proteins, and Cdc45 are the final components of the eukaryotic helicase, which is now active and able to participate in initiation of replication and progression of the replication forks away from the origin.

Identification of the components of the pre-RC and pre-IC takes us some distance toward understanding how genome replication is initiated, but it still leaves open the question of how replication is coordinated with other events in the cell cycle. Cell cycle control is a complex process, mediated largely by CDK proteins, which phosphorylate and activate enzymes and other proteins that have specific functions during the cell cycle. The activities of these CDKs change throughout the cell cycle, with minimal activity at the beginning of G1 phase and rapidly increasing activity during S phase (Figure 15.31). The CDK activity level is thought to influence the activities of the proteins whose functions are needed at different stages, thereby ensuring that the cell cycle progresses in an orderly manner. Most CDKs are present in the nucleus throughout the cell cycle, so they must themselves be subject to control. This control is exerted partly by proteins called cyclins, which vary in abundance at different stages of the cell cycle (Figure 15.32), partly by other protein kinases that activate the CDKs, and partly by inhibitory proteins. One of the main inhibitors is geminin, which accumulates during S, G2, and M phases and prevents the re-replication of genome regions that have already been replicated. It does this by binding to Cdt1 and hence ensuring that the MCM2–7 core helicase cannot be loaded onto origins that are present on the daughter DNA molecules. Along with many other proteins that are active during the late stages of the cell cycle, geminin is degraded during the anaphase period of mitosis by the anaphase-promoting complex/cyclosome (APC/C), which is a ubiquitin ligase that targets proteins for degradation by the proteasome (Section 13.3). The APC/C therefore remodels the nuclear proteome so the cell cycle can recommence in the daughter cells resulting from mitosis.
Replication origins do not all fire at the same time

Initiation of replication does not occur at the same time at all replication origins. Instead, during any particular cell cycle, some origins fire early in S phase and some later. Studies with S. cerevisiae have suggested that the pattern of origin firing is consistent from cell division to cell division, with euchromatin regions and the centromeres replicated early in S phase while heterochromatin and telomeres are replicated later. More recent work has confirmed that this general pattern is correct but has also indicated that there is little or no correspondence between the patterns of origin firing in different cells, suggesting that the pattern of firing is not inherited in any way.

How can origin firing be followed in living cells? One method involves labeling dividing cells with a brief pulse of a nucleotide analog, such as bromodeoxyuridine (BrdU), which is incorporated into growing polynucleotides in place of thymidine. The DNA that is synthesized during the pulse of labeling will contain BrdU nucleotides as well as T nucleotides, whereas the DNA made before and after the pulse will just contain T. The DNA is then extracted and combed onto a glass slide, so the molecules are obtained as linear fibers (Section 3.5). Treatment with an antibody specific for BrdU, this antibody carrying a fluorescent label, will reveal the genome regions that were being replicated at the time of the BrdU pulse. By comparison of molecules from cells at different stages in S phase, the pattern of origin firing can be followed. This technique has been used with Schizosaccharomyces pombe, which is a fission yeast, meaning that it divides by the cell splitting into two halves of equal size, rather than via small buds, as is the case with S. cerevisiae. The results show that the first origins to fire, at the beginning of S phase, are distributed randomly throughout the genome. As S phase continues, more origins fire, these forming clusters at various positions along the individual chromosomes. However, the positions of these clusters are not the same in different cells (Figure 15.33), showing that the pattern is not determined by the DNA sequences of individual initiation regions. Instead, it appears likely that the clusters represent regions of euchromatin, which are chosen at random as the starting points for replication of the genome. This hypothesis is supported by experiments with S. cerevisiae mutants that are unable to synthesize the RPD3 histone deacetylase. These cells, which are predicted to have a greater-than-normal degree of histone acetylation and hence a more open chromatin structure, display less control over replication origin firing, with early-firing origins now operating in parts of the genome that are not usually replicated until late in S phase.

Figure 15.33 Patterns of replication origin firing are not the same in different cells. Ten overlapping molecules from chromosome 2 of Schizosaccharomyces pombe are shown. The molecules were pulse-labeled with bromodeoxyuridine (BrdU) and combed into linear strands, and the label was detected with a BrdU-specific antibody. The clusters of active origins that are revealed are not the same in each molecule, indicating that the pattern of origin firing is different in different cells. The positions of the main clusters of fired origins are shown as pink bars on top of each DNA molecule. (From Kaykov A & Nurse P [2015] Genome Res. 25:391–401. With permission from Cold Spring Harbor Laboratory Press via CC BY 4.0.)
BrdU labeling also enables the migration rates of different replication forks in *S. pombe* to be estimated, from the lengths of the labeled tracts that are synthesized during the pulse period. The mean speed is 2.8 kb/min, but some forks move much more quickly, up to 11 kb/min for the most active ones (Figure 15.34). These figures are very similar to the migration rates measured for *S. cerevisiae* and estimated for human cells but are much slower than those for *E. coli*, which has to copy 116 kb of DNA/min in order to replicate its entire genome within the 20 min that elapsed between cell divisions when cultures are grown in rich medium. Fork migration rates are dependent on a number of factors, especially the presence of obstacles such as RNA polymerase complexes, which are bound to the DNA in order to transcribe genes, or damaged regions of the genome that must be repaired before they can be replicated. When a fork encounters such an obstacle, it might become stalled. After the blockage has been removed, the fork might recommence its journey, or it might remain stalled while a fork approaching from the other direction completes replication of the intervening region of DNA. Occasionally, however, both members of a pair of forks will stall permanently. This situation can be remedied by activation of additional replication origins within the region between the two stalled forks. This is possible, even in the presence of inhibitors such as geminin, because an excess of origins are always licensed during G1 phase. This means that the genome contains many activated but unused origins, some of which will remain unused even when the entire genome has been replicated.

**The cell has various options if the genome is damaged**

The cell cycle checkpoints are also important in preventing a damaged genome from being replicated. Damage can take many forms, including the dimerization of adjacent nucleotides, which is stimulated by UV irradiation (Section 16.1), and the presence of single- or double-strand breaks. If damage is detected, then the cell cycle will be arrested at the G1–S checkpoint while the DNA is repaired. There are also additional checkpoints within S phase that enable genome replication to be halted to repair damage that is detected when one or more replication forks become stalled, and the G2–M checkpoint ensures that postreplicative damage is corrected before mitosis takes place.

Cell cycle arrest occurs because of the activity of signal transduction pathways, activated by DNA damage, or indicators of damage, such as replication fork stalling. Two of these pathways are initiated by the ATM and ATR protein kinases, which either recognize damage directly or are activated by the detection proteins. ATM responds primarily to double-strand breaks in the genome while ATR responds to various types of damage, including single-strand breaks, that bring about replication fork arrest during S phase. The ATM pathway targets include the checkpoint kinase Chk2 as well as BRCA1, which when defective confers susceptibility to breast cancer. A second checkpoint kinase, Chk1, is activated by the ATR pathway. The checkpoint kinases act on cell cycle control proteins such as Cdc25, which regulate passage through the G2–M checkpoint. Phosphorylation of Cdc25 results in its degradation, so the cell cycle becomes arrested until the DNA damage is repaired.

If the damage is not excessive, then DNA repair processes are activated (Section 16.2). Alternatively, the cell may be shunted into the pathway of programmed cell death called apoptosis, the death of a single somatic cell as a result of DNA damage usually being less dangerous than allowing that cell to replicate its mutated DNA and possibly give rise to a tumor or other cancerous growth. In mammals, a central player in induction of cell cycle arrest and apoptosis is the protein called p53. This is classified as a tumor-suppressor protein because when this protein is defective, cells with damaged genomes can avoid the S-phase checkpoints and possibly proliferate into cancer. p53 is among the targets of both the ATM and ATR pathways. Once activated, it switches on transcription of a number of genes thought to be directly responsible for arrest and apoptosis, and represses expression of other genes that must be switched off to facilitate these processes.
SUMMARY

- In order to continue carrying out its function, the genome must replicate every time that a cell divides.

- Watson and Crick pointed out, when they first announced their discovery of the structure of DNA, that the specific base pairing that holds the two strands of the double helix together provides a means for accurate copying of each polynucleotide. They envisaged a semiconservative mode of replication in which each parent strand acts as a template for synthesis of a complementary daughter strand.

- The Meselson–Stahl experiment confirmed that DNA replicates by the semiconservative process, but there were still problems in understanding how the two strands of the helix were separated, especially in circular molecules that have little freedom to rotate.

- The discovery of DNA topoisomerases, which separate the strands of the double helix by repeated breakage and rejoining of one or both polynucleotides, solved the topological problem.

- No exceptions to the semiconservative mode of replication are known, though there are specialized versions such as displacement and rolling-circle replication.

- Initiation of genome replication occurs at discrete origins, which have been well characterized in bacteria and in yeast but are less clearly understood in higher eukaryotes.

- Once replication has been initiated, a pair of replication forks travel in opposite directions along the DNA.

- DNA polymerase can only synthesize DNA in the 5′→3′ direction, which means that although one strand, called the leading strand, can be replicated in a continuous fashion, the second, lagging strand has to be copied in short segments. These are called Okazaki fragments.

- DNA synthesis must be primed by an RNA polymerase.

- The replicating complex, called the replisome in bacteria, consists of the DNA polymerase enzyme along with ancillary proteins, such as the sliding clamp, which ensures that the connection between the polymerase and the DNA is secure but that the polymerase is still able to move along the DNA.

- Termination of replication occurs at specific regions within a bacterial chromosome but at less well-defined areas in eukaryotic chromosomes.

- Eukaryotic chromosomes require special processes to maintain their ends, as the replication process results in a gradual shortening of the telomeres. These are elongated by the telomerase enzyme, which has an RNA subunit that acts as the template for synthesis of new telomere repeat units.

- Replication of the genome must be coordinated with the cell cycle. This is achieved by a combination of regulatory proteins, many of which are active only at specific periods of the cell cycle.

- Assembly of the pre-replication complex at origins of replication is a critical step that is regulated to ensure that the genome is replicated just once per cell cycle.

- Once replication is underway, checkpoints during the synthesis phase respond to DNA damage in order to arrest or terminate genome replication.
SHORT ANSWER QUESTIONS

1. Prior to the Meselson–Stahl experiment it was not known if DNA replication is dispersive, semiconservative, or conservative. Describe the differences in the DNA contents of the daughter molecules resulting from these different modes of replication.

2. Outline the role of DNA topoisomerases in DNA replication.

3. Describe the mechanisms for (A) displacement replication and (B) rolling-circle replication.

4. Where and how does the DnaA protein bind at the origin of replication in *E. coli*?

5. What methods have been used to identify replication origins in eukaryotes?

6. List the key features of the bacterial and eukaryotic DNA polymerases that are involved in DNA replication.

7. How are Okazaki fragments joined together in (A) bacteria and (B) eukaryotes?

8. What is known about the termination of genome replication in *E. coli*? What proteins and sequences are involved in this process?

9. Why do the ends of linear chromosomes get shorter during successive rounds of DNA replication in eukaryotes?

10. How is telomerase activity regulated in eukaryotic cells?

11. Explain what is meant by the term origin licensing and describe the role of licensing in the cell cycle.

12. What general patterns have been observed regarding the timing of replication of different parts of the eukaryotic genome?

IN-DEPTH PROBLEMS

1. Discuss why the semiconservative mode of DNA replication was favored even before the Meselson–Stahl experiment was carried out.

2. Would it be possible to replicate the DNA molecules present in living cells if DNA topoisomerases did not exist?

3. Why is inactivation of the *Escherichia coli* polA gene, coding for DNA polymerase I, not lethal?

4. Construct a hypothesis to explain why all DNA polymerases require a primer in order to initiate synthesis of a new polynucleotide. Can your hypothesis be tested?

5. Our current knowledge of genome replication in eukaryotes is biased toward the events occurring at the replication fork. The next challenge is to convert this DNA-centered description of replication into a model that describes how replication is organized within the nucleus, addressing issues such as the role of replication factories and the processes used to avoid tangling of the daughter molecules. Devise a research plan to address one or more of these issues.
FURTHER READING

The history of research into genome replication


DNA topoisomerases


Termination of replication and the role of telomerase


Origins of replication


Control of genome replication


DNA polymerases and events at the replication fork


Termination of replication and the role of telomerase


MUTATIONS AND DNA REPAIR

Genomes are dynamic entities that change over time as a result of the cumulative effects of small-scale sequence alterations caused by mutation. A mutation is a change in the nucleotide sequence of a short region of a genome (Figure 16.1A). Many mutations are point mutations (also called simple mutations or single-site mutations) that replace one nucleotide with another. Point mutations are divided into two categories: transitions, which are purine-to-purine or pyrimidine-to-pyrimidine changes (A → G, G → A, C → T, or T → C), and transversions, which are purine-to-pyrimidine or pyrimidine-to-purine changes (A → C, A → T, G → C, G → T, C → A, C → G, T → A, or T → G). Other mutations arise from insertion or deletion of one or a few nucleotides.

Mutations result either from errors in DNA replication or from the damaging effects of mutagens, such as chemicals and radiation, which react with DNA and change the structures of individual nucleotides. All cells possess DNA repair enzymes that attempt to minimize the number of mutations that occur. These enzymes work in two ways. Some are prereplicative and search the DNA for nucleotides with unusual structures, these nucleotides being replaced before replication occurs; others are postreplicative and check newly synthesized DNA for errors, correcting any errors that they find (Figure 16.1B). A possible definition of mutation is therefore a deficiency in DNA repair.

Mutations can have dramatic effects on the cell in which they occur, a mutation in a key gene possibly resulting in a defective protein that could lead to death of the cell. Other mutations have a less significant impact on the phenotype of the cell, and many have none at all. As we will see in Chapter 18, all events that are not lethal have the potential to contribute to the evolution of the genome, but for this to happen, they must be inherited when the organism reproduces. With a single-celled organism such as a bacterium or yeast, all genome alterations that are not lethal or corrected are inherited by daughter cells and become permanent features of the lineage that descends from the original cell in which the alteration occurred. In a multicellular organism, only those events that occur in germ cells are relevant to genome evolution. Changes to the genomes of somatic cells are unimportant in an evolutionary sense, but they will have biological relevance if they result in a deleterious phenotype that affects the health of the organism.

In this chapter we will follow a logical progression, beginning with the causes of mutations and then moving on to the ways in which mutations are repaired.

16.1 THE CAUSES OF MUTATIONS

Mutations arise in two ways. Some mutations are spontaneous errors in replication that evade the proofreading function of the DNA polymerases that synthesize new polynucleotides at the replication fork (Section 15.3). These mutations are called mismatches because they are positions where the nucleotide that is inserted into the daughter polynucleotide does not match, by base pairing, the nucleotide at the corresponding position in the template DNA (Figure 16.2A). If the mismatch is not corrected in the daughter double helix, then one of the grand-daughter molecules produced during the next round of DNA replication will carry a permanent, double-stranded version of the mutation.

Other mutations arise because a mutagen has reacted with the parent DNA, causing a structural change that affects the base-pairing capability of the altered DNA molecule. A point mutation is shown but there are several other types of mutation, as described in the text. (B) DNA repair corrects mutations that arise as errors in replication and as a result of mutagenic activity.

Figure 16.1 Mutation and DNA repair.
(A) A mutation and DNA repair. A point mutation is shown but there are several other types of mutation, as described in the text. (B) DNA repair corrects mutations that arise as errors in replication and as a result of mutagenic activity.
Figure 16.2 Examples of mutations.
(A) An error in replication leads to a mismatch in one of the daughter double helices, in this case a T → C change because one of the As in the template DNA was miscopied. When the mismatched molecule is itself replicated, it gives one double helix with the correct sequence and one with a mutated sequence. (B) A mutagen has altered the structure of an A in the lower strand of the parent molecule, giving nucleotide X, which does not base-pair with the T in the other strand, so in effect, a mismatch has been created. When the parent molecule is replicated, X base-pairs with C, giving a mutated daughter molecule. When this daughter molecule is replicated, both granddaughters inherit the mutation.

Errors in replication are a source of point mutations
When considered purely as a chemical reaction, complementary base pairing is not particularly accurate. Nobody has yet devised a way of carrying out the template-dependent synthesis of DNA without the aid of enzymes, but if the process could be performed simply as a chemical reaction in a test tube then the resulting
polynucleotide would probably have point mutations at 5–10 positions out of every hundred. This represents an error rate of 5–10%, which would be completely unacceptable during genome replication. The template-dependent DNA polymerases that carry out DNA replication must therefore increase the accuracy of the process by several orders of magnitude. This improvement is brought about in two ways:

- A DNA polymerase operates a nucleotide selection process that dramatically increases the accuracy of template-dependent DNA synthesis. Exactly how this selection operates is not known, but it involves the nucleotide-binding part of the polymerase switching between an open and closed conformation, the closed conformation placing the selected nucleotide onto the template and checking that it base-pairs correctly. If the pairing is incorrect, then the nucleotide is rejected before it is attached to the 3’-end of the polynucleotide being synthesized.

- The accuracy of DNA synthesis is increased still further if the DNA polymerase has proofreading activity (Section 15.3), which means that it possesses a 3’ → 5’ exonuclease and so is able to remove an incorrect nucleotide that evades the nucleotide selection process and becomes attached to the 3’-end of the new polynucleotide (see Figure 2.7B).

*Escherichia coli* is able to synthesize DNA with an error rate of only 1 per 10⁷ nucleotide additions. Interestingly, these errors are not evenly distributed between the two daughter molecules, the lagging strand being replicated more accurately than the leading strand. The reason for this is not known but it probably relates to the efficiency with which errors in the two strands are repaired, rather than differences between the fidelity of DNA polymerase I, which is involved only in lagging-strand replication (Section 15.3), and that of DNA polymerase III, the main replicating enzyme.

Not all of the errors that occur during DNA synthesis can be blamed on the polymerase enzymes: sometimes an error occurs even though the enzyme adds the correct nucleotide, the one that base-pairs with the template. This is because each nucleotide base can occur as either of two alternative tautomers, structural isomers that are in dynamic equilibrium. For example, thymine exists as two tautomers, the keto and enol forms, with individual molecules occasionally undergoing a shift from one tautomer to the other. The equilibrium is biased very much toward the keto form, but every now and then the enol version of thymine occurs in the template DNA at the precise time that the replication fork is moving past. This will lead to an error, because the enol form of thymine base-pairs with G rather than A (Figure 16.3). The same problem can occur with adenine, where the rare imino tautomer of this base preferentially forms a pair with C, and with guanine, where the enol form pairs with thymine. After replication, the rare tautomer will inevitably revert to its more common form, leading to a mismatch in the daughter double helix.

As stated above, the error rate for DNA synthesis in *E. coli* is 1 in 10⁷. However, the overall error rate for replication of the *E. coli* genome is only 1 in 10¹⁰ to 1 in 10¹¹. The improvement compared with the polymerase error rate is the result of the mismatch repair system (Section 16.2) that scans newly replicated DNA for positions where the bases are unpaired and hence corrects the few mistakes that the replication enzymes make. This means that, on average, only one uncorrected replication error occurs every 2000 times that the *E. coli* genome is copied.

**Replication errors can also lead to insertion and deletion mutations**

Not all errors in replication are point mutations. Aberrant replication can also result in small numbers of extra nucleotides being inserted into the polynucleotide being synthesized or some nucleotides in the template not being copied. An insertion or deletion that occurs within a coding region might result in a frameshift mutation, which changes the reading frame used for translation of the
16.1 The Causes of Mutations

Chapter 16: Mutations and DNA Repair

There is a tendency to use the term frameshift to describe all insertions and deletions, but this is inaccurate, as inserting or deleting three nucleotides, or multiples of three, simply adds or removes codons or parts of adjacent codons without affecting the reading frame. Also, of course, many insertions and deletions occur outside of open reading frames, within introns or in the intergenic regions of a genome.

Insertion and deletion mutations can affect all parts of the genome but are particularly prevalent when the template DNA contains short repeated sequences, such as those found in microsatellites (Section 3.2). This is because repeated sequences can induce replication slippage, in which the template strand and its copy shift their relative positions so that part of the template is either copied twice or missed out. The result is that the new polynucleotide has a larger or smaller number, respectively, of the repeat units (Figure 16.5). This is the main reason why microsatellite sequences are so variable: replication slippage occasionally generates a new length variant, which can add to the collection of alleles already present in the population.

Figure 16.5 Replication slippage.
The diagram shows replication of a five-unit CA repeat microsatellite. Slippage has occurred during replication of the parent molecule, inserting an additional repeat unit into the newly synthesized polynucleotide of one of the daughter molecules. When this daughter molecule replicates, it gives a granddaughter molecule whose microsatellite array is one unit longer than that of the original parent.

protein specified by the gene (Figure 16.4). There is a tendency to use the term frameshift to describe all insertions and deletions, but this is inaccurate, as inserting or deleting three nucleotides, or multiples of three, simply adds or removes codons or parts of adjacent codons without affecting the reading frame. Also, of course, many insertions and deletions occur outside of open reading frames, within introns or in the intergenic regions of a genome.

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Replication slippage is probably also responsible, at least in part, for the trinucleotide repeat expansion diseases that have been discovered in humans in recent years. Each of these neurodegenerative diseases is caused by a relatively short series of trinucleotide repeats becoming elongated to two or more times...
16.1 The Causes of Mutations

For example, the human HTT gene contains the sequence 5’-CAG-3’ repeated between 6 and 29 times in tandem, coding for a series of glutamines in the protein product. In Huntington’s disease, this repeat expands to a copy number of 38–180, increasing the length of the polyglutamine tract and resulting in a dysfunctional protein. Several other human diseases are also caused by expansions of polyglutamine codons (Table 16.1). Some diseases associated with mental retardation result from trinucleotide expansions in the 5’-untranslated region of a gene, giving a fragile site, a position where the chromosome is likely to break. However, chromosome breakage is not the underlying cause of the disease. Instead, in both fragile X and fragile XE syndromes, the expansion results in aberrant methylation of a CpG island upstream of the affected gene, which is silenced, leading to loss of the protein product. Expansions involving intron and trailer regions are also known, most of these expansions affecting transcription of the gene or processing of the mRNA. An exception is the expansion in the 3’-untranslated region of the DMPK gene, which does not affect transcription or processing of the DMPK mRNA but does interfere with splicing of other RNAs. The explanation is not known, but it is possible that the expanded part of the DMPK mRNA either binds protein splicing factors, preventing the latter from acting on their normal target RNAs, or interferes in some way with the signaling pathways that control attachment of these factors to their targets. Finally, there are a few disease-causing mutations that involve expansions of longer sequences, such as progressive myoclonus epilepsy, caused by a \((\text{CTCGCGCGCGCG})_{2-3}\) to \((\text{CTCGCGCGCGCG})_{30+}\) expansion in the promoter region of the \(\text{EPM1}\) locus.

How triplet expansions are generated is not precisely understood. The size of the expansion is much greater than occurs with normal replication slippage, such as that seen with microsatellite sequences, and once the expansion reaches a certain length, it appears to become susceptible to further expansion in subsequent rounds of replication, so that the disease becomes increasingly severe in succeeding generations of an affected family. Expansions also occur between cell divisions, in cells that are not actively replicating their genomes. Initially, it was thought that expansion in nondividing cells might arise as a byproduct of double-strand break repair, but evidence to support this idea is lacking. It now seems likely that expansion can occur during excision repair, which involves removal of a segment of a polynucleotide containing a mutation, followed by resynthesis of the DNA strand to fill in the single-stranded gap that is formed (Section 16.2). If the

### Table 16.1 Examples of Human Trinucleotide Repeat Expansions

<table>
<thead>
<tr>
<th>Locus</th>
<th>Repeat sequence</th>
<th>Associated disease</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Polyglutamine expansions (all in coding regions of genes)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AR (exon 1)</td>
<td>(CAG)&lt;sub&gt;13-31&lt;/sub&gt;</td>
<td>(CAG)&lt;sub&gt;40&lt;/sub&gt;</td>
</tr>
<tr>
<td>ATN1 (exon 5)</td>
<td>(CAG)&lt;sub&gt;6-35&lt;/sub&gt;</td>
<td>(CAG)&lt;sub&gt;49-88&lt;/sub&gt;</td>
</tr>
<tr>
<td>ATXN1 (exon 8)</td>
<td>(CAG)&lt;sub&gt;6-39&lt;/sub&gt;</td>
<td>(CAG)&lt;sub&gt;41-83&lt;/sub&gt;</td>
</tr>
<tr>
<td>ATXN3 (exon 8)</td>
<td>(CAG)&lt;sub&gt;12-40&lt;/sub&gt;</td>
<td>(CAG)&lt;sub&gt;52-86&lt;/sub&gt;</td>
</tr>
<tr>
<td>HTT (exon 1)</td>
<td>(CAG)&lt;sub&gt;6-29&lt;/sub&gt;</td>
<td>(CAG)&lt;sub&gt;38-180&lt;/sub&gt;</td>
</tr>
<tr>
<td><strong>Fragile site expansions</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFF2 (5’-UTR)</td>
<td>(GCC)&lt;sub&gt;4-39&lt;/sub&gt;</td>
<td>(GCC)&lt;sub&gt;over 200&lt;/sub&gt;</td>
</tr>
<tr>
<td>FMR1 (5’-UTR)</td>
<td>(CGG)&lt;sub&gt;6-50&lt;/sub&gt;</td>
<td>(CGG)&lt;sub&gt;200-4000&lt;/sub&gt;</td>
</tr>
<tr>
<td><strong>Other expansions</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DMPK (3’-UTR)</td>
<td>(CTG)&lt;sub&gt;5-37&lt;/sub&gt;</td>
<td>(CTG)&lt;sub&gt;40-50&lt;/sub&gt;</td>
</tr>
<tr>
<td>FXN (intron 1)</td>
<td>(GAA)&lt;sub&gt;5-30&lt;/sub&gt;</td>
<td>(GAA)&lt;sub&gt;70-1000&lt;/sub&gt;</td>
</tr>
</tbody>
</table>

Abbreviation: UTR, untranslated region
excised segment includes part or all of a trinucleotide region, then slippage during the DNA synthesis step of the repair process could result in triplet expansion.

**Mutations are also caused by chemical and physical mutagens**

Many chemicals that occur naturally in the environment have mutagenic properties, and these have been supplemented in recent years with other chemical mutagens that result from human industrial activity. Physical agents such as radiation are also mutagenic. Most organisms are exposed to greater or lesser amounts of these various mutagens, and their genomes suffer damage as a result.

The definition of the term mutagen is a chemical or physical agent that causes mutations. This definition is important because it distinguishes mutagens from other types of environmental agent that cause damage to cells in ways other than by causing mutations (Table 16.2). There are overlaps between these categories (for example, some mutagens are also carcinogens), but each type of agent has a distinct biological effect. The definition of mutagen also makes a distinction between true mutagens and other agents that damage DNA without causing mutations, for example, by causing breaks in DNA molecules. This type of damage may block replication and cause the cell to die, but it is not a mutation in the strict sense of the term and the causative agents are therefore not mutagens.

Mutagens cause mutations in three different ways:

- Some mutagens act as **base analogs** and are mistakenly used as substrates when new DNA is synthesized at the replication fork.

- Other mutagens react directly with DNA, causing structural changes that lead to miscopying of the template strand when the DNA is replicated. These structural changes are diverse, as we will see when we look at individual mutagens.

- Some mutagens act indirectly on DNA. They do not themselves affect DNA structure, but instead they cause the cell to synthesize chemicals such as peroxides that have a direct mutagenic effect.

The range of mutagens is so vast that it is difficult to devise an all-embracing classification. We will therefore restrict our study to the most common types. For chemical mutagens, these are as follows:

- **Base analogs** are purine and pyrimidine bases that are similar enough to the standard bases of DNA to be incorporated into nucleotides when these are synthesized by the cell. The resulting unusual nucleotides can then be used as substrates for DNA synthesis during genome replication. For example, 5-bromouracil (5-bU; Figure 16.6A) has the same base-pairing properties as thymine, and nucleotides containing this base can be added to the daughter polynucleotide at positions opposite A in the template. The mutagenic effect arises because the equilibrium between the two tautomers of 5-bU is shifted more toward the rarer enol form than is the case with thymine. This means that during the next round of replication

<table>
<thead>
<tr>
<th>Agent</th>
<th>Effect on living cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carcinogen</td>
<td>Causes cancer, the neoplastic transformation of eukaryotic cells</td>
</tr>
<tr>
<td>Clastogen</td>
<td>Causes fragmentation of chromosomes</td>
</tr>
<tr>
<td>Mutagen</td>
<td>Causes mutations</td>
</tr>
<tr>
<td>Oncogen</td>
<td>Induces tumor formation</td>
</tr>
<tr>
<td>Teratogen</td>
<td>Results in developmental abnormalities</td>
</tr>
</tbody>
</table>
there is a relatively high chance of the polymerase encountering the enol form of 5-bU, which (like the enol form of thymine) pairs with G rather than A (Figure 16.6B). This results in a point mutation (Figure 16.6C).

2-Aminopurine acts in a similar way: it is an analog of adenine, with an amino tautomer that pairs with thymine and an imino tautomer that pairs with cytosine. The imino form of 2-aminopurine is more common than the imino form of adenine, so 2-aminopurine incorporation increases the frequency of T → C transitions.

- Deaminating agents also cause point mutations. A certain amount of base deamination (removal of an amino group) occurs spontaneously in genomic DNA molecules, and the rate is increased by chemicals such as nitrous acid, which is generated in the atmosphere from nitrogen and can accumulate in closed spaces such as poorly ventilated rooms. Nitrous acid deaminates adenine, cytosine, and guanine (thymine has no amino group and so cannot be deaminated). A second deaminating agent, sodium bisulfite, acts only on cytosine. Deamination of guanine is not mutagenic because the resulting base, xanthine, blocks replication when it appears in the template polynucleotide. Deamination of adenine gives hypoxanthine (Figure 16.7), which pairs with C rather than T, and deamination of cytosine gives uracil, which pairs with A rather than G. Deamination of these two bases therefore results in point mutations when the template strand is copied.

![Figure 16.6 5-Bromouracil and its mutagenic effect.](image)

![Figure 16.7 Hypoxanthine is a deaminated version of adenine. The nucleoside that contains hypoxanthine is called inosine.](image)
• **Alkylating agents** are a third type of mutagen that can give rise to point mutations. Chemicals such as ethylmethane sulfonate (EMS), dimethylnitrosamine, and methyl halides, the latter sometimes used as pesticides, add alkyl groups to nucleotides in DNA molecules. The effect of alkylation depends on the position at which the nucleotide is modified and the type of alkyl group that is added. Some methylations are not mutagenic, as they do not alter the base-pairing properties of the nucleotide. An example is the conversion of cytosine to 5-methylcytosine, which occurs extensively in vertebrate and plant genomes and is associated with gene silencing (Section 10.3). Other methylations do alter the base-pairing properties of the modified nucleotide and so lead to point mutations. There are also alkylations that block replication by forming cross-links between the two strands of a DNA molecule or simply by impeding progress of the replisome.

• **Intercalating agents** are usually associated with insertion mutations. The best-known mutagen of this type is ethidium bromide, which fluoresces when exposed to ultraviolet (UV) radiation and has been used to reveal the positions of DNA bands after agarose gel electrophoresis. Ethidium bromide and other intercalating agents are flat molecules that can slip between base pairs in the double helix, slightly unwinding the helix and hence increasing the distance between adjacent base pairs (Figure 16.8).

Now we move on to the most important types of physical mutagens, which are as follows:

• Ultraviolet radiation of wavelength 260 nm induces dimerization of adjacent pyrimidine bases, especially if these are both thymines (Figure 16.9A),
resulting in a **cyclobutyl dimer**. Other adjacent pyrimidine combinations also form dimers; the order of frequency with which this occurs is 5′-CT-3′ > 5′-TC-3′ > 5′-CC-3′. Purine dimers are much less common. UV-induced dimerization usually results in a deletion mutation when the modified strand is copied. Another type of UV-induced **photoproduct** is the *(6-4) lesion*, in which carbon 6 of one pyrimidine and carbon 4 of the adjacent pyrimidine become covalently linked (**Figure 16.9B**).

- Ionizing radiation has various effects on DNA depending on the type of radiation and its intensity. Point, insertion, and/or deletion mutations might arise, as well as more severe forms of DNA damage that prevent subsequent replication of the genome. Some types of ionizing radiation act directly on DNA, while others act indirectly by stimulating the formation of reactive molecules such as peroxides.

- Heat stimulates the water-induced cleavage of the β-N-glycosidic bond that attaches the base to the sugar component of the nucleotide (**Figure 16.10A**). This occurs more frequently with purines than with pyrimidines and results in an AP *(apurinic/apyrimidinic)* site, also called a **baseless site**. The sugar–phosphate that is left is unstable and rapidly degrades, leaving a gap if the DNA molecule is double-stranded (**Figure 16.10B**). This reaction is not normally mutagenic because cells have effective systems for repairing gaps (**Section 16.2**), which is reassuring when one considers that 10,000 AP sites are generated in each human cell per day.

**Figure 16.10** Mutagenic effect of heat. (A) Heat induces hydrolysis of a β-N-glycosidic bond, resulting in a baseless site in a polynucleotide. (B) Schematic representation of the effect of heat-induced hydrolysis on a double-stranded DNA molecule. The baseless site is unstable and degrades, leaving a gap in one strand.
16.2 REPAIR OF MUTATIONS AND OTHER TYPES OF DNA DAMAGE

Having studied the myriad ways in which the genome can become mutated, it is always reassuring to move quickly on to the equally myriad ways in which the genome can be repaired. In view of the thousands of damage events that genomes suffer every day, coupled with the errors that occur when the genome replicates, it is essential that cells possess efficient repair systems. Without these repair systems, a genome would not be able to maintain its essential cellular functions for more than a few hours before key genes became inactivated by DNA damage. Similarly, cell lineages would accumulate replication errors at such a rate that their genomes would become dysfunctional after a few cell divisions.

Most cells possess four different categories of DNA repair system (Figure 16.11):

- **Direct repair** systems, as the name suggests, act directly on damaged nucleotides, converting each one back to its original structure.

- **Excision repair** involves excision of a segment of the polynucleotide containing a damaged site, followed by resynthesis of the correct nucleotide sequence by a DNA polymerase.

- **Mismatch repair** corrects errors of replication, again by excising a stretch of single-stranded DNA containing the offending nucleotide and then repairing the resulting gap.

- **Break repair** is used to mend single- and double-strand breaks.

**Direct repair systems fill in nicks and correct some types of nucleotide modification**

Most types of DNA damage that are caused by chemical or physical mutagens can only be repaired by excision of the damaged nucleotide followed by resynthesis.

![Figure 16.11 Four categories of DNA repair system.](image-url)
of a new stretch of DNA, as shown in Figure 16.11B. Only a few types of damaged nucleotide can be repaired directly. These include nicks, which can be repaired by a DNA ligase if all that has happened is that a phosphodiester bond has been broken, without damage to the 5’-phosphate and 3’-hydroxyl groups of the nucleotides on either side of the nick (Figure 16.12). This is often the case with nicks resulting from the effects of ionizing radiation.

Some forms of alkylation damage are also directly reversible. This type of repair is carried out by enzymes that transfer the alkyl group from the nucleotide to their own polypeptide chains. Enzymes capable of doing this are known in many different organisms and include the Ada enzyme of E. coli, which is involved in an adaptive process that this bacterium is able to activate in response to DNA damage. Ada removes alkyl groups attached to the oxygen atoms at positions 4 and 6 of thymine and guanine, respectively, and can also repair phosphodiester bonds that have become alkylated. The transfer is irreversible, which means that Ada is an example of a suicide enzyme, an enzyme that is inactivated once it has carried out its biochemical reaction. However, the alkylated version of Ada now acts as a transcription factor that switches on the ada regulon, which includes genes for the Ada enzyme and three other repair proteins (Figure 16.13). Activation of these genes gives rise to the adaptive response. Eukaryotes also have alkylation repair enzymes, an example being human MGMT (O6-methylguanine-DNA methyltransferase) which, as its name suggests, removes alkyl groups from position 6 of guanine. The eukaryotic enzymes do not appear to affect transcription and are simply degraded once they have become alkylated.

The final type of direct repair system involves cyclobutyl dimers, which can be cleaved by a light-dependent direct system called photoreactivation. In E. coli, the process involves the enzyme called DNA photolyase (more correctly named deoxyribodipyrimidine photolyase). When stimulated by light with a wavelength between 300 and 500 nm, the enzyme binds to cyclobutyl dimers and converts them back to the original monomeric nucleotides. There are two types of photolyase, one type containing a folate cofactor and the other containing a flavin compound. Both types of cofactor capture light energy, which is used to transfer an electron to the cyclobutyl dimer, causing the latter to split into its monomers. Photoreactivation is a widespread but not universal type of repair: it is known in many but not all bacteria and also in quite a few eukaryotes, including some vertebrates, but it appears to be absent in humans and other placental mammals. A similar type of photoreactivation involves the (6–4) photoproduct photolyase and results in repair of (6–4) lesions. Neither E. coli nor humans have this enzyme, but it is possessed by a variety of other organisms.

**Base excision repairs many types of damaged nucleotide**

The direct types of damage reversal are important, but they form a very minor component of the DNA repair mechanisms of most organisms. Excision repair systems are much more prevalent and are able to correct a much broader range of mutations.

**Base excision**, which is the least complex of this type of repair system, is used to repair many modified nucleotides whose bases have suffered relatively minor damage resulting from, for example, exposure to alkylating agents or ionizing radiation. There are several versions of base excision repair, the main distinction being between short-patch repair, in which only the damaged nucleotide is replaced, and long-patch repair, which involves removal and resynthesis of up to 10 nucleotides, including the damaged one. The pathways are initiated by a DNA glycosylase, which cleaves the β-N-glycosidic bond between the damaged base and the sugar component of the nucleotide. This reaction has the same effect as the heat-induced creation of an AP site (see Figure 16.10A). Each DNA glycosylase has a limited specificity (Table 16.3), so the specificities of the glycosylases possessed by a cell determine the range of damaged nucleotides that can be repaired by this pathway. Most organisms are able to deal with deaminated bases such as uracil (deaminated cytosine) and hypoxanthine (deaminated adenine), oxidation

Figure 16.12 Repair of a nick by DNA ligase.

**Figure 16.13 Adaptive response stimulated by the Ada enzyme of E. coli.** Transfer of an alkyl group from a nucleotide to the Ada enzyme converts the latter into a transcription factor that activates the ada regulon. A regulon is a group of genes that are controlled by the same transcription factor or group of factors but are not linked together into a single operon. The ada gene codes for the Ada enzyme. The other genes in the regulon are alkA, which specifies a DNA glycosylase involved in base excision repair; alkB, which codes for a member of a different family of alkylation repair enzymes; and aidB, whose function is unknown.
products such as 5-hydroxycytosine and thymine glycol, and methylated bases such as 3-methyladenine and 7-methylguanine. Most of the DNA glycosylases involved in base excision repair are thought to diffuse along the minor groove of the DNA double helix in search of damaged nucleotides, but some may be associated with the replication enzymes.

A DNA glycosylase removes a damaged base by flipping the structure to a position outside the helix and then detaching it from the polynucleotide. This creates an AP (baseless) site. In the short-patch repair pathway, there are two ways in which this AP site can be converted into a single-nucleotide gap:

- An **AP endonuclease**, such as exonuclease III or endonuclease IV of *E. coli* or APE1 of humans, cuts the phosphodiester bond on the 5′-side of the AP site (Figure 16.14A). Some AP endonucleases might also remove the ribose from the AP site, as this is all that remains of the damaged nucleotide, but others lack this ability and so work in conjunction with a separate **phosphodiesterase** enzyme, which cleaves the phosphodiester bond between the sugar and the next nucleotide. Alternatively, in eukaryotes the ribose can be removed by a **lyase** activity, possessed by DNA polymerase β.

### Table 16.3 Examples of Mammalian DNA Glycosylases

<table>
<thead>
<tr>
<th>DNA glycosylase</th>
<th>Specific for</th>
</tr>
</thead>
<tbody>
<tr>
<td>MBD4</td>
<td>Uracil</td>
</tr>
<tr>
<td>MPG</td>
<td>Ethenoadenine, hypoxanthine, 3-methyladenine, 3-methylguanine, 7-methylguanine</td>
</tr>
<tr>
<td>NEIL1</td>
<td>Thymine glycol, formamidopyrimidine, 8-oxoguanine, 5-hydroxuracil, dihydroxuracil</td>
</tr>
<tr>
<td>NTHL1</td>
<td>5-Hydroxycytosine, 5-hydroxyuracil, formamidopyrimidine, thymine glycol</td>
</tr>
<tr>
<td>OGG1</td>
<td>Formamidopyrimidine, 8-oxoguanine</td>
</tr>
<tr>
<td>SMUG1</td>
<td>Uracil</td>
</tr>
<tr>
<td>UNG</td>
<td>Uracil, 5-hydroxyuracil</td>
</tr>
</tbody>
</table>

**Figure 16.14** Base excision repair. After removal of the damaged base by DNA glycosylase, the base excision repair process can follow one of three pathways. (A) An AP endonuclease cuts the phosphodiester bond at the 5′-side of the AP site. The ribose is then removed by the AP endonuclease, by a phosphodiesterase, or by the lyase activity of DNA polymerase β. (B) In the alternative short-patch pathway, the initial cut is made by a bifunctional DNA glycosylase, to the 3′-side of the ribose. The latter is then removed by a phosphodiesterase, the lyase activity of DNA polymerase β, or the 5′ → 3′ exonuclease activity of DNA polymerase I. (C) The long-patch pathway begins with a 5′ cut made by an AP endonuclease. The standard process for lagging-strand replication is then used to remove and resynthesize a strand of 2–10 nucleotides, the first of which is the AP site.
• If the DNA glycosylase is bifunctional, then it can make a cut at the 3'-side of the AP site, probably at the same time that it removes the damaged base. This cut is followed again by excision of the ribose by a phosphodiesterase, by DNA polymerase β, or in *E. coli* by the 5' → 3' exonuclease activity of DNA polymerase I (Figure 16.14B).

Whichever method is used, the resulting single-nucleotide gap is filled in by a DNA polymerase, using base pairing with the undamaged base in the other strand of the DNA molecule to ensure that the correct nucleotide is inserted. In *E. coli* the gap is filled by DNA polymerase I, and in mammals it is filled by DNA polymerase δ. After gap filling, the final phosphodiester bond is put in place by a DNA ligase.

In the long-patch pathway, the AP site created by DNA glycosylase activity is cut on the 5'-side by an AP endonuclease, but this is not followed by further cleavages. Instead, a process similar to lagging-strand replication (Section 15.3) is used to synthesize 2–10 new nucleotides of DNA, with the first nucleotide replacing the damaged site (Figure 16.14C). In bacteria, the synthesis is carried out by DNA polymerase I, which uses its 5' → 3' exonuclease activity to degrade the segment of polynucleotide that is replaced. In eukaryotes, DNA polymerase δ or ε displaces this segment as the new polynucleotide is made, and the resulting flap is cleaved by FEN1 endonuclease. Short-patch repair appears to be the main base excision pathway, the choice between the two pathways being influenced at least in part by the identity of the DNA glycosylase that initiates the process.

**Nucleotide excision repair is used to correct more extensive types of damage**

**Nucleotide excision repair** has a much broader specificity than the base excision system and is able to deal with more extreme forms of damage, such as intrastrand cross-links and bases that have become modified by attachment of large chemical groups. It is also able to correct cyclobutyl dimers by a dark repair process, providing those organisms that lack the photoreactivation system (such as humans) with a means of repairing this type of damage.

In nucleotide excision repair, a segment of single-stranded DNA containing the damaged nucleotide(s) is excised and replaced with new DNA. The process is therefore similar to base excision repair, except that it is not preceded by selective base removal and a longer stretch of polynucleotide is cut out. The best-studied example of nucleotide excision repair is the short-patch process of *E. coli*, so-called because the region of polynucleotide that is excised and subsequently patched is relatively short, usually 12 nucleotides in length. The use of the same terminology for base and nucleotide excision repair is unfortunate, especially as the short-patch version of nucleotide excision repair involves replacement of a segment that is similar in length to that removed during long-patch base excision repair.

Short-patch repair is initiated by a multienzyme complex called the **UvrABC endonuclease**, sometimes also referred to as the excinuclease. In the first stage of the process, a trimer comprising two UvrA proteins and one copy of UvrB attaches to the DNA at the damaged site. How the site is recognized is not known, but the broad specificity of the process indicates that individual types of damage are not directly detected and that the complex must search for a more general attribute of DNA damage such as distortion of the double helix. UvrA may be the component of the complex most involved in damage location because it dissociates once the site has been found and plays no further part in the repair process. Departure of UvrA allows UvrC to bind (Figure 16.15), forming a UvrBC dimer that cuts the polynucleotide either side of the damaged site. The first cut is made by UvrB at the fifth phosphodiester bond downstream of the damaged nucleotide, and the second cut is made by UvrC at the eighth phosphodiester bond upstream, resulting in the 12-nucleotide excision, although there is some variability, especially in the position of the UvrB cut site. The excised segment is then removed, usually as an intact oligonucleotide, by DNA helicase II (sometimes called UvrD), which...
presumably detaches the segment by breaking the base pairs holding it to the second strand. UvrC also detaches at this stage, but UvrB remains in place and bridges the gap produced by the excision. The bound UvrB is thought to prevent the single-stranded region that has been exposed from base-pairing with itself, but the role of UvrB could be to prevent this strand from becoming damaged, or possibly to direct the DNA polymerase to the site that needs to be repaired. As in base excision repair, the gap is filled in by DNA polymerase I and the last phosphodiester bond is synthesized by DNA ligase. E. coli also has a long-patch nucleotide excision repair system that involves Uvr proteins but differs in that the piece of DNA that is excised can be up to 2 kb in length. Long-patch repair has been less well studied and the process is not understood in detail, but it is presumed to work on more extensive forms of damage, possibly regions where groups of nucleotides, rather than just individual ones, have become modified.

In eukaryotes, there is a single nucleotide excision repair process that results in replacement of 24–32 nucleotides of DNA. The system is more complex than in E. coli, and the relevant proteins do not seem to be homologs of the Uvr family. In humans, at least 16 proteins are involved, several of these forming a complex that surveys the DNA for damage. The damaged site is then excised, with the downstream cut typically being made at the same position as in E. coli—the fifth phosphodiester bond—but with a more distant upstream cut, 22 nucleotides away, resulting in the longer excision. There is some variability in the cut positions, accounting for the range of fragment lengths that are replaced. Both cuts are made by endonucleases that attack single-stranded DNA specifically at its junction with a double-stranded region, indicating that before the cuts are made, the DNA around the damage site has been melted (Figure 16.16). The helicase activity for strand melting is provided by the XPB and XPD proteins, which are associated with transcription factor IIH (TFIIH), which stabilizes the resulting single-stranded region prior to its excision. The gap is then filled by DNA polymerase δ or ε. TFIIH is also one of the components of the RNA polymerase II initiation complex (Section 12.2). At first it was assumed that TFIIH simply had a dual role in the cell, functioning separately in both transcription and repair, but now it is thought that there is a more direct link between the two processes. This view is supported by the discovery of transcription-coupled repair, which repairs some forms of damage in the template strands of genes that are being actively transcribed. The only difference between transcription-coupled repair and the global nucleotide excision pathway described above is that the damage repaired by the transcription-coupled process is not detected by surveillance proteins but instead is signaled by the RNA polymerase stalling as it approaches the damaged site.

**Mismatch repair corrects replication errors**

Each repair system that we have looked at so far—direct, base excision, and nucleotide excision repair—recognizes and acts upon DNA damage caused by mutagens. These systems search for abnormal chemical structures such as modified nucleotides, cyclobutyl dimers, and helix distortions. They cannot, however, correct mismatches resulting from errors in replication because the mismatched nucleotide is not abnormal in any way, it is simply an A, C, G, or T that has been inserted at the wrong position. As these nucleotides look exactly like any other nucleotide, the mismatch repair system that corrects replication errors has to detect not the mismatched nucleotide itself but the absence of base pairing between the parent and daughter strands. Once it has found a non-base-paired position, the repair system excises part of the daughter polynucleotide and fills in the gap, in a manner similar to base and nucleotide excision repair.

The scheme described above leaves one important question unanswered. The repair must be made in the daughter polynucleotide because it is in this newly synthesized strand that the error has occurred; the parent polynucleotide has the correct sequence. How does the repair process know which strand is which? In E. coli the answer is that the daughter strand, at this stage, is undermethylated and can therefore be distinguished from the parent polynucleotide, which has a
full complement of methyl groups. *E. coli* DNA is methylated because of the activities of the **DNA adenine methylase** (*Dam*), which converts adenines to 6-methyl-adenines in the sequence 5′-GATC-3′, and the **DNA cytosine methylase** (*Dcm*), which converts cytosines to 5-methylcytosines in the sequences 5′-CCAGG-3′ and 5′-CCTGG-3′. These methylations are not mutagenic: the modified nucleotides have the same base-pairing properties as the unmodified versions. There is a delay between DNA replication and methylation of the daughter strand, and it is during this window of opportunity that the repair system scans the DNA for mismatches and makes the required corrections in the undermethylated daughter strand (Figure 16.17). *E. coli* has at least three mismatch repair systems, called long patch, short patch, and very short patch, the names once again indicating the relative lengths of the excised and resynthesized segments of DNA. The long-patch system replaces up to a kilobase or more of DNA and requires the *MutH*, *MutL*, and *MutS* proteins, as well as DNA helicase II, which we met during nucleotide excision repair. *MutS* recognizes the mismatch and *MutH* distinguishes the two strands by binding to unmethylated 5′-GATC-3′ sequences (Figure 16.18). The role of *MutL* is to coordinate the activities of the other two proteins so that *MutH* binds to unmethylated 5′-GATC-3′ sequences only in the vicinity of mismatch sites recognized by *MutS*. After binding, *MutH* cuts the phosphodiester bond immediately upstream of the G in the methylation sequence, and DNA helicase II detaches the single strand. There does not appear to be an enzyme that cuts the strand downstream of the mismatch; instead the detached single-stranded region is degraded by an exonuclease that follows the helicase and continues beyond the mismatch site. The gap is then filled in by DNA polymerase III and DNA ligase. Similar events are thought to occur during short-patch and very-short-patch mismatch repair, the difference being the specificities of the proteins that recognize the mismatch. The short-patch system, which results in excision of a segment less than 10 nucleotides in length, begins when *MutY* recognizes an A-G or A-C mismatch, and the very-short-patch repair system corrects G-T mismatches, which are recognized by the Vsr endonuclease.

Eukaryotes have homologs of the *E. coli* *MutS* and *MutL* proteins and their mismatch repair processes probably work in a similar way. The one difference is the absence of a homolog of *MutH*, which suggests that methylation might not be the method used to distinguish between the parent and daughter polynucleotides. Methylation has been implicated in mismatch repair in mammalian cells, but the DNA of some eukaryotes, including fruit flies and yeast, is not extensively methylated. It is therefore thought that in these organisms a different method must be used to identify the daughter strand. The most likely explanation is that the repair enzymes associate with the replisome in such a way that the parent and daughter strands can be distinguished as the latter are synthesized.

**Single- and double-strand breaks can be repaired**

A single-strand break in a double-stranded DNA molecule, as produced by some types of oxidative damage, does not present the cell with a critical problem as the double helix retains its overall intactness. The exposed single strand is coated with PARP1 proteins, which protect this intact strand from breaking and prevent it from participating in unwanted recombination events. The break is then filled in by the
enzymes involved in the excision repair pathways (Figure 16.19). Single-strand DNA breaks can also be repaired during replication, by a process that involves homologous recombination. To understand this version of DNA repair, we first need to study the molecular basis of recombination. We will therefore return to recombination repair in Chapter 17.

A double-strand break is more serious because this converts the original double helix into two separate fragments that have to be brought back together again in order for the break to be repaired. The two broken ends must be protected from further degradation, which could result in a deletion mutation appearing at the repaired break point. The repair processes must also ensure that the correct ends are joined: if there are two broken chromosomes in the nucleus, then the correct pairs must be brought together so that the original structures are restored. Experimental studies of mouse cells indicate that achieving this outcome is difficult, and if two chromosomes are broken, then misrepair resulting in hybrid structures occurs relatively frequently. Even if only one chromosome is broken, there is still a possibility that a natural chromosome end could be confused with a break and an incorrect repair might be made. This type of error is not unknown, despite the presence of shelterin proteins that mark the natural ends of chromosomes (Section 7.1).

Double-strand breaks are generated by exposure to ionizing radiation and some chemical mutagens, and breakages can also occur during DNA replication. They can be repaired by a system called nonhomologous end-joining (NHEJ). Progress in understanding NHEJ has been stimulated by studies of mutant human cell lines, which have resulted in the identification of various sets of genes involved in the process. These genes specify a multicomponent protein complex that directs a DNA ligase to the break (Figure 16.20A). The complex includes two copies of the Ku protein, one copy attaching to each broken DNA end. Ku proteins can only bind to cut ends, not to the internal regions of a DNA molecule, because the DNA molecule must fit into a loop formed by the association between the two subunits that make up each Ku protein (Figure 16.20B). Individual Ku proteins have an affinity for one another, which means that the two broken ends of the DNA molecule are brought into proximity. Ku binds to the DNA in association with the DNA-PKcs protein kinase, which activates a
third protein, XRCC4, that interacts with the mammalian DNA ligase IV, directing this repair protein to the double-strand break. NHEJ was originally thought to be restricted to eukaryotes, but genome annotations have uncovered bacterial homologs of the mammalian Ku proteins, and experimental studies have indicated that these act in conjunction with bacterial ligases in a simplified version of the double-strand break repair process.

**If necessary, DNA damage can be bypassed during genome replication**

If a region of the genome has suffered extensive damage, then it is conceivable that the repair processes will be overwhelmed. The cell then faces a stark choice between dying or attempting to replicate the damaged region, even though this replication may be error-prone and result in mutated daughter molecules. When faced with this choice, *E. coli* cells invariably take the second option, inducing one of several emergency procedures for bypassing sites of major damage.

The best studied of these bypass processes occurs as part of the SOS response, which enables an *E. coli* cell to replicate its DNA even though the template polynucleotides contain AP sites and/or cyclobutyl dimers and other photoproducts, resulting from exposure to chemical mutagens or UV radiation, that would normally block, or at least delay, the replication complex. Bypass of these sites requires construction of a mutasome, comprising DNA polymerase V (also called the UmuD′C complex, because it is a trimer made up of two UmuD′ proteins and one UmuC) and several copies of the RecA protein. RecA is primarily a single-strand binding protein that plays many roles in DNA repair and recombination. In this bypass system, RecA coats the damaged polynucleotide strands, enabling DNA polymerase V to displace DNA polymerase III and carry out error-prone DNA synthesis until the damaged region has been passed and DNA polymerase III can take over once again (Figure 16.21). DNA polymerase V is therefore an example of a translesion polymerase.

As well as acting as a single-strand binding protein that facilitates the mutasome bypass process, RecA also has a second function as an activator of the overall SOS response. The protein is stimulated by chemical signals (not yet identified) that indicate the presence of extensive DNA damage. In response, RecA cleaves a number of target proteins, directly or indirectly, including UmuD. RecA cleavage converts UmuD into its active form, UmuD′, and initiates the mutasome repair process. RecA also cleaves a repressor protein called LexA, switching on or increasing the expression of a number of genes normally repressed by LexA, including both the recA gene itself (leading to a 50-fold increase in RecA synthesis) and several other genes whose products are involved in DNA repair pathways. RecA also cleaves the cl repressor of λ bacteriophage, so if an integrated λ prophage is present in the genome, it can excise and leave the sinking ship (Section 14.3).

The SOS response is primarily looked on as the last best chance for a bacterium to replicate its DNA and hence survive under adverse conditions. However, the price of survival is an increased mutation rate because the mutasome does not repair damage, it simply allows a damaged region of a polynucleotide to be replicated. When it encounters a damaged position in the template DNA, the polymerase selects a nucleotide more or less at random, although with some preference for placing an A opposite an AP site: in effect, the error rate of the replication process increases. It has been suggested that this increased mutation rate is the purpose of the SOS response, mutation being in some way an advantageous response to extensive DNA damage, but this idea remains controversial.

For some time, the SOS response was thought to be the only damage-bypass process in bacteria, but we now appreciate that at least two other *E. coli* translesion polymerases act in a similar way, although with different types of damage. These are DNA polymerase II, which can bypass AP sites and nucleotide bases bound to mutagenic chemicals, such as deoxyguanine bound to 2-acetylaminofluorene, and DNA polymerase IV (also called DinB), which can replicate through a region of template DNA in which the two parent polynucleotides have become cross-linked. Translesion polymerases have also been discovered in eukaryotic cells.
cells. These include DNA polymerase η, which can bypass cyclobutyl dimers, and DNA polymerases ι and ζ, whose functions are less well understood.

**Defects in DNA repair underlie human diseases, including cancers**

The importance of DNA repair is emphasized by the number and severity of inherited human diseases that have been linked with defects in one of the repair processes. One of the best characterized of these is xeroderma pigmentosum, which results from a mutation in any one of several genes for proteins involved in nucleotide excision repair. Nucleotide excision is the only way in which human cells can repair cyclobutyl dimers and other photoproducts, so it is no surprise that the symptoms of xeroderma pigmentosum include hypersensitivity to UV radiation, patients suffering more mutations than normal upon exposure to sunlight and often developing skin cancer. Trichothiodystrophy is also caused by defects in nucleotide excision repair, but this is a more complex disorder that, although not involving cancer, usually includes problems with both the skin and nervous system.

A few diseases have been linked with defects in the transcription-coupled component of nucleotide excision repair. These include breast and ovarian cancers—the BRCA1 gene that confers susceptibility to these cancers codes for a protein that has been implicated in transcription-coupled repair—and Cockayne syndrome, a complex disease manifested by growth and neurologic disorders. Ataxia telangiectasia, the symptoms of which include sensitivity to ionizing radiation, results from defects in the ATM gene, which is involved in the damage-detection process. Other diseases that are associated with a breakdown in DNA repair are: Bloom's and Werner's syndromes, which are caused by inactivation of the RecQ DNA helicase, which has a role in NHEJ; the cancer-susceptibility syndrome called hereditary nonpolyposis colorectal cancer, which results from a defect in mismatch repair; and some types of spinocerebellar ataxia, which arise from defects in the pathway used to repair single-strand breaks. Understanding the genetic background to these disorders is important not just for devising therapies for managing the disease. The disorders can also provide novel information on the biochemical basis to DNA repair. An example is provided by Fanconi anemia, a rare disease first characterized in 1927 and with a frequency of only one case per 100,000 individuals. Patients suffering from this type of anemia have an increased susceptibility to chemicals that cause DNA cross-links. Recognition that the disease can arise as a result of mutations in at least 16 different genes has enabled the pathway that repairs this type of DNA damage to be elucidated.

**SUMMARY**

- A mutation is a change in the nucleotide sequence of a DNA molecule.
- A point mutation affects just a single nucleotide. Mutations can also arise by insertion or deletion of one or more adjacent nucleotides.
- Mutations can result from errors made during DNA replication. DNA polymerases have nucleotide selection and proofreading activities that maintain a high degree of accuracy, but these checking mechanisms can be evaded if an unusual tautomeric version of a nucleotide is present in the template.
- A second type of replication error, called slippage, can lead to insertion or deletion mutations.
- There are also many types of chemical or physical agent that can cause mutations.
- Some compounds act as base analogs and cause mutations after being mistaken for genuine nucleotides by the replication machinery.
• Deaminating and alkylating agents directly attack DNA molecules, and intercalating agents such as ethidium bromide slide between base pairs, causing insertions and deletions when the helix is replicated.

• UV radiation causes adjacent nucleotides to link together into dimers, and ionizing radiation and heat cause various types of damage.

• All cells possess DNA repair processes that enable many mutations to be corrected.

• Direct repair systems are uncommon but the few that are known correct some types of base damage including removal of UV-induced nucleotide dimers.

• Excision repair processes involve excision of a segment of a polynucleotide containing a damaged site, followed by resynthesis of the correct nucleotide sequence by a DNA polymerase.

• Mismatch repair corrects errors of replication, again by excising a stretch of single-stranded DNA containing the mutation and repairing the resulting gap.

• Nonhomologous end-joining is used to mend double-strand breaks.

• There are also processes for bypassing sites of DNA damage during replication, many of these acting as an emergency system for rescuing a genome that has become heavily mutated.

• Defects in DNA repair often result in disease, including several types of cancer.

**SHORT ANSWER QUESTIONS**

1. Describe how the mode of action of a DNA polymerase maximizes the accuracy with which that enzyme replicates DNA.

2. Outline the way in which tautomers induce errors in replication.

3. Explain what is meant by the term “replication slippage.”

4. How does the base analog 2-aminopurine produce mutations in DNA?

5. Describe the effect of UV irradiation on DNA structure.

6. How does heat affect the structure of DNA?

7. Describe the direct mutation repair processes that are known in bacterial and eukaryotic cells.

8. Outline the steps that occur during the base excision repair pathway.

9. Describe the key features of the nucleotide excision repair pathways of *E. coli*.

10. How are the parent and daughter strands distinguished during the mismatch repair process of *E. coli*?

11. What is the process by which double-strand breaks in DNA are repaired by the nonhomologous end-joining system?

12. What is the role of the RecA protein in the SOS response of *E. coli*?

**IN-DEPTH PROBLEMS**

1. Explain why a purine-to-purine or pyrimidine-to-pyrimidine point mutation is called a transition, whereas a purine-to-pyrimidine (or vice versa) change is called a transversion.
Chapter 16: Mutations and DNA Repair

2. What would be the anticipated ratio of transitions to transversions in a large number of mutations?

3. A mutation in a protein-coding gene can result in an alteration in the amino acid sequence of the protein product, possibly modifying or inactivating the function of that protein. An individual who inherits a mutated gene from one of their parents might therefore suffer from a genetic disease. However, not all such diseases have an immediate impact. Some are delayed onset and are only expressed later in the individual’s life. Others display nonpenetrance in some individuals, never being expressed. Devise mechanisms to explain how mutations can exhibit delayed onset or nonpenetration.

4. The bacterium *Deinococcus radiodurans* is highly resistant to radiation and to other physical and chemical mutagens. Discuss how these special properties of *D. radiodurans* might be reflected in its genome sequence.

5. Why do defects in DNA repair often lead to cancer?

**FURTHER READING**

**Causes of mutations**


**Trinucleotide repeat expansion diseases**


**Mismatch repair**


**Repair of DNA breaks**


Bypassing DNA damage

Repair and disease
**Recombination** is the term originally used by geneticists to describe the outcome of crossing over between pairs of homologous chromosomes during meiosis. Crossing over results in daughter chromosomes that have different combinations of alleles compared with their parent chromosomes (Section 3.3). In the 1960s, models were proposed for the molecular events that underlie crossing over, and it was realized that a key part of molecular recombination is the breakage and subsequent rejoining of DNA molecules. Biologists now use recombination to refer to a variety of processes that involve the breakage and reunion of polynucleotides. These include the following:

- **Homologous recombination**, also called **general** (or generalized) **recombination**, occurs between segments of DNA molecules that share extensive sequence homology. These segments might be present on different chromosomes or might be two parts of a single chromosome (Figure 17.1A). Homologous recombination is responsible for crossing over during meiosis and was initially studied in this context, but we now believe that its primary cellular role is in DNA repair.

- **Site-specific recombination** occurs between DNA molecules that have only short regions of sequence similarity, possibly just a few base pairs (Figure 17.1B). Site-specific recombination is responsible for the insertion of phage genomes, such as that of λ, into bacterial chromosomes.

![Figure 17.1 Two different types of recombination event.](image)
17.1 Homologous Recombination

The study of homologous recombination has presented two significant challenges for molecular biologists, neither of which has yet been fully met. The first challenge has been to describe the series of interactions, involving breakage and reunion of polynucleotides, that occur during recombination. The models for homologous recombination that have resulted from this work are described below. The second challenge relates to the fact that recombination is a cellular process that, like other cellular processes involving DNA (such as transcription and replication), is carried out and regulated by enzymes and other proteins. Biochemical studies have defined a series of related recombination pathways and have also revealed that homologous recombination underlies several important types of DNA repair, this repair function probably being more important to the cell (especially bacterial cells) than the capacity that homologous recombination provides for crossing over between chromosomes.

The Holliday and Meselson–Radding models for homologous recombination

Many of the breakthroughs in understanding homologous recombination were made by Robin Holliday, Matthew Meselson, and their colleagues in the 1960s and 1970s. This work resulted in a series of models that showed how breakage and reunion of DNA molecules could lead to the exchange of chromosome segments known to occur during crossing over. We will therefore begin our study of homologous recombination by examining these models.

The Holliday and Meselson–Radding models describe recombination between two homologous, double-stranded molecules with identical or nearly identical sequences. The central feature of these models is formation of a heteroduplex resulting from the exchange of polynucleotide segments between the two homologous molecules (Figure 17.3). The heteroduplex is initially stabilized by base pairing between each transferred strand and the intact polynucleotides of the recipient molecules, this base pairing being possible because of the sequence similarity between the two molecules. Subsequently the gaps are sealed by DNA ligase, giving a Holliday structure. This structure is dynamic, branch migration...
resulting in exchange of longer segments of DNA if the two helices rotate in the same direction.

Separation, or **resolution**, of the Holliday structure back into individual, double-stranded molecules occurs by cleavage across the branch point. This is the key to the entire process because the cut can be made in either of two orientations, as becomes apparent when the three-dimensional configuration, or **Chi form**, of the Holliday structure is examined (see Figure 17.3). These two cuts have very different results. If the cut is made from left to right across the Chi form (labeled horizontal resolution in Figure 17.3), then all that happens is that a short segment of polynucleotide, corresponding to the distance migrated by the branch of the Holliday structure, is transferred between the two molecules. On the other hand, an up–down cut (vertical resolution in Figure 17.3) results in **reciprocal strand exchange**, in which double-stranded DNA is transferred between the two molecules so that the end of one molecule is exchanged for the end of the other molecule. This is the DNA transfer seen in crossing over.

So far we have ignored one important aspect of this model. This is the way in which the two double-stranded molecules interact at the beginning of the process to produce the heteroduplex. In Holliday’s original scheme, the two molecules lined up with one another and single-strand nicks appeared at equivalent
positions in each helix. This produced free, single-stranded ends that could be exchanged, resulting in the heteroduplex (Figure 17.4A). This feature of the model was criticized because no mechanism could be proposed to ensure that the nicks occurred at precisely the same position on each molecule. The Meselson–Radding modification proposes a more satisfactory scheme whereby a single-strand nick occurs in just one of the double helices, the free end that is produced invading the unbroken double helix at the homologous position and displacing one of its strands, forming a D-loop (Figure 17.4B). Subsequent cleavage of the displaced strand at the junction between its single-stranded and base-paired regions produces the heteroduplex.

The double-strand break model for homologous recombination

Although the Holliday model for homologous recombination, either in its original form or as modified by Meselson and Radding, explained how crossing over could occur during meiosis, it had inadequacies that prompted the development of alternative schemes. In particular, it was thought that the Holliday model could not explain gene conversion, a phenomenon first described in yeast and fungi but now known to occur with many eukaryotes. In yeast, fusion of a pair of gametes results in a zygote that gives rise to an ascus containing four haploid spores whose genotypes can be individually determined (see Figure 14.15). If the gametes have different alleles at a particular locus, then under normal circumstances two of the spores will display one genotype and two will display the other genotype, but sometimes this expected 2:2 segregation pattern is replaced by an unexpected 3:1 ratio (Figure 17.5). This is called gene conversion because the ratio can only be explained by one of the alleles converting from one type to the other, presumably by recombination during the meiosis that occurs after the gametes have fused.

The double-strand break (DSB) model provides an opportunity for gene conversion to take place during recombination. According to this model, homologous recombination initiates not with a single-strand nick, as in the Meselson–Radding scheme, but with a double-strand cut that breaks one of the recombination partners into two pieces (Figure 17.6). After the double-strand cut, one strand in

Figure 17.5 Gene conversion. One gamete contains allele A and the other contains allele a. These fuse to produce a zygote that gives rise to four haploid spores, all contained in a single ascus. Normally, two of the spores will have allele A and two will have allele a, as shown on the left, but if gene conversion occurs, the ratio will be changed, possibly to 3A:1a as shown on the right.
each half of the molecule is shortened by exonuclease activity, so each end now has a 3' overhang. One of these overhangs invades the homologous DNA molecule in a manner similar to that envisaged by the Meselson–Radding scheme, setting up a Holliday junction that can migrate along the heteroduplex if the invading strand is extended by a DNA polymerase. To complete the heteroduplex, the other broken strand (the one not involved in the Holliday junction) is also extended. Note that both DNA syntheses involve extension of strands from the partner that suffered the double-strand cut, using as templates the equivalent regions of the uncut partner. This is the basis of gene conversion because it means that the polynucleotide segments removed from the cut partner have been replaced with copies of the DNA from the uncut partner. After ligation, the resulting heteroduplex has a pair of Holliday structures that can be resolved in a number of ways, some resulting in gene conversion and others giving a standard reciprocal strand exchange. An example leading to gene conversion is shown in Figure 17.6.

Although initially proposed as a mechanism for explaining gene conversion in yeast, the DSB model is now looked on as at least a close approximation to the way in which homologous recombination operates in all organisms. Acceptance of this model has come about for two reasons. First, in 1989 it was discovered that, during meiosis, chromosomes undergo double-strand breakages at 100–1000 times the rate seen in vegetative cells. The implication that formation of double-strand breaks is an inherent part of meiosis clearly favors the DSB model at the expense of schemes in which recombination is initiated by one or more single-strand nicks. The second factor leading to acceptance of the DSB model was the realization that homologous recombination is involved in DNA repair and specifically is responsible for repairing double-strand breaks that occur as aberrations in the replication process. The Holliday and Meselson–Radding models do not explain this aspect of homologous recombination, whereas double-strand break repair is implicit in the DSB model. We return to DNA repair after we have considered the biochemical basis of homologous recombination.

**RecBCD is the most important pathway for homologous recombination in bacteria**

Homologous recombination occurs in all organisms, but as with many aspects of molecular biology, our initial progress in understanding how the process is carried out in the cell was made with *Escherichia coli*. Of course, bacteria do not undergo meiosis, but crossing over by homologous recombination occurs following transfer of DNA from one bacterium to another and results in integration of the donated DNA into the chromosome of the recipient cell (see Figure 3.25A). The first breakthroughs in understanding the biochemistry of homologous recombination were made when mutation studies identified a number of *E. coli* genes that, when inactivated, give rise to defects in recombination, indicating that their protein products are involved in some way. Two distinct recombination systems have been described, the RecBCD and RecFOR pathways, with RecBCD apparently being the most important in bacteria.

In the RecBCD pathway, recombination is mediated by the **RecBCD complex**, which, as its name implies, is made up of three different proteins. Two of these, RecB and RecD, are helicases. To initiate homologous recombination, one copy of RecBCD attaches to the chromosome at a double-strand break. The DNA is unwound through the action of RecB, which travels along one strand in the 3' → 5' direction, and RecD, traveling 5' → 3' along the other strand. The RecB protein, as well as being a helicase, also has 3' → 5' exonuclease activity and so progressively degrades the strand it is traveling along, the one with the free 3'-end (Figure 17.7).

RecBCD progresses along the DNA molecule at a rate of approximately 1 kb per second until it reaches the first copy of the eight-nucleotide consensus sequence 5' - GCTGGTGG - 3' called the **chi (crossover hotspot initiator) site**, which occurs...
Figure 17.7 The RecBCD pathway for homologous recombination in E. coli. Translocation of the RecBCD complex along the DNA is accompanied by 3'→5' degradation of the upper strand, due to the exonuclease activity of RecB. When a chi site is encountered, the exonuclease activity is suppressed and the RecB endonuclease cleaves the lower strand, giving the 3'-overhang. If RecBCD translocates some distance along the DNA before reaching a chi site, then it is likely that the 5'-overhang shown in the second panel will occasionally be cleaved by the endonuclease, so that this overhang is never more than a few tens of base pairs in length.

The next step is establishment of the heteroduplex. This stage is mediated by the RecA protein, which forms a protein-coated DNA filament that is able to invade the intact double helix and set up the D-loop (Figure 17.8). An intermediate in formation of the D-loop is a triplex structure (see the third step in Figure 17.6), a three-stranded DNA helix in which the invading polynucleotide lies within the major groove of the intact helix and forms hydrogen bonds with the base pairs it encounters.

Branch migration is catalyzed by the RuvA and RuvB proteins, both of which attach to the branch point of the heteroduplex formed by invasion of the 3'-overhang into the partner molecule. X-ray crystallography studies suggest that RuvA is a tetramer of four identical proteins with one or two tetramers binding directly to the branch, forming a core to which two hexameric RuvB rings attach, one to either side (Figure 17.9). The resulting structure might act as a molecular motor, rotating the helices in the required manner so that the branch point moves. Branch migration does not appear to be a random process but instead stops preferentially at the sequence 5'-A/T TTG/C-3', where A/T and G/C denote that either of the two nucleotides can be present at the position indicated. This sequence occurs frequently in the E. coli genome, so presumably migration does not always halt at the first instance of the motif that is reached. When branch migration has ended, two RuvC proteins attach to the core of RuvA proteins, possibly displacing one RuvA tetramer if there are two present. RuvC is a resolvase that carries out the cleavage to resolve the Holliday structure, making a cut between the second T and the G/C of the recognition sequence.

Note that the above description provides no precise role for RecC in homologous recombination by the RecBCD pathway. X-ray crystallographic studies have revealed that the RecC protein comprises three structural domains, two of which are similar to the catalytic domains of the SF1 family of helicases and one of which is similar to a PD-(D/E)xK nuclease domain. This means that RecC has a structural relationship with RecB [which is an SF1 helicase and PD-(D/E)xK nuclease], but remarkably, key amino acids are absent from the RecC structure so that RecC has neither helicase nor nuclease activity. The nuclease-like domain retains the ability to make contacts with the DNA molecule, enabling RecC to form a loop through which one of the DNA strands is fed toward the RecD component. RecC may therefore stabilize the complex and help ensure that RecB and RecD are positioned correctly relative to the DNA. It is also possible that RecC has acquired a scanning function and is responsible for identifying the chi site and hence initiating the conformational change in RecBCD that leads to formation of the heteroduplex.
**E. coli** can also carry out homologous recombination by the RecFOR pathway

Mutants of *E. coli* that lack components of the RecBCD complex are still able to carry out homologous recombination, albeit with lowered efficiency. This is because the bacterium possesses at least one other homologous recombination pathway, called RecFOR. In normal *E. coli* cells, most homologous recombination takes place via RecBCD, but if this pathway is inactivated by mutation, the RecFOR system is able to take over.

The details of the RecFOR pathway are beginning to emerge, and the general mechanism appears to be similar to that described for RecBCD. The helicase activity for the RecFOR pathway is provided by RecQ, and the 5′-end of the strand is removed by RecJ, leaving a 3′-overhang that becomes coated in RecA proteins through the combined actions of RecF, RecO, and RecR. There is considerable interchangeability between the components of the RecBCD and RecFOR pathways, and it is thought that hybrid systems operate in some mutants that lack components of the standard processes. There are differences, however, as only the RecBCD pathway initiates recombination at the chi sites scattered around the *E. coli* genome, and only RecFOR is able to induce recombination between a pair of plasmids. RecFOR is also the primary pathway responsible for recombination repair of single-strand gaps resulting from replication of heavily damaged DNA.

As well as the RecBCD and RecFOR pathways, whose functions are to set up the heteroduplex structure, *E. coli* also has alternative means for carrying out the branch migration step. Mutants that lack RuvA or RuvB are still able to carry out homologous recombination because the function of RuvAB can also be provided by a helicase called RecG. It is not yet clear whether RuvAB and RecG are simply interchangeable or whether they are specific for different recombination scenarios. RuvC mutants are also able to carry out homologous recombination, suggesting that *E. coli* possesses other proteins that are able to resolve Holliday structures, but the identity of these protein(s) is unknown.

**Homologous recombination pathways in eukaryotes**

The double-strand break model for homologous recombination is thought to hold for all organisms, not just *E. coli*: recall that it was initially devised to explain gene conversion in *Saccharomyces cerevisiae*. The biochemical events underlying the process appear to be similar in all organisms, and a number of yeast proteins have been identified that carry out functions equivalent to those occurring during the RecBCD pathway in *E. coli*. In particular, two yeast proteins called RAD51 and DMC1 are the homologs of RecA of *E. coli*. Although specific roles for RAD51 and DMC1 are suspected, they are thought to work together or interchangeably in many homologous recombination events. This conclusion arises because mutants that lack one or other of these proteins have similar phenotypes, and the two proteins are found together at the same locations within nuclei.
that are undergoing meiosis. Proteins homologous to RAD51 and DMC1 are also known in other eukaryotes including humans.

One puzzling aspect of homologous recombination in eukaryotes has been the mechanism by which Holliday structures are resolved, because for many years, proteins homologous to *E. coli* RuvC had been sought but not found. In fact, RuvC is not universal in all bacteria, some species apparently using a totally different type of nuclease to resolve Holliday structures. The first human resolvase to be identified, MUS81, has homologs in both *S. cerevisiae* and *Schizosaccharomyces pombe*, but this protein acts differently than RuvC: although MUS81 can resolve Holliday structures, it does not give rise to crossovers. Eventually, in 2008, a protein called GEN1, or Yen1 in *S. cerevisiae*, was shown to be the functional equivalent of RuvC in eukaryotes. GEN1 is a member of the Rad1/XPG family of eukaryotic nucleases, whose other members include the FEN1 endonuclease that is involved in lagging-strand replication (Section 15.3), as well as nucleases with roles in mismatch and nucleotide excision repair (Section 16.2). None of these other enzymes is able to cleave Holliday junctions. The resolvase activity of GEN1 might be conferred in part by the presence of a structural motif called a chromodomain, which the other Rad1/XPG nucleases do not possess. A chromodomain does not itself have any DNA-binding ability, but the chromodomain within GEN1 is thought to help to position the protein onto the Holliday structure, providing GEN1 with its unique ability to make the cuts that give rise to crossovers.

The primary role of homologous recombination is thought to be DNA repair

The attention paid by geneticists to crossing over as a central feature of sexual reproduction inevitably biased the initial studies of homologous recombination toward events occurring during meiosis. An alternative role for homologous recombination became apparent when *E. coli* mutants defective for components of the RecBCD and other recombination pathways were first examined and shown to have deficiencies in DNA repair. Today we believe that the principal function of homologous recombination is in DNA repair, with its role in crossing over being of secondary importance in most cells.

Homologous recombination is particularly important as a repair process when breaks arise in daughter DNA molecules as a result of aberrations in replication. One such aberration can occur when the replication machinery is attempting to copy a segment of the genome that is heavily damaged, particularly in regions in which cyclobutyl dimers are prevalent. When a cyclobutyl dimer is encountered, the template strand cannot be copied and the DNA polymerase simply jumps ahead to the nearest undamaged region, where it restarts the replication process. The result is that one of the daughter polynucleotides has a gap (Figure 17.10). One way in which this gap could be repaired is by a recombination event that transfers the equivalent segment of DNA from the parent polynucleotide present in the second daughter double helix. The gap that is now present in the second double helix is refilled by a DNA polymerase, using the undamaged daughter polynucleotide within this helix as the template. In *E. coli*, this type of single-strand gap repair utilizes the RecFOR recombination pathway.

If the damaged site cannot be bypassed, then the daughter polynucleotide, rather than having a gap, will terminate (Figure 17.11). There are several ways in which this problem can be overcome. One possibility is that the replication fork stalls and reverses a short distance, so that a duplex is formed between the daughter polynucleotides. The incomplete polynucleotide is then extended by a DNA polymerase, using the undamaged daughter polynucleotide as a template. The replication fork then moves forward again, by a process equivalent to the branch migration step of homologous recombination. As a result, the damaged site is bypassed and replication can continue.

A more serious aberration occurs if one of the parent polynucleotides being replicated contains a single-strand nick. Now the replication process leads to a double-strand break in one of the daughter double helices, and the replication
fork is lost (Figure 17.12). The break can be repaired by a form of homologous recombination between the broken end and the second, undamaged molecule. In the scheme shown in Figure 17.12, the daughter polynucleotide at the double-strand break is extended via a strand-exchange reaction that enables it to use the other parent strand as a template. Branch migration followed by resolution of the Holliday structure then restores the replication fork.

17.2 SITE-SPECIFIC RECOMBINATION

A region of extensive homology is not a prerequisite for recombination: the process can also be initiated between two DNA molecules that have only very short sequences in common. This is called site-specific recombination, and it has been extensively studied because of the part that it plays during the infection cycle of bacteriophage λ.

Bacteriophage λ uses site-specific recombination during the lysogenic infection cycle

After injecting its DNA into an E. coli cell, bacteriophage λ can follow either of two infection pathways (Section 14.3). One of these, the lytic pathway, results in...
the rapid synthesis of λ coat proteins, combined with replication of the λ genome, leading to death of the bacterium and release of new phages within about 45 min of the initial infection. In contrast, if the phage follows the lysogenic pathway, new phages do not immediately appear. The bacterium divides as normal, possibly for many cell divisions, with the phage in a quiescent form called the prophage. Eventually, possibly as the result of DNA damage or some other stimulus, the phage becomes active again.

During the lysogenic phase, the λ genome becomes integrated into the E. coli chromosome. It is therefore replicated whenever the E. coli DNA is copied and is passed on to daughter cells just as though it was a standard part of the bacterium's genome. Integration occurs by site-specific recombination between the attachment or att sites, attP on the λ genome and attB on the E. coli chromosome. Each of these attachment sites has at its center an identical 15 bp core sequence referred to as O (Figure 17.13), flanked by variable sequences called B and B' in the bacterial genome and P and P' in the phage DNA. B and B' are quite short, just 4 bp each, meaning that attB covers just 23 bp of DNA, but P and P' are much longer, with the entire attP sequence spanning over 250 bp. Mutations in the core sequence inevitably lead to inactivation of the att site so that it can no longer participate in recombination, but mutations in the flanking sequences have a less severe consequence and only decrease the efficiency of recombination. If attB, the attachment site in the E. coli genome, is inactivated, then insertion of the λ DNA can occur at secondary sites that share some sequence similarity with the genuine attB locus. If a secondary site is being used, then the frequency of lysogeny is greatly reduced, with integration possibly occurring at less than 0.01% of the frequency observed with unmutated E. coli cells.

Because this is recombination between two circular molecules, the result is that one bigger circle is formed: in other words, the λ DNA becomes integrated into the bacterial genome (Figure 17.14). The recombination event is catalyzed by a specialized type I topoisomerase (Section 15.1) called integrase, a member of a diverse family of recombinases present in bacteria, archaea, and yeast. There are at least four binding sites for integrase within attP, as well as at least three sites for a second protein, the integration host factor or IHF. Together these proteins coat the phage attachment site. The integrase then makes a staggered, double-strand cut at equivalent positions in the λ and bacterial att sites. The two short, single-strand overhangs are then exchanged between the DNA molecules, producing a Holliday junction that migrates a few base pairs along the heteroduplex before being cleaved. This cleavage, providing that it is made in the appropriate orientation, resolves the Holliday structure in such a way that the λ DNA becomes inserted into the E. coli genome.

Figure 17.13 Core sequence of the att sites present in bacteriophage λ and in the E. coli chromosome. The pink line indicates the staggered cut made in each att site during integration and excision of the phage genome.

Figure 17.14 Integration of the bacteriophage λ genome into E. coli chromosomal DNA. Both λ and E. coli DNA have copies of the att site, each one comprising an identical central sequence, called O, and flanking sequences P and P' (for the phage att site) or B and B' (for the bacterial att site). Recombination between the O regions integrates the λ genome into the bacterial DNA.
Integration creates hybrid versions of the attachment sites, now called attR (which has the structure BOP⁺) and attL (whose structure is POB⁺). A second site-specific recombination between the two att sites, now both contained in the same molecule, reverses the original process and releases the λ DNA. This recombination is also catalyzed by the integrase but occurs in conjunction with a protein called excisionase, coded by the λ xis gene, rather than IHF. The functions of Xis and IHF in excision and integration, respectively, are probably quite different, and the two proteins should not be looked on as playing equivalent roles in the two processes. The key point is that the combination of integrase and excisionase is able to draw the attR and attL sites together in order to initiate the intramolecular recombination that excises the λ genome. After excision, the λ genome returns to the lytic mode of infection and directs synthesis of new phages.

Site-specific recombination is an aid in construction of genetically modified plants

The processes responsible for integration and excision of the λ genome are fairly typical of the strategies used by phages to establish lysogeny, though with some phages the molecular events are less complex than those seen with λ. Integration and excision of the bacteriophage P1 genome, for example, requires just a single enzyme, the Cre recombinase, which recognizes 34 bp target sites, called loxB and loxP, which are identical to one another and have no flanking sequences equivalent to B, B’, and so forth.

The simplicity of the P1 system has led to its utilization in genetic engineering projects in which site-specific recombination is a requirement. An important application has arisen in the technology used for the generation of genetically modified crops. One of the main areas of concern to emerge from the debate over genetically modified plants is the possible harmful effects of the marker genes used with plant cloning vectors. Most plant vectors carry a copy of a gene for kanamycin resistance (see Figure 2.33), enabling transformed plants to be identified during the cloning process. The kanB gene is bacterial in origin and codes for the enzyme neomycin phosphotransferase II. This gene and its enzyme product are present in all cells of an engineered plant. The fear that neomycin phosphotransferase might be toxic to humans has been allayed by tests with animal models, but there are still concerns that the kanB gene contained in a genetically modified foodstuff could be passed to bacteria in the human gut, making these resistant to kanamycin and related antibiotics, or that the kanB gene could be passed to other organisms in the environment, possibly resulting in damage to the ecosystem.

The fears surrounding the use of kanB and other marker genes have prompted biotechnologists to devise ways of removing these genes from plant DNA after the transformation event has been verified. One of the strategies makes use of Cre recombinase. To use this system, the plant is transformed with two cloning vectors. The first carries the gene being added to the plant along with its kanB selectable marker gene, the latter surrounded by the lox target sequences, and the second carries the Cre recombinase gene. After transformation, expression of the Cre gene therefore results in excision of the kanB gene from the plant DNA (Figure 17.15).

17.3 TRANPOSITION

Transposition results in the transfer of a segment of DNA from one position in the genome to another. The transposition process results in duplication of the target site, giving rise to a pair of direct repeats that flank the transposable element (Figure 17.16).

**Figure 17.16** Integrated transposable elements are flanked by short direct repeat sequences. This particular transposon is flanked by the tetranucleotide repeat 5’-CTGG-3’. Other transposons have different direct repeat sequences.
In Section 9.2 we examined the various types of transposable elements known in eukaryotes and prokaryotes and discovered that these could be broadly divided into three categories on the basis of their transposition mechanism:

- DNA transposons that transpose replicatively, with the original transposon remaining in place and a new copy appearing elsewhere in the genome (Figure 17.17A)
- DNA transposons that transpose conservatively, with the original transposon moving to a new site by a cut-and-paste process (Figure 17.17B)
- Retroelements, all of which transpose replicatively via an RNA intermediate

We will now examine the events responsible for each of these three types of transposition.

**Replicative and conservative transposition of DNA transposons**

A number of models for replicative and conservative transposition of DNA transposons have been proposed over the years, but most are modifications of a scheme originally outlined by Shapiro in 1979. According to this model, the replicative transposition of a bacterial element such as a Tn3-type transposon or a transposable phage (Section 9.2) is initiated by one or more endonucleases that make single-strand cuts on either side of the transposon and in the target site where the new copy of the element will be inserted (Figure 17.18). At the target
site, the two cuts are separated by a few base pairs, so that the cleaved double-stranded molecule has short 5’-overhangs. Ligation of these 5’-overhangs to the free 3’-ends either side of the transposon produces a hybrid molecule in which the original two DNAs—the one containing the transposon and the one containing the target site—are linked together by the transposable element flanked by a pair of structures resembling replication forks.

DNA synthesis at the two replication forks copies the transposable element and converts the initial hybrid into a cointegrate, in which the two original DNAs are still linked (Figure 17.19). Homologous recombination between the two copies of the transposon un couples the cointegrate, separating the original DNA molecule (with its copy of the transposon still in place) from the target molecule, which now contains a copy of the transposon. Replicative transposition has therefore occurred.

A modification of the process just described changes the mode of transposition from replicative to conservative (Figure 17.20). Rather than carrying out DNA synthesis, the hybrid structure is converted back into two separate DNA molecules simply by making additional single-strand nicks on either side of the transposon. This cuts the transposon out of its original molecule, transferring it into the target DNA.

**Retroelements transpose replicatively via an RNA intermediate**

From the human perspective, the most important retroelements are the retroviruses, which include the human immunodeficiency viruses that cause HIV/AIDS and various other virulent types. Most of what we know about retrotransposition...
refers specifically to retroviruses, although it is believed that other retrogene elements, such as retrotransposons of the Ty1/copia and Ty3/gypsy families, transpose by similar mechanisms.

The first step in retrotransposition is synthesis of an RNA copy of the inserted retroelement (Figure 17.21). The long terminal repeat (LTR) at the 5'-end of the element contains a TATA sequence that acts as a promoter for transcription by RNA polymerase II. Some retroelements also have enhancer sequences that are

![Figure 17.21 RNA and DNA replication during transposition of a retroelement.](image)

This diagram shows how an integrated retroelement is copied into a free double-stranded DNA version. The first step is synthesis of an RNA copy, which is then converted to double-stranded DNA by a series of events that involves two template switches, as described in the text.
thought to regulate the amount of transcription that occurs. Transcription continues through the entire length of the element, up to a polyadenylation sequence in the 3′-LTR.

The transcript now acts as the template for RNA-dependent DNA synthesis, catalyzed by a reverse transcriptase enzyme coded by part of the pol gene of the retroelement (see Figure 9.15). Because this is synthesis of DNA, a primer is required, and as during genome replication, the primer is made of RNA rather than DNA. During genome replication, the primer is synthesized de novo by a polymerase enzyme (see Figure 15.13), but retroelements do not code for RNA polymerases and so cannot make primers in this way. Instead they use one of the cell’s tRNA molecules as a primer, which one depending on the retroelement: the Ty1/copia family of elements always use tRNAMet as a primer, but other retroelements use different tRNAs.

The tRNA primer anneals to a site within the 5′-LTR (see Figure 17.21). At first glance this appears to be a strange location for the priming site, because it means that DNA synthesis is directed away from the central region of the retroelement and so results in only a short copy of part of the 5′-LTR. In fact, when the DNA copy has been extended to the end of the LTR, a part of the RNA template is degraded and the DNA overhang that is produced reanneals to the 3′-LTR of the retroelement; being a long terminal repeat, this has the same sequence as the 5′-LTR and so can base-pair with the DNA copy. DNA synthesis now continues along the RNA template. Note that the result is a DNA copy of the entire template, including the priming site: the template switching is, in effect, the strategy that the retroelement uses to solve the end-shortening problem, the same problem that chromosomal DNAs address through telomere synthesis (Section 15.4).

Completion of synthesis of the first DNA strand results in a DNA–RNA hybrid. The RNA is partially degraded by an RNase H enzyme, coded by another part of the pol gene. The RNA that is not degraded, usually just a single fragment attached to a short polypurine sequence adjacent to the 3′-LTR, primes synthesis of the second DNA strand, again by reverse transcriptase, which is able to act as both an RNA- and DNA-dependent DNA polymerase. As with the first round of DNA synthesis, second-strand synthesis initially results in a DNA copy of just the LTR, but a second template switch, to the other end of the molecule, enables the DNA copy to be extended until it is full length. This creates a template for further extension of the first DNA strand, so that the resulting double-stranded DNA is a complete copy of the internal region of the retroelement plus the two LTRs.

All that remains is to insert the new copy of the retroelement into the genome. It was originally thought that insertion occurred randomly, but it now appears that although no particular sequence is used as a target site, integration occurs preferentially at certain positions. Insertion involves removal of two nucleotides from the 3′-ends of the double-stranded retroelement by the integrase enzyme (coded by yet another part of pol). The integrase also makes a staggered cut in the genomic DNA so that both the retroelement and the integration site now have 5′-overhangs (Figure 17.22). These overhangs might not have complementary sequences but they still appear to interact in some way so that the retroelement becomes inserted into the genomic DNA. The interaction results in loss of the retroelement overhangs and filling in of the gaps that are left, which means that the target site becomes duplicated into a pair of direct repeats, one at either side of the inserted retroelement.

**SUMMARY**

- The term recombination was originally used to describe the outcome of crossing over between pairs of homologous chromosomes during meiosis. The term is now also used to refer to the molecular events that underlie this process.
- Homologous recombination occurs between segments of DNA molecules that share extensive sequence homology.
Chapter 17: Recombination and Transposition

- The initial models for homologous recombination envisaged the recombination process being initiated by nicks that occur in one or both of the double-stranded molecules, but it is now thought that the start point is a double-strand break in one of these molecules.

- Strand exchange leads to a heteroduplex structure that is resolved by cleavage, possibly leading to exchange of DNA segments or to gene conversion.

- *Escherichia coli* possesses at least two molecular pathways for homologous recombination. These are the RecBCD and RecFOR pathways.

- The RecBCD pathway involves the unwinding of one partner in the recombination by a pair of helicases, which attach to the double-strand break and progress along the molecule. At a recognition sequence called a chi site, the RecBCD complex initiates the strand exchange, with the RecA protein playing a central role in transfer of the invading strand into the intact double helix.

- Branch migration within the heteroduplex, and resolution of the structure, is catalyzed by the Ruv proteins.

- Homologous recombination pathways in eukaryotes are gradually being elucidated.

- Homologous recombination plays a major role in the repair of DNA strand breaks.

- Site-specific recombination does not require regions of extensive homology between the partner molecules. This type of recombination is responsible for insertion of bacteriophage genomes, such as the λ genome, into the host bacterial chromosome.

- Transposition occurs by recombination. With DNA transposons, the process can be either replicative or conservative, both modes occurring by a series of events initially described by Shapiro in 1979.

- Retroelements transpose via an RNA intermediate that is transcribed from the parent copy of the transposon. After copying into double-stranded DNA, the retroelement is reinserted into the host chromosome.

**SHORT ANSWER QUESTIONS**

1. What is the role of recombination in genome evolution?
2. How can the resolution of a Holliday structure yield two different results?
3. Describe how the double-strand break model explains how gene conversion occurs.
4. Describe the RecBCD pathway for homologous recombination in *E. coli*.
5. What are the distinctive features of the RecFOR pathway for homologous recombination in *E. coli*?
6. Outline our current knowledge of the molecular basis to homologous recombination in eukaryotes.
7. Describe the processes by which homologous recombination is used to repair breaks in DNA molecules.
8. What are the properties of the *attP* and *attB* sites that mediate integration of bacteriophage λ DNA into the *E. coli* genome?
9. Distinguish between the processes that lead to integration and excision of the bacteriophage λ DNA into and out of the *E. coli* genome.
10. Describe the model for conservative transposition.

11. How does the model for replicative transposition differ from that of conservative transposition?

12. How is the new copy of a retroelement inserted into a genome?

IN-DEPTH PROBLEMS

1. Determination of the structure of the RecBCD complex was looked on as a key step in understanding the molecular basis of homologous recombination. Explain why knowing the structure of this complex was so important.

2. Some *E. coli* strains that are used for propagating recombinant plasmids contain *recA* mutations. Why might *recA* defects be useful for researchers working with recombinant plasmids?

3. The Cre recombination system underlies one of the more controversial aspects of plant genetic engineering, the so-called terminator technology. This is one of the processes by which the companies who market genetically modified crops attempt to protect their financial investment by ensuring that farmers must buy new seed every year, rather than simply collecting seed from the crop and sowing this second-generation seed the following year. The terminator technology centers on the gene for ribosome inactivating protein (RIP). RIP blocks protein synthesis by cutting one of the ribosomal RNA molecules into two segments, which means that any cell in which RIP is active will quickly die. Use this information to deduce exactly how the terminator technology works.

4. Transposition can have deleterious effects on a genome. These effects go beyond the obvious disruption of gene activity that will occur if a transposable element takes up a new position that lies within the coding region of a gene. Some elements, notably retrotransposons, contain promoter and enhancer sequences that can modify the expression patterns of adjacent genes, and transposition often involves the creation of double-strand breaks. How might cells minimize these deleterious effects by preventing transposition from occurring?

5. Are there circumstances where transposition could be beneficial to a cell?

FURTHER READING

Models for homologous recombination


Molecular basis to homologous recombination


*Hybrid pathways involving parts of the RecBCD and RecFOR systems.*


**Role of homologous recombination in repair of DNA strand breaks**


**Site-specific recombination**


**Transposition**


Mutation and recombination provide the genome with the means to evolve, but we learn very little about the evolutionary histories of genomes simply by studying these events in living cells. Instead we must combine our understanding of mutation and recombination with comparisons between the genomes of different organisms in order to infer the patterns of genome evolution that have occurred. Clearly, this approach is imprecise and uncertain, but as we will see, it is based on a surprisingly large amount of hard data and we can be reasonably confident that, at least in outline, the picture that emerges is not too far from the truth.

In this chapter we will explore the evolution of genomes from the very origins of biochemical systems through to the present day. We will look at ideas regarding the RNA world, prior to the appearance of the first DNA molecules, and then examine how DNA genomes have gradually become more complex. We will examine the differences between the human and chimpanzee genomes in an attempt to identify the evolutionary changes that have occurred during the last 6 million years and which must, somehow, make us what we are. Finally, we will study how the diversity of genome sequences in modern-day populations is being used as a tool in research and in biotechnology.

18.1 GENOMES: THE FIRST 10 BILLION YEARS

Cosmologists believe that the universe began some 14 billion years ago with the gigantic primordial fireball called the Big Bang. Mathematical models suggest that, after about 4 billion years, galaxies began to fragment from the clouds of gas emitted by the Big Bang and that, within our own galaxy, the solar nebula condensed to form the Sun and its planets about 4.6 billion years ago (Figure 18.1). The early Earth was covered with water, and it was in this vast planetary ocean that the first biochemical systems appeared. Cellular life was well-established by the time land masses began to appear, some 3.5 billion years ago. We believe this to be the case because tiny microfossils of structures resembling bacteria have been discovered in 3.4 billion-year-old rocks from Australia (Figure 18.2). But cellular life was a relatively late stage in biochemical evolution, being preceded by self-replicating polynucleotides that were the progenitors of the first genomes. We must begin our study of genome evolution with these precellular systems.

The first biochemical systems were centered on RNA

The first oceans are thought to have had a similar salt composition to those of today, but the Earth’s atmosphere, and hence the dissolved gases in the oceans, was very different. The oxygen content of the atmosphere remained very low until photosynthesis evolved, and in the beginning the most abundant gases were probably methane and ammonia. The possibility that organic compounds important in biochemistry could be formed on the early Earth was first demonstrated in 1952, when experiments that recreated conditions in the ancient atmosphere showed that electrical discharges in a methane-ammonia mixture could result in...
synthesis of a range of amino acids, including alanine, glycine, valine, and several others found in proteins (Figure 18.3). Formation of ribonucleotides has been less easy to understand because of difficulties in envisaging how the first ribose sugars and nucleotide bases could be synthesized, but it has recently been demonstrated that complete ribonucleotides can be produced directly without prior synthesis of the sugar and base components. For example, pyrimidine nucleotides can be constructed from cyanamide, cyanoacetylene, glycolaldehyde, glyceraldehyde, and inorganic phosphate via arabinose amino-oxazoline and anhydronucleoside intermediates.

Once ribonucleotides had formed, their polymerization into RNA molecules would have been aided by base-stacking interactions (Section 1.1). These interactions would have been much too weak to hold nucleotides together for any length of time in the early ocean, but they might have been sufficient to stabilize structures that formed on solid surfaces such as clay particles or ice. Alternatively, base stacking could have been promoted by the repeated condensation and drying of droplets of water in clouds. The precise mechanism need not concern us: what is important is that it is possible to envisage purely geochemical processes that could lead to synthesis of polymeric RNA and that similar scenarios give plausible routes for polypeptide synthesis. It is the next steps that we must worry about. We have to go from a random collection of biomolecules to an ordered assemblage that displays at least some of the biochemical properties that we associate with life. These steps have never been reproduced experimentally, and our ideas are therefore based mainly on speculation tempered by a certain amount of computer simulation. One problem is that the speculations are unconstrained because the global ocean could have contained as many as $10^{10}$ biomolecules per liter and we
18.1 Genomes: The First 10 Billion Years

Progress in understanding the origins of cellular life was initially stalled by the apparent requirement that polynucleotides and polypeptides must work in harness in order to produce a self-reproducing biochemical system. This is because proteins are required to catalyze biochemical reactions but cannot carry out their own self-replication. Polynucleotides, on the other hand, can specify the synthesis of proteins and act as templates for their own self-replication, but it was thought that they could do neither without the aid of other proteins. It appeared that the biochemical system would have to spring fully formed from the random collection of biomolecules because any intermediate stage could not be perpetuated. Many of these problems were swept aside in the 1980s with the discovery of the first catalytic RNA molecules. It was quickly realized that ribozymes can catalyze biochemical reactions including: self-cleavage, as displayed by some modern-day viroid and virusoid genomes (see Figure 9.11); cleavage of other RNAs, as carried out by, for example, RNase P during pre-tRNA processing; and synthesis of peptide bonds, as performed by the rRNA component of the ribosome (Section 13.3).

In the test tube, synthetic RNA molecules have been shown to carry out other biologically relevant reactions such as synthesis of ribonucleotides, synthesis and copying of RNA molecules, and transfer of an RNA-bound amino acid to a second amino acid to form a dipeptide, in a manner analogous to the role of tRNA in protein synthesis. The discovery of these catalytic properties, and of various ways in which the activities of ribozymes might be regulated by riboswitches (Section 12.2), has solved the polynucleotide–polypeptide dilemma by showing that the first biochemical systems could have been centered entirely on RNA.

Various ideas about the RNA world have taken shape in recent years. We now envisage that RNA molecules initially replicated in a slow and haphazard fashion simply by acting as templates for binding of complementary nucleotides that polymerized spontaneously (Figure 18.4). This replication process would have been very inaccurate, so a variety of RNA sequences would have been generated, eventually leading to one or more with nascent ribozyme properties that were able to direct their own, more accurate self-replication. It is possible that a form of natural selection operated so that the most efficient replicating systems began to predominate, as has been shown to occur in experimental systems. Greater accuracy in replication would have enabled RNAs to increase in length without losing their sequence specificity, providing the potential for more sophisticated catalytic properties, possibly culminating in structures as complex as present-day ribosomal RNAs (see Figure 13.24).

To call the early RNAs genomes is a little fanciful, but the term protogenome has attractions as a descriptor for molecules that were self-replicating and able to direct simple biochemical reactions. These reactions might have included energy metabolism based, as today, on the release of free energy by hydrolysis of the phosphate–phosphate bonds in the ribonucleotides ATP and GTP, and the reactions might have become compartmentalized within lipid membranes, forming the first cell-like structures. There are difficulties in envisaging how long-chain unbranched lipids could form by chemical or ribozyme-catalyzed reactions, but once present in sufficient quantities they would have assembled spontaneously into membranes, possibly encapsulating one or more protogenomes. Within these early cells, the concentration of ribozymes would have been greater than in the unstructured external matrix, possibly giving the cells a selective advantage that enabled them to propagate. Inclusion in the cells of compounds such as citrate might have provided the first steps toward control of the ionic environment, as citrate chelates magnesium ions, which promote the activity of many of today’s ribozymes. The early cells could therefore have

**Figure 18.4 Copying of RNA molecules in the early RNA world.** Before the evolution of RNA polymerases, ribonucleotides that became associated with an RNA template would have had to polymerize spontaneously. This process would have been inaccurate, and many RNA sequences would have been generated.
provided the RNA protogenomes with an enclosed environment in which more controlled biochemical reactions could be carried out.

**The first DNA genomes**

How did the RNA world develop into the DNA world? The first major change was probably the development of protein enzymes, which supplemented, and eventually replaced, most of the catalytic activities of ribozymes. There are several unanswered questions relating to this stage of biochemical evolution, including the reason why the transition from RNA to protein occurred in the first place. Originally, it was assumed that the 20 amino acids in polypeptides provided proteins with greater chemical variability than the four ribonucleotides in RNA, enabling protein enzymes to catalyze a broader range of biochemical reactions, but this explanation has become less attractive as more and more ribozyme-catalyzed reactions have been demonstrated in the test tube. A more recent suggestion is that protein-mediated catalysis is more efficient because of the inherent flexibility of folded polypeptides compared with the greater rigidity of base-paired RNAs. Alternatively, enclosure of RNA protogenomes within membrane vesicles could have prompted the evolution of the first proteins, because RNA molecules are hydrophilic and must be given a hydrophobic coat, for instance, by attachment to peptide molecules, before being able to pass through or become integrated into a membrane.

The transition to protein-mediated catalysis demanded a radical shift in the function of the RNA protogenomes. Rather than being directly responsible for the biochemical reactions occurring in the early cell-like structures, the protogenomes became coding molecules whose main function was to specify the construction of the catalytic proteins. Whether the ribozymes themselves became coding molecules or coding molecules were synthesized by the ribozymes is not known, although the most persuasive theories about the origins of protein synthesis and the genetic code suggest that the latter alternative is more likely to be correct (Figure 18.5). Whatever the mechanism, the result was the paradoxical situation whereby the RNA protogenomes had abandoned their roles as enzymes, which they were good at, and taken on a coding function for which they were less well suited because of the relative instability of the RNA phosphodiester bond. A transfer of the coding function to the more stable DNA seems almost inevitable and would not have been difficult to achieve, reduction of ribonucleotides giving deoxyribonucleotides, which could then be polymerized into DNA copies of the RNA protogenomes by a reverse-transcriptase-catalyzed reaction (Figure 18.6). The replacement of uracil with its methylated derivative thymine probably conferred even more stability on the DNA polynucleotide, and the adoption of double-stranded DNA as the coding molecule was almost certainly prompted by the possibility of repairing DNA damage by copying the partner strand (Section 16.2).

According to this scenario, the first DNA genomes comprised many separate molecules, each specifying a single protein and each therefore equivalent to a

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**Figure 18.5** Two scenarios for evolution of the first coding RNA. A ribozyme could have evolved to have a dual catalytic and coding function (A), or a ribozyme could have synthesized a coding molecule (B). In both examples, the amino acids are shown attaching to the coding molecule via small adaptor RNAs, the presumed progenitors of today’s tRNAs.
single gene. The linking together of these genes into the first chromosomes, which could have begun before the transition to DNA, would have improved the efficiency of gene distribution during cell division, as it is easier to organize the equal distribution of a few large chromosomes than many separate genes. As with most stages in early genome evolution, several different mechanisms by which genes might have become linked have been proposed.

**How unique is life?**

If the experimental simulations and computer models are correct, then it is likely that the initial stages in biochemical evolution occurred many times in parallel in the oceans or atmosphere of the early Earth. It is therefore quite possible that life arose on more than one occasion, even though all present-day organisms appear to derive from a single origin. This single origin is indicated by the remarkable similarity between the basic molecular biological and biochemical mechanisms in bacterial, archaeal, and eukaryotic cells. To take just one example, there appears to be no obvious biological or chemical reason why any particular triplet of nucleotides should code for any particular amino acid, but the genetic code, although not universal, is virtually the same in all organisms that have been studied. If these organisms derived from more than one origin, then we would anticipate two or more very different codes.

If multiple origins are possible, but modern life is derived from just one origin, then at what stage did a single biochemical system begin to predominate? The question cannot be answered precisely, but the most likely scenario is that the predominant system was the first to develop the means to synthesize protein enzymes and therefore probably also the first to adopt a DNA genome. The greater catalytic potential and more accurate replication conferred by protein enzymes and DNA genomes would have given these cells a significant advantage compared to those still containing RNA protogenomes. The DNA–RNA–protein cells would have multiplied more rapidly, enabling them to outcompete the RNA cells for nutrients that, before long, would have included the RNA cells themselves.

What might have been the alternatives to life as we know it among the early biomolecules? One possibility is base pairing involving nucleotides other than the ones used in today’s versions of DNA and RNA. An interesting example is the pair formed by purine-2,6-dicarboxylate and 3-pyridine, which is coordinated by a Cu$^{2+}$ ion (Figure 18.7). Many natural DNA polymerases are able to add 3-pyridine to a growing polynucleotide if the template contains purine-2,6-dicarboxylate, indicating that the polymerizing capability of these enzymes is not limited to the canonical Watson–Crick base pairs. Equally interesting is the possibility that life forms based on informational molecules other than DNA or RNA might have occurred in the past, or might occur on a different planet. In particular, a pyranosyl nucleic acid, which is based on a six- rather than five-carbon sugar (Figure 18.8), might be a better choice than RNA for an early protogenome because the base-paired molecules that it forms are more stable. Other possibilities include threose nucleic acid, which uses four-carbon sugars, or a glycerol-derived nucleic acid, which, like peptide nucleic acid (see Figure 3.32), has a linear backbone that lacks sugar ring structures. None of these alternative nucleic acids is known in nature, but they can be synthesized in the laboratory and each one forms a stable double-stranded hybrid with RNA. A nucleic acid with two threose strands is also possible.
Prior to the discovery of pathways for the abiotic synthesis of ribonucleotides, these nucleic acid analogs were looked on as attractive precursors to the RNA world, in that their spontaneous synthesis might be more plausible than that of RNA. Today, the general consensus is that there is no need to postulate a precursor to RNA as the latter is likely to have evolved directly from the prebiotic soup.

18.2 EVOLUTION OF INCREASINGLY COMPLEX GENOMES

If we follow the planet’s history forward from the earliest bacterial cells, dating to 3.4 billion years ago, we see the first evidence for eukaryotes 2.7 billion years ago, in the form of sterols, thought to be characteristic features of eukaryotes but not prokaryotes, which have been detected in oil shales of this age. The earliest eukaryote visible in the fossil record is currently the organism called Bangiomorpha, a red alga from 1.2 billion years ago. Bangiomorpha is multicellular and reproduces sexually, so clearly it is a relatively advanced type, indicating that earlier eukaryotes remain undiscovered. Multicellular animals appeared around 640 million years ago, although there are enigmatic burrows suggesting that animals lived earlier than this. The Cambrian explosion, when invertebrate life proliferated into many novel forms, occurred 530 million years ago and ended with the disappearance of many of the novel forms in a mass extinction 490 million years ago. Since then, evolution has continued apace and with increasing diversification: the first terrestrial insects, animals, and plants were established by 350 million years ago, the dinosaurs had been and gone by the end of the Cretaceous Period, 65 million years ago, and the first hominins appeared a mere 4.5 million years ago.

Morphological evolution was accompanied by genome evolution. It is dangerous to equate evolution with progress, but it is undeniable that as we move up the evolutionary tree we see increasingly complex genomes. In this section we will explore how this complexity evolved.

Genome sequences provide extensive evidence of past gene duplications

One indication of genome complexity is gene number, which varies from less than 1000 in some bacteria to 20,000 in vertebrates such as humans. If we assume that the earliest genomes had fewer genes than the most complex modern ones, then we must ask how these gene numbers increased.
There are two fundamentally different ways in which new genes could be acquired by a genome: by duplicating some or all of the existing genes in the genome or by acquiring genes from other species. We have already discussed how prokaryotes acquire new genes by lateral transfer from other species (Section 8.2), and we will return to this topic later when we consider the less widespread ways in which eukaryotic genomes have been affected by lateral transfer. First we will focus on gene duplication, which is looked on as having played a central role in genome evolution in all types of organism.

The initial result of gene duplication will be two identical genes. Selective constraints will ensure that one of these genes retains its original nucleotide sequence, or something very similar to it, so that it can continue to provide the protein function that was originally supplied by the single gene copy before the duplication took place. It is possible that the same selective constraints will apply to the second gene, especially if the increase in the rate of synthesis of the gene product, made possible by the duplication, confers a benefit on the organism (Figure 18.9). More frequently, however, the second copy will confer no benefit and hence will not be subject to the same selective pressures and so will accumulate mutations at random. Evidence shows that the majority of new genes that arise by duplication acquire deleterious mutations that inactivate them, so that they become nonprocessed pseudogenes. Occasionally, however, mutations might not lead to inactivation of the gene but instead result in a new gene function that is useful to the organism (see Figure 18.9). In this case, the gene will be retained and the gene content of the genome will have increased.

The most cursory examination of genome sequences provides ample evidence that many genes have arisen by duplication events. The importance of the first scenario illustrated in Figure 18.9, where the increased amount of gene product resulting from a gene duplication is beneficial and stabilizes the duplication, is supported by the many examples of multigene families made up of genes with identical or near-identical sequences. The prime examples are the rRNA genes, which are multicopy in all but a few bacteria and have copy numbers of approximately 350 in the human genome and over 4000 in the pea genome. These multiple copies of identical genes presumably reflect the need for rapid synthesis of rRNAs at certain stages of the cell cycle. Note that the existence of these multigene families indicates not only that gene duplications have occurred in the past but also that there must be a molecular mechanism to ensure that the family members retain their identity over evolutionary time. This is called concerted evolution. If one copy of the family acquires an advantageous mutation, then it is possible for that mutation to spread throughout the family until all members possess it. The most likely way in which this can be achieved is by gene conversion, which, as described in Section 17.1, can result in the sequence of one copy of a gene being replaced with all or part of the sequence of a second copy. Multiple gene conversion events could therefore maintain identity among the sequences of the individual members of a multigene family, especially if those members are arranged in tandem arrays.

![Figure 18.9 Three scenarios for the outcome of a gene duplication.](image-url)
The third scenario in Figure 18.9 shows the duplicated gene accumulating mutations that give it a new, useful function. Again, multigene families provide many indications that such events have occurred frequently in the past. We have already seen that gene duplications in the globin gene families led to the evolution of new globin proteins that are used by animals at different stages in development (see Figure 7.19). All of the globin genes, both the α and β types, have related nucleotide sequences and hence form part of a single superfamily. This superfamily also includes genes specifying various other proteins that, like the blood globins, have the capacity to bind oxygen molecules. From the degrees of similarity displayed by pairs of genes in the superfamily, it is possible to deduce the pattern of gene duplications that gave rise to the genes we see today, and by applying the molecular clock to the data, which tells us how rapidly a pair of sequences will diverge over time, we can estimate how many millions of years ago each duplication took place. These analyses tell us that a duplication some 800 million years ago resulted in a pair of ancestral genes, one of which evolved into the modern gene for the brain protein neuroglobin and the other of which gave rise to all the other members of the superfamily (Figure 18.10). Some 250 million years later there was a second duplication on the path leading to the blood globins. One of the products of this duplication was a gene that, via another duplication, gave rise to myoglobin, which is active in muscle, and cytoglobin, which is present in many tissues but whose function is not yet understood. The proto-α and proto-β lineages split by a duplication that occurred 450 million years ago, and the duplications within the α- and β-globin gene families took place during the last 200 million years. Within this more recent time frame, it is possible to deduce not only the pattern of gene duplication but also some of the more detailed changes that have occurred within individual genes. Hence the events leading to the various groups of β-globin genes present in different mammals have been inferred (Figure 18.11).

Figure 18.10 Evolution of the globin gene superfamily of humans. The members of the superfamily are now on different chromosomes. The neuroglobin gene is on chromosome 14, the cytoglobin gene is on chromosome 17, and the myoglobin gene is on chromosome 22. The α-globin cluster is on chromosome 16 and the β-globin cluster is on chromosome 11. Relationships between the genes are inferred from the degree of nucleotide similarity between gene pairs, and the estimated rate at which mutations accumulate in human exons is used as a molecular clock to estimate the dates at which gene duplications occurred. Abbreviation: MYr, millions of years ago.
We observe similar patterns of evolution when we compare the sequences of other genes. The trypsin and chymotrypsin genes, for example, are related by a common ancestral gene that duplicated approximately 1500 million years ago. Both now code for proteases involved in protein breakdown in the vertebrate digestive tract: trypsin cuts other proteins at arginine and lysine amino acids, and chymotrypsin cuts at phenylalanines, tryptophans, and tyrosines. Genome evolution has therefore produced two complementary protein functions where originally there was just one.

Another striking example of gene evolution by duplication is provided by the homeotic selector genes, the key developmental genes responsible for specification of the body plans of animals. As described in Section 14.3, *Drosophila melanogaster* has a single cluster of homeotic selector genes, called HOM-C, which consists of eight genes, each containing a homeodomain sequence coding for a DNA-binding motif in the protein product (see Figure 14.37). These eight genes, as well as other homeodomain genes in *Drosophila*, are believed to have arisen by a series of gene duplications that began with an ancestral gene that existed about 1000 million years ago. The functions of the modern genes, each specifying the identity of a different segment of the fruit fly, give us a tantalizing glimpse into how gene duplication and sequence divergence could, in this case, have been the underlying processes responsible for increasing the morphological complexity of the series of organisms in the evolutionary lineage leading to *Drosophila*. If we then move further up the evolutionary tree, we see that most vertebrates have four Hox gene clusters (see Figure 14.37), each a recognizable copy of the *Drosophila* cluster, with sequence similarities between genes in equivalent positions. The implication is that in the vertebrate lineage there were two duplications, not of individual Hox genes but of the entire cluster (Figure 18.12). So far, no intermediate species with two Hox clusters has been found, but there are well-known examples of vertebrates with more than four. Teleosts, a group of ray-finned fish that are probably the most diverse group of vertebrates, with a vast range of different variations of the basic body plan, have seven or eight Hox

![Figure 18.11](image-url) Evolution of the mammalian β-globin genes. This scheme shows the events that are believed to have given rise to the various β-globin gene clusters seen in different mammalian species. (From Tagle DA, Stanhope MJ, Siemieniak DR et al. [1992] Genomics 13:741-760. With permission from Elsevier.)
clusters, presumed to have arisen by duplication of the set of four, followed by the loss of one cluster in the ancestor of those fish that have just seven clusters. A further duplication in the salmonid lineage has resulted in 13 Hox clusters in the Atlantic salmon *Salmo salar* and the rainbow trout *Oncorhynchus mykiss*.

**A variety of processes could result in gene duplication**

Genome annotations provide extensive evidence that gene duplications have occurred in the past. How did these duplications arise? There are several possibilities, including the following:

- **Unequal crossing over** is a recombination event that is initiated by similar nucleotide sequences that are not at identical places in a pair of homologous chromosomes. As shown in Figure 18.13A, the result of unequal crossing over can be duplication of a segment of DNA in one of the recombination products.

- **Unequal sister chromatid exchange** occurs by the same mechanism as unequal crossing over but involves a pair of chromatids from a single chromosome (Figure 18.13B).

- **DNA amplification** is sometimes used in this context to describe duplication of segments of DNA in bacteria and other haploid organisms, arising by unequal recombination between the two daughter DNA molecules in a replication bubble (Figure 18.13C).

Each of these three processes leads to tandem duplications, in which the two duplicated segments lie adjacent to one another in the genome. This is the pattern seen with many multigene families, but it is not the only possibility (Section 7.3). Family members are not always co-located: for example, in the human genome there are five genes for the metabolic enzyme aldolase, each on a different chromosome. These copies might have once been present as a tandem array and become dispersed as part of large-scale genome reorganizations, but it is also possible that the distant locations are a consequence of a different type of duplication process, one that occurs by retrotransposition, in a manner similar to that thought to lead to formation of processed pseudogenes (see Figure 7.20). A processed pseudogene arises when the mRNA copy of a gene is converted into cDNA and reinserted into the genome. The resulting structure is a pseudogene because it lacks a promoter sequence, which is absent from the mRNA. But what if the mRNA copy becomes inserted adjacent to a novel promoter (Figure 18.14)? Now it might become active by subverting this promoter for its own use. Gene duplicates that arise in this way are called retrogenes.

How do we identify retrogenes when a genome is being annotated? A distinctive feature of a retrogene is that it lacks any introns that were present in the parent copy of the gene, as these are not present in the mRNA. The discovery within an annotation of a pair of homologous genes, one containing introns and one lacking introns, would therefore be an indication that the latter might be a retrogene. However, when we discussed the possible activity of those nonprocessed pseudogenes that are still transcribed from their original promoters (Section 7.3), we learned that transcription and translation *per se* do not provide sufficient evidence for a function to be assigned to a pseudogene. It must also be established that natural selection is acting in a positive way on that gene; otherwise it has to be concluded that expression is merely fortuitous and is of no long-term benefit to the organism. The same issue relates to retrogenes. Expression due to insertion of the retrocopy adjacent to a novel promoter is an indication that the gene might be functional, but this has to be confirmed by establishing that positive selection is acting upon it. When these issues are taken into account, the best estimates are that there are 600–700 retrogenes in the human genome, a small number of which, 25 or so, are orphan retrogenes, whose parent copies have been lost, the retrogene now bearing sole responsibility for providing the protein function.
Whole-genome duplication is also possible

The processes described above give rise to relatively short DNA duplications, perhaps a few tens of kilobases in length. Are larger duplications possible? It seems unlikely that duplication of entire chromosomes has played any major role in genome evolution, because we know that duplication of individual human chromosomes, resulting in a cell that contains three copies of one chromosome and two copies of all the others (the condition called trisomy), is either lethal or results in a genetic disease, such as Down syndrome, and similar harmful effects have been observed in artificially generated trisomic mutants of *Drosophila*. Probably the resulting increase in copy numbers for some genes but not others leads to an imbalance of the gene products and disruption of the cellular biochemistry.

The harmful effects of trisomy do not mean that duplication of the entire set of chromosomes in a nucleus must be discounted. Genome duplication can occur if an error during meiosis leads to the production of gametes that are diploid rather than haploid (Figure 18.15). If two diploid gametes fuse, then the result will be a type of autopolyploid, in this case, a tetraploid cell whose nucleus contains four copies of each chromosome. Autopolyploidy, as with other types of polyploidy, is not uncommon, especially among plants. Autopolyploids are often viable because each chromosome still has a homologous partner and so can form a bivalent during meiosis. This allows an autopolyploid to reproduce successfully, but it generally prevents interbreeding with the original organism from which it was
derived. This is because a cross between, for example, a tetraploid and diploid would give a triploid offspring that would not itself be able to reproduce because one full set of its chromosomes would lack homologous partners (Figure 18.16). Autopolyploidy is therefore a mechanism by which speciation can occur, a pair of species usually being defined as two organisms that are unable to interbreed. The generation of new plant species by autopolyploidy has in fact been observed, notably by Hugo de Vries, one of the rediscoverers of Mendel’s experiments. During his work with evening primrose, *Oenothera lamarckiana*, de Vries isolated a tetraploid version of this normally diploid plant, which he named *Oenothera gigas*. Autopolyploidy among animals is less common, especially in those with two distinct sexes, possibly because of problems that arise if a nucleus possesses more than one pair of sex chromosomes.

Autopolyploidy does not lead directly to an increase in genome complexity because the initial product is an organism that simply has extra copies of every gene, rather than any new genes. It does, however, provide the potential for increased complexity because the extra genes are not essential to the functioning of the cell and so can undergo mutational change without harming the viability of the organism. With this in mind, is there any evidence that whole-genome

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**Figure 18.15** The basis of autopolyploidization. The normal events occurring during meiosis are shown, in abbreviated form, on the left. On the right, an aberration has occurred between prophase I and prophase II, and the pairs of homologous chromosomes have not separated into different nuclei. The resulting gametes will be diploid rather than haploid.

**Figure 18.16** Autopolyploids cannot interbreed successfully with their parents. Fusion of a diploid gamete produced by the aberrant meiosis shown in Figure 18.15 with a haploid gamete produced by normal meiosis leads to a triploid nucleus, one that has three copies of each homologous chromosome. During prophase I of the next meiosis, two of these homologous chromosomes will form a bivalent but the third will have no partner. This has a disruptive effect on the segregation of chromosomes during anaphase and usually prevents meiosis from reaching a successful conclusion. This means that gametes are not produced and the triploid organism is sterile. Note that the bivalent could have formed between any two of the three homologous chromosomes, not just between the pair shown in the diagram.
duplication has been important in the large-scale acquisition of new genes during the evolutionary histories of present-day genomes.

From what we understand about the way in which genomes change over time, we might anticipate that evidence for whole-genome duplication would be quite difficult to obtain. Many of the extra gene copies resulting from genome duplication would decay to the extent that they are no longer visible in the DNA sequence. Those genes that are retained, because their duplicated function is useful to the organism or because they have evolved new functions, should be identifiable, but it would be difficult to distinguish whether they have arisen by duplication of the entire genome or by duplication of much smaller segments. For a genome duplication to be signaled, the following criteria have to be met:

- Molecular clock analyses should indicate that a substantial fraction of the duplicated genes in a genome arose simultaneously.
- Despite postduplication rearrangements, sets of duplicated genes should still occur in blocks, and there should be synteny (similar gene orders) in different copies of each duplicated block.

The first evidence for whole-genome duplication in the evolutionary ancestry of eukaryotes was obtained during the *Saccharomyces cerevisiae* sequencing project. As parts of the sequence were assembled, it became clear that the genome contained syntenic blocks. When the sequence was complete, homology analysis (Section 5.1) was carried out with every yeast gene tested against every other yeast gene. To be considered descendants of a duplication event, two genes had to display at least 25% identity when the predicted amino acid sequences of their protein products were compared. About 800 gene pairs were identified in this way, 376 of which could be placed in 55 duplicate sets, each of these sets containing at least three genes in the same order, possibly with other genes interspersed between them, the sets altogether covering half the genome. These sets could have arisen by duplication of segments rather than the entire genome, but if this was the case, then it might be anticipated that some of the genes would have been duplicated more than once. The fact that there were just two copies of each gene, and never three or four, therefore supported the notion that the copies arose by whole-genome duplication. This possibility became more certain when the complete genome sequences of other yeast species were obtained. Comparisons between the genomes of *S. cerevisiae*, *Kluyveromyces waltii*, and *Ashbya gossypii* have been particularly informative. These three species shared a common ancestor that lived over 100 million years ago, previous to the time of the genome duplication event inferred from the homology analysis. If that duplication had indeed occurred in the lineage leading to *S. cerevisiae*, then it would be anticipated that this species would have duplicated copies of many genes present as singletons in the *K. waltii* and *A. gossypii* genomes. This turns out to be the case, this new analysis suggesting that some 10% of the genes in the modern *S. cerevisiae* genome derive from a whole-genome duplication that occurred just under 100 million years ago.

Equivalent work has now been carried out with other genomes, using more sophisticated computational tools to detect syntenic blocks of duplicated genes and incorporating molecular clock studies to date when duplications occurred. These analyses have uncovered evidence of two whole-genome duplications early in the evolutionary history of vertebrates, some 550 and 450 million years ago (Figure 18.17). A subsequent duplication on the lineage leading to teleost fish occurred about 310 million years ago, and more recently the protosalmonid genome duplicated 80–100 million years ago. Detailed analysis of these two more recent duplications has enabled the events following the duplication to be inferred, suggesting that during the 60 million years following the teleost duplication, 60% of the duplicated genes were lost, reverting these gene pairs back to singletons. Since then, there has been a continued but much slower loss of duplicates. Whole-genome duplications have been equally important during plant evolution. Comparisons between the *Arabidopsis thaliana* sequence and other plant genomes suggest that there have been at least three rounds of genome duplication in the lineage leading to *A. thaliana*, since the origins of monocotyledonous

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**Figure 18.17** Genome duplications in the vertebrate evolutionary lineage. The four genome duplications believed to have occurred in the vertebrate evolutionary lineage are shown. Note that the timing of these duplications explains the different numbers of Hox clusters found in flies, vertebrates, teleosts, and salmonids (see Figure 18.12). Abbreviation: MYr, millions of years ago.
plants some 150 million years ago, and that there have been several other duplications in other flowering plant lineages (Figure 18.18).

**Smaller duplications can also be identified in the human genome and other genomes**

Although the most recent whole-genome duplication in the human lineage occurred about 450 million years ago, the human genome has not been quiescent in the intervening period. Indeed, the opposite is true. One surprise arising from detailed analysis of the human genome sequence is the discovery that there has been extensive and frequent duplication of short segments of the genome in the relatively recent past. These are called **segmental duplications** or **low-copy repeats**. They are usually defined as duplications between 1 and 400 kb in length, with greater than 90% sequence identity, that are repeated up to 50 times in the genome, though often they occur as just two copies. Although segmental duplications sometimes contain repetitive DNA elements, such as long and short interspersed nuclear elements (LINEs and SINEs) or human endogenous retroviruses (HERVs) (Section 9.2), these are not themselves classified as segmental duplications because they have high copy numbers.

The impact of segmental duplication on the structure of the human genome is illustrated in **Figure 18.19**, which depicts the duplication events that have affected chromosome 1. When the parameters are set at >5 kb and >90% identity, many repeats are detected, some with two or more copies confined to chromosome 1, and others with duplicates on other chromosomes. Even at a much higher stringency of >40 kb and >99% identity, there are still multiple intrachromosomal duplications within chromosome 1 as well as interchromosomal duplications involving 11 other chromosomes. Duplications occur more frequently in the vicinity of the centromeres and in the subtelomeric regions, but there are still many repeats away from these areas.

The distribution of segmental duplications in chromosome 1 is fairly typical of the human genome as a whole, but there are significant differences when duplications in the mouse genome are examined. The proportions of the human and mouse genomes involved in segmental duplications are about the same, but
in the mouse genome the majority of duplicates form tandem repeats, whereas most human duplicates are located at a distant site in the same chromosome or on an entirely different chromosome. The human genome appears to be unusual in this regard: the tandem organization of segmental duplicates is more typical of vertebrates as a whole. A second, and particularly interesting, difference is that the human genome possesses a higher proportion of duplicates with >98% sequence identity than are present in the mouse (Figure 18.20). We have already learned that the similarity between a pair of duplicated sequences decreases over time due to the random accumulation of mutations. The high degree of similarity between the human duplicates therefore suggests that there was a recent burst of segmental duplication in the lineage leading to the human genome during the recent evolutionary past. Comparisons with segmental duplications in the chimpanzee genome suggest that this burst of activity began before the divergence of the great apes from their common ancestor and may have resulted in duplication of some genes involved in brain development, contributing to the special features of primate evolution.

In one regard, segmental duplications have a potentially deleterious impact on a genome. Recombination between a pair of intrachromosomal duplicates could result in deletion of the region between the repeated sequences, deletions generated in this way possibly giving rise to genetic disease. An increasing number of disorders are known to have originated in this way. An example is Prader–Willi syndrome, an incurable disorder associated with developmental and nutritional problems, the latter often leading to obesity. In about 70% of cases,
Prader–Willi syndrome is caused by deletion of a segment of the paternal version of chromosome 13, due to recombination between a pair of segmental duplicates. The deleted region contains various protein-coding and noncoding RNA genes, several of which are maternally imprinted. Deletion of the active, paternal versions of these genes therefore leads to a loss of function. Which gene or genes in the deleted region is directly responsible for the disease is not yet known, but attention is currently focused on the \textit{SNORD116} locus, which contains 29 copies of a small nucleolar RNA (snoRNA) gene. Patients who still have the region between the duplicates but have smaller deletions that remove the \textit{SNORD116} locus display many of the symptoms of Prader–Willi syndrome, suggesting that the snoRNA genes play a central role in the disease.

\textbf{Both prokaryotes and eukaryotes acquire genes from other species}

Duplication events are not the only way of adding new genes to a genome. Both prokaryotes and eukaryotes can also acquire genes from other species by lateral
gene transfer. In bacteria and archaea, lateral gene transfer is so frequent that it has a major impact on the gene catalogs of individual genomes, the resulting sharing of genes leading to a blurring of the distinction between prokaryotic species (Section 8.2).

The prevalence of lateral gene transfer in prokaryotes is due, at least in part, to the ease with which many prokaryotic species can take up DNA from their environment. Some species have cell membrane proteins specifically for this purpose. Eukaryotes do not have equivalent mechanisms for DNA uptake, and it is therefore not surprising that lateral gene transfer has been much less important during the evolution of eukaryotic genomes. Exactly how important the process has been is not clear: several reports of bacterial genes in eukaryotic genomes were subsequently ascribed to contamination of eukaryotic DNA preparations with bacteria, the latter contributing sequence reads that were mistakenly included in the eukaryotic genome assembly. Conversely, exclusion of bacterial reads from a eukaryotic assembly, simply on the grounds that these reads might be contaminants, risks missing genuine cases of gene transfer from bacteria to eukaryotes.

One example where the reported lateral transfer of genes into eukaryotic genomes is likely to be authentic concerns a group of plant-parasitic roundworms, including the potato cyst nematode *Globodera pallida*. In order to feed on sucrose provided by the host plant, these animals synthesize enzymes that degrade root cell walls and suppress the plant defense response. Several components of this specialized biochemical repertoire appear to have been acquired by the nematodes via lateral gene transfer from other species that inhabit the root biosphere. The genes for the cell-wall-degrading enzymes, for example, have closest similarity with homologous genes in the soil bacterium *Ralstonia*, and one gene for sucrose utilization is also probably bacterial in origin. Other plant-parasitic nematodes appear to have obtained cellulase genes from soil fungi.

A more extreme form of lateral gene transfer is well-documented in plant species. We have already seen how autoploidy can result in genome duplication in plants (see Figure 18.15). **Allopolyploidy**, which results from interbreeding between two different species, is also common and, like autoploidy, can result in a viable hybrid. Usually, the two species that form the allopolyploid are closely related and have many genes in common, but each parent will possess a few novel genes or at least distinctive alleles of shared genes. For example, the bread wheat *Triticum aestivum* is a hexaploid that arose by allopolyploidization between cultivated emmer wheat, *Triticum turgidum*, which is a tetraploid, and a diploid wild grass, *Aegilops squarrosa*. The wild grass nucleus contained novel alleles for the high-molecular-weight glutenin genes which, when combined with the glutenin alleles already present in emmer wheat, resulted in the superior properties for breadmaking displayed by the hexaploid wheats. Allopolyploidization can therefore be looked upon as a combination of genome duplication and interspecies gene transfer.

**Genome evolution also involves rearrangement of existing genes**

Duplications do not have to be of entire genes in order to have an impact on the genetic content of a genome. Duplications of gene segments containing one or more exons can alter the coding specificity of existing genes and even create entirely new genes. Rearrangements of this type can result in novel protein functions because most proteins are made up of structural domains, each domain comprising a segment of the polypeptide chain and hence encoded by a contiguous series of nucleotides (Figure 18.21). **Domain duplication** occurs when the

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**Figure 18.21** Each structural domain is an individual unit in a polypeptide chain and is coded by a contiguous series of nucleotides. In this simplified example, each α-helix and β-sheet in the polypeptide is looked upon as an individual structural domain. In reality, most structural domains comprise two or more secondary structural units.
gene segment coding for a structural domain is duplicated by unequal crossing over, replication slippage, or one of the other methods that we have considered for duplication of DNA sequences (Figure 18.22A). Duplication could result in the structural domain being repeated in the protein, which might itself be advantageous, for example by making the protein more stable. The duplicated domain might also change over time as its coding sequence becomes mutated, leading to a modified structure that might provide the protein with a new activity. Note that domain duplication causes the gene to become longer. Gene elongation appears to be a general consequence of genome evolution, as the genes of higher eukaryotes are longer, on average, than those of lower organisms. Alternatively, domain shuffling combines segments coding for structural domains from completely different genes to form a new coding sequence that specifies a hybrid or mosaic protein, one that would have a novel combination of structural features and might provide the cell with an entirely new biochemical function (Figure 18.22B).

Implicit in these models of domain duplication and shuffling is the need for the relevant gene segments to be separated so that they can themselves be rearranged and shuffled. This requirement has led to the attractive suggestion that exons might code for structural domains. With some proteins, duplication or shuffling of exons does seem to have resulted in the structures seen today. An example is provided by the α2 type I collagen gene of vertebrates, which codes for one of the three polypeptide chains of collagen. Each of the three collagen polypeptides has a highly repetitive sequence made up of repeats of the tripeptide glycine-X-Y, where X is usually proline and Y is usually hydroxyproline (Figure 18.23). The chicken α2 type I gene is split into 52 exons, 42 of which cover the part of the gene coding for the glycine-X-Y repeats. Within this region, each exon encodes a set of complete tripeptide repeats. The number of repeats per exon varies but is 5 (in 5 exons), 6 (23 exons), 11 (5 exons), 12 (8 exons), or 18 (1 exon). Clearly this gene could have evolved by duplication of exons leading to repetition of the structural domains.

Domain shuffling is illustrated by tissue plasminogen activator (TPA), a protein found in the blood of vertebrates that is involved in the blood clotting...
response. The TPA gene has four exons, each coding for a different structural domain (Figure 18.24). The upstream exon codes for a finger module that enables the TPA protein to bind to fibrin, a fibrous protein found in blood clots that activates TPA. This exon appears to be derived from a second fibrin-binding protein, fibronectin, and is absent from the gene for a related protein, urokinase, which is not activated by fibrin. The second TPA exon specifies a growth-factor domain that has apparently been obtained from the gene for epidermal growth factor and may enable TPA to stimulate cell proliferation. The last two exons code for kringle structures, which TPA uses to bind to fibrin clots; these kringle exons come from the plasminogen gene.

Type I collagen and TPA provide elegant examples of gene evolution but, unfortunately, the clear links that they display between structural domains and exons are exceptional and are rarely seen with other genes. Many other genes appear to have evolved by duplication and shuffling of segments, but in these the structural domains are coded by segments of genes that do not coincide with individual exons or even groups of exons. Domain duplication and shuffling still occur but presumably in a less precise manner and with many of the rearranged genes having no useful function. Despite being haphazard, the process clearly works, as indicated by, among other examples, the number of proteins that share the same DNA-binding motifs (Section 11.2). Several of these motifs probably evolved de novo on more than one occasion, but it is clear that in many cases the nucleotide sequence coding for the motif has been transferred to a variety of different genes.

One possible mechanism for moving gene segments around a genome is in association with transposable elements. The transposition of a LINE-1 element (Section 9.2) can occasionally result in a short piece of the adjacent DNA being transferred along with the transposon, a process called 3’-transduction because the transferred segment is located at the 3’-end of the element. LINE-1 elements are sometimes found in introns, so 3’-transduction could conceivably move downstream exons to new sites in a genome. Movement of exons and other gene segments might also be brought about by DNA transposons called Mutator-like transposable elements (MULEs), which are found in many eukaryotes but are especially common in plants. MULEs often contain within their DNA sequence segments of genes captured from the host genome. Transposition of a MULE would therefore move the captured segments to a new location. MULEs can collect segments of different genes as they travel around a genome, assembling new hybrid genes as they go. MULEs therefore provide an attractive way of driving gene evolution, but there are still several unanswered questions about their impact. In particular, it is not yet clear how frequently gene segments are able to escape from MULEs.
There are competing hypotheses for the origins of introns

Ever since introns were discovered in 1977, there has been debate about their origins. Many of the early ideas were influenced by the exon theory of genes, which suggests that introns were formed when the first DNA genomes were constructed, soon after the end of the RNA world. These genomes would have contained many short genes, each derived from a single coding RNA molecule and each specifying a very small polypeptide, perhaps just a single structural domain. These polypeptides would probably have had to associate together into larger multidomain proteins in order to produce enzymes with specific and efficient catalytic mechanisms (Figure 18.25). To aid the synthesis of a multidomain enzyme, it would have been beneficial for the enzyme’s individual polypeptides to become linked into a single protein, such as we see today. It is envisaged that this was achieved by splicing together the transcripts of the relevant minigenes, a process that was aided by rearranging the genome so that groups of minigenes specifying the different parts of individual multidomain proteins were positioned next to each other. In other words, the minigenes became exons and the DNA sequences between them became introns.

According to the exon theory of genes and other introns early hypotheses, all genomes originally possessed introns. But we know that bacterial genomes do not have GU-AG introns, so if these hypotheses are correct, then we must assume that for some reason introns became lost from the ancestral bacterial genome at an early stage in its evolution. The alternative introns late hypothesis avoids this problem by proposing that, to begin with, no genes had introns, these structures invading eukaryotic nuclear genomes and subsequently proliferating into the numbers seen today.

One reason the debate regarding the origin of GU-AG introns has continued for over 40 years is because evidence in support of the early and late hypotheses has been difficult to obtain and has often been ambiguous. A study of the vertebrate globin genes illustrates the problem. Initially it was concluded that a globin protein comprises four structural domains, the first corresponding to exon 1 of the globin gene, the second and third to exon 2, and the fourth to exon 3 (Figure 18.26). This pattern matches the expectations of the exon theory of genes. The prediction that there should be globin genes with another intron that splits the second and third domains was found to be correct when the leghemoglobin gene of soybean was shown to have an intron at exactly the expected position. Unfortunately, as globin genes were sequenced from other species, more introns were discovered, and most of these were located at positions that do not correspond to junctions between the protein domains.

There is still no consensus regarding the origins of introns, but the availability today of genome sequences from multiple species does at least enable greater amounts of data to be applied to the problem. Comparisons of intron positions in homologous genes from different species have been particularly informative. These have shown that there is a significant correlation between the positions of introns in fungi, plants, and animals, with over 25% of introns having the same position in genes from at least two of these three groups of organisms. Importantly, intron positions in fungi, plants, and animals are also shared with basal eukaryotes, ones whose lineages diverged from other eukaryotes at a very early evolutionary stage and which root back to the base of the eukaryotic phylogenetic tree. Examples are the excavates, a group of single-celled flagellated organisms including Trichomonas vaginalis, which causes the sexually transmitted disease
called trichomoniasis (Figure 18.27). The genomes of excavates contain introns, and many of those introns are at the same positions as in other eukaryotes. The implication is that the genome of the last eukaryotic common ancestor (LECA) contained multiple introns and that many of those introns have been retained at the same positions in the genes of modern-day organisms.

It has also been discovered that the components of the spliceosome, the structure that removes introns from eukaryotic pre-mRNA (Section 12.4), are very similar in different groups of eukaryotes, suggesting that the splicing process evolved at an early stage in evolution and has not changed greatly since then. On the basis of these various strands of evidence, many researchers now favor a hypothesis for the origin of introns that is intermediate between the early and late schemes. The absence of GU-AG introns in prokaryotes is looked on as a major difficulty for the exon theory of genes, and the very earliest genomes are therefore considered to have lacked introns of this type. Those early genomes might, however, have contained ancestors of the retroelements called group II introns, which are present in organelle genomes and are known in a few prokaryotes. The splicing pathway for group II introns is very similar to that of GU-AG introns, but group II introns do not require a spliceosome, as they are ribozymes and so are able to self-splice. The group II intron sequence encodes a reverse transcriptase, which enables the excised intron to be copied into DNA and to reinsert at a second position in its host genome, by a process called retrohoming. The model now emerging is one in which group II introns were present in the genomes of the endosymbionts that formed the precursors of mitochondria and chloroplasts in the very first eukaryotic cells (Section 8.3). These introns escaped from their organelles and invaded the early eukaryotic nucleus, where they proliferated by retrohoming. The resulting structures evolved into the large numbers of GU-AG introns that were present in LECA. The ability of modern group II introns to transpose by retrohoming in human cells in experimental systems indicates that the proliferation of group II introns in early eukaryotic nuclear genomes was possible. Other group II introns are thought to have evolved into non-LTR retroelements such as LINEs. Introns are therefore neither early nor late: their genesis accompanied the origins of the first eukaryotic cells.

The evolution of the epigenome

In this overview of the evolution of genome complexity, we have so far concentrated solely on the DNA sequence features of genomes. We must not forget that, in eukaryotic cells, the activity of the genome is inextricably linked with the pattern of histone modifications, nucleosome positioning, and DNA methylation that is present in different functional domains (Chapter 10). These patterns determine which parts of the genome are accessible to the RNA polymerases that transcribe genes into RNA and, in particular, determine which regions of the genome are
packaged into heterochromatin and hence are silenced in a particular cell or cell lineage. The combination of processes including histone modification, nucleosome positioning, and DNA methylation that regulate genome expression are sometimes referred to as the epigenome, reflecting the role that these processes play in the genome’s response to environmental and other stimuli that give rise to epigenetic effects. Epigenetic effects are changes in phenotype that occur not because of alterations in the genome sequence but because of alterations in the way in which the genome is expressed.

Evolutionary geneticists are starting to ask how the epigenome evolved or, to phrase the question slightly differently, how the genome evolved its ability to respond to external and internal signals through the use of chromatin modification as a means of silencing those genes that are not needed in a particular tissue or at a particular developmental stage. So far, much of the work in this area has centered on comparative epigenomics, which explores the extent to which the equivalent regions of two different genomes display the same pattern of chromatin modification, the resulting data being used to infer the extent to which epigenomic effects are programmed by the DNA sequence. It has already been established that, within a single genome, both copies of a segmental duplication have similar modification patterns. In the human genome, for example, the pattern of methylation within CpG islands tends to be retained after duplication of the segment containing that island, even when the daughter copy has been inserted at a new genomic location distant from the genes whose expression the CpG island normally influences (Figure 18.28). This holds true even if the sequence surrounding the island in the daughter duplicate has begun to degrade. Similar patterns of histone modification and nucleosome positioning are also seen in duplicated regions. The implication is that, at least in a single genome, the DNA sequence plays a major role in determining the pattern of chromatin modification.

When genomes of different species are compared, the analysis becomes more complicated because of the possibility that equivalent genome regions have nonidentical functions in different organisms. In a comparison of human, chimpanzee, bonobo, gorilla, and orangutan genomes, many examples were found of homologous genes with different methylation patterns, but these included genes with developmental or neurological functions, whose altered methylation patterns might reflect differences in the associated physiological processes in the different primate species. Broader-based studies employing phylogenetic methods have shown that the degree of relatedness between pairs of species as inferred from their methylation patterns is similar to the degree of relatedness as measured by DNA sequence comparisons. A correlation between genome methylation and genome sequence is therefore evident, even if that correlation may be less easy to discern when short regions of the genome are examined.

Understanding the evolution of the epigenome is challenging, but it is a challenge worth tackling because it provides one of the keys to linking the evolution of genomes with the evolution of phenotypes. Because of the ease with which DNA sequences can now be obtained, we still tend to think of a genome predominantly as a string of As, Cs, Gs, and Ts, displayed in a genome browser with the interesting sequence features highlighted. In order to move on to the next era of genome biology, we must increasingly study the genome as a component of the cell and organism in which it resides. Describing the way in which the epigenome has evolved alongside the genome is one of the first steps toward the establishment of this new and more integrated approach to genome biology.

### 18.3 Genomes: The Last 6 Million Years

Bishop Samuel Wilberforce once famously asked Thomas Huxley, one of Charles Darwin’s supporters, if his descent from a monkey was on his mother or father’s side. The answer is both: humans and chimpanzees are descended from a common ancestor that lived around 6 million years ago. Since the split, the human lineage has embraced two genera, *Australopithecus* and *Homo*, and a number of species, not all of which were on the direct line of descent to *Homo sapiens*.
The result is us, a novel species in possession of what are, at least to our eyes, important biological attributes that make us very different from all other animals. So how different are we from the chimpanzees?

The human genome is very similar to that of the chimpanzee

A draft of the chimpanzee genome sequence was completed in 2005. Initial comparisons with the human genome revealed the extensive similarity expected for two species that diverged just 6 million years ago. The degree of nucleotide sequence identity within the coding DNA is greater than 98.5%, with 29% of the genes in the human genome coding for proteins whose amino acid sequences are identical to the sequences of their counterparts in chimpanzees. Gene order is almost the same in the two genomes, and the chromosomes have very similar appearances. At this level, the most dramatic difference is that human chromosome 2 is two separate chromosomes in chimpanzees (Figure 18.30), so chimpanzees, as well as other apes, have 24 pairs of chromosomes whereas humans have just 23 pairs.

Even in the noncoding regions of the human and chimpanzee genomes, the nucleotide identity is rarely less than 97%. This figure does, however, imply a greater degree of similarity than is the case, in part because it refers to the alignment between the sequences and does not take account of the presence of indels. These are positions where one genome contains a segment of DNA that is absent in the second genome, and which could be an insertion in one genome or a deletion in the other (Figure 18.31). Because of indels, about 1.5% of both the human and chimpanzee genomes are made up of DNA sequence that is unique and not possessed by the other species, but most of the individual indels are quite short, and few if any are likely to have a significant impact on the function of the genome in which they reside. There are also differences in repetitive DNA content, with 5000 unique retroelement insertions in the human genome and 2500 unique insertions in chimpanzee, with the species-specific patterns of Alu insertions being particularly distinctive. These differences are interesting for what they tell us about the evolution of the repetitive DNA content of a genome, and they also mean that the human and chimpanzee genomes can potentially be affected by homologous recombination events in different ways, but as with indels, the retroelement insertions are unlikely to be the Rosetta Stone that explains why humans are special.

For several years, progress in understanding the unique features of the human genome was frustratingly slow. This trend is now starting to reverse, and although we are still some way from knowing how our genome specifies the distinctive human characteristics, there are several open lines of research that are being actively pursued. One of these concerns the gene for the FOXP2 transcription factor. Defects in this protein result in the human disability called dysarthria, characterized by a difficulty in articulating speech, suggesting that the FOXP2 gene...
might be involved in the human ability for language. There are two amino acid differences between the FOXP2 proteins in humans and chimpanzees, and there is evidence that there has been positive selection of the gene in the human lineage. The latter is an important observation because genes under positive selection are likely to be ones that have contributed to the recent phenotypic evolution of the species. Further indications of the importance of FOXP2 were obtained from studies of mice whose own FOXP2 genes had been humanized by directed mutations that introduced the amino acid substitutions coded in the human gene. These mice had increased neuron growth in the striatum of the basal ganglia, the part of the forebrain associated with speech in humans. Mice use a range of acoustic vocalizations to communicate with one another: most remarkably, the mice containing the humanized FOXP2 gene generated patterns of ultrasonic vocalizations not recorded from other mice.

Other investigations are suggesting that partial and complete gene duplications might have a role in specifying the human phenotype. In particular, a protein domain called DUF1220 has undergone amplification in the human genome. This domain comprises 65 amino acids that are coded by two adjacent exons. The exon pair has a copy number of 272 in the human genome, compared with 126 in chimpanzee and just one in mouse. DUF1220 domains are mainly found in the neuroblastoma breakpoint family of genes, some members of which have been duplicated in the human genome and some of which have undergone expansion because of intragenic amplification of the DUF1220 exons. The functions of the members of the gene family are not known, but they have been associated with brain size and cortical neuron number.

Segmental duplications have also been implicated in the generation of human-specific genome features that might be relevant to brain development. The SRGAP2 gene, which is involved in cortical development, has undergone two consecutive duplications in the human lineage, giving rise to daughter and granddaughter copies of the parent gene. The granddaughter duplicate codes for a truncated version of the SRGAP2 protein, which can form a heterodimer with the full-length product of the parent gene. Unlike the homodimers formed by two complete SRGAP2 proteins, the heterodimer is nonfunctional. Formation of the heterodimer therefore sequesters full-length SRGAP2 copies, modulating the activity of the protein. The segmental duplication giving rise to the truncated granddaughter gene is thought to have occurred between 2 and 3 million years ago, just before the emergence of the Homo genus (see Figure 18.29), at a period when the human brain began to increase in size.

As well as changes to the DNA sequence, alterations in gene expression patterns could quite possibly underlie the unique capabilities of the human genome. At least 100 genes, and possibly many more, give rise to human-specific transcripts via novel alternative promoters or splicing pathways. Characterization of these and other changes in gene expression is less advanced than the research described above, but is likely to make an increasing contribution to our understanding of the species-specific features of the human genome during the next few years.

**Paleogenomics is helping us understand the recent evolution of the human genome**

One of the latest advances in genomics research is the ability to obtain sequences from extinct species by analysis of the ancient DNA fragments that are sometimes preserved in bones and other remains. We have already seen how ancient DNA has been used to obtain a complete sequence of the genome of Neanderthals, the extinct hominins who inhabited many parts of Europe and Asia between 200,000
and 30,000 years ago (Section 4.4). Neanderthals are, arguably, a subspecies of *H. sapiens*, and it is therefore no surprise that the Neanderthal genome is almost identical with our own genome, with 99.7% nucleotide similarity. What can such a closely related genome tell us about human evolution?

The most remarkable insight into our past that has been revealed by this branch of paleogenomics is that our ancestors interbred with Neanderthals. Evidence of interbreeding was obtained when comparisons were made between the Neanderthal genome and the genomes of present-day humans from Europe and Africa. If there had been no interbreeding, then we would expect the genomes of modern Europeans and Africans to display identical degrees of divergence when compared with the Neanderthal genome. In fact, the divergence between the Neanderthal and modern European genomes is slightly less than that between the genomes of Neanderthals and modern Africans, suggesting that some Neanderthal DNA has found its way into the genomes of modern Europeans. This finding indicates that there was a small amount of interbreeding between Neanderthals and *H. sapiens* during the 15,000 years or so that they were co-resident in Europe.

Ancient DNA sequencing has also identified an Asian version of Neanderthals, called the Denisovans. The Denisovan genome sequence gives additional evidence for interbreeding with *H. sapiens*, in this case specifically with the ancestors of modern inhabitants of Oceania. The most recent estimates are that 1.5–2.1% of the DNA of modern humans from outside Africa is of Neanderthal origin, and 3.0–6.0% of the genomes of modern inhabitants of Oceania is derived from Denisovans (Figure 18.32). There is also evidence for interbreeding between Neanderthals and Denisovans and between Denisovans and an unidentified extinct type of human.

Has the influx of Neanderthal and Denisovan DNA had any significant effect on the human genome? Of course, the three genomes have the same genes, but it is possible that Neanderthals and/or Denisovans possessed alleles that were not present in the ancestral human population, and these alleles might have been acquired by humans via interbreeding. One intriguing possibility is that the ability of people from Tibet to withstand low-oxygen conditions, such as found in the high altitudes of the Tibetan plateau, was inherited from Denisovans. This hypothesis is prompted by the discovery that an allele of the *EPAS1* gene that is common in Tibetans is absent in human genomes from all other parts of the world. The *EPAS1* protein is a transcription factor that controls the expression of genes for proteins that help the body tolerate anoxic conditions. The allele possessed by Tibetans, although absent in non-Tibetans, is present in the Denisovan genome, raising the possibility that its presence in Tibetans is the outcome of interbreeding between Denisovans and *H. sapiens* in the distant past.

### 18.4 Genomes Today: Diversity in Populations

We conclude our survey of genome evolution by exploring the snapshot of evolutionary time represented by the present day. Genomes continue to evolve through mutation and recombination, creating new variants of the DNA sequence and new arrangements of different sequence units. One outcome of this continuing process is that the genomes of most species display intraspecific variability. The human genome, for example, is not a single DNA sequence. Every individual, with the possible exception of identical twins, possesses his or her own unique genome sequence, the uniqueness determined by the identities of the nucleotides present at each of the 10 million or so single-nucleotide polymorphism (SNP) positions in the genome, as well as the repeat lengths of each of the 3 million simple sequence length polymorphisms (SSLPs). A small minority of these personal variations fall in genes and so specify the phenotype of the individual, generating the characteristic features that distinguish one person from another. The vast majority of the polymorphisms are silent, but all of the variations, whether in coding or intergenic regions, provide an evolutionary record of the genome. Comparisons between those variations enable the relationships between individuals and
groups of individuals to be inferred. In this section we will explore how the diversity of genome sequences is used in research and in biotechnology. We will examine three case studies, on the origin of HIV/AIDS, the prehistoric migrations of *H. sapiens*, and the breeding of new varieties of crop plants.

**The origins of HIV/AIDS**

The global epidemic of human immunodeficiency virus infection and acquired immune deficiency syndrome (HIV/AIDS) has touched everyone's lives. AIDS is the final stage of infection by human immunodeficiency virus 1 (HIV-1), a retrovirus (Section 9.1) that attacks cells involved in the immune response. The suppression of the immune system that occurs in patients who progress to this final stage increases the risk of opportunistic infections and tumor growth that can lead to death.

The demonstration in the early 1980s that HIV-1 is responsible for AIDS was quickly followed by speculation about the origin of the disease. Speculation centered around the discovery that similar immunodeficiency viruses are present in primates such as the chimpanzee, sooty mangabey, mandrill, and various monkeys. These simian immunodeficiency viruses (SIVs) are not pathogenic in their normal hosts, but it was thought that if one had become transferred to humans, then within this new species the virus might have acquired new properties, such as the ability to cause disease and to spread rapidly through the population.

Retrovirus genomes accumulate mutations relatively quickly because reverse transcriptase, the enzyme that copies the RNA genome contained in the virus particle into the DNA version that integrates into the host genome (see Section 9.1), lacks an efficient proofreading activity and so tends to make errors when it carries out RNA-dependent DNA synthesis. This means that retrovirus genomes that diverged quite recently display sufficient nucleotide dissimilarity for an evolutionary analysis to be carried out.

Various methods can be used to deduce the evolutionary relationships between genomes. If the genome sequences are relatively short, as is the case with retroviruses, and they have not been rearranged by recombination, then a **phylogenetic tree** is usually constructed. The tree comprises a set of external nodes, each representing one of the genome sequences, linked by branches to internal nodes representing ancestral genome sequences (Figure 18.33). The lengths of the branches indicate the degrees of difference between the sequences represented by the nodes. The tree is generated by aligning the various sequences that are being examined (Figure 18.34) and then converting this multiple alignment into numerical data that can be analyzed mathematically in order to produce the tree. The simplest method involves conversion of the sequence data into a **distance matrix**, which is a table showing the evolutionary distances between all pairs of sequences, calculated from the number of nucleotide differences between each pair. The matrix is used by the tree-building software to establish the lengths of the branches connecting pairs of sequences in the tree that is drawn.

What do we discover when phylogenetic analysis is applied to HIV and SIV genome sequences? The tree that is obtained has a number of interesting features (Figure 18.35). First, it shows that different samples of HIV-1 have slightly different sequences, with the samples as a whole forming a tight cluster, almost a starlike
pattern, that radiates from one end of the tree. This starlike topology implies that the global AIDS epidemic began with a very small number of viruses, perhaps just one, which have spread and diversified since entering the human population. The closest relative to HIV-1 among primates is the SIV of chimpanzees, the implication being that this virus jumped across the species barrier between chimpanzees and humans and initiated the HIV/AIDS epidemic. However, this epidemic did not begin immediately: a relatively long, uninterrupted branch links the center of the HIV-1 radiation with the internal node leading to the relevant SIV sequence, suggesting that, after transmission to humans, HIV-1 underwent a latent period when it remained restricted to a small part of the global human population, presumably in Africa, before beginning its rapid spread to other parts of the world. Other primate SIVs are less closely related to HIV-1, but one, the SIV from sooty mangabey, clusters in the tree with the second human immunodeficiency virus, HIV-2. It appears that HIV-2 was transferred to the human population independently of HIV-1 and from a different simian host. HIV-2 is also able to cause HIV/AIDS but has not, as yet, become globally epidemic.

An intriguing addition to the HIV/SIV tree was made in 1998, when the RNA of an HIV-1 isolate from a blood sample taken in 1959 from an African male was sequenced. The RNA was highly fragmented and only a short sequence could be obtained, but this was sufficient for the sample to be placed on the phylogenetic tree (see Figure 18.35). The sequence, called ZR59, attaches to the tree by a short branch that emerges from near the center of the HIV-1 radiation. The positioning indicates that the ZR59 sequence represents one of the earliest versions of HIV-1 and shows that the global spread of HIV-1 was already underway by 1959. A later and more comprehensive analysis of HIV-1 sequences has suggested that the spread began in the period between 1915 and 1941, with a best estimate of 1931. Pinning down the date in this way has enabled epidemiologists to begin an investigation of the historic and social conditions that might have been responsible for the start of the HIV/AIDS epidemic.

The first migrations of humans out of Africa

Our second case study illustrating the ways in which intraspecific genome diversity is used in research concerns the prehistoric migrations that resulted in the expansion of *H. sapiens* from our evolutionary homeland in Africa to our current global distribution. Paleoanthropologists refer to our species as **anatomically modern humans**, a term that sidesteps debates about the precise evolutionary relationships between us and our extinct ancestors. The oldest fossils identified as belonging to anatomically modern humans were found at Omo in Ethiopia and date to 195,000 years ago. Other early fossils have been found at Herto, also in Ethiopia, from 160,000 years ago, at Laetoli in Tanzania from 120,000 years ago, and at Border Cave, South Africa, from about 110,000 years ago. The earliest fossil remains of modern humans outside Africa were discovered in the Qafzeh and Skhul caves near Nazareth in modern Israel. These fossils are dated to 90,000–100,000 years ago, but the people who lived in the caves at that time are unlikely to have been part of the first major migration of humans into Eurasia. The caves were reoccupied by Neanderthals at a later date, and modern humans were not seen there again for some time. It seems that the Qafzeh and Skhul people either were part of a failed attempt at colonization or merely represent an extension of the African population into Asia during a warm period.
Initial attempts to follow the early migrations of humans out of Africa made use of mitochondrial DNA. In mammals such as humans, the molecular clock for mitochondrial DNA is faster than that for DNA in the nucleus, probably because mitochondria lack many of the DNA repair systems that operate in the nucleus, enabling a greater proportion of the mutations that occur to be retained, each of these retained mutations resulting in a substitution, a permanent change in the nucleotide sequence. This means that within the human population there are a variety of mitochondrial DNA sequences, which on the basis of shared substitutions can be divided into about 275 haplogroups. Each haplogroup can then be further subdivided into haplotypes, all of which have the characteristic substitutions that define the haplogroup, along with private substitutions that distinguish each haplotype (Figure 18.36). The date when a haplogroup originated can be deduced by estimating its coalescence time. The reasoning is that the greater the diversity among the haplotypes within the haplogroup, then the larger the number of substitutions that has occurred, and the more ancient the coalescence time.

Mitochondrial DNA analysis has suggested that the first modern human migration out of Africa left from Ethiopia, further south than the area of modern Suez that forms the physical link between Africa and Asia. The basis for this hypothesis is as follows. All the mitochondrial DNA haplogroups that are known in the modern human population can be linked together in a phylogenetic tree displaying their sequence relationships. Within this tree, all of the haplogroups that are common today in Africa are clustered together, with just two links connecting them with the remainder of the tree. These two links are between haplogroup L3 on the African side and M and N on the non-African side. Coalescence analyses suggest that the M and N haplogroups originated 60,000–70,000 years ago. We believe that at that time the L3 haplogroup was present mainly in East Africa. The most direct route from East Africa to Asia is across the Bab-el-Mandeb strait, at the entrance of the Red Sea, to the southern coast of Arabia (Figure 18.37). We therefore conclude that the initial migration out of Africa was from Ethiopia to southern Arabia and occurred some 60,000–70,000 years ago.

Mitochondrial DNA studies therefore provide a model for the first human migrations from Africa, but to what extent is this a true interpretation of the past? The mitochondrial genome is just a small part of the human DNA, and its evolutionary pattern is unusual in that it is inherited solely through the maternal line and does not recombine with paternal DNA during reproduction. Hypotheses based on mitochondrial DNA have been criticized as narratives that may or may not give a true indication of the past evolutionary history of a species or population. So do studies of the complete human genome confirm or contradict the scheme suggested by mitochondrial DNA? This is a rapidly developing area of research, with genome sequences being collected from more and more individuals from different parts of the world, and increasingly sophisticated methods being developed for the analysis of these sequences. While the data sets and methodologies are still reaching maturity, it is inevitable that different projects will yield conflicting results. But the growing evidence suggests that the single wave of migration out of Africa around 60,000–70,000 years ago indicated by the mitochondrial studies might be correct. One recent project compared over 4 million SNPs in genomes from Australia, Africa, Europe, and East Asia. For each SNP, data from primate
The diversity of plant genomes is an aid in crop breeding

The genomes of plant species also display intraspecific variability. Geneticists have used this variability to study the migrations of plants, in particular the expansion of species from the glacial refugia that they occupied during the Ice Ages. These refugia were relatively warm geographical areas where plants, insects, and animals were able to survive the harsh conditions of the glaciations, prior to recolonizing the adjacent regions when the climate improved. In Europe, there were refugia in Iberia, Italy, and the Balkans, and in North America refugia are believed to have been present in the coastal regions of Alaska and British Columbia.

Population genomics is proving particularly important in research with crop plants. Agriculture began independently in at least three parts of the world, Mesoamerica, Southwest Asia, and Southeast Asia, at about the same time, 10,000 years ago. In the Fertile Crescent of Southwest Asia, early farmers grew barley, einkorn and emmer wheat, lentils, peas, chickpeas, and bitter vetch. Barley and wheat cultivation then entered southeast Europe about 9000 years ago and, over the next 3000 years, spread through the continent by two principal routes. One route followed the Danube and Rhine valleys through central Europe and into the north European plain, and the second was a coastal route around Italy and Iberia to northwest Europe. By 6000 years ago, agriculture had reached all but the most northern parts of the continent. This human-driven spread of agriculture exposed crops to a variety of environmental conditions, many of which were quite different from those present in the natural ranges of the wild progenitor species in Southwest Asia. These environmental conditions included not only changes to the climate but also exposure to new insect and fungal pests, as well as growth on new types of soil. The barley and wheat genomes evolved by adapting to the local environments in which the plants were grown, resulting in what botanists call landraces, populations of crop plants that are locally adapted and which were grown throughout Europe until the twentieth century, when they gradually were replaced with the products of modern crop breeding. The landraces were not entirely lost, however, as ardent plant collectors scoured the continent, collecting seeds that have been propagated in germplasm collections, from which samples of these landraces can be obtained today.

The adaptations displayed by landraces are now being utilized in breeding programs that aim to improve modern barley and wheat cultivars so that these are better able to withstand the changing climate. A problem in many of these breeding programs is that the genes that specify desirable traits, such as tolerance to drought or resistance to pests, are unidentified. This means that it is not possible to use DNA cloning techniques to transfer the interesting genes from one plant to another. Crop improvement therefore still depends very much on conventional breeding methods, in which two plants with different characteristics are crossed in the hope that at least some of the progeny will inherit the combination of desirable traits from the parents. This approach works, but it is a lengthy process that typically involves thousands of seedlings being generated whose characteristics are often not known until they have been grown to maturity and subjected to a
range of physiological and biochemical tests. Genomics is now speeding up the process by making possible the approach called marker-associated selection. Although the genes for a trait might be unknown, SNPs whose positions in the genome are close to the relevant genes can usually be identified by an association study, which searches for SNP alleles that are consistently present in plants with a particular trait (Figure 18.39). Being closely linked to the genes of interest, these SNPs cosegregate with those genes when different plants are crossed. This means that the SNP alleles can be used as proxies for the genes, using an approach called marker-assisted selection. A cross is carried out, and when the seedlings are obtained, leaf samples are taken, DNA is prepared, and the diagnostic SNPs are typed by DNA chip technology or some other appropriate high-throughput method (Section 3.2). Those seedlings that contain SNP alleles associated with the desirable traits of the two parents are retained, and those that do not display the diagnostic SNP pattern are discarded. The time-consuming growth and testing procedures can then be carried out just with that subset of seedlings that is likely to be of value.

**SUMMARY**

- It is thought that the first polynucleotides to evolve, several billion years ago, were made of RNA rather than DNA.
- These RNA molecules probably combined a self-replicating ability with some enzymatic activity, and it is possible that enclosure of these in simple lipid envelopes gave rise to the progenitors of the first cells.
- DNA probably evolved as a more stable version of the initial RNA protogenomes.
- Life forms based on coding molecules other than DNA and RNA are unknown but are thought to be possible.
- Gene duplication is an important event that can result in a genome acquiring new genes. This is illustrated by the globin gene superfamily, which arose...
by a series of gene duplications whose pattern and timing can be inferred by making comparisons between the sequences of the globin genes in existence today.

- Duplication events have also played an important role during evolution of the homeotic selector genes that specify the body plan of eukaryotes. These duplication events can be linked to whole-genome duplications that have occurred in the past.
- Smaller duplications, 1–400 kb in length, have occurred regularly in the recent evolution of the human genome.
- Lateral gene transfer results in the acquisition of genes from other species. This has been a regular event in the evolution of prokaryotic genomes but is much less common in eukaryotes, except possibly in plants, which can form new polyploids by fusion of gametes from related species.
- Duplications and rearrangements of gene segments have resulted in new combinations of exons. Exons can also be transported around a genome by attachment to transposable elements.
- The origins of introns are unclear, but current evidence suggests that introns proliferated in nuclear genomes shortly after the evolution of the first eukaryotic cell.
- The epigenome has also evolved over time, but we are only just beginning to understand the processes that were involved.
- Six million years ago the human and chimpanzee lineages diverged. The genomes of humans and chimpanzees still display considerable sequence similarities, with many genes giving rise to identical protein products. It is thought that point mutations in key genes, segmental duplications, and changes in gene expression patterns are responsible for the features of the human genome that give rise to the particular characteristics of our species.
- The ability to sequence ancient DNA from preserved fossils has revealed that Homo sapiens interbred with Neanderthals and Denisovans.
- The diversity of genome sequences within a species enables recent evolutionary events to be studied. The origins of HIV/AIDS have been deduced from phylogenetic studies of HIV and SIV genome sequences.
- The diversity of modern human genome sequences enables past migrations of human populations to be mapped.
- Plant breeders attempt to combine desirable characteristics that have evolved in different populations of a single crop species, for example to develop high-yielding crops that are better able to withstand climate change.

### SHORT ANSWER QUESTIONS

1. How might some of the amino acids, nucleotide bases, and sugars have been synthesized before life evolved?
2. Provide a timeline for the evolution of living organisms from the formation of the Earth to the appearance of the first hominins.
3. Describe the evidence indicating that the content of the human genome has been influenced by past gene duplication events.
4. Outline the mechanisms by which gene duplication might occur.
5. Describe the evidence indicating that the content of the human genome has been influenced by past whole-genome duplication events.
6. How have segmental duplications affected the genomes of primates?
7. Give examples of polyploidization in plants.
8. In what ways can new genes arise by rearrangement of existing genes?
9. What is the exon theory of genes? How does this theory compare with modern theories for the origin of introns?
10. What are the key differences between the human and chimpanzee genomes?
11. What has ancient DNA sequencing revealed regarding the relationships between *Homo sapiens* and extinct types of human?
12. Describe how genome sequencing has been used to study the origin of HIV/AIDS.

**IN-DEPTH PROBLEMS**

1. Are the examples of domain duplication and domain shuffling given on pp.445–447 special cases or are they representative of genome evolution in general?
2. One of the initial publications of the draft human genome sequence (International Human Genome Sequencing Consortium [2001] Initial sequencing and analysis of the human genome. *Nature* 409: 860–921) suggested that between 113 and 223 human genes might have been acquired from bacteria by lateral gene transfer. Subsequently it was concluded that this interpretation is incorrect and these genes are not bacterial in origin. What was the evidence that supported lateral transfer of these genes and why was this evidence subsequently discounted?
3. To what extent do you believe it will be possible to determine the genetic basis to the special attributes of humans from comparisons between the genome sequences of humans and other primates?
4. How reliable are molecular clocks?
5. What is the potential of ancient DNA in studies of human evolution?

**FURTHER READING**

**The RNA world and the origins of genomes**


**Gene, genome, and segmental duplications**


Evolution of the epigenome


Humans and other primates


Paleogenomics and past human migrations


Origins of HIV/AIDS

Glossary

2-Aminopurine A base analog that can cause mutations by replacing adenine in a DNA molecule.
2’-Deoxyribose The sugar component of a deoxyribonucleotide.
2 μm Plasmid A plasmid found in the yeast Saccharomyces cerevisiae and used as the basis for a series of cloning vectors.
-25 Box A component of the bacterial promoter.
3’-OH terminus The end of a polynucleotide that terminates with a hydroxyl group attached to the 3’-carbon of the sugar.
3’-Transduction Transfer of a segment of genomic DNA from one place to another caused by movement of a LINE element.
3’-Untranslated region The untranslated region of an mRNA upstream of the termination codon.
3’→5’ Exonuclease An exonuclease activity that removes nucleotides sequentially from the 3’-end of a polynucleotide.
30 nm Fiber A relatively unpacked form of chromatin consisting of a possibly helical array of nucleosomes in a fiber approximately 30 nm in diameter.
454 Sequencing A next-generation method that makes use of the pyrosequencing procedure.
5-Bromouracil (5-bU) A base analog that can cause mutations by replacing thymine in a DNA molecule.
5’-P terminus The end of a polynucleotide that terminates with a mono-, di-, or triphosphate attached to the 5’-carbon of the sugar.
5’-Untranslated region The untranslated region of an mRNA downstream of the termination codon.
5’→3’ Exonuclease An exonuclease activity that removes nucleotides sequentially from the 5’-end of a polynucleotide.
(6–4) Lesion A dimer of two adjacent pyrimidine bases in a polynucleotide, formed by ultraviolet irradiation.
(6–4) Photoproduction photolyase An enzyme involved in photoreactivation repair.
7SK RNA A component of a protein–RNA complex that is indirectly involved in the control of transcription elongation.
7SL RNA A component of the eukaryotic signal recognition particle.
A-DNA A structural configuration of the double helix, which is present but not common in cellular DNA.
Ab initio gene prediction Identification of putative genes by ORF scanning of a DNA sequence.
ABC model A model for the genetic control of the identity of the whorls in a flower.
Ac/Ds transposon A DNA transposon of maize.
Acceptor arm Part of the structure of a tRNA molecule.
Acceptor site The splice site at the 3’-end of an intron.
Accessory genome The component of a prokaryotic genome comprising all those genes not present in the core genome.
Acridine dye A chemical compound that causes a frameshift mutation by intercalating between adjacent base pairs of the double helix.
Acylation The attachment of a lipid side chain to a polypeptide.
Ada enzyme An Escherichia coli enzyme that is involved in the direct repair of alkylation mutations.
ada regulon The set of genes that is switched on by the Ada enzyme.
Adaptor A synthetic, double-stranded oligonucleotide used to attach sticky ends to a blunt-ended molecule.
Adenine A purine base found in DNA and RNA.
Adenosine deaminase acting on RNA (ADAR) An enzyme that edits various eukaryotic mRNAs by deaminating adenosine to inosine.
Adenylate cyclase The enzyme that converts ATP to cyclic AMP.
Affinity chromatography A column chromatography method that makes use of a ligand that binds to the molecule being purified.
Agarose gel electrophoresis Electrophoresis carried out in an agarose gel and used to separate DNA molecules between 100 bp and 50 kb in length.
Alarmone One of the stringent response activators, ppGpp and pppGpp.
Alkaline phosphatase An enzyme that removes phosphate groups from the 5’-ends of DNA molecules.
Alkylating agent A mutagen that acts by adding alkyl groups to nucleotide bases.
Allele One of two or more alternative forms of a gene.
Allele frequency The frequency of an allele in a population.
Allele-specific oligonucleotide (ASO) hybridization The use of an oligonucleotide probe to determine which of two alternative nucleotide sequences is contained in a DNA molecule.
Allopolyploidy The result of interbreeding between two different species, giving a polyploid nucleus derived from fusion between gametes of those two species.
Alphoid DNA The tandemly repeated nucleotide sequences located in the centromeric regions of human chromosomes.
Alternative exons An alternative splicing scenario where the mRNA contains either of a pair of exons, but not both at the same time.
Alternative polyadenylation The use of two or more different sites for polyadenylation of an mRNA.
**Alternative promoter** One of two or more different promoters acting on the same gene.

**Alternative site selection** An alternative splicing scenario where the usual donor or acceptor site is ignored and a second site used in its place.

**Alternative splicing** The production of two or more mRNAs from a single pre-mRNA by joining together different combinations of exons.

**Alu** A type of SINE found in the genomes of humans and related mammals.

**Alu-PCR** A clone fingerprinting technique that uses PCR to detect the relative positions of Alu sequences in cloned DNA fragments.

**Amino acid** One of the monomeric units of a protein molecule.

**Amino-terminus** The end of a polypeptide that has a free amino group.

**Aminoacyl or A site** The site in the ribosome occupied by the aminoacyl-tRNA during translation.

**Aminoacyl-tRNA synthetase** An enzyme that catalyzes the aminoacylation of one or more tRNAs.

**Aminoacylation** Attachment of an amino acid to the acceptor arm of a tRNA.

**Amplification refraction mutation system (ARMS test)** A technique for SNP typing, in which PCR is directed by a pair of primers, one covering the position of the SNP.

**Analytical protein array** A type of protein array that is used in protein profiling.

**Anaphase-promoting complex/cyclosome (APC/C)** A ubiquitin ligase that directs degradation of proteins that are active late in the cell cycle, so that these proteins are not carried over to the daughter cells resulting from mitosis.

**Anatomically modern humans** Fossil members of our own species.

**Ancestral allele** The version of an allele that is possessed by the common ancestor of a group of organisms.

**Ancient DNA** DNA preserved in ancient biological material.

**Annealing** Attachment of an oligonucleotide primer to a DNA or RNA template.

**Antibody array** A protein array that carries a series of antibodies.

**Anticodon** The triplet of nucleotides, at positions 34–36 in a tRNA molecule, that base-pairs with a codon in an mRNA molecule.

**Anticodon arm** Part of the structure of a tRNA molecule.

**Antigen** A substance that elicits an immune response.

**Antitermination** A bacterial mechanism for regulating the termination of transcription.

**Antiterminator protein** A protein that attaches to bacterial DNA and mediates antitermination.

**AP (apurinic/apyrimidinic) site** A position in a DNA molecule where the base component of the nucleotide is missing.

**AP endonuclease** An enzyme involved in base excision repair.

**Apoptosis** Programmed cell death.

**Archaea** One of the two main groups of prokaryotes, mostly found in extreme environments.

**Artificial gene synthesis** Construction of an artificial gene from a series of overlapping oligonucleotides.

**Ascospore** One of the haploid products of meiosis in an ascomycete such as the yeast *Saccharomyces cerevisiae*.

**Ascus** The structure that contains the four ascospores produced by a single meiosis in the yeast *Saccharomyces cerevisiae*.

**Attenuation** A process used by some bacteria to regulate expression of an amino acid biosynthetic operon in accordance with the levels of the amino acid in the cell.

**AU-AC intron** A type of intron found in eukaryotic nuclear genes: the first two nucleotides in the intron are 5’-AU-3’ and the last two are 5’-AC-3’.

**Autonomous consensus sequence (ACS)** An 11 bp subdomain of a yeast origin of replication that is part of the origin recognition sequence.

**Autonomously replicating sequence (ARS)** A DNA sequence, particularly from yeast, that confers replicative ability on a nonreplicative plasmid.

**Autopolyploid** A polyploid nucleus derived from fusion of two gametes from the same species, neither of which is haploid.

**Autoradiography** The detection of radioactively labeled molecules by exposure of an X-ray-sensitive photographic film.

**Autosome** A chromosome that is not a sex chromosome.

**Auxotroph** A mutant microorganism that can grow only when supplied with a nutrient that is not needed by the wild type.

**Avidin** A protein, from egg white, that has a high binding affinity for biotin.

**B chromosome** A chromosome possessed by some individuals in a population, but not all.

**B-DNA** The commonest structural conformation of the DNA double helix in living cells.

**Backtracking** The reversal of an RNA polymerase a short distance along its DNA template strand.

**Bacterium** One of the two main groups of prokaryotes.

**Bacterial artificial chromosome (BAC)** A high-capacity cloning vector based on the F plasmid of *Escherichia coli*.

**Bacteriophage** A virus that infects a bacterium.

**Bacteriophage P1 vector** A high-capacity cloning vector based on bacteriophage P1.

**Bait** One of a set of oligonucleotides used to capture particular DNA fragments during target enrichment.

**Barcode deletion strategy** A method that has been developed for the large-scale screening of deletion mutations in *Saccharomyces cerevisiae*.
**Barr body** The highly condensed chromatin structure taken up by an inactivated X chromosome.

**Basal promoter** The position within a eukaryotic promoter where the initiation complex is assembled.

**Basal rate of transcription initiation** The number of productive initiations of transcription occurring per unit time at a particular promoter.

**Base analog** A compound whose structural similarity to one of the bases in DNA enables it to act as a mutagen.

**Base excision repair** A DNA repair process that involves excision and replacement of an abnormal base.

**Basepair** The hydrogen-bonded structure formed by two complementary nucleotides. When abbreviated to bp, the shortest unit of length for a double-stranded DNA molecule.

**Base pairing** The attachment of one polynucleotide to another, or one part of a polynucleotide to another part of the same polynucleotide, by base pairs.

**Base ratio** The ratio of A to T, or G to C, in a double-stranded DNA molecule. Chargaff showed that the base ratios are always close to 1.0.

**Base stacking** The hydrophobic interactions that occur between adjacent base pairs in a double-stranded DNA molecule.

**Baseless site** A position in a DNA molecule where the base component of the nucleotide is missing.

**Basic helix-loop-helix** A DNA-binding domain.

**Beads-on-a-string** An unpacked form of chromatin consisting of nucleosome beads on a string of DNA.

**Biobank** A collection of biological material such as blood samples, donated following informed consent by patients and volunteers, that is often used to study the genetic basis to inherited disease.

**Biochemical profiling** The study of metabolomes.

**Bioinformatics** The use of computer methods in studies of genomes.

**Biolistics** A means of introducing DNA into cells that involves bombardment with high-velocity microprojectiles coated with DNA.

**Biological information** The information contained in the genome of an organism and which directs the development and maintenance of that organism.

**Biotechnology** The use of living organisms, often but not always microbes, in industrial processes.

**Biotin** A molecule that can be incorporated into dUTP and used as a nonradioactive label for a DNA probe.

**Biotinylation** Attachment of a biotin label to a DNA or RNA molecule.

**Bivalent** The structure formed when a pair of homologous chromosomes lines up during meiosis.

**BLAST** An algorithm frequently used in homology searching.

**Blunt end** An end of a double-stranded DNA molecule where both strands terminate at the same nucleotide position with no single-stranded extension.

**Bootstrap analysis** A method for inferring the degree of confidence that can be assigned to a branch point in a phylogenetic tree.

**Bootstrap value** The statistical value obtained by bootstrap analysis.

**Bottleneck** A temporary reduction in the size of a population.

**Bottom-up proteomics** A version of proteomics in which proteins are broken into peptides by treatment with a sequence-specific protease, such as trypsin, prior to mass spectrometry.

**Branch** A component of a phylogenetic tree.

**Branch migration** A step in the Holliday model for homologous recombination, involving exchange of polynucleotides between a pair of recombining double-stranded DNA molecules.

**Break repair** A process for the repair of single- or double-strand breaks in a DNA molecule.

**Bubble-seq** A method used to identify the positions of replication origins in an eukaryotic genome sequence.

**Buoyant density** The density possessed by a molecule or particle when suspended in an aqueous salt or sugar solution.

**Buoyant density centrifugation** A centrifugation method used to separate molecules or structures on the basis of their buoyant densities.

**C-terminal domain (CTD)** A component of the largest subunit of RNA polymerase II, important in the activation of the polymerase.

**C-terminus** The end of a polypeptide that has a free carboxyl group.

**C-value paradox** The nonequivalence between genome size and gene number that is seen when comparisons are made between some eukaryotes.

**Candidate gene** A gene, identified by experimental means, that might be a disease-causing or disease-susceptibility gene.

**Cap analysis gene expression (CAGE)** A method for the rapid acquisition of RNA-seq data.

**Cap binding complex** The complex that makes the initial attachment to the cap structure at the beginning of the scanning phase of eukaryotic translation.

**CAP site** A DNA binding site for the catabolite activator protein.

**Cap structure** The chemical modification at the 5'-end of most eukaryotic mRNA molecules.

**Capillary electrophoresis** Polyacrylamide gel electrophoresis carried out in a thin capillary tube, providing high resolution.

**Capping** Attachment of a cap to the 5'-end of a eukaryotic mRNA.

**Capsid** The protein coat that surrounds the DNA or RNA genome of a virus.
Carboxyl-terminus | The end of a polypeptide that has a free carboxyl group.

Cas9 endonuclease | A programmable nuclease that is directed to its target site by a 20-nucleotide guide RNA.

Cascade | A pathway comprising a series of proteins or other molecules, which passes a signal from the cell surface to genes and other targets within a cell.

Catabolite activator protein | A regulatory protein that binds to various sites in a bacterial genome and activates transcription initiation at downstream promoters.

Catabolite repression | The means by which extracellular glucose levels dictate whether genes for sugar utilization are switched on or off in bacteria.

cDNA | A double-stranded DNA copy of an mRNA molecule.

cDNA capture or cDNA selection | Repeated hybridization probing of a pool of cDNAs with the objective of obtaining a subpool enriched in certain sequences.

Cell cycle | The series of events occurring in a cell between one cell division and the next.

Cell cycle checkpoint | A period before entry into S or M phase of the cell cycle, a key point at which regulation is exerted.

Cell senescence | The period in a cell lineage when the cells are alive but no longer able to divide.

Cell transformation | The alteration in morphological and biochemical properties that occurs when an animal cell is infected by an oncogenic virus.

Cell-free protein-synthesizing system | A cell extract containing all the components needed for protein synthesis and able to translate added mRNA molecules.

centiMorgan | The unit used to describe the distance between two genes on a chromosome. 1 cM is the distance that corresponds to a 1% probability of recombination in a single meiosis.

Centromere | The constricted region of a chromosome that is the position at which the pair of chromatids is held together.

Chain-termination method | A DNA sequencing method that involves enzymatic synthesis of polynucleotide chains that terminate at specific nucleotide positions.

Chaperonin | A multisubunit protein that forms a structure that aids the folding of other proteins.

Chemical degradation sequencing | A DNA sequencing method that involves the use of chemicals that cut DNA molecules at specific nucleotide positions.

Chemical modification | Modification of a protein or RNA by addition of novel chemical groups.

Chemical shift | The change in the rotation of a chemical nucleus, used as the basis of NMR.

Chemiluminescent marker | A chemiluminescent chemical group incorporated into or attached to a molecule and whose chemiluminescent emissions are subsequently used to detect and follow that molecule during a biochemical reaction.

Chi (crossover hotspot initiation) site | A repeated nucleotide sequence in the Escherichia coli genome that is involved in the initiation of homologous recombination.

Chi (γ) form | An intermediate structure seen during recombination between DNA molecules.

Chimera | An organism composed of two or more genetically different cell types.

ChiP-on-chip or ChiP-chip | A microarray-based version of chromatin immunoprecipitation sequencing.

Chloroplast | One of the photosynthetic organelles of a eukaryotic cell.

Chloroplast genome | The genome present in the chloroplasts of a photosynthetic eukaryotic cell.

Chromatid | The arm of a chromosome.

Chromatin | The complex of DNA and histone proteins found in chromosomes.

Chromatin immunoprecipitation sequencing (ChIP-seq) | A method for identifying the positions where individual DNA-binding proteins attach to a genome.

Chromosome | One of the DNA–protein structures that contains part of the nuclear genome of a eukaryote. Less accurately, the DNA molecule(s) that contain(s) a prokaryotic genome.

Chromosome conformation capture (3C) | A method for identifying regions of chromosomes that are located close to one another in the nucleus.

Chromosome painting | A version of fluorescent in situ hybridization in which the hybridization probe is a mixture of DNA molecules, each specific for different regions of a single chromosome.

Chromosome theory | The theory, first propounded by Sutton in 1903, that genes lie on chromosomes.

Chromosome walking | A technique that can be used to construct a clone contig by identifying overlapping fragments of cloned DNA.

Cis-displacement | Movement of a nucleosome to a new position on a DNA molecule.

Class switching | A process that results in a complete change in the type of immunoglobulin synthesized by a B lymphocyte.

Cleavage and polyadenylation specificity factor (CPSF) | A protein that plays an ancillary role during polyadenylation of eukaryotic mRNAs.

Cleavage stimulation factor (CstF) | A protein that plays an ancillary role during polyadenylation of eukaryotic mRNAs.

Clone | A group of cells that contain the same recombinant DNA molecule.

Clone contig | A collection of clones whose DNA fragments overlap.
**Clone contig approach** A genome sequencing strategy in which the molecules to be sequenced are broken into manageable segments, each a few hundred kb or few Mb in length, which are sequenced individually.

**Clone fingerprinting** Any one of several techniques that compare cloned DNA fragments in order to identify ones that overlap.

**Clone library** A collection of clones, possibly representing an entire genome, from which individual clones of interest are obtained.

**Cloning vector** A DNA molecule that is able to replicate inside a host cell and therefore can be used to clone other fragments of DNA.

**Closed promoter complex** The structure formed during the initial step in assembly of the transcription initiation complex. The closed promoter complex consists of the DNA polymerase and/or accessory proteins attached to the promoter, before the DNA has been opened up by breakage of base pairs.

**Cloverleaf** A two-dimensional representation of the structure of a tRNA molecule.

**Clumped regularly interspaced short palindromic repeats (CRISPRs)** A type of bacterial repetitive DNA made up of 20–50 bp sequences found in tandem arrays, with each pair of repeats separated by a spacer of similar length but with a unique sequence.

**Coalescence time** An estimate of the time that has elapsed since a haplogroup first came into existence, based on the degree of divergence of the haplotypes in that haplogroup.

**Coding RNA** An RNA molecule that codes for a protein; an mRNA.

**Co-dominance** The relationship between a pair of alleles that both contribute to the phenotype of a heterozygote.

**Codon** A triplet of nucleotides coding for a single amino acid.

**Codon bias** Refers to the fact that not all codons are used equally frequently in the genes of a particular organism.

**Codon-anticodon recognition** The interaction between a codon on an mRNA molecule and the corresponding anticodon on a tRNA.

**Cohesin** The protein that holds sister chromatids together during the period between genome replication and nuclear division.

**Cohesive end** An end of a double-stranded DNA molecule where there is a single-stranded extension.

**Co-immunoprecipitation** Isolation of all the members of a protein complex using an antibody specific for just one of those proteins.

**Cointegrate** An intermediate in the pathway resulting in replicative transposition.

**Column chromatography** A method for separation of compounds that makes use of a resin contained in a column.

**Comorbidity** The tendency for patients suffering from one disease to display symptoms associated with other diseases.

**Comparative epigenomics** Examination of the extent to which the equivalent regions of two different genomes display the same pattern of chromatin modification.

**Comparative genomics** A research strategy that uses information obtained from the study of one genome to make inferences about the map positions and functions of genes in a second genome.

**Competent** Refers to a culture of bacteria that have been treated, for example, by soaking in calcium chloride, so that their ability to take up DNA molecules is enhanced.

**Complementary** Refers to two nucleotides or nucleotide sequences that are able to base-pair with one another.

**Complementary DNA (cDNA)** A double-stranded DNA copy of an mRNA molecule.

**Complex A** The prespliceosome complex.

**Complex B** The pre catalytic spliceosome.

**Complex E** The first protein-RNA complex formed during splicing of a GU-AG intron.

**Composite transposon** A DNA transposon comprising a pair of insertion sequences flanking a segment of DNA usually containing one or more genes.

**Concatemer** A DNA molecule made up of linear genomes or other DNA units linked head to tail.

**Concerted evolution** The evolutionary process that results in the members of a multigene family retaining the same or similar sequences.

**Conjugation** Transfer of DNA between two bacteria that come into physical contact with one another.

**Conjugation mapping** A technique for mapping bacterial genes by determining the time it takes for each gene to be transferred during conjugation.

**Consensus sequence** A nucleotide sequence that represents an average of a number of related but nonidentical sequences.

**Conservative replication** A hypothetical mode of DNA replication in which one daughter double helix is made up of the two parental polynucleotides and the other is made up of two newly synthesized polynucleotides.

**Conservative transposition** Transposition that does not result in copying of the transposable element.

**Constitutive heterochromatin** Chromatin that is permanently in a compact organization.

**Context-dependent codon reassignment** Refers to the situation whereby the DNA sequence surrounding a codon changes the meaning of that codon.

**Contour clamped homogeneous electric fields (CHEF)** An electrophoresis method used to separate large DNA molecules.

**Conventional pseudogene** A gene that has become inactive because of the accumulation of mutations.

**COOH-terminus** The end of a polypeptide that has a free carboxyl group.

**Core genome** The component of a bacterial pan-genome that contains the set of genes possessed by all members of the species.
Core octamer  The central component of a nucleosome, made up of two subunits each of histones H2A, H2B, H3, and H4, around which DNA is wound.

Core promoter  The position within a eukaryotic promoter where the initiation complex is assembled.

cos site  One of the cohesive, single-stranded extensions present at the ends of the DNA molecules of certain strains of λ phage.

Cosmid  A high-capacity cloning vector consisting of the λ cos site inserted into a plasmid.

Co-transduction  Transfer of two or more genes from one bacterium to another via a transducing phage.

Co-transformation  Uptake of two or more genes on a single DNA molecule during transformation of a bacterium.

CpG island  A GC-rich DNA region located upstream of approximately 56% of the genes in the human genome.

Crossing over  The exchange of DNA between chromosomes during meiosis.

Cryptic splice site  A site whose sequence resembles an authentic splice site and which might be selected instead of the authentic splice site inserted into a plasmid.

Cyanelle  A photosynthetic organelle that resembles an ingested cyanobacterium.

Cyclic AMP (cAMP)  A modified version of AMP in which the 5'- and 3'-carbons.

Cyclic AMP response element (CRE)  The binding site for the CAMP response element-binding (CREB) protein.

Cyclic AMP response element-binding (CREB) protein  A transcription factor that responds to elevated levels of cyclic AMP by binding to the CRE (cyclic AMP response element) sequence.

Cyclin  A regulatory protein whose abundance varies during the cell cycle and which regulates biochemical events in a cell-cycle-specific manner.

Cyclobutyl dimer  A dimer between two adjacent pyrimidine bases in a polynucleotide, formed by ultraviolet irradiation.

Cys2His2 finger  A type of zinc-finger DNA-binding domain.

Cytochemistry  The use of compound-specific stains, combined with microscopy, to determine the biochemical content of cellular structures.

Cytokines  Proteins involved in cell–cell signaling.

Cytosine  One of the pyrimidine bases found in DNA and RNA.

D arm  Part of the structure of a tRNA molecule.

D-loop  An intermediate structure formed in the Meselson–Radding model for homologous recombination. Also, a region of approximately 500 bp where the double helix is disrupted by the presence of an RNA molecule base-paired to one of the DNA strands, and which acts as the origin for the displacement mode of replication.

Dark repair  A type of nucleotide excision repair process that corrects cyclobutyl dimers.

De Bruijn graph  A computational approach, based on a mathematical concept for identifying overlaps between strings of symbols, used by some types of sequence assembler.

De novo methylation  Addition of methyl groups to new positions on a DNA molecule.

De novo sequencing  A strategy in which a genome sequence is assembled solely by finding overlaps between individual sequence reads.

Deadenylation-dependent decapping  A process for degradation of eukaryotic mRNAs that is initiated by removal of the poly(A) tail.

Deaminating agent  A mutagen that acts by removing amino groups from nucleotide bases.

Defective retrovirus  A retrovirus whose genome contains a cellular gene, which replaces part or all of a retrovirus gene, so the virus is unable to replicate without the use of proteins from other retroviruses.

Degeneracy  Refers to the fact that the genetic code has more than one codon for most amino acids.

Degradosome  A multienzyme complex responsible for degradation of bacterial mRNAs.

Delayed-onset mutation  A mutation whose effect is not apparent until a relatively late stage in the life of the mutant organism.

Deletion mutation  A mutation resulting from deletion of one or more nucleotides from a DNA sequence.

Denaturation  Breakdown by chemical or physical means of the noncovalent interactions, such as hydrogen bonding, that maintain the secondary and higher levels of structure of proteins and nucleic acids.

Dendrogram  A tree that is drawn to indicate the relationships between, for example, a group of transcriptomes.

Density gradient centrifugation  A technique in which a cell fraction is centrifuged through a dense solution, in the form of a gradient, so that individual components are separated.

Deoxyribonuclease  An enzyme that cleaves phosphodiester bonds in a DNA molecule.

Derived allele  An allele that arises in a population by mutation of an existing allele.

Development  A coordinated series of transient and permanent changes that occurs during the life history of a cell or organism.

Diauxie  The phenomenon whereby a bacterium, when provided with a mixture of sugars, uses up one sugar before beginning to metabolize the second sugar.

Dicer  The ribonuclease that plays a central role in RNA interference.

Dideoxynucleotide  A modified nucleotide that lacks the 3'-hydroxyl group and so terminates strand synthesis when incorporated into a polynucleotide.

Differential centrifugation  A technique that separates cell components by centrifuging an extract at different speeds.
**Differentiation** The adoption by a cell of a specialized biochemical and/or physiological role.

**Dihybrid cross** A sexual cross in which the inheritance of two pairs of alleles is followed.

**Dimer** A protein or other structure that comprises two subunits.

**Diol** A compound containing two hydroxyl groups.

**Diploid** Describes a nucleus that has two copies of each chromosome.

**Direct readout** The recognition of a DNA sequence by a binding protein that makes contacts with the outside of a double helix.

**Direct repair** A DNA repair system that acts directly on a damaged nucleotide.

**Direct repeat** A nucleotide sequence that is repeated twice or more frequently in a DNA molecule.

**Directed acyclic graph (DAG)** A device used to give a hierarchical categorization of the molecular function.

**Discontinuous gene** A gene that is split into exons and introns.

**Disease module** A set of proteins that when defective give rise to the same genetic disease, often co-located in a protein interaction map.

**Dispersive replication** A hypothetical mode of DNA replication in which both polynucleotides of each daughter double helix are made up partly of parental DNA and partly of newly synthesized DNA.

**Displacement replication** A mode of DNA replication that involves continuous copying of one strand of the helix, the second strand being displaced and subsequently copied after synthesis of the first daughter strand has been completed.

**Distance matrix** A table showing the evolutionary distances between all pairs of nucleotide sequences in a data set.

**Disulfide bridge** A covalent bond linking cysteine amino acids on different polypeptides or at different positions on the same polypeptide.

**DNA** Deoxyribonucleic acid, one of the two forms of nucleic acid in living cells; the genetic material for all cellular life forms and many viruses.

**DNA adenine methylase (Dam)** An enzyme involved in methylation of *Escherichia coli* DNA.

**DNA binding** A type of conformational change introduced into a DNA molecule by a binding protein.

**DNA chip** A high-density array of DNA molecules used for parallel hybridization analyses.

**DNA cloning** Insertion of a fragment of DNA into a cloning vector, and subsequent propagation of the recombinant DNA molecule in a host organism.

**DNA cytosine methylase (Dcm)** An enzyme involved in methylation of *Escherichia coli* DNA.

**DNA glycosylase** An enzyme that cleaves the β-N-glycosidic bond between a base and the sugar component of a nucleotide as part of the base excision and mismatch repair processes. The name is a misnomer and should be DNA glycolyase, but the incorrect usage is now embedded in the literature.

**DNA gyrase** A type II topoisomerase of *Escherichia coli*.

**DNA labeling** Attachment of a radioactive, fluorescent, or other marker to a DNA molecule.

**DNA ligase** An enzyme that synthesizes phosphodiester bonds as part of DNA replication, repair, and recombination processes.

**DNA marker** A DNA sequence that exists as two or more readily distinguished versions and which can therefore be used to mark a map position on a genetic, physical, or integrated genome map.

**DNA methylation** Refers to the chemical modification of DNA by attachment of methyl groups.

**DNA methyltransferase** An enzyme that attaches methyl groups to a DNA molecule.

**DNA photolyase** A bacterial enzyme involved in photoreactivation repair.

**DNA polymerase** An enzyme that synthesizes DNA.

**DNA polymerase I** The bacterial enzyme that completes synthesis of Okazaki fragments during genome replication.

**DNA polymerase II** A bacterial DNA polymerase involved in DNA repair.

**DNA polymerase III** The main DNA replicating enzyme of bacteria.

**DNA polymerase α** The enzyme that primes DNA replication in eukaryotes.

**DNA polymerase γ** The enzyme responsible for replication of the mitochondrial genome.

**DNA polymerase δ** The enzyme responsible for replication of the lagging DNA strand in eukaryotes.

**DNA polymerase ε** The enzyme responsible for replication of the leading DNA strand in eukaryotes.

**DNA repair** The biochemical processes that correct mutations arising from replication errors and the effects of mutagenic agents.

**DNA replication** Synthesis of a new copy of the genome.

**DNA sequencing** The technique for determining the order of nucleotides in a DNA molecule.

**DNA topoisomerase** An enzyme that introduces or removes turns from the double helix by breakage and reunion of one or both polynucleotides.

**DNA transposon** A transposon whose transposition mechanism does not involve an RNA intermediate.

**DNA tumor virus** A virus with a DNA genome, able to cause cancer after infection of an animal cell.

**DNA unwinding element (DUE)** The AT-rich component of a bacterial origin of replication; the position at which helix melting occurs.

**DNA-binding motif** The part of a DNA-binding protein that makes contact with the double helix.

**DNA-binding protein** A protein that attaches to a DNA molecule.
DNA-dependent DNA polymerase  An enzyme that makes a DNA copy of a DNA template.
DNA-dependent RNA polymerase  An enzyme that makes an RNA copy of a DNA template.
DNase I hypersensitive site  A short region of eukaryotic DNA that is relatively easily cleaved with deoxyribonuclease I, possibly coinciding with positions where nucleosomes are absent.
Domain  A segment of a polypeptide that folds independently of other segments; also the segment of a gene coding for such a domain.
Domain duplication  Duplication of a gene segment coding for a structural domain in the protein product.
Domain shuffling  Rearrangement of segments of one or more genes, with each segment coding for a structural domain in the gene product, to create a new gene.
Dominant  The allele that is expressed in a heterozygote.
Donor site  The splice site at the 5’-end of an intron.
Double helix  The base-paired double-stranded structure that is the natural form of DNA in the cell.
Double heterozygote  A nucleus that is heterozygous for two genes.
Double homozygote  A nucleus that is homozygous for two genes.
Double restriction  Digestion of DNA with two restriction endonucleases at the same time.
Double-stranded  Comprising two polynucleotides attached to one another by base pairing.
Double-strand break (DSB) model  A model of the events occurring during homologous recombination.
Double-stranded RNA binding domain (dsRBD)  A common type of RNA-binding domain.
Downstream  Toward the 3’-end of a polynucleotide.
Draft sequence  An incomplete chromosome or genome sequence, typically containing some errors, gaps, and ambiguity about the order and/or orientation of some sequence contigs.
Duplicated pseudogene  A nonprocessed pseudogene that is the decayed version of one copy of a duplicated gene.
E or exit site  A position within a bacterial ribosome to which a tRNA moves immediately after deacylation.
EC number  A four-part number describing the activity of an enzyme in accordance with the nomenclature set by the International Union of Biochemistry and Molecular Biology.
Edge  A line used to link pairs of interacting proteins in a protein interaction map.
Electroendosmosis  The motion of a liquid, such as the buffer in a gel, induced by an electric field.
Electron density map  A plot of the electron density at different positions within a molecule, deduced from an X-ray diffraction pattern.

**Electron microscopy**  A version of microscopy in which visualization of the sample is provided by a beam of electrons.
**Electrophoresis**  Separation of molecules on the basis of their net electrical charge.
**Electrospray ionization**  An ionization method used during mass spectrometry whereby a high voltage is applied to a liquid to create an aerosol.
**Electrostatic interactions**  Ionic bonds that form between charged chemical groups.
**Elution**  The unbinding of a molecule from a chromatography column.
**Embryonic stem (ES) cell**  A totipotent cell from the embryo of a mouse or other organism.
**End-labeling**  The attachment of a radioactive or other label to one end of a DNA or RNA molecule.
**End modification**  The chemical alteration of the end of a DNA or RNA molecule.
**End-modification enzyme**  An enzyme used in recombinant DNA technology that alters the chemical structure at the end of a DNA molecule.
**Endogenous retrovirus (ERV)**  An active or inactive retroviral genome integrated into a host chromosome.
**Endonuclease**  An enzyme that breaks phosphodiester bonds within a nucleic acid molecule.
**Endosymbiont theory**  A theory that states that the mitochondria and chloroplasts of eukaryotic cells are derived from symbiotic prokaryotes.
**Enhancer**  A regulatory sequence that increases the rate of transcription of a gene or genes located some distance away in either direction.
**Ensembl**  An online genome browser.
**Epigenetic effects**  Changes in phenotype that occur not because of alterations in the genome sequence but because of alterations in the way in which the genome is expressed.
**Epigenome**  The combination of processes including histone modification, nucleosome positioning, and DNA methylation that regulate genome expression.
**Episome**  A plasmid that is able to integrate into the host cell’s chromosome.
**Episome transfer**  Transfer between cells of some or all of a bacterial chromosome by integration into a plasmid.
**Ethidium bromide**  A type of intercalating agent that causes mutations by inserting between adjacent base pairs in a double-stranded DNA molecule.
**Ethylmethane sulfonate (EMS)**  A mutagen that acts by adding alkyl groups to nucleotide bases.
**Euchromatin**  Regions of a eukaryotic chromosome that are relatively uncondensed, thought to contain active genes.
**Eukaryote**  An organism whose cells contain membrane-bound nuclei.
**Eulerian pathway**  A pathway through a graph that visits each edge just once, used by some sequence assemblers.
to identify the correct sequence in regions of a genome containing repetitive DNA.

**Excision repair** A DNA repair process that corrects various types of DNA damage by excising and resynthesizing a region of polynucleotide.

**Exit or E site** A position within a bacterial ribosome to which a tRNA moves immediately after deacylation.

**Exome** The sequences of all of the exons in a genome.

**Exon** A coding region within a discontinuous gene.

**Exon skipping** Aberrant splicing, or an alternative splicing scenario, in which one or more of the exons are omitted from the spliced RNA.

**Exon theory of genes** An introns early hypothesis that holds that introns were formed when the first DNA genomes were constructed.

**Exon trapping** A method, based on cloning, for identifying the positions of exons in a DNA sequence.

**Exon–intron boundary** The nucleotide sequence at the junction between an exon and an intron.

**Exonic splicing enhancer (ESE)** A nucleotide sequence that plays a positive regulatory role during splicing of GU-AG introns.

**Exonic splicing silencer (ESS)** A nucleotide sequence that plays a negative regulatory role during splicing of GU-AG introns.

**Exonuclease** An enzyme that removes nucleotides from the ends of a nucleic acid molecule.

**Exosome** A multiprotein complex involved in degradation of mRNA in eukaryotes.

**Expressed sequence tag (EST)** A cDNA that is sequenced in order to gain rapid access to the genes in a genome.

**Expression proteomics** The methodology used to identify the proteins in a proteome.

**External node** The end of a branch in a phylogenetic tree, representing one of the organisms or DNA sequences being studied.

**Extrachromosomal gene** A gene in a mitochondrial or chloroplast genome.

**Extremophile** An organism that is able to live in an environment whose physical and/or chemical conditions are hostile to other organisms.

**F plasmid** A fertility plasmid that directs conjugal transfer of DNA between bacteria.

**Facultative heterochromatin** Chromatin that has a compact organization in some but not all cells, thought to contain genes that are inactive in some cells or at some periods of the cell cycle.

**FEN1** The flap endonuclease involved in replication of the lagging strand in eukaryotes.

**Fertile Crescent** The region of Southwest Asia where barley and wheat are thought to have been cultivated.

**Fiber-FISH** A specialized form of FISH that enables high marker resolution.

**Field inversion gel electrophoresis (FIGE)** An electrophoresis method used to separate large DNA molecules.

**Filamentous** One of the capsid structures of a bacteriophage or virus.

**Finished sequence** A chromosome or genome sequence that is almost complete, but typically still has some unsequenced gaps between contigs and an average of up to one error per \(10^4\) nucleotides.

**Flap endonuclease (FEN1)** An enzyme involved in replication of the lagging strand in eukaryotes.

**Flow cytometry** A method for the separation of chromosomes.

**Fluorescence recovery after photobleaching (FRAP)** A technique used to study the mobility of nuclear proteins.

**Fluorescent in situ hybridization (FISH)** A technique for locating markers on chromosomes by observing the hybridization positions of fluorescent labels.

**Fluorescent marker** A fluorescent chemical group incorporated into or attached to a molecule and whose fluorescent emissions are subsequently used to detect and follow that molecule during a biochemical reaction.

**Flush end** An end of a double-stranded DNA molecule where both strands terminate at the same nucleotide position with no single-stranded extension.

**fMet** N-formylmethionine, the modified amino acid carried by the tRNA that is used during the initiation of translation in bacteria.

**Foldback RNA** The precursor RNA molecules that are cleaved to produce microRNAs.

**Folding pathway** The series of events, involving partially folded intermediates, that results in an unfolded protein attaining its correct three-dimensional structure.

**Footprinting** A range of techniques used for locating bound proteins on DNA molecules.

**Föhrster resonance energy transfer (FRET)** Energy transfer between two molecules, resulting in quenching of dye when a reporter probe is used.

**Forward genetics** The conventional approach to genetics, where the researcher starts with a phenotype and attempts to discover the gene or genes responsible for that phenotype.

**Forward sequence** One of the two directions in which a double-stranded DNA molecule can be sequenced.

**Fosmid** A high-capacity vector carrying the F plasmid origin of replication and a \(\lambda\) cos site.

**Fourier transform ion cyclotron resonance (FT-ICR) mass analyzer** A mass analyzer that incorporates an ion trap that captures individual ions and further excites them within a cyclotron, so they accelerate along an outward spiral, the vector of this spiral revealing the \(m/z\) ratio.

**Fourth-generation sequencing** An approach to DNA sequencing in which the sequence is read directly without copying that DNA molecule in any way.

**Fragile site** A position in a chromosome that is prone to breakage, possibly because it contains an expanded trinucleotide repeat sequence.
Fragment ion  An ion resulting from fragmentation of a molecule during the ionization phase of mass spectrometry.

Frameshift mutation  A mutation resulting from insertion or deletion of a group of nucleotides that is not a multiple of three and which therefore changes the frame in which translation occurs.

Functional domain  A region of eukaryotic DNA containing a set of genes that are subject to a similar expression pattern, that expression pattern set by the identities of regulatory sequences also present in the domain.

Functional RNA  RNA that has a functional role in the cell; that is, RNAs other than mRNA.

Fusion protein  A protein that consists of a fusion of two polypeptides, or parts of polypeptides, normally coded by separate genes.

G-protein  A small protein that binds either a molecule of GDP or GTP, the replacement of GDP with GTP activating the protein.

G1–S checkpoint  A cell cycle checkpoint that a cell must pass before it is able to replicate its DNA.

G2–M checkpoint  A cell cycle checkpoint that can only be passed when a cell is ready to enter mitosis.

Gamete  A reproductive cell, usually haploid, that fuses with a second gamete to produce a new cell during sexual reproduction.

Gap 1 or G1 phase  The first gap period of the cell cycle.

Gap 2 or G2 phase  The second gap period of the cell cycle.

Gap genes  Developmental genes that play a role in establishing positional information within the Drosophila embryo.

Gap period  One of two intermediate periods within the cell cycle.

GATA zinc finger  A type of zinc-finger DNA-binding domain.

GC content  The percentage of nucleotides in a genome that are G or C.

Gel electrophoresis  Electrophoresis performed in a gel so that molecules of similar electrical charge can be separated on the basis of size.

Gel retardation analysis  A technique that identifies protein-binding sites on DNA molecules by virtue of the effect that a bound protein has on the mobility of the DNA fragments during gel electrophoresis.

Gel stretching  A technique for preparing restricted DNA molecules for optical mapping.

GenBank  An online repository of DNA sequences.

Gene  A DNA segment containing biological information and hence coding for an RNA and/or polypeptide molecule.

Gene cloning  Insertion of a fragment of DNA, containing a gene, into a cloning vector, and subsequent propagation of the recombinant DNA molecule in a host organism.

Gene conversion  A process that results in the four haploid products of meiosis displaying an unusual segregation pattern.

Gene desert  A region of genome in which there are few if any genes.

Gene duplication  The duplication of a gene to give two daughter copies, which initially will have identical nucleotide sequences, but whose sequences might subsequently change due to mutation.

Gene expression  The series of events by which the biological information carried by a gene is released and made available to the cell.

Gene flow  The transfer of a gene from one organism to another.

Gene fragment  A gene relic consisting of a short isolated region from within a gene.

Gene ontology (GO)  A scheme for describing gene function.

Gene space  A version of the barley genome, comprising the sequences of the vast majority of barley genes anchored on to a detailed genome map.

Gene superfamily  A group of two or more evolutionarily related multigene families.

Gene therapy  A clinical procedure in which a gene or other DNA sequence is used to treat a disease.

General recombination  Recombination between two homologous double-stranded DNA molecules.

General transcription factor (GTF)  A protein or protein complex that is a transient or permanent component of the initiation complex formed during eukaryotic transcription.

Genes-within-genes  Refers to a gene whose intron contains a second gene.

Genetic code  The rules that determine which triplet of nucleotides codes for which amino acid during protein synthesis.

Genetic linkage  The physical association between two genes that are on the same chromosome.

Genetic mapping  The use of genetic techniques to construct a genome map.

Genetic marker  A gene that exists as two or more readily distinguished alleles and whose inheritance can therefore be followed during a genetic cross, enabling the map position of the gene to be determined.

Genetic profile  The banding pattern revealed after electrophoresis of the products of PCRs directed at a range of microsatellite loci.

Genetic redundancy  The situation that occurs when two genes in the same genome perform the same function.

Genetics  The branch of biology devoted to the study of genes.

Genome  The entire genetic complement of a living organism.

Genome annotation  The process by which the genes, control sequences, and other interesting features are identified in a genome sequence.

Genome browser  A software package or online system for display of an annotated genome sequence.
<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genome expression</td>
<td>The series of events by which the biological information carried by a genome is released and made available to the cell.</td>
</tr>
<tr>
<td>Genome map</td>
<td>A chart showing the positions of genetic and/or physical markers in a genome.</td>
</tr>
<tr>
<td>Genome resequencing</td>
<td>Sequencing of multiple versions of a genome that has already been sequenced, in order to study the sequence variations that occur within a species or within populations of that species.</td>
</tr>
<tr>
<td>Genomewide association study (GWAS)</td>
<td>A method that attempts to identify all of the markers, from all over the genome, that are associated with a disease.</td>
</tr>
<tr>
<td>Genomewide repeat</td>
<td>A sequence that recurs at many dispersed positions within a genome.</td>
</tr>
<tr>
<td>Genomic imprinting</td>
<td>Inactivation by methylation of a gene on one of a pair of homologous chromosomes.</td>
</tr>
<tr>
<td>Genotype</td>
<td>A description of the genetic composition of an organism.</td>
</tr>
<tr>
<td>Gigabase pair</td>
<td>1,000,000 kb; 1,000,000,000 bp.</td>
</tr>
<tr>
<td>Glacial refugia</td>
<td>Geographical regions occupied by various species during the Last Glacial Maximum, enabling those species to survive the glaciations and eventually recolonize adjacent regions following the warming of the planet.</td>
</tr>
<tr>
<td>Glycan</td>
<td>The oligosaccharide at a single glycosylated position in a glycoprotein.</td>
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<tr>
<td>Glycosylation</td>
<td>The attachment of sugar units to a polypeptide.</td>
</tr>
<tr>
<td>Greedy algorithm</td>
<td>A computational approach, based on making the most logical choice at each step in an iterative process, used by some types of sequence assembler.</td>
</tr>
<tr>
<td>Green fluorescent protein</td>
<td>A protein that is used to label other proteins and whose gene is used as a reporter gene.</td>
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<tr>
<td>Group I intron</td>
<td>A type of intron found mainly in organelle genes.</td>
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<tr>
<td>Group II intron</td>
<td>A type of intron found in organelle genes.</td>
</tr>
<tr>
<td>Group III intron</td>
<td>A type of intron found in organelle genes.</td>
</tr>
<tr>
<td>GTPase-activating protein (GAP)</td>
<td>A protein that inactivates a G-protein by stimulating it to convert its GTP molecule to a GDP molecule.</td>
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<tr>
<td>GU-AG intron</td>
<td>The commonest type of intron in eukaryotic nuclear genes. The first two nucleotides of the intron are 5′-GU-3′ and the last two are 5′-AG-3′.</td>
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<tr>
<td>Guanine</td>
<td>One of the purine bases found in DNA and RNA.</td>
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<tr>
<td>Guanine methyltransferase</td>
<td>The enzyme that attaches a methyl group to the 5′-end of a eukaryotic mRNA during the capping reaction.</td>
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<tr>
<td>Guanine nucleotide exchange factor (GEF)</td>
<td>A protein that activates a G-protein by replacing its GDP molecule with a GTP molecule.</td>
</tr>
<tr>
<td>Guanylyl transferase</td>
<td>The enzyme that attaches a GTP to the 5′-end of a eukaryotic mRNA at the start of the capping reaction.</td>
</tr>
<tr>
<td>Hairpin</td>
<td>A stem–loop structure made up of a base-paired stem and non-base-paired loop, which can form in a single-stranded polynucleotide that contains an inverted repeat.</td>
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<tr>
<td>Half-life</td>
<td>The time needed for half the atoms or molecules in a sample to decay or be degraded.</td>
</tr>
<tr>
<td>Hammerhead</td>
<td>An RNA structure with ribozyme activity that is found in some viruses.</td>
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<tr>
<td>Haplogroup</td>
<td>One of the major sequence classes of mitochondrial DNA present in the human population.</td>
</tr>
<tr>
<td>Haploid</td>
<td>Describes a nucleus that has a single copy of each chromosome.</td>
</tr>
<tr>
<td>Haploinsufficiency</td>
<td>The situation where inactivation of a gene on one of a pair of homologous chromosomes results in a change in the phenotype of the mutant organism.</td>
</tr>
<tr>
<td>Haplotype</td>
<td>An individual mitochondrial DNA sequence.</td>
</tr>
<tr>
<td>Head-and-tail</td>
<td>One of the capsid structures of a bacteriophage.</td>
</tr>
<tr>
<td>Helicase</td>
<td>An enzyme that breaks base pairs in a double-stranded DNA molecule.</td>
</tr>
<tr>
<td>Helix-turn-helix motif</td>
<td>A common structural motif for attachment of a protein to a DNA molecule.</td>
</tr>
<tr>
<td>Heterochromatin</td>
<td>Chromatin that is relatively condensed and is thought to contain DNA that is not being transcribed.</td>
</tr>
<tr>
<td>Heteroduplex</td>
<td>A DNA–DNA or DNA–RNA hybrid.</td>
</tr>
<tr>
<td>Heteroduplex analysis</td>
<td>Transcript mapping by analysis of DNA–RNA hybrids with a single-strand-specific nuclease such as S1.</td>
</tr>
<tr>
<td>Heterogenous nuclear ribonucleoproteins (hnRNPs)</td>
<td>A broad group of RNA–protein complexes that play several roles in the nucleus, most of which involve binding to RNAs.</td>
</tr>
<tr>
<td>Heterogenous nuclear RNA (hnRNA)</td>
<td>The nuclear RNA fraction that comprises unprocessed transcripts synthesized by RNA polymerase II.</td>
</tr>
<tr>
<td>Heterozygosity</td>
<td>The probability that a person chosen at random from the population will be heterozygous for a particular marker.</td>
</tr>
<tr>
<td>Heterozygous</td>
<td>A diploid nucleus that contains two different alleles for a particular gene.</td>
</tr>
<tr>
<td>Hexaploid</td>
<td>An auto- or allopolyploid with three diploid genome copies.</td>
</tr>
<tr>
<td>Hibernation promotion factor</td>
<td>A protein involved in the inactivation of surplus ribosomes in <em>Escherichia coli</em>.</td>
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<tr>
<td>Hierarchical clustering</td>
<td>A method for analyzing transcriptomes based on comparisons between the expression levels of pairs of genes.</td>
</tr>
<tr>
<td>Hierarchical shotgun sequencing</td>
<td>A DNA sequencing strategy that involves a pre-sequencing phase during which the genome is broken into large fragments, which are cloned and each sequenced individually by the shotgun method.</td>
</tr>
<tr>
<td>High mobility group (HMG) box</td>
<td>A DNA-binding domain.</td>
</tr>
<tr>
<td>High-performance liquid chromatography (HPLC)</td>
<td>A column chromatography method with many applications in biochemistry.</td>
</tr>
<tr>
<td>Histone</td>
<td>One of the basic proteins found in nucleosomes.</td>
</tr>
<tr>
<td>Histone acetylation</td>
<td>Modification of chromatin structure by attachment of acetyl groups to core histones.</td>
</tr>
</tbody>
</table>
Histone acetyltransferase (HAT) An enzyme that attaches acetyl groups to core histones.

Histone code The hypothesis that the pattern of chemical modification on histone proteins influences various cellular activities.

Histone deacetylase (HDAC) An enzyme that removes acetyl groups from core histones.

Histone-like nucleoid structuring protein (H-NS) A nucleoid protein that binds specifically to AT-rich regions, which are thought to be present at the boundaries of the supercoiled loops of a bacterial chromosome.

 Holliday structure An intermediate structure formed during recombination between two DNA molecules.

Holocentric chromosome A chromosome that does not have a single centromere but instead has multiple kinetochores spread along its length.

Homeodomain A DNA-binding motif found in many proteins involved in developmental regulation of gene expression.

Homeotic mutation A mutation that results in the transformation of one body part into another.

Homeotic selector gene A gene that establishes the identity of a body part such as a segment of the Drosophila embryo.

Homologous chromosomes Two or more identical chromosomes present in a single nucleus.

Homologous recombination Recombination between two homologous double-stranded DNA molecules; that is, ones that share extensive nucleotide sequence similarity.

Homologous sequences DNA sequences that share a common evolutionary ancestor.

Homology searching A technique in which genes with sequences similar to that of an unknown gene are sought, the objective being to gain an insight into the function of the unknown gene.

Homopolymer tailing The attachment of a sequence of identical nucleotides (for example, AAAAA) to the end of a nucleic acid molecule; usually referring to the synthesis of single-stranded homopolymer extensions on the ends of a double-stranded DNA molecule.

Homozygous A diploid nucleus that contains two identical alleles for a particular gene.

Hoogsteen base pairs Base pairs that involve the same combinations (A-T and G-C) as Watson–Crick base pairs but are held together by hydrogen bonds between different groups.

Horizontal gene transfer Transfer of a gene from one species to another.

Hormone response element A nucleotide sequence upstream of a gene that mediates the regulatory effect of a steroid hormone.

Housekeeping protein A protein that is continually expressed in all or at least most cells of a multicellular organism.

Hsp70 chaperones A family of proteins that bind to hydrophobic regions in other proteins in order to aid their folding.

HU family A family of nucleoid proteins that have some amino acid sequence similarity with the eukaryotic histone H2B.

Hub A protein that has many interactions within a protein interaction map.

Human Genome Project The publicly funded project responsible for one of the draft human genome sequences and which continues to study the functions of human genes.

Hybrid dysgenesis The event that occurs when females from laboratory strains of Drosophila melanogaster are crossed with males from wild populations, the offspring resulting from such crosses being sterile and having chromosomal abnormalities and other genetic malfunctions.

Hybridization The attachment to one another, by base pairing, of two complementary polynucleotides.

Hybridization probe A labeled nucleic acid molecule used as a probe to identify complementary or homologous molecules to which it base-pairs.

Hybridize Base pairing between nucleic acid molecules that may or may not have entirely complementary sequences.

Hydrogen bond A weak electrostatic attraction between an electronegative atom such as oxygen or nitrogen and a hydrogen atom attached to a second electronegative atom.

Hydrophobic effects Chemical interactions that result in hydrophobic groups becoming buried inside a protein.

Icosahedral One of the capsid structures of a bacteriophage or virus.

Illumina sequencing A next-generation sequencing method utilizing reversible terminator sequencing of fragments immobilized on a slide.

Immobilized metal affinity chromatography A type of chromatography for purification of phosphorylated proteins.

Immunocytochemistry A technique that uses antibody probing to locate the position of a protein in a tissue.

Immunoelectron microscopy An electron microscopy technique that uses antibody labeling to identify the positions of specific proteins on the surface of a structure such as a ribosome.

Immunofluorescence microscopy A microscopy method that utilizes a fluorescently labeled antibody to visualize the location of particular proteins within a cell.

Immunoglobulin fold A DNA-binding domain made up of three loops emerging from a barrel-shaped β-sheet.

Immunoscreening The use of an antibody probe to detect a polypeptide synthesized by a cloned gene.

Imprint control element A DNA sequence, found within a few kb of clusters of imprinted genes, that mediates the methylation of the imprinted regions.

In vitro mutagenesis Techniques used to produce a specified mutation at a predetermined position in a DNA molecule.

In vitro packaging Synthesis of infective λ phages from a preparation of λ proteins and a concatemer of λ DNA molecules.
**Incomplete dominance** Refers to a pair of alleles, neither of which displays dominance, the phenotype of a heterozygote being intermediate between the phenotypes of the two homozygotes.

**Indel** A position in an alignment between two DNA sequences where an insertion or deletion has occurred.

**Inducer** A molecule that induces expression of a gene or operon by binding to a repressor protein and preventing the repressor from attaching to the operator.

**Inducible operon** An operon that is switched on by an inducer molecule.

**Induction** (1) Of a gene: the switching on of the expression of a gene or group of genes in response to a chemical or other stimulus. (2) Of λ phage: the excision of the integrated form of λ and accompanying switch to the lytic mode of infection in response to a chemical or other stimulus.

**Informational problem** The problem tackled by the early molecular biologists concerning the nature of the genetic code.

**Inherited disease** A disease caused by a defect in a gene.

**Initiation codon** The codon, usually but not exclusively 5′-AUG-3′, found at the start of the coding region of a gene.

**Initiation complex** The complex of proteins that initiates transcription. Also the complex that initiates translation.

**Initiation factor** A protein that plays an ancillary role during initiation of translation.

**Initiation of transcription** The assembly upstream of a gene of the complex of proteins that will subsequently copy the gene into RNA.

**Initiation region** A region of eukaryotic chromosomal DNA within which replication initiates at positions that are not clearly defined.

**Initiator (Inr) sequence** A component of the RNA polymerase II core promoter.

**Initiator tRNA** The tRNA, aminoacylated with methionine in eukaryotes or N-formylmethionine in bacteria, that recognizes the initiation codon during protein synthesis.

**Inosine** A modified version of adenosine, sometimes found at the wobble position of an anticodon.

**Insertion mutation** A mutation that arises by insertion of one or more nucleotides into a DNA sequence.

**Insertion sequence** A short transposable element found in bacteria.

**Insertion vector** A λ vector constructed by deleting a segment of nonessential DNA.

**Insertional inactivation** A cloning strategy whereby insertion of a new piece of DNA into a vector inactivates a gene carried by the vector.

**Insulator** A segment of DNA that acts as the boundary point between two functional domains.

**Integrase** A type I topoisomerase that catalyzes insertion of the λ genome into *Escherichia coli* DNA.

**Integron** A set of genes and other DNA sequences that enables plasmids to capture genes from bacteriophages and other plasmids.

**Interactome** The whole set of molecular interactions occurring in a cell.

**Intercalating agent** A compound that can enter the space between adjacent base pairs in a double-stranded DNA molecule, often causing mutations.

**Interferon** A type of cytokine.

**Interferon γ-stimulated gene response (GAS) element** One type of DNA binding site for STAT dimers.

**Interferon-stimulated response element (ISRE)** One type of DNA binding site for STAT dimers.

**Intergenic DNA** The regions of a genome that do not contain genes.

**Internal node** A branch point within a phylogenetic tree, representing an organism or DNA sequence that is ancestral to those being studied.

**Internal ribosome entry site (IRES)** A nucleotide sequence that enables the ribosome to assemble at an internal position in some eukaryotic mRNAs.

**Interphase** The period between cell divisions.

**Interspersed repeat** A sequence that recurs at many dispersed positions within a genome.

**Interspersed repeat element PCR (IRE-PCR)** A clone fingerprinting technique that uses PCR to detect the relative positions of genomewide repeats in cloned DNA fragments.

**Intramolecular base pairing** Base pairing that occurs between two parts of the same DNA or RNA polynucleotide.

**Intrinsic terminator** A position in bacterial DNA where termination of transcription occurs without the involvement of Rho.

**Intron** A noncoding region within a discontinuous gene.

**Intron retention** An alternative splicing scenario in which an intron that is usually spliced out of the pre-mRNA is retained in the final mRNA.

**Intron splicing enhancer (ISE)** A nucleotide sequence that plays a positive regulatory role during splicing of GU-AG introns.

**Intron splicing silencer (ISS)** A nucleotide sequence that plays a negative regulatory role during splicing of GU-AG introns.

**Introns early** The hypothesis that introns evolved relatively early and are gradually being lost from eukaryotic genomes.

**Introns late** The hypothesis that introns evolved relatively late and are gradually accumulating in eukaryotic genomes.

**Inverted repeat** Two identical nucleotide sequences repeated in opposite orientations in a DNA molecule.

**Ion-exchange chromatography** A method for separating molecules according to how tightly they bind to electrically charged particles present in a chromatographic matrix.

**Ion torrent sequencing** A next-generation sequencing method that reads a sequence by detection of the hydrogen ions that are released every time a nucleotide is incorporated into the growing strand.
Ion-sensitive field effect transistor (ISFET)  The component of an ion torrent sequencer that detects the hydrogen ions that are released during strand synthesis.

IRES trans-acting factors (ITAFs)  Proteins that regulate the usage of internal ribosome entry sites.


Isoaccepting tRNAs  Two or more tRNAs that are charged with the same amino acid.

Isobaric labeling  The use of labeled tags that have equal mass but during mass spectrometry give rise to reporter fragment ions that are differentially labeled.

Isochore model  A model that proposes that a eukaryotic genome is a mosaic of DNA segments, each of which has a uniform base composition that differs from that of the adjacent segments.

Isoelectric focusing  Separation of proteins in a gel that contains chemicals that establish a pH gradient when the electrical charge is applied.

Isoelectric point  The position in a pH gradient where the net charge of a protein is zero.

Isoforms  The products of the alternative splicing pathways of a single gene.

Isopycnic centrifugation  A centrifugation method used to separate molecules or structures on the basis of their buoyant densities.

Isotope  One of two or more atoms that have the same atomic number but different atomic mass.

Isotope-coded affinity tag (ICAT)  Markers, containing normal hydrogen and deuterium atoms, used to label individual proteomes.

JAK/STAT pathway  A relatively noncomplex type of signal transduction pathway found in many vertebrates.

Janus kinase (JAK)  A type of kinase that plays an intermediary role in some types of signal transduction involving STATs.

Junk DNA  One interpretation of the intergenic DNA content of a genome.

K homology (KH) domain  An RNA-binding domain.

Karyogram  The entire chromosome complement of a cell, with each chromosome described in terms of its appearance at metaphase.

Kilobase pair (kb)  1000 base pairs.

Kinase receptor  A type of cell surface receptor that has kinase activity.

Kinase-associated receptors  A type of cell surface receptor that works in conjunction with a protein that has kinase activity.

Kinetochore  The part of the centromere to which spindle microtubules attach.

Klenow polymerase  A DNA polymerase enzyme, obtained by chemical modification of Escherichia coli DNA polymerase I, used primarily in chain-termination DNA sequencing.

k-mers  Sequence reads of length k.

Knockout mouse  A mouse that has been engineered so that it carries an inactivated gene.

Kornberg polymerase  The DNA polymerase I enzyme of Escherichia coli.

Lac selection  A means of identifying recombinant bacteria containing vectors that carry the lacZ gene. The bacteria are plated on a medium that contains an analog of lactose that gives a blue color in the presence of β-galactosidase activity.

Lactose operon  The cluster of three genes that codes for enzymes involved in utilization of lactose by Escherichia coli.

Lactose repressor  The regulatory protein that controls transcription of the lactose operon in response to the presence or absence of lactose in the environment.

Lagging strand  The strand of the double helix that is copied in a discontinuous fashion during genome replication.

Lambda (λ)  A bacteriophage that infects Escherichia coli, derivatives of which are used as cloning vectors.

Landrace  Populations of crop plants that are locally adapted and which were grown by farmers until replaced by the products of modern breeding programs during the twentieth century.

Lariat  Refers to the lariat-shaped intron RNA that results from splicing a GU-AG intron.

Last eukaryotic common ancestor (LECA)  The archaic organism from which all modern DNA eukaryotes are descended.

Latent period  The period between injection of a phage genome into a bacterial cell and the time when cell lysis occurs.

Lateral gene transfer  Transfer of a gene from one species to another.

Leader segment  The untranslated region of an mRNA upstream of the initiation codon.

Leading strand  The strand of the double helix that is copied in a continuous fashion during genome replication.

Lectin  A plant or animal protein with specific sugar-binding properties.

Leucine zipper  A dimerization domain commonly found in DNA-binding proteins.

Ligase  An enzyme that synthesizes phosphodiester bonds as part of DNA replication, repair, and recombination processes.

LINE (long interspersed nuclear element)  A type of genomewide repeat, often with transposable activity.

LINE-1  One type of human LINE.

Linkage  The physical association between two genes that are on the same chromosome.

Linkage analysis  The procedure used to assign map positions to genes by genetic crosses.

Linkage disequilibrium  The situation where a particular combination of alleles at linked loci occurs more frequently or less frequently than expected in a population.

Linkage group  A group of genes that display linkage. With eukaryotes, a single linkage group usually corresponds to a single chromosome.
**Linker** A synthetic, double-stranded oligonucleotide used to attach sticky ends to a blunt-ended molecule.

**Linker DNA** The DNA that links nucleosomes—the string in the beads-on-a-string model for chromatin structure.

**Linker histone** A histone, such as H1, that is located outside of the nucleosome core octamer.

**Linking number** The number of times one strand crosses the other in a circular molecule.

**Lod score** A statistical measure of linkage as revealed by pedigree analysis.

**Long intergenic noncoding RNA (lincRNA)** A long noncoding RNA that is located entirely within an intergenic region.

**Long noncoding RNA (lncRNA)** A noncoding RNA longer than 200 nucleotides in length.

**Long terminal repeat (LTR)** A repeated DNA sequence found at the ends of some retroelements.

**Low-copy repeat** A duplication between 1 and 400 kb in length, with greater than 90% sequence identity, that is repeated up to 50 times in the genome.

**LTR element** A type of genomewide repeat typified by the presence of long terminal repeats (LTRs).

**Lyase** An enzyme that breaks chemical bonds by processes other than oxidation and hydrolysis.

**Lysis** The disruption of a bacterial cell by lysozyme, such as occurs at the end of the infection cycle of a lytic bacteriophage.

**Lysogenic infection cycle** The type of bacteriophage infection that involves integration of the phage genome into the host DNA molecule.

**Lysozyme** A protein used to destabilize the bacterial cell wall prior to DNA purification.

**Lytic infection cycle** The type of bacteriophage infection that involves lysis of the host cell immediately after the initial infection, with no integration of the phage DNA molecule into the host genome.

**M13 bacteriophage** A bacteriophage that infects Escherichia coli, and derivatives of which are used as cloning vectors.

**Macrochromosome** One of the larger gene-deficient chromosomes seen in the nuclei of chickens and various other species.

**MADS box** A DNA-binding domain found in several transcription factors involved in plant development.

**Magnetic tweezer** A set of magnets whose positions and field strengths can be varied to move magnetic particles, such as magnetic beads, in a controlled manner to study the mechanical properties of biomolecules.

**Maintenance methylation** Addition of methyl groups to positions on newly synthesized DNA strands that correspond with the positions of methylation on the parent strand.

**Major groove** The larger of the two grooves that spiral around the surface of the B-form of DNA.

**MAP (mitogen-activated protein) kinase or MAPK/ERK pathway** An important signal transduction pathway found in many organisms.

**Map unit** A unit used to describe the distance between two genes on a chromosome, now superseded by the centiMorgan.

**Mapping reagent** A collection of DNA fragments spanning a chromosome or the entire genome and used in STS mapping.

**Mass analyzer** The component of a mass spectrometer that measures the mass-to-charge ratios of the ions that are being studied.

**Mass spectrometry** An analytical technique in which ions are separated according to their mass-to-charge ratios.

**Mass-to-charge ratio** The basis to separation of ions by mass spectrometry.

**Massively parallel** A high-throughput sequencing strategy in which many individual sequences are generated in parallel.

**Massively parallel array** An array of DNA fragments immobilized in a format suitable for next-generation DNA sequencing.

**Maternal-effect gene** A Drosophila gene that is expressed in the parent and whose mRNA is subsequently injected into the egg, after which it influences development of the embryo.

**Mating type** The equivalent of male and female for a eukaryotic microorganism.

**Mating-type switching** The ability of yeast cells to change from the a to α mating type, or vice versa, by gene conversion.

**Matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF)** A type of mass spectrometry used in proteomics.

**Matrix-assisted selection** A DNA screening method that enables individuals with a particular characteristic to be identified from their possession of DNA markers associated with that characteristic.

**Mediator** A protein complex that forms a contact between various activators and the C-terminal domain of the largest subunit of RNA polymerase II.

**Megabase pair (Mb)** 1000 kb; 1,000,000 bp.

**Meiosis** The series of events, involving two nuclear divisions, by which diploid nuclei are converted to haploid gametes.

**Melting temperature (Tm)** The temperature at which the two strands of a double-stranded DNA molecule or base-paired hybrid detach as a result of complete breakage of hydrogen bonding.

**Meselson–Stahl experiment** The experiment that showed that cellular DNA replication occurs by the semiconservative process.

**Messenger RNA (mRNA)** The transcript of a protein-coding gene.

**Metabolic engineering** The process by which changes are made to the genome by mutation or recombinant DNA techniques in order to influence the cellular biochemistry in a predetermined way.
Metabolic flux The rate of flow of metabolites through the network of pathways that make up the cellular biochemistry.

Metabolic labeling A labeling method that involves growing cells in the presence of labeled nutrients.

Metabolome The complete collection of metabolites present in a cell under a particular set of conditions.

Metabolomics The study of metabolomes.

Metagenomics Studies of the mixture of genomes present in a particular habitat.

Metaphase chromosome A chromosome at the metaphase stage of cell division, when the chromatin takes on its most condensed structure and features such as the banding pattern can be visualized.

Methyl-CpG-binding protein (MeCP) A protein that binds to methylated CpG islands and may influence acetylation of nearby histones.

MGMT (O6-methylguanine-DNA methyltransferase) An enzyme involved in the direct repair of alkylation mutations.

Microarray A low-density array of DNA molecules used for parallel hybridization analysis.

Microbiome The microorganisms that live on or within the human body.

Microchromosome One of the shorter gene-rich chromosomes seen in the nuclei of chickens and various other species.

MicroRNA (miRNA) A class of short RNAs involved in regulation of gene expression in eukaryotes, and which act by a pathway similar to RNA interference.

Microsatellite A type of simple sequence length polymorphism comprising tandem copies of, usually, di-, tri-, or tetranucleotide repeat units. Also called a short tandem repeat (STR).

Miniature inverted repeat transposable element (MITE) A general term for the truncated relic of a DNA transposon.

Minigene The name given to the pair of exons carried by a cloning vector used in the exon-trapping procedure.

Minisatellite A type of simple sequence length polymorphism comprising tandem copies of repeats that are a few tens of nucleotides in length. Also called a variable number of tandem repeats (VNTR).

Minor groove The smaller of the two grooves that spiral around the surface of the B-form of DNA.

Miscoding lesion A sequence error caused by the chemical alteration of a nucleotide resulting from the partial degradation of an ancient DNA molecule.

Mismatch A position in a double-stranded DNA molecule where base pairing does not occur because the nucleotides are not complementary; in particular, a non-base-paired position resulting from an error in replication.

Mismatch repair A DNA repair process that corrects mismatched nucleotide pairs by replacing the incorrect nucleotide in the daughter polynucleotide.

Mitochondrial genome The genome present in the mitochondria of a eukaryotic cell.

Mitochondrion One of the energy-generating organelles of eukaryotic cells.

Mitosis The series of events that results in nuclear division.

Mitosis or M phase The stage of the cell cycle when mitosis or meiosis occurs.

Mobile phase The movable phase in a chromatography system, usually a liquid in which the compounds have been dissolved or a gas in which they have been vaporized.

Model organism An organism that is relatively easy to study and hence can be used to obtain information that is relevant to the biology of a second organism that is more difficult to study.

Modification assay A range of techniques used for locating bound proteins on DNA molecules.

Modification interference A technique used to identify nucleotides involved in interactions with a DNA-binding protein.

Modification protection A technique used to identify nucleotides involved in interactions with a DNA-binding protein.

Molecular beacon A method, based on dye quenching, used to type SNPs.

Molecular biologist A person who studies the molecular life sciences.

Molecular chaperone A protein that helps other proteins to fold.

Molecular clock A device based on the inferred mutation rate that enables times to be assigned to the branch points in a gene tree.

Molecular combing A technique for preparing restricted DNA molecules for optical mapping.

Molecular evolution The gradual changes that occur in genomes over time as a result of the accumulation of mutations and structural rearrangements resulting from recombination and transposition.

Molecular ion An ion formed during peptide mass fingerprinting: \([M+H]^+\) and \([M-H]^–\), where M is the peptide.

Molecular life sciences The area of research comprising molecular biology, biochemistry, and cell biology, as well as some aspects of genetics and physiology.

Molten globule An intermediate in protein folding formed by the rapid collapse of a polypeptide into a compact structure with slightly larger dimensions than the final protein.

Monogenic A characteristic that is specified by a single gene.

Monohybrid cross A sexual cross in which the inheritance of one pair of alleles is followed.

Multicopy A gene, cloning vector, or other genetic element that is present in multiple copies in a single cell.

Multicysteine zinc finger A type of zinc-finger DNA-binding domain.

Multidimensional protein identification technique (MudPIT) A technique that combines various
chromatography methods in order to isolate intact protein complexes.

**Glossary**

**Multigene family** A group of genes, clustered or dispersed, with related nucleotide sequences.

**Multiple alignment** An alignment of three or more nucleotide sequences.

**Multiple alleles** The different alternative forms of a gene that has more than two alleles.

**Mutagen** A chemical or physical agent that can cause a mutation in a DNA molecule.

**Mutagenesis** Treatment of a group of cells or organisms with a mutagen as a means of inducing mutations.

**Mutant** A cell or organism that possesses a mutation.

**Mutasome** A protein complex that is constructed during the SOS response of *Escherichia coli*.

**Mutation** An alteration in the nucleotide sequence of a DNA molecule.

**Mutator-like transposable element (MULE)** A type of DNA transposon that is able to capture exons and other gene segments.

**N50 size** A measure of the degree of completeness of a genome sequence.

**N-linked glycosylation** The attachment of sugar units to an asparagine in a polypeptide.

**N-terminus** The end of a polypeptide that has a free amino group.

**Nanopore sequencing** A fourth-generation sequencing method.

**Next-generation sequencing** A collection of DNA sequencing methods, each involving a massively parallel strategy.

**NG50 size** A measure of the degree of completeness of a genome sequence.

**NH2-terminus** The end of a polypeptide that has a free amino group.

**Nick** A position in a double-stranded DNA molecule where one of the polynucleotides is broken as a result of the absence of a phosphodiester bond.

**Nitrogenous base** One of the purines or pyrimidines that form part of the molecular structure of a nucleotide.

**Nonhomologous end-joining (NHEJ)** Another name for the double-strand break repair process.

**Node** The depiction of a protein in a protein interaction map.

**Nonchromatin region** The space separating the chromosome territories within a nucleus.

**Noncoding RNA** An RNA molecule that does not code for a protein.

**Nonpenetrance** The situation whereby the effect of a mutation is never observed during the lifetime of a mutant organism.

**Nonpolar** A hydrophobic (water-hating) chemical group.

**Nonprocessed pseudogene** A gene that has become inactive because of the accumulation of mutations.

**Northern blotting** The transfer of RNA from an electrophoresis gel to a membrane prior to northern hybridization.

**Northern hybridization** A technique used for detection of a specific RNA molecule against a background of many other RNA molecules.

**Nuclear genome** The DNA molecules present in the nucleus of a eukaryotic cell.

**Nuclear lamina** A network of filaments on the internal side of the nuclear membrane.

**Nuclear magnetic resonance (NMR) spectroscopy** A technique for determining the three-dimensional structure of large molecules.

**Nuclear matrix** A proteinaceous scaffold-like network that is thought to permeate the nucleus.

**Nuclear receptor superfamily** A family of receptor proteins that bind hormones as an intermediate step in the modulation of genome activity by these hormones.

**Nuclease** An enzyme that degrades a nucleic acid molecule.

**Nuclease protection experiment** A technique that uses nuclease digestion to determine the positions of proteins on DNA or RNA molecules.

**Nucleic acid** The term first used to describe the acidic chemical compound isolated from the nuclei of eukaryotic cells. Now used specifically to describe a polymeric molecule comprising nucleotide monomers, such as DNA and RNA.

**Nucleic acid hybridization** Formation of a double-stranded hybrid by base pairing between complementary polynucleotides.

**Nucleoid** The DNA-containing region of a prokaryotic cell.

**Nucleoid-associated proteins** The protein component of a bacterial nucleoid.

**Nucleolus** The region of the eukaryotic nucleus in which rRNA transcription occurs.

**Nucleoside** A purine or pyrimidine base attached to a five-carbon sugar.

**Nucleosome** The complex of histones and DNA that is the basic structural unit in chromatin.

**Nucleosome remodeling** A change in the conformation of a nucleosome, which is associated with a change in access to the DNA with which the nucleosome is associated.

**Nucleotide** A purine or pyrimidine base attached to a five-carbon sugar, to which a mono-, di-, or triphosphate is also attached; the monomeric unit of DNA and RNA.

**Nucleotide excision repair** A repair process that corrects various types of DNA damage by excising and resynthesizing a region of a polynucleotide.

**Nucleus** The membrane-bound structure of a eukaryotic cell in which the chromosomes are contained.

**O-linked glycosylation** The attachment of sugar units to a serine or threonine in a polypeptide.
Okazaki fragment  One of the short segments of RNA-primed DNA synthesized during replication of the lagging strand of the double helix.

Oligonucleotide  A short, synthetic, single-stranded DNA molecule.

Oligonucleotide hybridization analysis  The use of an oligonucleotide as a hybridization probe.

Oligonucleotide ligation assay (OLA)  A technique for SNP typing, which depends on ligation of two oligonucleotides that anneal adjacent to one another, with one covering the position of the SNP.

Oligonucleotide-directed mutagenesis  An in vitro mutagenesis technique in which a synthetic oligonucleotide is used to introduce a predetermined nucleotide alteration into the gene to be mutated.

Oncogene  A gene that, when defective, can give rise to cancer.

One-step growth curve  A single infection cycle for a lytic bacteriophage.

Oocyte  An unfertilized female egg cell.

Open promoter complex  A structure formed during assembly of the transcription initiation complex consisting of the RNA polymerase and/or accessory proteins attached to the promoter, after the DNA has been opened up by breakage of base pairs.

Open reading frame (ORF)  A series of codons starting with an initiation codon and ending with a termination codon; the part of a protein-coding gene that is translated into protein.

Operator  The nucleotide sequence to which a repressor protein binds to prevent transcription of a gene or operon.

Operon  A set of adjacent genes in a bacterial genome, transcribed from a single promoter and subject to the same regulatory regime.

Optical mapping  A technique for the direct visual examination of restricted DNA molecules.

ORF scanning  Examination of a DNA sequence for open reading frames in order to locate the genes.

Origin licensing  The construction of pre-replication complexes on replication origins.

Origin of replication  A site on a DNA molecule where replication initiates.

Origin recognition complex (ORC)  A set of proteins that binds to the origin recognition sequence.

Origin recognition sequence  A component of a eukaryotic origin of replication.

Orphan retrogene  A retrogene whose parent copy has been lost.

Orthogonal field alternation gel electrophoresis (OFAGE)  An electrophoresis system, in which the field alternates between pairs of electrodes set at an angle of 45°, used to separate large DNA molecules.

Orthologous  Refers to homologous genes located in the genomes of different organisms.

Overlap graph  The output of a sequence assembler, comprising overlapping sequence reads.

Overlapping genes  Two genes whose coding regions overlap.

P element  A DNA transposon of Drosophila.

P1-derived artificial chromosome (PAC)  A high-capacity vector that combines features of bacteriophage P1 vectors and bacterial artificial chromosomes.

p300/CBP  A protein complex that is able to modify histone proteins and so affect chromatin structure and nucleosome positioning.

PacBio sequencing  A version of single-molecule real-time DNA sequencing.

Pair-rule genes  Developmental genes that establish the basic segmentation pattern of the Drosophila embryo.

Paired-end reads  Mini-sequences from the two ends of a single cloned fragment.

Paleogenomics  The study of the genomes of extinct species.

Pan-genome concept  The concept that views a bacterial genome as a combination of a core and accessory genome.

Paralogous  Refers to two or more homologous genes located in the same genome.

Paranemic  Refers to a helix whose strands can be separated without unwinding.

Pararetrovirus  A viral retroelement whose encapsidated genome is made of DNA.

Parental genotype  The genotype possessed by one or both of the parents in a genetic cross.

Partial linkage  The type of linkage usually displayed by a pair of genetic and/or physical markers on the same chromosome, the markers not always being inherited together because of the possibility of recombination between them.

Partial restriction  Digestion of DNA with a restriction endonuclease under limiting conditions so that not all restriction sites are cut.

Pedigree  A chart showing the genetic relationships between the members of a human family.

Pedigree analysis  The use of pedigree charts to analyze the inheritance of a genetic or DNA marker in a human family.

Pentose  A sugar comprising five carbon atoms.

Peptide bond  The chemical link between adjacent amino acids in a polypeptide.

Peptide mass fingerprinting  Identification of a protein by examination of the mass spectrometric properties of peptides generated by treatment with a sequence-specific protease.

Peptide nucleic acid (PNA)  A polynucleotide analog in which the sugar-phosphate backbone is replaced by amide bonds.

Peptidyl or P site  The site in the ribosome occupied by the tRNA attached to the growing polypeptide during translation.

Peptidyl transferase  The enzyme activity that synthesizes peptide bonds during translation.
**Personalized medicine**  The use of individual genome sequences to make accurate diagnoses of a person’s risk of developing a disease, and the use of that person’s genetic characteristics to plan effective therapies and treatment regimes.

**Phage**  A virus that infects a bacterium.

**Phage display**  A technique for identifying proteins that interact with one another.

**Phage display library**  A collection of clones carrying different DNA fragments, used in phage display.

**Phenotype**  The observable characteristics displayed by a cell or organism.

**Philadelphia chromosome**  An abnormal chromosome resulting from a translocation between human chromosomes 9 and 22; a common cause of chronic myeloid leukemia.

**Phosphate group**  One of the components of a nucleotide.

**Phosphodiester bond**  The chemical link between adjacent nucleotides in a polynucleotide.

**Phosphodiesterase**  A type of enzyme that can break phosphodiester bonds.

**Phosphorylase**  An enzyme that adds a phosphate group to another molecule.

**Photobleaching**  A component of the FRAP technique for studying protein mobility in the nucleus.

**Photolithography**  A technique that uses pulses of light to construct an oligonucleotide from light-activated nucleotide substrates.

**Photolyase**  An *Escherichia coli* enzyme involved in photoreactivation DNA repair.

**Photoproduct**  A modified nucleotide resulting from treatment of DNA with ultraviolet radiation.

**Photoreactivation**  A DNA repair process in which cyclobutyl dimers and (6–4) photoproducts are corrected by a light-activated enzyme.

**Phylogenetic tree**  A tree depicting the evolutionary relationships between a set of DNA sequences, species, or other taxa.

**Physical mapping**  The use of molecular biology techniques to construct a genome map.

**Plus**  A structure involved in bringing a pair of bacteria together during conjugation; possibly the tube through which DNA is transferred.

**Piwi protein**  A type of protein that forms a complex with piRNA to form structures that regulate gene expression during various developmental processes.

**Piwi-interacting RNA (piRNA)**  A type of snRNA, 25–30 nucleotides in length, that associates with piwi proteins.

**Plant GDB**  An online plant genome browser.

**Plaque**  A zone of clearing on a lawn of bacteria caused by lysis of the cells by infecting bacteriophages.

**Plasmid**  A usually circular piece of DNA often found in bacteria and some other types of cell.

**Plectonemic**  Refers to a helix whose strands can only be separated by unwinding.

**Point centromere**  A type of centromere, found in *Saccharomyces cerevisiae*, which does not contain repetitive DNA and instead is defined by a single-copy sequence.

**Point mutation**  A mutation that results from a single nucleotide change in a DNA molecule.

**Polar**  A hydrophilic (water-loving) chemical group.

**Poly(A) polymerase**  The enzyme that attaches a poly(A) tail to the 3’-end of a eukaryotic mRNA.

**Poly(A) tail**  A series of A nucleotides attached to the 3’-end of a eukaryotic mRNA.

**Polyacrylamide gel electrophoresis**  Electrophoresis carried out in a polyacrylamide gel and used to separate DNA molecules between 10 and 1500 bp in length.

**Polyadenylate-binding protein**  A protein that aids poly(A) polymerase during polyadenylation of eukaryotic mRNAs, and which plays a role in maintenance of the tail after synthesis.

**Polyadenylation**  The addition of a series of As to the 3’-end of a eukaryotic mRNA.

**Polycomb group (PCG)**  A group of proteins that induce localized formation of heterochromatin.

**Polycomb response element**  The DNA recognition sequences for Polycomb group proteins.

**Polymer**  A compound made up of a long chain of identical or similar units.

**Polymerase chain reaction (PCR)**  A technique that results in exponential amplification of a selected region of a DNA molecule.

**Polynucleotide**  A single-stranded DNA or RNA molecule.

**Polynucleotide kinase**  An enzyme that adds phosphate groups to the 5’-ends of DNA molecules.

**Polypeptide**  A polymer of amino acids.

**Polyploidy**  Having two or more diploid genome copies.

**Polypeptide**  A translation product consisting of a series of linked proteins that are processed by proteolytic cleavage to release the mature proteins.

**Polypyrimidine tract**  A pyrimidine-rich region near the 3’-end of a GU-AG intron.

**Positional effect**  Refers to the different levels of expression that result after insertion of a gene at different positions in a eukaryotic genome.

**Post-spliceosome complex**  The immediate product of the splicing reaction for a GU-AG intron, which dissociates into the spliced mRNA and the intron lariat.

**POU domain**  A DNA-binding motif found in a variety of proteins.

**Precatalytic spliceosome**  An intermediate in the splicing pathway for a GU-AG intron, the immediate precursor of the spliceosome.

**Pre-mRNA**  The primary transcript of a protein-coding gene.

**Pre-replication complex (pre-RC)**  A protein complex that is constructed at a eukaryotic origin of replication and enables initiation of replication to occur.
Pre-RNA  The initial product of transcription of a gene or group of genes, subsequently processed to give the mature transcript(s).

Pre-rRNA  The primary transcript of a gene or group of genes specifying tRNA molecules.

Pre-tRNA  The primary transcript of a gene or group of genes specifying tRNA molecules.

Preinitiation complex  (1) The structure comprising the small subunit of the ribosome and the initiator tRNA plus ancillary factors that forms the initial association with the mRNA during protein synthesis. (2) The structure that forms at the core promoter of a gene transcribed by RNA polymerase II. (3) The activated version of a pre-replication complex.

Prepriming complex  A complex of proteins formed during initiation of genome replication in bacteria.

Prespliceosome complex  An intermediate in the splicing pathway for a GU-AG intron.

Pribnow box  A component of the bacterial promoter.

Primary structure  The sequence of amino acids in a polypeptide.

Primary transcript  The initial product of transcription of a gene or group of genes, subsequently processed to give the mature transcript(s).

Primase  The RNA polymerase enzyme that synthesizes RNA primers during bacterial DNA replication.

Primer  A short oligonucleotide that is attached to a single-stranded DNA molecule in order to provide a start point for strand synthesis.

Primosome  A protein complex involved in genome replication.

Prion  An unusual infectious agent that consists purely of protein.

Processed pseudogene  A pseudogene that results from the integration into the genome of a reverse-transcribed copy of an mRNA.

Processivity  Refers to the amount of DNA synthesis that is carried out by a DNA polymerase before dissociation from the template.

Programmable nuclease  A nuclease that can be directed to a specific site in a genome.

Prokaryote  An organism whose cells lack a distinct nucleus.

Proliferating cell nuclear antigen (PCNA)  An accessory protein involved in genome replication in eukaryotes.

Promiscuous DNA  DNA that has been transferred from one organelle genome to another.

Promoter  The nucleotide sequence, upstream of a gene, to which RNA polymerase binds in order to initiate transcription.

Promoter clearance  The completion of successful initiation of transcription that occurs when the RNA polymerase moves away from the promoter sequence.

Promoter escape  The stage in transcription during which the polymerase moves away from the promoter region and becomes committed to making a transcript.

Proofreading  The 3′→5′ exonuclease activity possessed by some DNA polymerases, which enables the enzymes to replace a misincorporated nucleotide.

Prophage  The integrated form of the genome of a lysogenic bacteriophage.

PROSITE  An online protein structure database.

Protease  An enzyme that degrades protein.

Proteasome  A multisubunit protein structure that is involved in the degradation of other proteins.

Protein  The polymeric compound made of amino acid monomers.

Protein array  A microarray comprising immobilized proteins.

Protein electrophoresis  Separation of proteins in an electrophoresis gel.

Protein engineering  Various techniques for making directed alterations in protein molecules, often to improve the properties of enzymes used in industrial processes.

Protein folding  The adoption of a folded structure by a polypeptide.

Protein interaction map  A map showing the interactions between all or some of the proteins in a proteome.

Protein profiling  The methodology used to identify the proteins in a proteome.

Protein–protein cross-linking  A technique that links together adjacent proteins in order to identify proteins that are positioned close to one another in a structure such as a ribosome.

Proteome  The collection of functioning proteins synthesized by a living cell.

Proteomics  A variety of techniques used to study proteomes.

Protogenome  An RNA genome that existed during the RNA world.

Protomer  One of the protein subunits in a bacteriophage or virus capsid.

Protoplast  A cell from which the cell wall has been completely removed.

Pseudogene  An inactivated and hence nonfunctional copy of a gene.

PSI-BLAST  A modified and more powerful version of the BLAST algorithm.

Pulse labeling  A brief period of labeling carried out at a defined period during the progress of an experiment.

Punctuation codon  A codon that specifies either the start or the end of a gene.

Purine  One of the two types of nitrogenous base found in nucleotides.

Pyrimidine  One of the two types of nitrogenous base found in nucleotides.

Pyrosequencing  A novel DNA sequencing method in which addition of a nucleotide to the end of a growing polynucleotide is detected directly by conversion of the released pyrophosphate into a flash of chemiluminescence.
**Quadrupole mass analyzer** A mass spectrometer in which the mass analyzer has four magnetic rods placed parallel to one another, surrounding a central channel through which the ions must pass.

**Quantitative PCR** A PCR method that enables the number of DNA molecules in a sample to be estimated.

**Quantitative trait locus (QTL)** A region of a genome, which possibly contains several genes, that controls a variable trait.

**Quaternary structure** The structure resulting from the association of two or more polypeptides.

**Radiation hybrid** A collection of rodent cell lines that contain different fragments of a second genome; it is constructed by a technique involving irradiation and used as a mapping reagent, for example in studies of the human genome.

**Radioactive marker** A radioactive atom incorporated into a molecule and whose radioactive emissions are subsequently used to detect and follow that molecule during a biochemical reaction.

**Radiolabeling** The technique for attaching a radioactive atom to a molecule.

**Random genomic sequences** Sequence-tagged sites obtained by sequencing random pieces of cloned genomic DNA.

**Rapid amplification of cDNA ends (RACE)** A PCR-based technique for mapping the end of an RNA molecule.

**Read** A single sequence from the output of a next-generation sequencing run.

**Reading frame** A series of triplet codons in a DNA sequence.

**Real-time PCR** A modification of the standard PCR technique in which synthesis of the product is measured as the PCR proceeds through its series of cycles.

**RecA** An *Escherichia coli* protein involved in homologous recombination.

**RecBCD complex** An enzyme complex involved in homologous recombination in *Escherichia coli*.

**Recessive** The allele that is not expressed in a heterozygote.

**RecFOR pathway** A pathway for homologous recombination in *Escherichia coli*.

**Reciprocal strand exchange** The exchange of DNA between two double-stranded molecules, occurring as a result of recombination, such that the end of one molecule is exchanged for the end of the other molecule.

**Recognition helix** An α-helix in a DNA-binding protein that is responsible for recognition of the target nucleotide sequence.

**Recombinant** A progeny member that possesses neither of the combinations of alleles displayed by the parents.

**Recombinant DNA molecule** A DNA molecule created in the test tube by ligating pieces of DNA that are not normally joined together.

**Recombinant DNA technology** The techniques involved in the construction, study, and use of recombinant DNA molecules.

**Recombinant genotype** A genotype not possessed by either of the parents in a genetic cross.

**Recombinant plasmid** A plasmid that contains an inserted piece of DNA.

**Recombinase** A diverse family of enzymes that catalyze site-specific recombination events.

**Recombination** A large-scale rearrangement of a DNA molecule.

**Recombination frequency** The proportion of recombinant progeny arising from a genetic cross.

**Recombination hotspot** A region of a chromosome where crossovers occur at a higher frequency than the average for the chromosome as a whole.

**Recombination repair** A DNA repair process that involves excising and replacing DNA segments.

**Reference genome** An existing genome sequence that is used to aid assembly of the reads obtained by next-generation sequencing of a related genome.

**Reflectron** An ion mirror used in some types of mass spectrometer; also used to denote a mass spectrometer that contains an ion mirror.

**Regional centromere** A typical eukaryotic centromere, associated with a region of repetitive DNA.

**Renaturation** The return of a denatured molecule to its natural state.

**Replicative DNA** A DNA sequence that is repeated two or more times in a DNA molecule or genome.

**Replicative DNA fingerprinting** A clone fingerprinting technique that involves determining the positions of genomewide repeats in cloned DNA fragments.

**Replicative DNA PCR** A clone fingerprinting technique that uses PCR to detect the relative positions of genomewide repeats in cloned DNA fragments.

**Repetitive extragenic palindromic (REP) sequences** Type of bacterial repetitive DNA made up of motifs, most of which are 20–35 bp in length, that occur singly or in arrays.

**Replacement vector** A λ vector designed so that insertion of new DNA is by replacement of part of the nonessential region of the λ DNA molecule.

**Replication factor C (RFC)** A multisubunit accessory protein involved in eukaryotic genome replication.

**Replication factory** A discrete, immobile region that contains all the relevant proteins for DNA replication and through which the DNA threads as it is being replicated.

**Replication foci** Discrete, immobile regions that contain all the relevant proteins for DNA replication and through which the DNA threads as it is being replicated.

**Replication fork** The region of a double-stranded DNA molecule that is being opened up to enable DNA replication to occur.

**Replication mediator protein (RMP)** A protein responsible for detachment of single-strand binding proteins during genome replication.

**Replication origin** A site on a DNA molecule where replication initiates.
Replication protein A (RPA) The main single-strand binding protein involved in replication of eukaryotic DNA.

Replication slippage An error in replication that leads to an increase or decrease in the number of repeat units in a tandem repeat such as a microsatellite.

Replicative transposition Transposition that results in copying of the transposable element.

Replisome A complex of proteins involved in genome replication.

Reporter gene A gene whose phenotype can be assayed and which can therefore be used to determine the function of a regulatory DNA sequence.

Reporter probe A short oligonucleotide that gives a fluorescent signal when it hybridizes with a target DNA.

Repressible operon An operon that is switched off by a co-repressor molecule.

Resin A chromatography matrix.

Resolution Separation of a pair of recombining double-stranded DNA molecules.

Resolvase A protein capable of resolving a Holliday structure.

Resonance frequency The energy difference between the α and β spin states of a nucleus.

Restriction endonuclease An enzyme that cuts DNA molecules at a limited number of specific nucleotide sequences.

Restriction fragment length polymorphism (RFLP) A restriction fragment whose length is variable because of the presence of a polymorphic restriction site at one or both ends.

Restriction mapping Determination of the positions of restriction sites in a DNA molecule by analyzing the sizes of restriction fragments.

Restriction pattern The set of fragments obtained after digestion of a DNA molecule with a restriction endonuclease, referring to the pattern of bands obtained after separation of the fragments by gel electrophoresis.

Retroelement A genetic element that transposes via an RNA intermediate.

Retrogene A gene duplicate that arises by insertion of a pseudogene adjacent to the promoter of an existing gene.

Retrohoming A process during which an excised intron, comprising single-stranded RNA, inserts directly into an organelle genome prior to being copied into double-stranded DNA.

Retron The commonest type of bacterial retroelement.

Retroposon A retroelement that does not have LTRs.

Retrotansposition Transposition via an RNA intermediate.

Retrotansposon A genomewide repeat with a sequence similar to an integrated retroviral genome and possibly with retrotansposition activity.

Retrovirus A virus with an RNA genome that integrates into the genome of its host cell.

Reverse genetics The strategy by which the function of a gene is identified by mutating that gene and identifying the phenotypic change that results.

Reverse-phase liquid chromatography (RPLC) A column chromatography method that separates proteins according to their degree of surface hydrophobicity.

Reverse sequence One of the two directions in which a double-stranded DNA molecule can be sequenced.

Reverse transcriptase A polymerase that synthesizes DNA on an RNA template.

Reverse transcriptase PCR (RT-PCR) PCR in which the first step is carried out by reverse transcriptase, so RNA can be used as the starting material.

Reversible terminator sequencing A DNA sequencing method in which the sequence is read by detection of the fluorescent label attached to each nucleotide that is added to a growing polynucleotide.

Rho A protein involved in termination of transcription of some bacterial genes.

Rho-dependent terminator A position in bacterial DNA where termination of transcription occurs with the involvement of Rho.

Ribonucleic acid, one of the two forms of nucleic acid.

Ribonuclease An enzyme that degrades RNA.

Ribose The sugar component of a ribonucleotide.

Ribosomal protein One of the protein components of a ribosome.

Ribosomal RNA (rRNA) The RNA molecules that are components of ribosomes.

Ribosome One of the protein–RNA assemblies on which translation occurs.

Ribosome binding site The nucleotide sequence that acts as the attachment site for the small subunit of the ribosome during initiation of translation in bacteria.

Ribosome modulation factor A protein involved in the inactivation of surplus ribosomes in Escherichia coli.

Riboswitch A segment of an mRNA that can bind a small molecule, attachment of this molecule affecting the translation or processing of the mRNA.

Ribozyme An RNA molecule that has catalytic activity.

RNA Ribonucleic acid, one of the two forms of nucleic acid in living cells; the genetic material for some viruses.

RNA editing A process by which nucleotides not coded by a gene are introduced at specific positions in an RNA molecule after transcription.

RNA interference (RNAi) An RNA degradation process in eukaryotes.

RNA polymerase An enzyme that synthesizes RNA on a DNA or RNA template.

RNA polymerase I The eukaryotic RNA polymerase that transcribes ribosomal RNAs.

RNA polymerase II The eukaryotic RNA polymerase that transcribes protein-coding and snRNA genes.
RNA polymerase III The eukaryotic RNA polymerase that transcribes tRNA and other short genes.

RNA recognition domain An RNA-binding domain.

RNA sequencing (RNA-seq) Next-generation sequencing of RNA.

RNA silencing An RNA degradation process in eukaryotes.

RNA transcript An RNA copy of a gene.

RNA world The early period of evolution when all biological reactions were centered on RNA.

RNA-dependent DNA polymerase An enzyme that makes a DNA copy of an RNA template; a reverse transcriptase.

RNA-dependent RNA polymerase An enzyme that makes an RNA copy of an RNA template.

RNA-induced silencing complex (RISC) A complex of proteins that cleaves and hence silences an mRNA as part of the RNA interference pathway.

Rolling-circle replication A replication process that involves continual synthesis of a polynucleotide, which is rolled off of a circular template molecule.

S phase The stage of the cell cycle when DNA synthesis occurs.

S value The unit of measurement for a sedimentation coefficient.

S1 nuclease An enzyme that degrades single-stranded DNA or RNA molecules, including single-stranded regions in predominantly double-stranded molecules.

Satellite DNA Repetitive DNA that forms a satellite band in a density gradient.

Satellite RNA An RNA molecule some 320–400 nucleotides in length that does not encode its own capsid proteins, instead moving from cell to cell within the capsid of a helper virus.

Scaffold A series of sequence contigs separated by sequence gaps.

Scaffold/matrix attachment regions (S/MARs) Nucleotide sequences, within chromosomes, that bind to the proteins of the nuclear matrix.

Scanning A system used during initiation of eukaryotic translation, in which the preinitiation complex attaches to the 5′-terminal cap structure of the mRNA and then scans along the molecule until it reaches an initiation codon.

Second messenger An intermediate in a certain type of signal transduction pathway.

Secondary structure The conformations, such as the α-helix and β-sheet, taken up by a polypeptide.

Sedimentation analysis The centrifugal technique used to measure the sedimentation coefficient of a molecule or structure.

Sedimentation coefficient The value used to express the velocity at which a molecule or structure sediments when centrifuged in a dense solution.

Segment polarity genes Developmental genes that provide greater definition to the segmentation pattern of the Drosophila embryo that is established by the action of the pair-rule genes.

Segmental duplication A duplication between 1 and 400 kb in length, with greater than 90% sequence identity, that is repeated up to 50 times in the genome.

Segmented genome A virus genome that is split into two or more DNA or RNA molecules.

Selectable marker A gene carried by a vector and conferring a recognizable characteristic on a cell containing the vector or a recombinant DNA molecule derived from the vector.

Selfish DNA DNA that appears to have no function and apparently contributes nothing to the cell in which it is found.

Semiconservative replication The mode of DNA replication in which each daughter double helix is made up of one polynucleotide from the parent and one newly synthesized polynucleotide.

Sequence assembler A software package that converts sequence reads into contigs.

Sequence assembly Assembly of the many short reads obtained by next-generation sequencing into a contiguous DNA sequence.

Sequence contig A contiguous DNA sequence obtained as an intermediate in a genome sequencing project.

Sequence coverage or sequence depth The average number of reads that cover each nucleotide position in a DNA sequence obtained by a next-generation method.

Sequence-tagged site (STS) A DNA sequence that is unique in the genome.

Sequence-tagged site (STS) content mapping A clone fingerprinting technique.

Sequencing by oligonucleotide ligation and detection (SOLID) A next-generation sequencing method in which the sequence is deduced by hybridization of a series of oligonucleotides whose sequences are complementary to that of the template.

Sequence library A set of DNA fragments that have been immobilized on a solid support in such a way that multiple sequencing reactions can be carried out side by side in a massively parallel array format.

Serial analysis of gene expression (SAGE) A method for studying the composition of a transcriptome.

Serine–threonine kinase receptors A type of cell surface receptor that has serine–threonine kinase activity.

Sex cell A reproductive cell; a cell that divides by meiosis.

Sex chromosome A chromosome that is involved in sex determination.

Shelterin A structure, comprising telomere-binding proteins, that protects the telomeres from degradation by nuclease enzymes and mediates the enzymatic activity that maintains the length of each telomere during DNA replication.

Shine–Dalgarno sequence Another name for the prokaryotic ribosome binding site.

Short interfering RNA (siRNA) An intermediate in the RNA interference pathway.
**Short nascent strand (SNS) sequencing** A method used to identify the positions of replication origins in a eukaryotic genome sequence.

**Short noncoding RNA (snRNA)** Noncoding RNAs less than 200 nucleotides in length.

**Short tandem repeat (STR)** A type of simple sequence length polymorphism comprising tandem copies of, usually, di-, tri-, or tetranucleotide repeat units. Also called a microsatellite.

**Shotgun method** A genome sequencing strategy in which the molecules to be sequenced are randomly broken into fragments that are then sequenced individually.

**Shotgun proteomics** A version of proteomics in which a mixture of proteins is fragmented.

**Shuttle vector** A vector that can replicate in the cells of more than one organism (for example, in Escherichia coli and in yeast).

**Signal peptide** A short sequence at the N-terminus of some proteins that directs the protein across a membrane.

**Signal transduction** Control of cellular activity, including genome expression, via a cell surface receptor that responds to an external signal.

**Silencer** A regulatory sequence that reduces the rate of transcription of a gene or genes located some distance away in either direction.

**Simple sequence length polymorphism (SSLP)** An array of repeat sequences that display length variations.

**SINE (short interspersed nuclear element)** A type of genomewide repeat, typified by the Alu sequences found in the human genome.

**Single-copy DNA** A DNA sequence that is not repeated elsewhere in the genome.

**Single-molecule real-time sequencing** A third-generation DNA sequencing method that uses an advanced optical system to observe the addition of individual nucleotides to a growing polynucleotide.

**Single-nucleotide polymorphism (SNP)** A point mutation that is carried by some individuals of a population.

**Single-strand binding protein (SSB)** One of the proteins that attach to single-stranded DNA in the region of the replication fork, preventing base pairs forming between the two parent strands before they have been copied.

**Single-stranded** A DNA or RNA molecule that comprises just a single polynucleotide.

**Site-directed hydroxyl radical probing** A technique for locating the position of a protein in a protein–RNA complex, such as a ribosome, by making use of the ability of Fe(II) ions to generate hydroxyl radicals, which cleave nearby RNA phosphodiester bonds.

**Site-directed mutagenesis** Techniques used to produce a specified mutation at a predetermined position in a DNA molecule.

**Site-specific recombination** Recombination between two double-stranded DNA molecules that have only short regions of nucleotide sequence similarity.

**Small Cajal body-specific RNA (scaRNA)** A snoRNA associated with a Cajal body.

**Small nuclear ribonucleoprotein (snRNP)** A structure involved in the splicing of GU-AG and AU-AC introns and in other RNA processing events, comprising one or two snRNA molecules complexed with proteins.

**Small nuclear RNA (snRNA)** A type of short eukaryotic RNA molecule involved in the splicing of GU-AG and AU-AC introns and in other RNA processing events.

**Small nucleolar RNA (snoRNA)** A type of short eukaryotic RNA molecule involved in chemical modification of rRNA.

**SOLiD** A next-generation sequencing method in which the DNA sequence is deduced by hybridization of a series of oligonucleotides whose sequences are complementary to that of the template.

**Solid phase** The immobile phase in a chromatography system.

**Solution hybridization** Hybridization between nucleic acid molecules carried out in a solution.

**Somatic cell** A nonreproductive cell; a cell that divides by mitosis.

**Sonication** A procedure that uses ultrasound to cause random breaks in DNA molecules.

**Sorting sequence** An amino acid sequence that directs a protein to an organelle such as the nucleus or mitochondrion, or might specify that the protein is to be secreted from the cell.

**SOS response** A series of biochemical changes that occurs in Escherichia coli in response to damage to the genome and other stimuli.

**Southern hybridization** A technique used for detection of a specific restriction fragment against a background of many other restriction fragments.

**Speckle** A nuclear structure associated with mRNA splicing.

**Spliceosome** The protein–RNA complex involved in splicing GU-AG or AU-AC introns.

**Splicing** The removal of introns from the primary transcript of a discontinuous gene.

**Splicing code** A hypothetical code that would explain the impact on a splicing pathway of the various interactions that can occur between enhancers, silencers, and their binding proteins.

**Splicing pathway** The series of events that converts a discontinuous pre-mRNA into a functional mRNA.

**Spm element** A DNA transposon of maize.

**Spontaneous mutation** A mutation that arises from an error in replication.

**SR protein** A protein that plays a role in splice-site selection during splicing of GU-AG introns.

**STAT (signal transducer and activator of transcription)** A type of protein that responds to binding of an extracellular signaling compound to a cell surface receptor by activating a transcription factor.

**Stem cell** A progenitor cell that divides continually throughout the lifetime of an organism.
**Stem–loop structure** A structure made up of a base-paired stem and non-base-paired loop, which can form in a single-stranded polynucleotide that contains an inverted repeat.

**Steroid hormone** A type of extracellular signaling compound.

**Steroid receptor** A protein that binds a steroid hormone after the latter has entered the cell, as an intermediate step in modulation of genome activity.

**Sticky end** An end of a double-stranded DNA molecule where there is a single-stranded extension.

**Streptavidin** A protein, from the bacterium *Streptomyces avidinii*, that has a high binding affinity for biotin.

**Stringent response** A biochemical and genetic response initiated in *Escherichia coli* when the bacterium encounters poor growth conditions such as low levels of essential amino acids.

**Strong promoter** A promoter that directs a relatively large number of productive initiations per unit time.

**STS mapping** A physical mapping procedure that locates the positions of sequence-tagged sites (STSs) in a genome.

**Stuffer fragment** A DNA fragment contained within a λ vector that is replaced by the DNA to be cloned.

**Substitution** A point mutation that escapes the repair processes and results in a permanent change in a DNA sequence.

**Sugar pucker** Alternative conformations of a sugar ring structure.

**Suicide enzyme** An enzyme that is inactivated once it has carried out its biochemical reaction.

**SUMO** A protein related to ubiquitin.

**Superciling** A conformational state in which a double helix is overwound or underwound so that superhelical coiling occurs.

**Surveillance mechanism** A process that identifies mRNAs that lack a termination codon, or have a termination codon at an unexpected position, and which therefore should be degraded.

**Syncytium** A cell-like structure comprising a mass of cytoplasm and many nuclei.

**Synteny** Refers to a pair of genomes in which at least some of the genes are located at similar map positions.

**Synthesis or S phase** The phase of the cell cycle when DNA replication takes place.

**Systems biology** An approach to biology that attempts to link metabolic pathways and subcellular processes with genome expression.

**T-DNA** The portion of the Ti plasmid that is transferred to the plant DNA.

**T4 polynucleotide kinase** An enzyme that adds phosphate groups to the 5’-ends of DNA molecules.

**TAF- and initiator-dependent cofactor (TIC)** A type of protein involved in initiation of transcription by RNA polymerase II.

**Tandem affinity purification (TAP)** A method for isolating protein complexes that makes use of a test protein with a C-terminal extension that binds to calmodulin.

**Tandem mass spectrometry** A type of mass spectrometry that uses two or more mass analyzers linked in series.

**Tandem repeat** Direct repeats that are adjacent to each other.

**Tandemly repeated DNA** DNA sequence motifs that are repeated head to tail.

**Target enrichment** A method for enriching a next-generation DNA sequencing library for fragments derived from particular genes of interest.

**TATA box** A component of the RNA polymerase II core promoter.

**TATA-binding protein (TBP)** A component of the general transcription factor TFII D, the part that recognizes the TATA box of the RNA polymerase II promoter.

**Tautomeric shift** The spontaneous change of a molecule from one structural isomer to another.

**Tautomers** Structural isomers that are in dynamic equilibrium.

**TBP-associated factor (TAF)** One of several components of the general transcription factor TFII D that play ancillary roles in recognition of the TATA box.

**Telomerase** The enzyme that maintains the ends of eukaryotic chromosomes by synthesizing telomeric repeat sequences.

**Telomere** The end of a eukaryotic chromosome.

**Telomere-binding protein (TBP)** A protein that binds to and regulates the length of a telomere.

**Temperate bacteriophage** A bacteriophage that is able to follow a lysogenic mode of infection.

**Template** The polynucleotide that is copied during a strand-synthesis reaction catalyzed by a DNA or RNA polymerase.

**Template-dependent DNA polymerase** An enzyme that synthesizes DNA in accordance with the sequence of a template.

**Template-dependent DNA synthesis** Synthesis of a DNA molecule on a DNA or RNA template.

**Template-dependent RNA polymerase** An enzyme that synthesizes RNA in accordance with the sequence of a template.

**Template-dependent RNA synthesis** Synthesis of an RNA molecule on a DNA or RNA template.

**Template-independent DNA polymerase** An enzyme that synthesizes DNA without the use of a template.

**Template-independent RNA polymerase** An enzyme that synthesizes RNA without the use of a template.

**Terminal deoxynucleotidyl transferase** An enzyme that adds one or more nucleotides to the 3’-end of a DNA molecule.

**Termination codon** One of the three codons that mark the position where translation of an mRNA should stop.
**Terminator sequence** One of several sequences on a bacterial genome involved in termination of genome replication.

**Territory (chromosome)** The region of a nucleus occupied by a single chromosome.

**Tertiary structure** The structure resulting from folding the secondary structural units of a polypeptide.

**Test cross** A genetic cross between a double heterozygote and a double homozygote.

**Thermal cycle sequencing** A DNA sequencing method that uses PCR to generate chain-terminated polynucleotides.

**Thermostable** Able to withstand high temperatures.

**Third-generation sequencing** Methods in which DNA sequencing is carried out in real time.

**Thermostable** Able to withstand high temperatures.

**Thymine** One of the pyrimidine bases found in DNA.

**Ti plasmid** The large plasmid found in those *Agrobacterium tumefaciens* cells able to direct crown gall formation on certain species of plants.

**Tiling array** A collection of oligonucleotide probes, each targeting a different position along a chromosome or a part of a chromosome.

**Tm** Melting temperature.

**Tn3-type transposon** A type of DNA transposon that does not have flanking insertion sequences.

**Top-down proteomics** A version of proteomics in which individual proteins are directly examined by mass spectrometry.

**Topological problem** Refers to the need to unwind the double helix in order for DNA replication to occur, and the difficulties that the resulting rotation of the DNA molecule would cause.

**Topologically associated domain (TAD)** A contiguous segment of chromatin folded into coils and loops.

**Topology** The branching pattern of a phylogenetic tree.

**Totipotent** Refers to a cell that is not committed to a single developmental pathway and can hence give rise to all types of differentiated cell.

**Trailer segment** The untranslated region of an mRNA downstream of the termination codon.

**Trans-displacement** Transfer of a nucleosome from one DNA molecule to another.

**Transcript** An RNA copy of a gene.

**Transcript-specific regulation** Regulatory mechanisms that control protein synthesis by acting on a single transcript or a small group of transcripts coding for related proteins.

**Transcription** The synthesis of an RNA copy of a gene.

**Transcription factor** A protein that activates or represses the initiation of transcription.

**Transcription initiation** The assembly, upstream of a gene, of the complex of proteins that will subsequently copy the gene into RNA.

**Transcription-coupled repair** A nucleotide excision repair process that results in repair of the template strands of genes.

**Transcriptome** The entire mRNA content of a cell.

**Transduction** Transfer of bacterial genes from one cell to another by packaging in a phage particle.

**Transduction mapping** The use of transduction to map the relative positions of genes in a bacterial genome.

**Transfection** The introduction of purified phage DNA molecules into a bacterial cell.

**Transfer RNA (tRNA)** A small RNA molecule that acts as an adaptor during translation and is responsible for decoding the genetic code.

**Transfer-messenger RNA (tmRNA)** A bacterial RNA involved in protein degradation.

**Transformant** A cell that has become transformed by the uptake of naked DNA.

**Transformation** The acquisition by a cell of new genes by the uptake of naked DNA.

**Transformation mapping** The use of transformation to map the relative positions of genes in a bacterial genome.

**Transforming principle** The compound, now known to be DNA, responsible for transformation of an avirulent *Streptococcus pneumoniae* bacterium into a virulent form.

**Transgenic mouse** A mouse that carries a cloned gene.

**Transition** A point mutation that replaces a purine with another purine, or a pyrimidine with another pyrimidine.

**Translation** The synthesis of a polypeptide, the amino acid sequence of which is determined by the nucleotide sequence of an mRNA in accordance with the rules of the genetic code.

**Translational efficiency** The rate at which proteins are synthesized from an mRNA.

**Translesion polymerase** A DNA polymerase that can carry out error-prone replication of a damaged region of DNA.

**Translocation** (1) The attachment of a segment of one chromosome to another chromosome. (2) The movement of a ribosome along an mRNA molecule during translation.

**Transposable element** A genetic element that can move from one position to another in a DNA molecule.

**Transposable phage** A bacteriophage that transposes as part of its infection cycle.

**Transposase** An enzyme that catalyzes transposition of a transposable genetic element.

**Transposition** The movement of a genetic element from one site to another in a DNA molecule.

**Transposon** A genetic element that can move from one position to another in a DNA molecule.

**Transposon tagging** A gene-isolation technique that involves inactivation of a gene by movement of a transposon into its coding sequence, followed by the use of a transposon-specific hybridization probe to isolate a copy of the tagged gene from a clone library.

**Transversion** A point mutation that involves a purine being replaced by a pyrimidine, or vice versa.

**Treble clef finger** A type of zinc finger.
**Trinucleotide repeat expansion disease** A disease that results from the expansion of an array of trinucleotide repeats in or near a gene.

**Triplex** A DNA structure comprising three polynucleotides.

**Trisomy** The presence of three copies of a homologous chromosome in a nucleus that is otherwise diploid.

**Trithorax group (trxG)** A group of proteins that maintain an open chromatin state in the regions of active genes.

**Truncated gene** A gene relic that lacks a segment from one end of the original, complete gene.

**Tudor domain** A five-stranded β-sheet structure encoded by a sequence of approximately 60 amino acids that binds to methylated arginine and/or lysine amino acids contained in other proteins.

**Tus (terminator utilization substance) protein** The protein that binds to a bacterial terminator sequence and mediates termination of genome replication.

**Two-dimensional gel electrophoresis** A method for separation of proteins used especially in studies of the proteome.

**Type 0 cap** The basic cap structure, consisting of 7'-methylguanosine attached to the 5'-end of an mRNA.

**Type 1 cap** A cap structure comprising the basic 5'-terminal cap plus an additional methylation of the ribose of the second nucleotide.

**Type 2 cap** A cap structure comprising the basic 5'-terminal cap plus methylation of the riboses of the second and third nucleotides.

**Tyrosine kinase receptor** A type of cell surface receptor that has tyrosine kinase activity.

**Tyrosine kinase-associated receptor** A type of cell surface receptor that works in conjunction with a protein that has tyrosine kinase activity.

**TrpCarm** Part of the structure of a tRNA molecule.

**U-RNA** A uracil-rich nuclear RNA molecule including the snRNAs and snoRNAs.

**Ubiquitin** A 76-amino-acid protein that, when attached to a second protein, acts as a tag directing that protein for degradation.

**Ubiquitin-receptor protein** A protein that directs ubiquitinated proteins into the proteasome.

**Ubiquitination** The attachment of ubiquitin to a protein.

**UCSC Genome Browser** An online genome browser.

**Unequal crossing over** A recombination event that results in duplication of a segment of DNA.

**Unequal sister chromatid exchange** A recombination event that results in duplication of a segment of DNA.

**Unit factor** Mendel's term for a gene.

**Unit transposon** A Tn3-type transposon.

**Unitary pseudogene** A nonprocessed pseudogene that arises by decay of a gene that is not a member of a gene family and whose function is lost as a result of the inactivation of the gene.

**Universal primer** A sequencing primer that is complementary to the part of the vector DNA immediately adjacent to the point into which new DNA is ligated.

**Untranslated region (UTR)** The parts of an mRNA, upstream and downstream of the open reading frame, that are not translated into protein.

**Upstream** Toward the 5′-end of a polynucleotide.

**Upstream control element** A component of an RNA polymerase I promoter.

**Upstream promoter element** Components of a eukaryotic promoter that lie upstream of the position where the initiation complex is assembled.

**Upstream regulatory sequence** A regulatory sequence, usually a binding site for a transcription factor, found upstream of a gene.

**Uracil** One of the pyrimidine bases found in RNA.

**UvrABC endonuclease** A multienzyme complex involved in the short-patch repair process of *Escherichia coli*.

**V loop** Part of the structure of a tRNA molecule.

**van der Waals forces** A particular type of attractive or repulsive noncovalent bond.

**Variable number of tandem repeats (VNTR)** A type of simple sequence length polymorphism comprising tandem copies of repeats that are a few tens of nucleotides in length. Also called a minisatellite.

**Vault RNA** A type of snRNA found in protein–RNA complexes called vaults, which are found in most eukaryotic cells but whose functions are not known.

**Vegetative cell** A nonreproductive cell; a cell that divides by mitosis.

**Viral retroelement** A virus whose genome replication process involves reverse transcription.

**Viroid** An RNA molecule 240–375 nucleotides in length that contains no genes and never becomes encapsidated, spreading from cell to cell as naked DNA.

**Virulent bacteriophage** A bacteriophage that follows the lytic mode of infection.

**Virus** An infective particle, composed of protein and nucleic acid, that must parasitize a host cell in order to replicate.

**Virusoid** An RNA molecule some 320–400 nucleotides in length that does not encode its own capsid proteins, instead moving from cell to cell within the capsid of a helper virus.

**Weak promoter** A promoter that directs relatively few productive initiations per unit time.

**Wild type** A gene, cell, or organism that displays the typical phenotype and/or genotype for the species and is therefore adopted as a standard.

**Winged helix–turn–helix** A type of DNA-binding domain.

**X inactivation** Inactivation by methylation of most of the genes on one copy of the X chromosome in a female nucleus.

**X-ray crystallography** A technique for determining the three-dimensional structure of a large molecule.
X-ray diffraction  The diffraction of X-rays that occurs during their passage through a crystal.

X-ray diffraction pattern  The pattern obtained after diffraction of X-rays through a crystal.

Yeast two-hybrid system  A technique for identifying proteins that interact with one another.

Z-DNA  A conformation of DNA in which the two polynucleotides are wound into a left-handed helix.

Zero-mode waveguide  A nanostructure that enables individual molecules to be observed.

Zinc finger  A common structural motif for attachment of a protein to a DNA molecule.

Zoo-blotting  A technique that attempts to determine if a DNA fragment contains a gene by hybridizing that fragment to DNA preparations from related species, on the basis that genes have similar sequences in related species and so give positive hybridization signals, whereas the regions between genes have less similar sequences and so do not hybridize.

Zygote  The cell resulting from fusion of gametes.

α-helix  One of the commonest secondary structural conformations taken up by segments of polypeptides.

β-N-glycosidic bond  The linkage between the base and sugar of a nucleotide.

β-sheet  One of the commonest secondary structural conformations taken up by segments of polypeptides.

β-turn  A sequence of four amino acids, the second usually glycine, that causes a polypeptide to change direction.

γ-complex  A component of DNA polymerase III comprising subunit γ in association with δ, δ', χ, and ψ.

π–π interactions  The hydrophobic interactions that occur between adjacent base pairs in a double-stranded DNA molecule.
INDEX

Note: The index covers the main text but not the glossary or the end-of-chapter questions and reading lists. Acronyms have usually been preferred to their expansions as entries. Unmodified page references indicate text discussion that may be accompanied by relevant Figures or Tables: the suffix F or T distinguishes pages where only a Figure or Table is relevant.

A
A (aminoacyl) sites 310–11
A-T base pairs 9, 80, 122, 212
A-U base pairs 271, 309F
*ab initio* gene prediction 119, 122–3
Ac/Ds elements 214
acceptor sites (3′-splice sites) 280, 282–3
acetylation see histones
ACS (autonomously consensus sequences) 365
activation coefficients 324
activator proteins
catabolite activator protein 270–1, 336
RNA synthesis 266, 268, 270–3, 275
acylation of proteins 320T, 321
ADA complexes (yeast) 228–9
Ada enzyme/ada regulon (*E. coli*) 399
adaptors
linkers and 37
next-generation sequencing 93–4
tRNAs in protein synthesis 19
adenine
3-methlyadenine 400
decay and 14, 126, 129, 212, 278
structure 5F
adenosine 3′-phosphosulfate 144
adenylate cyclase 248F, 270
*Agilops squarrosa* 445
affinity chromatography 129–30, 299F, 304–5, 321
age-related macular degeneration 152
agriculture 57, 194, 457
*Agrobacterium tumefaciens* 49–50, 183, 184T
AIDS (acquired immune deficiency syndrome) see HIV/AIDS
*Aluropoda melanoleuca* (giant panda) 111–13
alarmones 311
aldolase genes 173–4, 438
alkaline phosphatase 36, 38
alkylating agents 396, 399
alleles
ancestral and derived 457
defective, in breast cancers 151
defined 58
multiple, in HLA system 59
relation to phenotype 63
segregation 64
allo lactose 269, 331
allopolyploidy 445
α-helices
death domain 172
in HTH motifs 249
in leucine zipper 254
in secondary protein structure 16, 17F
in zinc finger motifs 137, 250
alphoid DNA 160, 176
ALS (amyotrophic lateral sclerosis) 212
alternative exons 282
alternative promoters 267–8, 285, 331, 452
alternative site selection 282
alternative splicing pathways 169, 279, 282–5, 298, 452
Alu elements 211T, 212–13, 451
amino acids
additional amino acids 19
biosynthesis response to stress 311
chemical modification 320–1
codons corresponding 20F
histone modification 230–1, 232T
and homology searching 136–7, 139
the Miller-Urey experiment 430F
peptide analysis with mass spectrometry 297–8
in primary protein structure 16
and protein diversity 17–19
in the recognition helix 254
sequence and protein folding 315–18
structures and abbreviations 18
2-amino purine 395
amoebas
*Amoeba dubia* 166T, 167
*Paulinella* 196
*Pelomyxa* 195
ampicillin resistance gene 43–4, 48–9F
analytical protein arrays 300–1
anatomically modern humans 455
anchor cells in differentiation 347–8
Angelman syndrome 235
annealing, in DNA copying and PCR 30, 39–41
Antennapedia complex (ANT-C) 249T, 350–1
anti- and syn-conformations, DNA 9
antibiotic resistance
  deletion cassettes 140
  IS elements 213
  lateral gene transfer 193
  plasmids 184
antibodies
  labeling 147
  polyclonal antibodies 300, 301F
antigens, and immunoglobulin diversity 339–41
Antirrhinum 352
antisense transcripts 236, 260, 261F, 277
antiterminator proteins 342
AP (apurinic/apyrimidinic) sites 397, 399–401, 405
APC/C (anaphase-promoting complex/cyclosome) 381
apolipoprotein B, human 15
apoptosis
  cancer cells 324
  death domain 172
  eIF-4G cleavage 313
  following genome damage 383
  PANDA gene 261F
Arabidopsis thaliana
  centromeres 159–60
  DNA transfer 195
  gene catalog 171
  genome sequenced 162
  genome size and gene numbers 166T, 168, 171
  heat stress response 287
  mitochondrial genome 195, 197T, 199T
  plant development model 252, 352
  S/MARs in 223
  transcriptome 287, 288F
  whole-genome duplication 441–2
archaea
  DNA topoisomerases 363
  formerly regarded as extremophiles 181
  gene transfer with bacteria 193
  nucleoids 183
  proteasomes in 314
  RNA polymerases 264
  use of pyrrolysine 21
arginine, methylation and citrullination 230
Argonaute protein family 142F; 277–8
ARMS (amplification refractory mutation system) test 63
ARS (autonomously replicating sequences) 365–6
artificial gene synthesis 148
ascorbid acid (Vitamin C) 174
ascus and ascospores 338, 414
Ashbya gossypii 441
ataxias 393T, 406
attachment sites (att sites)
  DNA binding proteins 241–9
  sequence binding proteins 244
  site-specific recombination 420–1
attenuation 272, 273F
AU-AC introns 279
Australopithecus 450, 451F
autocatalysis 188
automation
  chain-termination sequencing 91
  optical mapping 78
autophagy 324–5
autopolyploidy 439–40, 445
autoradiography 36
auxotrophs 73
Avery, Oswald et al 3, 4F, 43
avidin 130, 299F
see also streptavidin
B
B chromosomes 158
BAC (bacterial artificial chromosomes) vectors 47T, 48, 78, 83, 104, 109–10, 113, 114F
Bacillus spp. 266
  B. anthracis 192
  B. subtilis 181, 204, 343–6
backtracking 265
bacteria
  DNA transfer methods 73
  gene location 120
  gene transfer with archaea 193
  genetic mapping 73–4
  homologous recombination in 415–17
  with non-circular genomes 183–6
  RNA synthesis and degradation 264, 268–72
  and the species concept 191
see also E. coli
bacteriophages
  capsid structures 203–5
  Cas endonuclease inactivation 187
  as cloning vectors 44–7, 48, 126, 301, 421
  genomes of 203–6
  Hershey–Chase experiment 3, 4F
  lambda (λ) 44–8, 99, 100F, 205, 301, 342–3, 405, 419–21
  as lytic or lysogenic 205–6, 342, 419–20
  M13 126, 127F, 148, 203, 204T, 301
  MS2 203–4
  P1 47T, 48, 421
  φX174 204
  prophages 205–6, 342–3, 405, 420
  replication strategies 205–6, 402
  rolling-circle replication 364
  SPO1 204
  T series (T1-T7) 38, 205
cancers
  breast cancer 150–1, 286, 324–5, 383, 406
  colon cancer 175
  DNA repair defects 406
  and the transcriptome 286–7
candidate genes 151–2
cap binding complexes 312–13
cap structures, mRNAs 130
capillary electrophoresis
  chain-termination sequencing 88–9, 91
  genetic profiling 177F
  SNP analysis 63
  STR typing 60F, 61
capping, 5'-end, RNA 14
caps, chromosome 378
capsid structures 203–5
carcinogens 394
Cas9 endonuclease
  DNA cleavage 144F
  phage inactivation 187
  as a programmable nuclease 143
cascade pathways, signal transduction 334–5
catabolite activator protein 270–1, 336
catalysis see enzymes; ribozymes
CD4 protein 207
CDEI-III elements 160
CDKs (cyclin-dependent kinases) 381
cDNAs (complementary DNAs)
  cDNA capture/selection 129
  cDNA library sequencing 129
  preparation 82
  RNA-seq 114, 124
  use of reverse transcriptases 31, 126
cell-cell signaling 301, 346–8
cell cycle
  protein degradation 314
  stalling 229
  synchronization of DNA replication 380
cell cycle checkpoints 380, 383
cell division, types of nuclear division 65
cell fates, primary and secondary 347–8
cell fusion, Sendai virus or polyethylene glycol 82
cell senescence 378–9
cell surface receptors, signal transmission 332–3
cell transformation and cancer 207–9, 224F
  by importing signaling compounds 330–2
  protein content 19
  RNA content 12–13
  somatic and sex cells 1
  cellular genes, in retroviruses 208
  cellular life
    biochemical systems preceding 429
    origins 431
  cellular response, to external signals 330–6
  CENP-A protein 160
  CENP-B protein 224
centiMorgan (cM) unit 68
centromeres
  in chromosome maps 75F, 81F
  DNA-protein interactions 159–61
  introduced 158
  in metaphase 79
  in mitosis 66, 67F
  regional and point 160
  stability 224, 234
  tandem repeats 176
  tandem repeats at 176
centromeric DNA 160
CEPH (Centre d'Études du Polymorphisme Humaine) 72
chain-termination sequencing
  applications 99
  choice of polymerase 89–90
  forward and reverse sequencing 90–1
  human genome 109
  method outlined 87–9
  strengths and limitations 91–2
  thermal cycle sequencing 90–1
chaperones 266, 316–18, 324–5
chaperonins 316–18
Chargaff, Erwin 7, 8F
charge, on amino acids 17, 19
Chase, Martha 3, 4F
checkpoint kinases 383
checkpoints, cell cycle 380, 383
chemical and physical mutagens 389, 394–7
chemical degradation sequencing 91
chemical modification of proteins 315, 320–1
chemical shifts 243
chemiluminescence 36, 95, 96F
chi (crossover hotspot initiator) forms/sites 413, 415–17
chimeras 141
chimpanzees 125F, 135, 443, 450–2, 454–5
ChiP-chip 248
ChiP-seq (chromatin immunoprecipitation sequencing) 247, 248F, 334
chip technology 62, 128, 152
Chlamydomonas reinhardtii 195–6, 197T, 199T
chloroplasts
  genome discovery and characteristics 195–8, 199F
  RNA polymerases 265
chromatids
  crossovers in meiosis 66, 68–9
  gene inactivation effects 145F
  of metaphase chromosomes 158
  unequal sister chromatid exchange 438, 439F
chromatin
  arrangement of protein complexes 156–7
  constitutive heterochromatin 158T, 159F
  DNA packaging into 220–2
euchromatin and heterochromatin 221
modification and DNA sequence 450
structural changes and differentiation 336–7
chromatin loops 225
chromatin-remodeling complexes 230, 233
chromatophores 196
chromosomes 157
chromids 185–6
chromodomains 418
chromosomes
caps 378
denaturing with formamide 79F
DNA attachment 222–3
dye references 84
homologous chromosomes 66–7, 68F, 70–1, 107F
meiosis 65–70, 74
micro- and macrochromosomes 158, 224
and the nuclear genome 155–61
prokaryotic ‘chromosomes’ 181–2
range of numbers 155, 451
size, compared with DNA 155–6
TADs (topologically associated domains) 225–7
translocations 224, 377F, 416
chromosome banding 79, 158, 159F, 161–2
chromosome conformation capture (3C) 224–5, 247
chromosome painting 221, 223
chromosome territories 223–5
chromosome theory 3
chromosome walking 105
chylomicrons 15
chymotrypsin, evolution 437
circular DNA
organellar genomes 196–7
prokaryotic genomes 181
replication problem 358, 361, 363–4, 373, 379F
cis-displacement, nucleosomes 233
citrate 322F, 431
citrullination 230
clamp loaders 370
class switching, immunoglobulins 340–1
climate change 113, 457
cloning contigs 104–5, 107, 109–10, 113–14
cloning technique 105–7
clone libraries
choice of vector 47–8
chromosome-specific libraries 83
from gene cloning 42–3
in genome sequencing 47T, 99, 100F, 112
as a mapping reagent 82–4, 107
cloned DNA, chain-termination sequencing 91
cloning vectors
adeno-associated viruses 144
for bacteria, yeasts, fungi and plants 49
bacteriophages 44–7, 301
for gene overexpression 144
insertion and replacement vectors 45–6
for longer fragments 47–8
plasmids as 41–4
shuttle vectors 49, 50F
Clostridium spp. 194
cloverleaf structures, RNA 122
co-dominance 64, 71
co-immunoprecipitation 305
coalescence times, haplogroups 456
Cockayne syndrome 406
coding RNA, mRNAs as 12
codon bias 121–2, 146
codons
average gene length 120
context-dependent reassignment 21
initiation codons 20, 119F, 311
introduced 19
termination codons 20–1, 119F, 121F, 276
cohesins 227, 375
cohesive ends (sticky ends) 32, 33F, 34, 37, 38F, 41
co-integrate formation 423
collagen
α2 type I collagen gene 446–7
post-translational hydroxylation 320T
structural protein 22
Collins, Francis 110
column chromatography 129, 294–8, 300
combinatorial-mode RNAs 287
combinatorial screening 105, 106F, 107
comorbidities 308
comparative epigenomics 450
comparative genomics
functional protein interactions 305–6
in gene prediction 123–4
complex A 280, 281F
complex B 280
complex E 280, 281F, 282
composite transposons 213
computer use
  gene function analysis 135–9
  in genome annotation 119–24
  in genome sequencing 55–6, 79, 96–7, 103, 262, 441
  in mass spectrometry 297, 324
  modeling early biology 430, 432, 441
  sequence databases 131
  X-ray diffraction studies 242
  see also software
centration gradients 347–9
concerted evolution 435
condensins 227
confocal microscopy 147
conjugation, in bacteria 73, 192
consensus sequences
  autonomous consensus sequence (ACS) 365
  chi sites 415
genome activity changes in 329, 341–52
plants 352
developmental processes, C. elegans 346–50
diagnosis
personalized medicine and 92
and the transcriptome 286
use of PCR 41
diauxic growth 271
Dicer ribonuclease 142F, 277–8
Dictyostelium discoideum 166T, 167
dideoxynucleotides 88–90
differentiation
B. subtilis sporulation 345–6
Caenorhabditis elegans 346–50
Drosophila melanogaster 348–50
gene activity changes 329, 336–41
protein degradation in 314
dihydrouracil 15F
dimerization, DNA-binding proteins 254
dinoflagellates 196
diols 130
dioxetane 36
diploid cells
chromosomes 66
distinguished from haploid 1
direct readout 252–3
directed mutagenesis 147–9
disease genes see genetic diseases; inherited diseases
disease modules, protein interaction maps 307, 308F
dispensive model, DNA replication 359, 361
displacement replication 363–4
distance matrices, phylogenetics 454
disulfide bonds/bridges 17, 315–16, 318, 339
DMS (dimethyl sulfate) 244–6, 247F
DNA amplification 438, 439F
DNA bending 253
DNA binding motifs 249–52, 352, 447
DNA binding proteins
proximal and distant binding sites 274
special features 249–52
structure and attachment sites 241–9
zinc fingers as 137
DNA chip technology 62, 128, 152
DNA cloning
ESTs as DNA clones 82
longer fragments 47–8
organisms other than E. coli 48–50
positional effect 226, 227F
process and applications 41–50
as a recombinant DNA technology 28, 41
DNA damage
bypassing during replication 405–6
double strand breaks 229, 368T, 383, 393, 403–5
single strand breaks 361, 383, 403–6
see also DNA repair; nicks
DNA (deoxyribonucleic acid)
A-form 10, 11F, 253
anti- and syn-conformations 9
available manipulations 27F
B-form 9, 10, 11F, 252–3
biosynthesis 7F
cellular genomes as 1
chemical differences from RNA 12F
chromosome packaging 155–6
conformational variations 253
discovery 2
end-shortening 375, 376F, 379, 425
genes as 4
packaging 221–2
probable first appearance 432–3
survival of ancient DNA 110–11
T-DNAs 10, 49, 50F
Z-form 10, 11F, 253
see also cDNAs; double helix; recombinant DNA;
repetitive DNA sequences
DNA-dependent DNA polymerases 367, 425
DNA-dependent RNA polymerases 12
DNA fragments, membrane protein capture 193
DNA glycosylases 399–401
DNA gyrase 182, 363
DNA labeling, with E. coli polymerase I 30
DNA ligases
activity 29F, 37
bacterial promoter 268T
3C method 224, 225F
damage repair 401–5
DNA cloning 41
DNA ligase IV 405
function 37–8
homologous recombination 412
joining Okazaki fragments 371–2
nick repair 399
in SOLI D sequencing 96
DNA markers
for gene mapping 59–61
linkage analysis 68
DNA methylation 230, 234–6, 449–50
DNA methyltransferases 234, 236, 337, 399
DNA polymerases
activity and role 29F, 367–70
chain-termination sequencing 87–9
direction of synthesis 6, 29
DNA-dependent 367, 425
exonuclease activity 30–1, 367, 371, 391
Kornberg/Klenow polymerase (E. coli DNA polymerase I) 30–1, 90
limitations 369–70
novel nucleotides and 433
nucleotide selection by 391
RNA-dependent 31, 367, 425
DNA polymerase 31T, 39, 63, 90–1, 126
DNA polymerase α 369, 370F, 371T, 372, 377
DNA polymerase β 368T, 400–1
DNA polymerase δ 368T, 369, 371T, 372, 401–2
DNA polymerase ε 368–9, 371, 401
DNA polymerase γ 368T, 369
DNA polymerase η 368T, 406
DNA polymerase I 5′→3′ exonuclease activity 82F, 371, 372F, 400F
DNA polymerase II 368, 405
DNA polymerase III 368–71, 372F, 391, 403, 405
DNA polymerase IV 405
DNA polymerase V 405
DNA polymerases ζ, θ, λ, μ, ν, κ 368T
DNA-protein interactions in centromeres and telomeres 159–61
DNA repair defects and human disease 406
direct repair systems 398–9
excision repair systems 399–402
four repair categories examined 398–406
halting replication 383
HMG box domain 251–2
homologous recombination in 412, 415, 418–19
mismatch repair systems 402–3
prereplicative and postreplicative enzymes 389
strand break repair systems 403–5
DNA replication base pairing and 9
bypassing DNA damage 405–6
daughter strand identification 402, 403
dispersive, conservative and semiconservative models 359–61
error correction 369
initiation phase 364–7
link to evolution 357
polymerase limitations 369–70
regulation in eukaryotes 380–3
termination 372–9
topology of 357–64
DNA sequencing see chain-termination sequencing, genome sequencing, next-generation sequencing
DNA synthesis error rates 391
template-dependent DNA synthesis 8–9
DNA topoisomerases DNA gyrase and reverse gyrase 182, 363
DNA replication role 358, 361–3, 370, 374–5
and the topological problem 361
Type I 361–2
Type II 361–2, 374–5
Type IV 374
DNA transfer, eukaryotic 195
DNA transposons 163, 166, 211T, 213–15
DnaA protein 365
DnaB helicase 268F, 365, 373–4
DNase I in footprinting hypersensitive sites 245
hypersensitive sites 232–3
dNTPs (deoxynucleotide triphosphates) 88
domain duplications 445–7
domain model, E. coli nucleoid 183
domain shuffling 446
dominant phenotypes, distinguished from recessive 64
donor sites (5′-splice sites) 279–80, 283F
double helix, DNA complicating replication 358–9
importance of discovery 4, 6, 8
leading and lagging strands 369
major and minor grooves 11
melting 62F, 365–6, 402
structural flexibility 9–11
double restriction, genetic mapping 76
double-strand breaks DNA topoisomerases 362
double-strand break model, homologous recombination 414–16
repairs 393, 398, 403–5, 415
Down syndrome 439
DPE (downstream promoter element) 267
Drosophila melanogaster (fruit fly) alternative splicing 284–5
bicoid gene/protein 252, 348–50
copia retroelement 211
Dicer ribonuclease 278
differentiation 348–50
DNA end-shortening 379
embryonic development 146, 348
gene catalog 171
genetic mapping in 58, 68
genome sequencing 92, 284–5
half pint gene 285
insulator sequences 226
operons 189
origin of GO system 139
P element transposon 142, 215
partial linkage example 67
piRNAs in 259
S/MARs in 223
sex determination 282–3
TADs 225
tudor domain 138
drug resistance 324–5
D17S1321/1325 STRs 151
dsRBD (double-stranded RNA binding domain) 252
dsRNA (double stranded RNA) 142, 143, 204T, 306T
DUE (DNA unwinding elements) 364–5
DUF1220 domain 452
dye quenching 40, 62
dyes see fluorescence; staining
dystrophin gene, human 265, 267, 268F

E

E. coli

core genome and pan-genome 192
DNA polymerase I (Kornberg/Klenow polymerases)

30–1, 90
DNA synthesis errors 391
flagellum 323–4
genome sequencing 74
genome supercoiling 182
genome transfer by conjugation 73
homologous recombination 415–17
initiation of replication 364
K12 strain 184, 185T, 186–7, 189, 191
lateral gene transfer 192
metabolic flux analysis 322–3
nucleoid 181–2
number of protein-coding genes 135
O157:H7 strain 191
plasmids as cloning vectors 43–4, 48
promoter sequences 266, 268T
protein-protein interactions 305–6
replication termination 373–4
repetitive DNA 213
ribosome numbers 308
sample genome segment 120F, 122F
E (exit) site 310
EcoP151 endonuclease 130, 131F
EcoRI restriction endonuclease 32, 59, 75, 76F
EGF (epidermal growth factor) gene/receptor 333F, 447
eIF initiation factors 312–14
electroendosmosis (EEO) 34
electron density maps, X-ray diffraction 242
electron microscopy
cell nucleus 220–3
chromatin 156, 157F
immunocytochemistry 147
organelles 195–6
ribosomes 309–10
electrophoresis, principles 34
see also capillary electrophoresis; gel electrophoresis
electrospray ionization 298
Ellis, Emory 205
EMS (ethylmethane sulfonate) 396
end-modification
enzyme activity 29F, 38
RNA 4
end-shortening problem 375, 376F, 379, 425
C2’- and C3’-endo configurations 10
endogenous promoters 277F
endogenous retroviruses (ERVs) 211–12, 442
endogenous transposons 210–11
endonucleases
Cas9 143, 144F, 187
FEN1 (flap endonuclease) 372, 401, 418
initiation of replicative transposition 422
programmable nucleases 143, 187
restriction endonucleases 32–4, 100, 130, 224, 225F, 366F, 367
RNAse E, H and P 276, 425, 431
endoplasmic reticulum 260, 308, 318, 325F, 336
endospores, Bacillus subtilis 344
endosymbiosis 190, 195–6, 449
enhancers
binding sites within 274
in retroelements 424
splicing enhancers 275, 281–4, 330–1, 334
enol tautomers 391, 394–5
Ensembl genome browser 131
environmental agents causing cell damage 394T
environmental stress, transcriptome effects 285, 287
enzymes
for DNA manipulation 28–38
as proteins 22
replacing ribozymes 432
enzyme activity, GO nomenclature 170
Enzyme Commission (EC) numbers 138
EPAS1 transcription factor 453
epigenome evolution 449–50
episome transfer 73
epitopes 301
error correction, in DNA replication 369
error rate
DNA polymerases 391
finished genomes 107
RNA polymerases 265, 276
ES (embryonic stem) cells 141
Escherichia coli see E. coli
ESTs (expressed sequence tags) 82, 109
ethidium bromide 35, 40F, 125F, 396
euchromatin
in cell division 382
distinguished from heterochromatin 221
human genome 109–10
organization 222–3
replication priority 382
eukaryotes
DNA polymerases 368
DNA topoisomerases 362–3
evidence for first appearance 434
homologous recombination 417–18
last common ancestor 449
lateral gene transfer 444–5
proteome remodeling 312–13
regulation of DNA replication 380–3
replication termination 374–5
RNA end modification 14
RNA splicing pathways 279–80
RNA synthesis and degradation 272–5
eukaryotic genomes
DNA transposons 214
nuclear genome organization 161–7
organelar 195–9
sequencing projects 109–15
shotgun sequencing 102–4
see also nuclear genomes
eukaryotic virus genomes 206–7
Eulerian pathways 103
evolution
concerted evolution 435
evolutionary relics 174–6
homeotic selector genes 351
homologous genes and 135–6
and homologous sequences 123
multigene families 174, 435
whole-genome duplication 439–42
see also genome evolution
excavates 448–9
excinuclease 401
excision repairs
base excision repairs 399–401
nucleotide excision repairs 401–2
short-patch and long-patch 399–401
excisionase 421
exome sequencing 108
exons
in the human genome 164
and pre-RNA splicing 13
whether coding for domains 446
exon-intron boundaries 121–2, 124, 126–7, 129, 280, 298
see also intron-exon
exon skipping 281–3
exon theory of genes 448
exon trapping 127
exonic splicing enhancers (ESE) 281
exonic splicing silencers (ESS) 281
exonucleases 32, 400
exonuclease activity
direction 30, 367
DNA polymerases 30–1, 90, 367, 371, 391
proofreading 368
exosome 276, 278
expression pattern and gene function 146–9
expression proteomics see protein profiling
extinct proteomes see protein profiling
extinct species, genome sequences 41, 110–11, 452–3
extrachromosomal genes 195
extremophiles, archaea formerly seen as 181

F
F plasmid, *E. coli* 48
facultative heterochromatin 222
family studies
gene mapping 59, 70–1
pedigree analysis 71–2, 150–1
Fanconi anemia 406
FEN1 (flap endonuclease) 372, 401, 418
ferritin 22, 313
fetal hemoglobin 173
fiber-FISH 79
filamentous capsid structures 203–4, 206, 301, 302F
finished genomes, error rates 107
fish, gene duplication 437–8, 441
FISH (fluorescent in situ hybridization) 79
5’ or 5’-P terminus 6
5’-RNA end capping 14
flagella, *E. coli* 323–4
flow cytometry 83, 84F
fluorescence
in chain termination sequencing 88–9
dye quenching 40, 62
FISH 79
in flow cytometry 83, 84F
FRAP 221
FRET 40
GFP 147F, 220, 222, 224
labels, in immunocytochemistry 147
labels, polyclonal antibodies 301
labels, reversible terminator sequencing 95
labels, via Kornberg polymerase 30
labels, with alkaline phosphatase 36
markers in next-generation sequencing 97
markers in PCR 40–1, 61
markers in protein profiling 299, 303
see also dye quenching; FRET
fluorescence microscopy 62, 77, 220
immunofluorescence microscopy 222
flush ends (blunt ends) 32, 33F, 37, 38F, 46F
foldback RNAs 278
folding pathways 316, 317F
footprinting 245, 246F
formaldehyde 224, 247
formamide 79–80
forward and reverse genetics 150
forward and reverse sequencing 90–1
fosmids 47T, 48, 110
454 sequencing 96
fourth-generation sequencing 97–8
FOX2 transcription factors 451–2
fragile sites 393
fragment ions 298–300
frameshift mutations 391–2
Franklin, Rosalind 6–7
FRAP (fluorescence recovery after photobleaching) 221
FRET (Förster resonance energy transfer) 40
fruit flies see Drosophila
FT-ICR (Fourier transform ion cyclotron resonance) 298
functional domains, TADs as 225
functional RNAs 12

G
G-band 158T, 159F, 161–2
G-C base pairs 9, 80, 122
G-proteins 334
G-protein receptors 172
gag, pol and env genes 207, 211–12
Gallo, Robert 207

gametes
autopolyploidyation 439, 440F
direct examination 69
in gene conversion 414
introduced 1
knockout mice 141–2
meiosis 66–7, 68F, 69–70, 71F
S. cerevisiae 338
γ complex 370
gap genes 349–50
GAPs (GTPase-activating proteins) 334
GAS (interferon γ-stimulated gene response) elements 334
GC repeats 10
GC-rich regions 80, 84, 158T, 161–2
GCN5 protein 228–9
GEFs (guanine nucleotide exchange factors) 334, 335F
gel electrophoresis
agarose gel electrophoresis 34–5, 40, 59–61, 75, 99, 125
denaturing gel electrophoresis 246F, 247
heteroduplex analysis 126
limitations 61
PCR monitoring 40
polyacrylamide gel electrophoresis (PAGE) 34, 87, 88F, 91, 294–7
two-dimensional 294, 295F, 297, 299
variety of sncRNAs 259
see also capillary electrophoresis
gel retardation 244, 247
gel stretching 77, 78F
geminin 381, 383
GenBank database 131
gene catalogs 169–72, 191, 196–8
gene cloning see DNA cloning
gene clusters
homeotic gene clusters 351, 437

operon-like 189
for 5S rRNA 173
see also globin
gene conversion
coordinated evolution 435
Holliday model of recombination and 414–15
S. cerevisiae mating type determination 338–9, 412, 417

gene densities 158, 162, 165–6, 167T, 187, 190
gene deserts 162
gene duplication
domain duplications 445–7
human phenotype 452
multigene families 173, 436–8
possible mechanisms 438
segmental duplications 442, 445, 452
whole-genome duplication 439–42
gene expression
CAGE 130, 131F
cell specific and time-dependent 128
expression pattern and gene function 146–9
gene overexpression 139, 144–6
as genome expression 219
globin family 173
nucleosome remodeling and 231–3
small regulatory RNAs 15–16, 259
tissue specific 125, 234
gene families see multigene
gene flow and the species concept 191
gene function
annotation terminology 138–8
assignment by expression pattern and protein product 146–9
assignment by inactivation 139–44
assignment by overexpression 139, 144–6
computer analysis 135–9
conventional genetic analysis 149–52
directed mutagenesis 147–9
homology analysis 135–7
gene inactivation
investigation of gene function 139–44
knockout mice 141–2
phenotypic effects 145–6
and protein-protein interactions 306–7
using homologous recombination 141–2
gene overexpression 139, 144–6
gene prediction 119, 122–4
see also genome annotation
gene regulation, lactose operons 189
gene silencing
by DNA methylation 234
histone deacetylation in 229
lysine methylation in 230
PcG proteins 337
silent cassettes 338
X inactivation 235
gene space 113–15, 124
gene transfer 192–5, 444–5
general recombination see homologous recombination
general transcription factor (GTF) 229, 273–4
see also TFIIA etc.
genetic analysis, forward and reverse genetics 150–1
intron theory 448
maternal-effect genes 348–50
numbers of, and genomic complexity 434–8
numbers of, in bacteriophages 204–5
numbers of, in prokaryotes 189–92
overall numbers and functions 167–76
partial linkage 65
protein-coding, numbers 135, 162, 164, 168
protein-coding regions as ORFs 119
strain-specific 191
truncated genes and gene fragments 123F, 175
whether protein or DNA 3–4
genetic association studies 458F
incremented origin of biological systems 429–34
origin of complexity 434–50
primate evolution 450–3
genome expression
DNA binding proteins in 241
DNA modification and 234–6
gene expression as 219
miRNA regulation 278
nucleosome modifications 228–33
translation and transcription in 2
genome browsers 131
genome damage/repair see DNA damage; DNA repair
genome evolution
duplication events 435–44
epigenome evolution 449–50
gene rearrangements 445–9
from germ cell mutations 389
origin of biological systems 429–34
genome expression
DNA binding proteins in 241
DNA modification and 234–6
gene expression as 219
miRNA regulation 278
nucleosome modifications 228–33
translation and transcription in 2
genome maps see genetic mapping
genome rearrangements 210, 339–41
genome reduction 196
genome replication see DNA replication
genome resequencing 92, 97, 101, 113, 129
genome sequencing
alteration in gene conversion 338
categories of method 87
chemical degradation sequencing 91
E. coli 74–5, 186–8
error rates 107
eukaryotic genomes 102–4, 109–15
extinct species 41, 110–11, 452–3
genetic maps in 55–7, 74
of giant panda 111–13
H. influenzae 91, 99–101
Mycoplasma genitalium 100
of plants 57, 109, 113–15, 124, 159
of prokaryotes 100–1, 181
real-time sequencing 97
S. cerevisiae 74–5
sequence assembly 55, 56F, 98–110
shotgun method 55–6, 99–104, 109–12
whether necessary 107–9
see also chain-termination; next-generation

genome sizes
bacterial 91–2, 99, 135
bacteriophages 44, 204
crop plants 57, 113
eukaryotes 165–6T, 167
eukaryotic organelles 195–8
eukaryotic viruses 206–7
human 1, 92, 102
prokaryotic 189–92
and the species concept in prokaryotes 190–2
yeasts 146, 164–5

genomes
bacteriophages 203–206
chloroplast 195–198
core and accessory prokaryote 191–2
eukaryotic nuclear 1, 155–177
extinct species 41
metagenomes 194
mitochondrial 1, 195–198
origin of DNA genomes 432–3
population diversity 453–8
prokaryotic, genetic features 186–95
prokaryotic, physical features 181–6
response to external signals 330–6
viruses 203–209
whole-genome duplication 439–42
genomewide RNA mapping 127–31
genomewide scans 247
genomic imprinting 235, 261, 444
germ cell mutations, leading to evolution 389
GFP (green fluorescent protein) 147F, 220, 222, 224
giant panda (Ailuropoda melanoleuca) genome 111–13
Giemsa stain 158T, 161
glass slide method 93–4

globin gene superfamily
α-globin gene cluster 174
β-globin gene cluster 161, 174, 232, 437
δ-globin gene 174
evolution in humans/mammals 436, 437F
gene duplication 136, 173–4, 436
as multigene example 173–4
myoglobin and β-globin as paralogs 136

Globodera pallida 445

glutathione peroxidase 21
glycans 320–1
β-N-glycosidic bonds 5, 9–10, 397, 399
O-glycosidic bonds 138, 139F
glycosylation 320–1
GO (gene ontology) nomenclature 139, 169
Gossypium hirsutum and G. barbadense. 83
greedy algorithms 102
group II introns 449
GRP78 chaperone 324–5
GU-AC introns 279, 448–9

guanine
7-methyl- 14, 15F, 130, 400
methylation 244
structure 5F
guide RNAs 143–4, 187
GWAS (genomewide association studies) 151–2

H
H-NS (histone-like nucleoid structuring) protein 183

Haemophilus influenzae

genome sequencing 91–2, 99
HindIII restriction endonuclease 22T, 32, 34

hairpins see stem-loop structures

half-life, RNAs 269F, 275

haploid cells, distinguished from diploid 1

haplotypes/haplogroups 456

HAT (hypoxanthine, aminopterin, and thymidine) medium 83

HATs (histone acetyltransferases) 228–9, 233
HDACs (histone deacetylases) 229–30, 234, 235F, 338, 382

head-and-tail capsid structures 203F, 204

heat shock
bacterial response 266
eukaryotic response 274, 284, 287–8, 312, 317
mutagenic effect of heat 397F

Helianthus annuus 57

helicases
DNA helicase II 401, 403
in DNA replication 365–6, 370, 372–4, 380–1, 401
nanopore sequencing 98
RecBCD pathway/complex 415–18
RecFOR pathway 417
RecQ helicase 406
RNA helicase B 276

Helicobacter pylori 191, 306

hemoglobin, fetal and maternal 173
hemophilia 150T, 213
hereditary diseases see inherited diseases
Hershey, Alfred 3, 4F

HERVs (human ERVs) 211–12, 442

heterochromatin
Barr bodies as 235
consortial 110, 221–2
distinguished from euchromatin 221
facultative 222
PcG proteins and 337
replication priority 382
heteroduplex analysis 126–7
heteroduplex formation
in homologous recombination 412–17
in site-specific recombination 420
heterozygosity
  distinguished from homozygosity 63–4
  double and triple heterozygotes 70
  in genome sequences 107
hexaploid genomes 57
hibernation promotion factors 311
hierarchical classification, annotation terminology 138–9
hierarchical clustering 263
hierarchical shotgun approach 104–7
high mobility group (HMG) box domain 251
high-throughput functional analysis 141
high-throughput mapping 78
high-throughput sequencing 129
  see also next-generation
Hinf I restriction endonuclease 32, 33T, 34
histidine biosynthesis 305–6
histones, introduced 156
histone acetylation, and genome expression 228–9
histone acetyltransferases (HATs) 228–9, 233
histone code 230–1
histone deacetylases (HDACs) 229–30, 234, 235F, 338, 382
histone deacetylation
  and firing of replication origins 382
  and genome repression 229–30
  and X inactivation 236
histone H2A 230, 236, 337
histone H2B 183, 230, 233
histone H3
  centromeric replacements 160
  covalent modification 230–1
  lysine methylation 230–1, 236, 337
  in the nucleosome 156–7
histone modification
  in epigenome evolution 449–50
  other than acetylation 230–1, 232T
HIV/AIDS (human immunodeficiency virus/acquired immune deficiency syndrome)
  HIV-1 and HIV-2 455
  origins 454–5
  polyprotein use 319
  retrovirus causation 207, 423, 454
HLA (human leukocyte antigen) system 58–9
hnRNAs (heterogeneous nuclear RNA) 14
hnRNPs (heterogenous nuclear ribonucleoproteins) 282
Holliday model/structures (Robin Holliday) 412–20
holocentric chromosomes 158
homeodomain proteins 249T, 250, 252, 350–2, 437
homeotic mutants 350
homeotic selector genes 337, 350–2, 437
Homo genus 110, 450, 451F, 452, 453F
Homo neanderthalensis 110, 452–453
homogeneous domains 162
homologous chromosomes 66–7, 68F, 70–1, 107F
homologous genes
  with differing functions 137
  and evolution 135
homologous recombination
  distinguished from site-specific recombination 411–12
  in DNA repair 412, 415, 418–19
  double-strand break model 414–17
  eukaryotes 417–18
  following site-directed mutagenesis 148–9
  gene inactivation by 140–2
  Holliday model/structures 412–20
  LTR origins 211
  Meselson–Radding modification 412, 414–15
  models and pathways 412–19
  RecBCD pathway 415–16
  reporter genes 147F
YIp5 vector 49
homologous sequences, and evolution 123, 441
homology searching
  and gene function 136–7, 139
  in gene prediction 123
  protein–protein interactions 305F
homopolymer tailing 37, 38F
  poly(A) tails 14, 126, 129, 212, 278
homozygosity
  distinguished from heterozygosity 63–4
  double and triple homozygotes 70–1
Hoogsteen base pairs 80
horizontal resolution of Holliday structures 413
hormone response elements 331–2
housekeeping genes 234
housekeeping proteins 19
Hox gene clusters 351, 437–8, 441F
HPLC (high-performance liquid chromatography) 296, 298
HPRT (hypoxanthine phosphoribosyltransferase) 83
hsp70 genes/Hsp70 chaperones 226, 233, 316–18
HTH (helix–turn–helix) binding motif/HTH family 249–50, 254, 269, 350
HU family proteins 183
hubs, protein interaction maps 306–7
humans
  anatomically modern 455
  embryonic development models 351
  globin gene evolution 436
  number of cells and cell types 329, 341
  number of different immunoglobulins 339
  number of origins of replication 364
  out-of-Africa migration 455–7
  protein interaction maps 306–7
  ribosome numbers 308
telomere sequences 377T
human endogenous retroviruses (HERVs) 211–12, 442
human evolution 450–3
human genetic studies
  blood typing in 58
  pedigree analysis 71–2, 150–1
  radiation hybrid mapping 83
human genome
  browser illustration 131F
centromeric DNA 160
chimpanzee genome and 451–2
chromosome lengths 156
DNA transposons 214
duplication events 442–4
exons and interspersed repeats 164, 212
gene distribution pattern 162
mitochondrial genome 196–7, 199T
number of base-pairs 1
number of baseless sites 397
number of IncRNAs 260
number of protein-coding and noncoding genes 168–9
  number of SNPs 107, 453
S/MARs in 223
sample segment of 200kb 163–4, 212
transposable elements 211T
Human Genome Project (HGP) 48, 92, 109–10
Human Genome Sequencing Consortium 110
human karyogram 158, 159F, 161, 165
human lifetime, number of mitosis 66
human metabolome 323
human microbiome 194–5
humidity and DNA structure 10
Huntington’s disease 150T, 393
hybrid dysgenesis 215
hybridization
  introduced 36
  northern hybridization 125
  oligonucleotide hybridization analysis 61–2, 96F
  in PCR 40
  radiation hybrids 82
  solution hybridization technique 62
  Southern hybridization 35, 36F, 59, 60F, 100, 107
hybridization probes/tests 36, 79, 100, 105, 125
hydrogen bonding
  acceptors and donors, DNA 253
  affinity chromatography 304
  base pairing 8, 80
dehydrins 287
  secondary protein structure 16
tertiary protein structure 17, 315
hydrophilicity
  dehyadrins 287
RNA 432
hydropobicity
  of chaperones 317
molten globules 316
  in protein-DNA interactions 253–4
  in protein structure 17
  in protein transport 198
  in RPLC 295
hypoxanthine 15F, 83, 395, 399, 400T

I
ICAT (isotope-coded affinity tags) 299, 300F
Ice Ages 110, 457
ICF (immunodeficiency, centromere instability, and facial anomalies) 234
icosahedral capsid structures 203, 204T, 206
Igf2 gene 235
IHF (integration host factor)/IHF motifs 249T, 420–1
IIA^Glc protein 270
Illumina sequencing 95–6, 111–13, 129
immediate-early genes 342–3
immobilization methods, next-generation sequencing 93–4
immobilized metal ion affinity chromatography 321
immune system, in prokaryotes 187
immunocytchemistry 146–7
immunoelectron microscopy 310
immunofluorescence microscopy 222
immunoglobulin fold 334
immunoglobulins
  classes and class switching 340–1
diversity through genome rearrangement 339–41, 412
immunoprecipitation 247, 305, 334
imprint control elements 235
imprinting, genomic 235, 261, 444
in vitro packaging 45–6, 48
in vitro (site-directed) mutagenesis 147–9
incomplete dominance 64
indels 451, 452F
independent-mode RNAs 287
inducers of operons 269
induction of lytic infection 206
infectious proteins 209
informational problem 19
infrared spectroscopy 322
inheritance, principles of 63–5
inherited diseases
  DNA repair defects 406
  forward genetics and 150–1
  interspersed elements and 213
  monogenic 150
  polygenic traits 151–2
  see also genetic diseases
initiation codons 20, 119F, 311
initiation factors/complexes 311–13
initiation phase, DNA replication 364–7
initiation regions/sites 366–7, 382
initiator (Inr) sequence 267, 273
insertion cloning vectors 45–6
insertion mutations 389, 391–4, 396
insertion sequences (IS) 187, 213–14
inselional inactivation 43
insulator sequences 226–7, 274
insulin, activation by proteolytic cleavage 318
integrases 342, 420–1, 425
integrative eukaryotic viruses 207
interactomes 306
intercalating agents 35, 396
interferons 143, 333–4
intergenic regions/DNA 120, 123, 125, 163, 164F, 167, 186
interleukin 1β gene 331
interleukins 6 and 10 333–4
internal primers 91, 93, 100F
International Human Genome Sequencing Consortium 110
interphase
DNA replication during 380
interphase chromosome territories 223F
30 nm fiber 157
interspersed repeat element PCR (IRE-PCR) 107
interspersed repeat sequences
distributed by transposition 210
fruit fly genome 166
as one form of repetitive DNA 176–7
types of 163–7
in various eukaryotic genomes 165–7
see also LINEs;LTRs;SINEs; transposons
intramolecular base pairing 12, 122, 129
intrinsic terminator sequences 271, 272F
intron-exon boundaries 121, 247
see also exon-intron
intron retention 282
intronic splicing enhancers (ISE) 281
intronic splicing silencers (ISS) 281
introns
absent from retrogenes 438
group II introns 449
GU-AC 448
GU-AC and AU-AC 279
and ORF scanning 120–1
origin 448
and pre-RNA splicing 13, 279–80
scarcity in prokaryotes 188
various eukaryotic genomes 165–7, 197
introns early and introns late hypotheses 448
ion-exchange chromatography 295
ion torrent method 96–7
IPTG (isopropyl thiogalactoside) 44
IRE-PCR (interspersed repeat element PCR) 107
IRES (internal ribosome entry sites) 312–13
iron-response elements 313
ISFET (ion-sensitive field effect transistor) 96
isobaric labeling 299, 300F
isochore model 161–2
isoelectric focusing/isoelectric point 294–6
isoforms, RNA 282, 285–6
isopycnic centrifugation 161, 162F
isotope studies
Hershey-Chase experiment 3, 4F
ICAT 299–300
Meselson–Stahl experiment 359
NMR 243, 275
ISRE (interferon-stimulated response element) 334
ITAFS (IRES trans-acting factors) 313
J
Jacob, François 268
JAK/STAT pathway 333–4
Janus kinases (JAK) 333
K
K-islands 191
k-mers 102–3
kanamycin resistance 50F, 141F, 213, 421
karyograms
fruit fly 165–7
human 158, 159F, 161, 165
KH (K homology) domain 252
kinase or kinase-associated receptors 332
kinetochores 160
Klebsiella pneumoniae 266
Klenow polymerases 31, 90
Kluyveromyces waltii 441
knockdown mice 143
knockout mice 49, 142–3
Koch, Robert 190
Kornberg, Arthur 30, 368
Kornberg polymerase see DNA polymerase I
Ku proteins 404–5
L
Lac selection 44, 48
Lactobacillus helveticus H10 189
lactoferrin 330–1
lactose operon 268, 269F, 331
lactose permease 188, 305–6
lactose repressor 250, 269–70
lacZ gene 43, 47
lacZ’ gene 43–4, 50F
lagging strand, DNA replication 369–72, 374–5, 376F, 391, 400F, 401, 418
lambda (λ) see bacteriophages
landraces 457
lariat structures 279–80
latent periods 205, 208, 455
lateral gene transfer 192–4, 444–5
leading strand, DNA replication 369–70, 374, 376F, 391
LECA (last eukaryotic common ancestor) 449
lectins 321
leghemoglobin 448
leucine zippers 254
leukemia viruses 208
leukemias
chronic myeloid leukemia 224
and the transcriptome 286–7
life, origins of 429–33
ligases see DNA ligases
light-activated oligonucleotide synthesis 128
LIN-32 signaling compound 347
LINEs (long interspersed nuclear elements) 163, 166, 211T, 212–13
LINE-1 elements 211–12, 447
LINE-1 transposition 212–13
linear genomes 183–6, 196
linkage analysis
different organisms 68–71
limitations 74–5
STS mapping and 81–2, 84
linkage disequilibrium 151
linkage groups 68
linker DNA, between nucleosomes 157, 231, 369
linker histones 157, 221
linkers and adaptors 37, 38F
linkers region, ICAT 299F
linking number, DNA topoisomerases 361–2
lncRNAs (long noncoding RNAs) 13, 254, 257, 260–2, 265T
lod scores 72
long-patch repairs 399–403
long products, PCR 39
Lou Gehrig’s disease 212
low-copy repeats (segmental duplications) 442–4, 450, 452
LTR (long terminal repeat) elements 163, 165–7, 210–13, 424–5
lyases 138, 399–400
lymphocytes, B and T 339–40
lysine acetylation see histone acetylation
lysine carbamoylation 320T, 321
lysine methylation
histone H3, lysines 4 and 9 230–1, 236
histone H3, lysines 9 and 27 337
lysine ubiquitination 230, 314, 337
lysogenic infection cycles 44, 45F, 205–6, 342–3, 419–20
lytic infection cycles 44, 45–6F, 47, 205–6, 342–3, 419–21
M
M13 bacteriophage 126, 127F, 148, 203, 204T, 301
m/z (mass-to-charge) ratio 297–9, 300F, 321
MacLeod, Colin 3
MADS box binding domain 352
magnesium ions 77–8, 431
magnetic beads 96, 109, 130, 373
magnetic tweezers 373, 374F
maintenance methylation 234
maize (Zea mays)
genetic code anomaly 21F
genome 166–7, 197T
mobile genetic elements 211, 214–15
major groove, DNA double helix 11, 249–54
MALDI-TOF (matrix-assisted laser desorption ionization time-of-flight) 297–8
MAP kinase (mitogen-activated protein) 334–5, 338, 348
map units, genetic mapping 68
marker-assisted/associated selection 57, 61, 69, 458
marker genes, plant cloning 421
mass analyzers 298–300
mass spectrometry 293–4, 296–300, 305, 322
tandem mass spectrometry 298–300
mass-to-charge ratio (m/z) 297–9, 300F, 321
massively parallel arrays 93
see also next-generation sequencing
maternal-effect genes 348–51
mating type switching 338–9, 412
McCarty, Maclyn 3
McClintock, Barbara 214–15
MeCPs (methyl-CpG-binding proteins) 234, 235F
mediator proteins
replication mediators 366, 370
role 274
two-hybrid system 302–4
meiosis
chromosomes in 65–70, 74
distinguished from mitosis 65–6
double-strand break frequency 415
gene inactivation effects 145T
melittin and promelittin 318
melting, of the double helix 62F, 365–6, 402
membrane proteins, DNA capture 193
membrane transport, signaling compounds 330
Mendel, Gregor 27, 55, 58, 63–4, 65F
Mendelian inheritance 150–1
Mendel’s laws 64–5F
Meselson, Matthew 359, 412
Meselson-Radding model/modification 412, 414–15
Meselson-Stahl experiment 359, 360F, 361
messenger RNA see mRNAs
metabolic engineering 323
metabolic flux 322–3
metabolic labeling 300
metabolomes 322–3
metagenomes and metagenomics 194
metal ion uptake 331
metallic beads 93–4
metaphase chromosomes
chromosome features 157, 158F
DNA packaging 221
and FISH 79
metaphase (meiosis) 67F
metaphase (mitosis) 66F
metastases 262, 286
Methanosarcina spp. 194
3-methyladenine 400
7-methylguanosine 14, 15F, 130, 400
methaspartate cycle 193–4
methylation
of arginine and lysine 138, 230
of bases 80, 244, 400
maintenance and de novo methylation 234
patterns and species relatedness 450
of RNA 15
*Methylobacterium* spp. 194
MGMT (O6-methylguanine–DNA methyltransferase) 399
microarray analysis
chip-on-chip method 248
protein profiling 300–1
SNP association studies 458
tiling arrays 128–9, 258, 262
transcriptome 258, 262–3, 284, 287
microbiomes
  giant panda 113
  human 194–5
microchromosomes 158
*Micrococcus* spp. 21
microfluidic devices 78–80
microfossils 429, 430F
microsatellite DNA 60–1, 71, 72F, 164F, 176–7
  see also STRs
microsomes see ribosomes
microtiter techniques 62, 106F
Miescher, Friedrich 2–3
Miller, Stanley 430F
minigenes 127
minisatellite DNA 60, 176–7, 375
minor groove, DNA double helix 11, 252–3, 273, 400
miRNAs (micro RNAs) 15–16, 175, 259, 260F, 264, 278
mismatch mutations 389
mismatch repair system 391, 398, 402–3, 406, 418
MITEs (miniature inverted repeat transposable elements) 214
mitochondrial genome
  discovery and characteristics 195–8
  human migration and 456
  nonstandard genetic code 20–1
  replication 363, 369
  size and gene content for several species 1, 197, 199T
mitochondrial RNA polymerases 265
mitosis
  in the cell cycle 380
  distinguished from meiosis 65–6
  in a human lifetime 66
  phases 66F
mobile genetic elements
  in prokaryotes 187
  replicative and conservative transposition 210–15
  see also transposons
mobile phase, column chromatography 294
model organisms
  *Arabidopsis thaliana* as 352
  *Caenorhabditis elegans* as 142
  development processes 341
  *Drosophila melanogaster* as 348
  the mouse as 141–2
  modification interference 246–7
  modification protection assays 244–5, 246F, 247
  molecular beacons 62
  molecular biology, development of 27
  molecular chaperones 256, 316–18, 324
  molecular clock analyses 436, 441, 456
  molecular combing 77–8
  molecular ions 297–9
  molecular phylogenetics 135
  molecular rotors 361
  molten globule model 316
  Monod, Jacques 268, 271
  monogenic inherited diseases 150
  Montagnier, Luc 207
  Morgan, Thomas Hunt 65–9, 215
  mouse
    FOXP2 gene effects 452
    knockout mice 49, 142–3
    as a model organism 141–2
    nervous system development 351
    segmental duplications 442–3
    transgenic mice 144
mRNAs (messenger RNAs)
  cap structures 130
  degradation by RNAi 142, 278
  fraction of total RNA 257–8
  half-lives 276
  initiation complex scanning 312–13
  pre-mRNA splicing 13–14, 279–80
  regulating translation of 313–14
role 12
MudPIT (multidimensional protein identification technique) 305
MULEs (*Mutator*-like transposable elements) 447
Mullis, Kary 38
multicellular animals, evolution 434
multicopy cloning vectors 144
multicysteine zinc fingers 250
multigene families
  examples and characteristics 172–4
  gene duplication and 173, 435
  gene superfamilies 174
  paralogous genes 136
  problems with hybridization assays 125, 129–30
multipartite bacterial genomes 183–6
multiple alleles 59
multiple sclerosis 307, 308F
multiprotein complexes 228–9, 270, 304–5
multisubunit proteins 17, 264, 274, 368, 375, 448F
muntjac deer 155
muscular dystrophy 150T, 267
mutagens, chemical and physical 389, 394–7, 405–6
mutasome 405
<table>
<thead>
<tr>
<th>Term</th>
<th>Page(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>mutations</td>
<td>389–97</td>
</tr>
<tr>
<td>causes</td>
<td>389–97</td>
</tr>
<tr>
<td>insertions and deletions</td>
<td>389, 391–4</td>
</tr>
<tr>
<td>mitochondrial DNA</td>
<td>456</td>
</tr>
<tr>
<td>point mutation types</td>
<td>389, 390F</td>
</tr>
<tr>
<td>retrovirus mutation rates</td>
<td>454</td>
</tr>
<tr>
<td>Mycoplasma spp.</td>
<td>21</td>
</tr>
<tr>
<td>M. genitalium</td>
<td>100, 190</td>
</tr>
<tr>
<td>myoglobin</td>
<td>135–6, 436</td>
</tr>
<tr>
<td>Myrmecia pilosula</td>
<td>155</td>
</tr>
<tr>
<td>N</td>
<td></td>
</tr>
<tr>
<td>N50 and NG50 sizes</td>
<td>108, 112–13</td>
</tr>
<tr>
<td>N-CoR protein</td>
<td>275</td>
</tr>
<tr>
<td>N-linked glycosylation</td>
<td>320, 321F</td>
</tr>
<tr>
<td>N-termini, polypeptides</td>
<td>16</td>
</tr>
<tr>
<td>nanopore sequencing</td>
<td>98</td>
</tr>
<tr>
<td>Nasuia deltocephalnicola</td>
<td>189T, 190</td>
</tr>
<tr>
<td>natural selection</td>
<td></td>
</tr>
<tr>
<td>and pseudogene function</td>
<td>175–6, 438</td>
</tr>
<tr>
<td>in the RNA world</td>
<td>431</td>
</tr>
<tr>
<td>sequence similarity effects</td>
<td>123</td>
</tr>
<tr>
<td>Neanderthal genome</td>
<td>110–11, 452–3, 457</td>
</tr>
<tr>
<td>negative supercoiling</td>
<td>182, 365, 367</td>
</tr>
<tr>
<td>Neisseria meningitidis</td>
<td>188</td>
</tr>
<tr>
<td>nematodes see Caenorhabditis elegans; Globodera pallida</td>
<td></td>
</tr>
<tr>
<td>neuroglobin</td>
<td>436</td>
</tr>
<tr>
<td>Neurospora crassa</td>
<td>195</td>
</tr>
<tr>
<td>neutrons, in NMR</td>
<td>243</td>
</tr>
<tr>
<td>next-generation sequencing</td>
<td></td>
</tr>
<tr>
<td>de Bruijn graphs</td>
<td>102–3</td>
</tr>
<tr>
<td>methods</td>
<td>95–7</td>
</tr>
<tr>
<td>sequence assembly</td>
<td>102–104</td>
</tr>
<tr>
<td>sequencing library preparation</td>
<td>92–4</td>
</tr>
<tr>
<td>third- and fourth-generation</td>
<td>97–8</td>
</tr>
<tr>
<td>NHEJ (nonhomologous end-joining)</td>
<td>143, 404–6</td>
</tr>
<tr>
<td>nicks</td>
<td></td>
</tr>
<tr>
<td>direct repair</td>
<td>398–9</td>
</tr>
<tr>
<td>homologous recombination</td>
<td>413–15, 418</td>
</tr>
<tr>
<td>piperidine nicking</td>
<td>246, 247F</td>
</tr>
<tr>
<td>rolling-circle replication</td>
<td>364</td>
</tr>
<tr>
<td>transposition</td>
<td>423</td>
</tr>
<tr>
<td>nitrogenous bases</td>
<td></td>
</tr>
<tr>
<td>base ratios</td>
<td>7–8</td>
</tr>
<tr>
<td>in DNA structure</td>
<td>5</td>
</tr>
<tr>
<td>tautomers</td>
<td>391, 394T, 395</td>
</tr>
<tr>
<td>NMR (nuclear magnetic resonance) spectroscopy</td>
<td></td>
</tr>
<tr>
<td>DNA binding proteins</td>
<td>241, 243, 249</td>
</tr>
<tr>
<td>metabolome characterization</td>
<td>322</td>
</tr>
<tr>
<td>NOESY (nuclear Overhauser effect spectroscopy)</td>
<td>243</td>
</tr>
<tr>
<td>non-LTR retroelements</td>
<td>212–13, 379, 449</td>
</tr>
<tr>
<td>nonchromatin regions</td>
<td>224</td>
</tr>
<tr>
<td>noncoding RNAs</td>
<td></td>
</tr>
<tr>
<td>categories</td>
<td>12–13</td>
</tr>
<tr>
<td>half-lives</td>
<td>276</td>
</tr>
<tr>
<td>locating genes</td>
<td>122–3</td>
</tr>
<tr>
<td>as proportion of the transcriptome</td>
<td>15–16, 257</td>
</tr>
<tr>
<td>short and long</td>
<td>13, 257</td>
</tr>
<tr>
<td>see also IncRNAs; rRNAs; snRNAs; tRNAs</td>
<td></td>
</tr>
<tr>
<td>nonprocessed pseudogenes</td>
<td>174–5</td>
</tr>
<tr>
<td>nuclear genomes</td>
<td></td>
</tr>
<tr>
<td>chromosomal location and arrangement</td>
<td>155–61</td>
</tr>
<tr>
<td>DNA transfer with organelles</td>
<td>195</td>
</tr>
<tr>
<td>gene arrangement in</td>
<td>161–7</td>
</tr>
<tr>
<td>repetitive DNA</td>
<td>176–7</td>
</tr>
<tr>
<td>size</td>
<td>165F, 166T</td>
</tr>
<tr>
<td>nuclear lamina</td>
<td>222</td>
</tr>
<tr>
<td>nuclear matrix</td>
<td>222–3, 225</td>
</tr>
<tr>
<td>nuclear receptor superfamily</td>
<td>249T, 251, 332</td>
</tr>
<tr>
<td>nuclease protection experiments</td>
<td>156–7, 309</td>
</tr>
<tr>
<td>nuclease activity</td>
<td>29F</td>
</tr>
<tr>
<td>characteristics of useful nucleaseases</td>
<td>32F</td>
</tr>
<tr>
<td>S1</td>
<td>32F, 126, 127F</td>
</tr>
<tr>
<td>in yeast and humans</td>
<td>418</td>
</tr>
<tr>
<td>see also endonucleases; exonucleases</td>
<td></td>
</tr>
<tr>
<td>nucleic acids see DNA; RNA</td>
<td></td>
</tr>
<tr>
<td>nucleic acid analogs</td>
<td>433–4</td>
</tr>
<tr>
<td>nucleic-acid binding proteins</td>
<td></td>
</tr>
<tr>
<td>peptide NMR studies</td>
<td>243</td>
</tr>
<tr>
<td>see also DNA binding; RNA-binding</td>
<td></td>
</tr>
<tr>
<td>nucleoid-associated proteins</td>
<td>183</td>
</tr>
<tr>
<td>nucleoids</td>
<td>181–3, 185, 220</td>
</tr>
<tr>
<td>nucleolus</td>
<td>220–1, 223F</td>
</tr>
<tr>
<td>snoRNAs</td>
<td>259, 264, 265T, 444</td>
</tr>
<tr>
<td>nucleosides</td>
<td>5</td>
</tr>
<tr>
<td>nucleosomes</td>
<td></td>
</tr>
<tr>
<td>centromeric</td>
<td>160</td>
</tr>
<tr>
<td>and DNA replication</td>
<td>367, 369</td>
</tr>
<tr>
<td>introduced</td>
<td>156–7</td>
</tr>
<tr>
<td>modifications and genome expression</td>
<td>228–33</td>
</tr>
<tr>
<td>nucleosome remodeling</td>
<td>231–3</td>
</tr>
<tr>
<td>positioning</td>
<td>336, 449–50</td>
</tr>
<tr>
<td>nucleosome remodeling complexes</td>
<td>230, 233</td>
</tr>
<tr>
<td>nucleotides</td>
<td></td>
</tr>
<tr>
<td>abiotic formation</td>
<td>430</td>
</tr>
<tr>
<td>as DNA constituents</td>
<td>1, 4–8</td>
</tr>
<tr>
<td>nucleotide excision repair</td>
<td>401–2, 418</td>
</tr>
<tr>
<td>nucleus</td>
<td></td>
</tr>
<tr>
<td>internal structure</td>
<td>219–27</td>
</tr>
<tr>
<td>protein migration in</td>
<td>221</td>
</tr>
<tr>
<td>nutrient starvation</td>
<td>266, 311, 344</td>
</tr>
<tr>
<td>O</td>
<td></td>
</tr>
<tr>
<td>O-islands</td>
<td>191</td>
</tr>
<tr>
<td>O-linked glycosylation</td>
<td>320, 321F</td>
</tr>
<tr>
<td>O sequences</td>
<td>420</td>
</tr>
<tr>
<td>odds of linkage (lod scores)</td>
<td>72</td>
</tr>
<tr>
<td>Okazaki fragments</td>
<td>369–72, 375, 376–7F, 379F, 419F</td>
</tr>
</tbody>
</table>
oligonucleotide-directed mutagenesis 148
oligonucleotide hybridization analysis 61–2, 96F
oligonucleotide ligation assay (OLA) 62–3, 96
oligonucleotides, as primers 30
oncogenes 208F, 209
one-dimensional gel electrophoresis 296
one-step growth curve 205
oocytes 1, 138, 143
operator locus 269, 343
operons
  *E. coli* flagellum 323
  inducible and repressible operons 189
  lactose operon example 120, 188–9
  numbers of 187–9
  protein–protein interactions 306
  ribosomal protein genes 313
  threonine synthesis 187
  tryptophan synthesis 188F, 189, 272, 273F, 323
optical mapping 77–80
ORC (origin recognition complex) 365–7, 380
ORFs (open reading frames)
  assembly from gene fragments 340
  assigning functions 141, 192F
  function 119–24
  in genome browsers 131
  as protein coding regions 119–22, 168
  reporter gene replacement 146–7
ORF scanning 119–22, 186
organelles
  DNA transfer with nuclei 195
  genomes 195–9
  see also chloroplasts; mitochondrial
organic compounds, formation on the early Earth 429–30
organisms see species
organs, extent of gene expression 258
origin licensing 380
origin recognition complex (ORC) 365–7, 380
origin recognition sequence 365
origin(s) of life 429–33
origins of replication
  in bacterial plasmids 43–4, 48–9, 50F
  firing patterns 382–3
  in higher eukaryotes 366–7
  role, and numbers of 364
  S/MARs and 223
orphan retrogenes 438
orthologous genes 135
*Oryza sativa* (rice) 197, 199F
overlap graphs 102
overlapping genes 204–5
*Oxytricha* 377T

**P**
p53 protein 261F, 383
p300/CBP protein 228, 336
P element transposons 142, 215
P (peptidyl) site 310
P-TEFb protein 260
PacBio sequencing 98
PACs (P1-derived artificial chromosomes) 48
PAGE (polyacrylamide gel electrophoresis) 34, 87, 88F, 91, 294–7
pair-rule genes 349–51
paired-end reads 103, 104F, 112
paleogenomics 111, 452–3
palindromic sequences 187, 260, 271
  see also CRISPR
pan-genome concept 191–2
PANDA gene/RNA 261F
paralogous genes 135F, 136
*Paramecium* 196
pararetroviruses 207
partial linkage 65, 68–9
partial restriction, genetic mapping 76–7
party hubs 307
pathogenicity 184, 213
Pauling, Linus 6
pBIN19 cloning vector 50
pBR322 cloning vector 43
PcG (polycomb group) proteins 337
PCNA (proliferating cell nuclear antigen) 369
PCR (polymerase chain reaction)
  chromosome walking 105
  IRE-PCR 107
  limitations 41
  in next-generation sequencing 94
  in physical genome mapping 81
  process and applications 38–41
  quantitative PCR 41, 258
  real-time PCR 40, 62
  as a recombinant DNA technology 28
  in RFLP, SSLP and STR typing 59–61
  RT-PCR 126–7
  in site-directed mutagenesis 148–9
  thermal cycle sequencing 90–1
pedigree analysis
  genetic mapping by 71–2
  inherited breast cancer 150–1
*Pelomyxa* 195
pentoses 5, 433
peptide analysis, by mass spectrometry 297–8
peptide bonds 16, 308, 431
peptide nucleic acids (PNAs) 80, 433
peroxides 313, 394, 397
personalized medicine 92
phage display method/libraries 301–4
phages see bacteriophages
phenotypes
  alleles and 63
  changes, as clues to gene function 140
chromids as 185–6
as cloning vectors 28F, 41–4, 99
distinguishing recombinant plasmids 43–4
episome transfer 73
Saccharomyces cerevisiae 2μm plasmid 49
Plasmodium falciparum 78, 166T, 197, 199T
plectonemic, double helix as 358
pleiohomeotic (PHO) proteins 337
PNAs (peptide nucleic acids) 80, 433
point centromeres 160
point mutations
arising from replication errors 390–1
caued by mutagens 395–6
as origins of SNPs 61
as transitions and transversions 389
pol gene 27, 211–12, 425
polarities of amino acids 19
poly(A) tails 14, 126, 129, 212, 278
polyclonal antibodies 300, 301F
polycomb response elements 337
polyethylene glycol 82
polygenic traits 151–2
polyglutamine 393
polynucleotide phosphorylase (PNPase) 276
polynucleotide structure 6F
polypeptides, in primary protein structure 16
polyploidy 439–40, 445
polyproteins 207, 211–12, 318–19
polypyrimidine tracts 279–80, 282
positional effect, DNA cloning 226, 227F
positional information
Caenorhabditis elegans 346–8
Drosophila melanogaster 348–50
post-translational processing 270, 278, 315, 320, 446F
postspliceosome complex 280
POU domain 249T, 250, 275
ppGpp and ppGpp 311
Frader–Willi syndrome 235, 443–4
pre-replication complex (pre-RC) 380–1
pre-RNA splicing 13–14, 278–9
preinitiation complexes (pre-ICs) 273–4, 312, 381
prepriming complexes 365
prespliceosome complex 280
prespores 344–6
primary protein structure 16
primases 369, 370F, 371, 377
primates
evolution 443, 450–3
simian immunodeficiency viruses 454–5
primers
chain-termination sequencing 88
definition 30–1
DNA replication 369
internal primers 91, 93, 100F
PCR 39–41, 59, 60F, 61, 63, 105

Index 511
preparing cDNA 82F
retrotransposition 425
universal primers 91
prions 209
prioritized-mode RNAs 288
processed pseudogenes 174–5
processivity, of DNA polymerases 90
programmable nucleases 143, 187
progressive myoclonus epilepsy 393
prokaryotes
bacteria and archaea 181
DNA topoisomerases 362–3
DNA transposons 213–14
genome sizes and gene numbers 189–92
‘immune system’ in 187
species concept 190–2
prokaryotic genomes
genetic features 186–95
numbers of operons 187–9
physical features 181–6
shotgun sequencing 100–1
promiscuous DNA 195–6
promoter sequences
alternative promoters 267–8, 285, 331, 452
core and basal promoters 266–7, 273
endogenous promoters 277F
and genetic switching 343
strong promoters 268
transcription start points and 266–8
upstream promoter elements 266–7, 274–5
proofreading 30, 368, 391
proopiomelanocortin 318–19
prophages 205–6, 342–3, 405, 420
prophase (meiosis) 66, 67–8F
prophase (mitosis) 66F
PROSITE database 137
proteases 3, 4F, 31, 294
proteasomes 314–15, 381
protection assays 244–5, 246F, 247
proteins
activation by cleavage 318–19
chemical modification 315, 320–1
digestion by trypsin and chymotrypsin 437
DNA-binding 137
functional diversity 22
glycosylation 320–1
housekeeping proteins 19
origins of protein-based catalysis 432–3
post-translational processing 270, 278, 315, 320, 446F
structural diversity 17–19
protein arrays, in protein profiling 300–1, 303
protein classification 138–9
protein-DNA interactions see DNA binding
protein domains
death domain 172
and gene catalogs 172
and gene function 137–8
genomic specification 172
subdomain folding 316
Tudor domains 137–8
whether coded for by exons 446
zinc finger domains 137, 172F, 250–1, 332
protein folding 243, 315–18, 324–5
protein interaction maps 306–7
protein kinase A 336
protein products and gene function 146–9
protein profiling
alternatives 300, 303
chemical modification in 321
comparing proteome compositions 299–301
defined 293
identification stage 297–9
separation stage 294–7
top-down and bottom-up approaches 293
using protein arrays 300–1
protein-protein binding domains 335
protein-protein interactions
cross-linking 309
identifying 301–8
protein-RNA complexes 259–60
protein structure
chemical modification 19
four levels 16–17
immunoglobulins 339–40
NMR spectroscopy 243
X-ray crystallography 241–3
proteolytic cleavage
activation by 318–19
alternative patterns 318–19
proteomes
comparing compositions 299–301
component synthesis and degradation 308–15
compositional studies 293–301
defined 293
downsizing 311–12
genome expression and 2
and genome sizes 169
link to transcriptome 19–20, 314
metabolome, systems biology and 322–5
protein processing influence 315–21
and protein structure 16–19
proteome remodeling 312–13
proteomes 431–3
protoplasts 49, 83
protozoa, telomere sequences 377T
PSE (proximal sequence element) 267
pseudogenes
duplicated and unitary 174
as evolution relics 174–6, 435
processed and nonprocessed 174–5, 438
PTENP1 175–6
recombinant DNA technology 43–4
recombination events
changing meaning 411
discovery 66
double recombination 70
gene inactivation 140
interspersed repeat origins 177
lod analysis 72
use of reference genomes 101
see also crossing over; homologous recombination;
site-specific recombination
recombination frequency 68, 70–2
recombination hotspots 69, 74
RecQ helicase 406, 417
reference genomes 101, 110, 129–30
reflectrons 297
regional centromeres 160
regulatory proteins, response to signaling compounds 331
remediation of contaminated land 194
REP (repetitive extragenic palindromic) sequences 187, 260
repetitive DNA fingerprints 106F, 107
repetitive DNA PCR 106F, 107
repetitive DNA sequences
barley gene space 114
centromeric regions 160
dinucleotide and trinucleotide repeats 177
direct repeats from transposition 421, 425
distinguished from low-copy repeats 442
in humans and chimpanzees 451
inducing replication slippage 392
as interspersed and tandem repeats 176–7
in prokaryotes 187
shotgun sequencing errors 56, 102–4
trinucleotide repeats 392
see also transposable elements
replacement cloning vectors 45
replication errors
causing point mutations 390–1
insertion and deletion mutations 391–4
mismatch repair 402–3
replication factories/foci 374
replication forks
initiation and progression 365–6
migration rates 383
overwinding 361
processes around 367–72
semiconservative replication 363–4
stalling 383, 418
replication mediator proteins (RMPs) 366, 370
replication origins see origins of replication
replication slippage 177, 392–3, 446
replisomes 371, 373–4, 396, 403
reporter fragment ions 300

Pseudomonas syringae DC3000 189
PSI-BLAST (position-specific iterated BLAST) 137
PstI restriction endonuclease 33–4
PTENPI pseudogene 175–6
pUC8 plasmid 43–4, 47
pulse labeling 275–6, 382
punctuation codons 20
purine, 2-amino- 395
purine-2,6-dicarboxylate 433
purine bases
adenine and guanine as 5
transitions between 389
pyridine 433
pyrimidine bases
dimerization 396
polypurimidine tracts 279–80, 282
transitions between 389
pyrosequencing 95–7
pyrrolysine 19, 21, 320

Q
QTLs (quantitative trait loci) 57, 61, 113
quadrupole mass analyzers 298
quaternary protein structure 17

R
RACE (rapid amplification of cDNA ends) 126
Radding, C. M. see Meselson–Radding
radiation hybrids 82–3, 109
radiolabeling 3, 4F
see also isotope studies
Ralstonia 445
random genomic sequences 82
reactive oxygen compounds 313
read depth 92F
read lengths 96–8
reading frames
and genome browsers 131
and ORF scanning 119, 120F, 121
overlapping genes 204–5
see also ORFs
real-time PCR 40, 62
real-time sequencing 97
RecA protein 405, 416–17
RecBCD pathway/complex 415–18
receptor activity, GO 170
receptor proteins, transmembrane 332–3
recessive and dominant phenotypes 64
RecFOR pathway 415, 417–18
reciprocal strand exchange 413, 415
Reclinomonas americana 197, 199T
recognition helix 11, 250, 254, 275F
recombinant DNA technology
development 28
distinguishing recombinant plasmids 43–4
see also DNA cloning; PCR
recombinants 420–1
recognition helix
Reclinomonas americana 11
reciprocal strand exchange
RecFOR pathway 415, 417–18
recognition helix 11, 250, 254, 275F
recombinant DNA technology
development 28
distinguishing recombinant plasmids 43–4
see also DNA cloning; PCR
recombinases 420–1
recombination events
changing meaning 411
discovery 66
double recombination 70
gene inactivation 140
interspersed repeat origins 177
lod analysis 72
use of reference genomes 101
see also crossing over; homologous recombination;
site-specific recombination
recombination frequency 68, 70–2
recombination hotspots 69, 74
RecQ helicase 406, 417
reference genomes 101, 110, 129–30
reflectrons 297
regional centromeres 160
regulatory proteins, response to signaling compounds 331
remediation of contaminated land 194
REP (repetitive extragenic palindromic) sequences 187, 260
repetitive DNA fingerprints 106F, 107
repetitive DNA PCR 106F, 107
repetitive DNA sequences
barley gene space 114
centromeric regions 160
dinucleotide and trinucleotide repeats 177
direct repeats from transposition 421, 425
distinguished from low-copy repeats 442
in humans and chimpanzees 451
inducing replication slippage 392
as interspersed and tandem repeats 176–7
in prokaryotes 187
shotgun sequencing errors 56, 102–4
trinucleotide repeats 392
see also transposable elements
replacement cloning vectors 45
replication errors
causing point mutations 390–1
insertion and deletion mutations 391–4
mismatch repair 402–3
replication factories/foci 374
replication forks
initiation and progression 365–6
migration rates 383
overwinding 361
processes around 367–72
semiconservative replication 363–4
stalling 383, 418
replication mediator proteins (RMPs) 366, 370
replication origins see origins of replication
replication slippage 177, 392–3, 446
replisomes 371, 373–4, 396, 403
reporter fragment ions 300
reporter genes
  E. coli flagellum operons 324
  in the two-hybrid system 302–3
reporter probes 40, 62
repressor proteins, RNA synthesis 258, 268–73
resolution, Holliday structures 413
resolvases 214F, 416, 418
resonance frequencies 243
respiratory chain/complex 194, 197F, 198, 199T
restriction endonucleases 32–4, 100, 130, 224, 225F, 366F, 367
restriction mapping 75–9, 125F
restriction patterns 106
retinoblastomas 150T, 229
retroelements
  group II introns 449
  human and chimpanzee 451
  LTR elements 163–4F, 165–7, 210–11
  non-LTR elements 212–13, 379, 449
  reverse transcriptase 368T
  transposition mechanism 422–5
  viral retroelements 207, 210–11
  with and without LTRs 210
retrogenes 438, 439F
retrohoming 449
retroons 213
retroposons 212, 379
retrottransposition 210, 423–4, 438
retrotransposons 210F; 211–12, 259, 424
retroviruses
  and cancer 207–9
  in HIV/AIDS 207, 423, 454
  human endogenous retroviruses 212, 442
  mutation rate 454
  reverse transcriptases 31
  RNA intermediate 423–4
  structure and replication strategies 206F, 207–9
reverse transcriptases
  in cDNA preparation 31, 126
  as DNA polymerases 31, 367
  and early DNA genomes 432
  isolation of 207
  RT-PCR 126–7
  in telomeras 376
reversible terminator sequencing 95, 97, 129
RFC (replication factor C) 371
RFLPs (restriction fragment length polymorphisms) 59–61, 68–9, 75, 109, 150–1
rheumatoid arthritis 212, 307, 308F
Rho-dependent terminators 271–2
Rhodobacter spp. 194
Rhodopirellula baltica 189
ribbon-helix-helix motif 251
ribonuclease structure 315
ribose, possible alternatives 433–4
  see also deoxyribose
ribosomal RNAs see rRNAs
ribosomes
  A, P and E sites 310
  bacterial and eukaryotic 308–9
  binding sites 310–13
  inactivation 311–12
  introduced 13
  in protein synthesis 19, 308–10
  stalling 272–3, 311
  X-ray crystallography 310–11
riboswitches 272, 431
ribozymes 431–2, 449
rice (Oryza sativa) 197, 199F
RISC (RNA-induced silencing complex) 277–8
RMPs (replication mediator proteins) 370
RNA (ribonucleic acid)
  chemical differences from DNA 12F
  chemical modification 15
  cloverleaf structure 122
  coding and noncoding 12
  degradation, nonspecific 275–6
  double stranded (dsRNA) 142, 143, 204T, 277, 306T
  end modification 14
  enzyme activity 209
  foldback RNAs 278
  guide RNAs 143–4, 187
  IncRNAs (long noncoding RNAs) 13, 254, 257, 260–2, 265T
  miRNAs (micro RNAs) 15–16, 175, 259, 260F, 264, 278
  and the origin of proteins 432
  piRNAs (piwi-interacting RNAs) 138, 259, 264, 265T
  precursor molecules 13–14
  scaRNAs (small Cajal body-specific RNAs) 259
  7SK and 7SL RNAs 212, 260, 265T
  siRNAs (short interfering RNAs) 142F, 259, 260F, 264, 265T, 277
  snRNAs (short noncoding RNAs) 13–15, 122, 257, 259–60, 264
  snoRNAs (small nucleolar RNAs) 259, 264, 265T, 444
  snRNAs (small nuclear RNAs) 259, 264, 265T, 267
  synthesis and degradation 263–75
  and the transcriptome 11–16
  transfer-messenger RNA 260
  as transposition intermediate 423–5
  vault RNAs 259–60
  see also mRNAs; rRNAs; tRNAs
RNA-binding proteins
  in the exosome 276
  motifs 252
  RNA-dependent DNA polymerases 31, 367, 425
  RNA-dependent RNA polymerases 12
  RNA editing 15, 20
  RNA genomes
    HIV 454
    plant viruses 206
    RNA protogenomes 431–3
RNA interference (RNAi)
- as defense against viruses 276-8
- in genomics research 142-3
  - small non-coding RNAs in 138, 259
RNA mapping
- genomewide 127-31
- transcript mapping 124-7, 284
RNA polymerases
- during B. subtilis sporulation 345
- CTD phosphorylation 274, 281
- DNA- and RNA-dependent 12, 264
- error rate 265
- mitochondrial polymerases 265
- poly(A) polymerase 14
- rate of RNA synthesis 265
RNA polymerase I 264, 265T, 266
RNA polymerase II 212, 264-5, 267, 273-5, 278, 281-2, 285
RNA polymerase III 212, 264, 265T, 267
RNA polymerases IV and VI 264
RNA recognition domains 252
RNA-seq
- barley genome 114, 124
- transcript analysis 129-30, 262-3
RNA silencing see RNA interference
RNA splicing 13-14, 138, 220, 276
RNA world concept 431-2, 434, 448
RNAi (RNA interference) 138, 142-3, 259, 277
RNases
- RNase E 276
- RNase H 425
- RNase P 431
rolling-circle replication 364
Rous sarcoma virus 208
RPA (replication protein A) 370
RPLC (reverse-phase liquid chromatography) 295, 299
rRNAs (ribosomal RNAs)
- bacterial pre-rRNA 14
- characteristics 13-15
- gene cluster for 5S rRNA 173
- gene family as evidence of duplication 435
- nucleolar synthesis 220
RT-PCR (reverse transcriptase PCR) 126-7
Rubisco (ribulose-bisphosphate carboxylase) 321
S
S/MARs (scaffold/matrix attachment regions) 223, 225
Saccharomyces spp. 124
Saccharomyces cerevisiae
- biochemical markers 58T
- copper uptake 331
- gene catalog 171
- gene inactivation 140-1
- genome annotation 123-4
- genome characteristics 164-5, 196-8, 199T
- genome duplication 441
- genome sequencing 74-5
- haploid colonies 69
- mating type determination 338-9, 412
- 2μm plasmid 49
- origins of replication 364, 365-6
- point centromeres 160
- prions of 209
- protein interaction maps 306
- in the two-hybrid system 302
- Ty element 211
- typical phenotype screens 145T, 146
SAGA complex 228-9
Salmonella typhimurium 190, 192, 213
sample genome segments
- Drosophila 165, 167F
- E. coli 120F, 122F, 187
- human genome 163-4, 212
- S. cerevisiae 164, 167F
Sanger, Frederick 20, 87
see also chain-termination
satellite DNA
- minisatellites and microsatellites 176-7
- tandem repeats as 176
satellite RNAs 209
Sau3AI restriction endonuclease 33-4
scaffolds 103-4, 108, 112, 129
scaffold/matrix attachment regions (S/MARs) 223, 225
scRNA (small Cajal body-specific RNAs) 259
Schizosaccharomyces pombe 168T, 382, 418
SDS (sodium dodecyl sulfate) 294, 296
second messengers 336
secondary protein structure 16
segment polarity genes 349-51
segmental duplications 442-4, 450, 452
segmented body plans 348
segmented genomes 204
segregation, allelic 64
selectable markers 43
selenocysteine 19, 21, 22F, 320
self-splicing 188
semiconservative model, DNA replication 359-61, 363-4
Sendai virus 82
sequence assembly 55, 56F, 98-110
sequence contigs 99, 103, 107, 114
sequence depth see coverage
sequence inspection 119
sequence similarity 129, 135-9, 173-4, 183, 207
- between homologous genes 123, 135
sequence-specificity
- DNA binding proteins 241, 244, 249, 251, 253F, 273, 331, 366, 373
- DNA endonucleases 32T
- proteases 294
sequencing-by-synthesis methods 96-7
sequencing libraries 92-4, 103, 107-9, 112-14
serine–threonine kinase receptors 332
7SK and 7SL RNAs 212, 260, 265T
sex cells, distinguished from somatic 1
sex determination in Drosophila 282–3
sex-specific alternative splicing 285
sexual reproduction in Bangiomorpha 434
Shapiro model (J. A. Shapiro) of transposition 422
shelterins 161, 404
short-patch and very-short-patch repairs 399–401, 403
short products, PCR 39–40
shotgun proteomics 296, 298, 305
shotgun sequencing
eukaryotic genomes 102–4
giant panda genome 111
in the HGP 109–10
hierarchical shotgun approach 104–7
limitations 55–6
prokaryotic genomes 100–1
sequence assembly 98–104
shuttle vectors 49, 50F
signal peptides 318, 319F, 331
signal recognition particles 260
signal transduction pathways
C. elegans differentiation 348
cascade pathways 334–5
importance of phosphorylation 321
MAP kinase 334–5, 338
from receptor to genome 332–6
using second messengers 336
signal transmission across membranes
by importing signaling compounds 330–2
by receptor proteins 332–3
silencer elements 275, 281–4, 330, 338
silent cassettes 338
similar-mode RNAs 287
Sin3 complex 229–30, 234
SINEs (short interspersed nuclear elements) 163, 166, 211T, 212–13
single-molecule realtime sequencing 97
single-strand break repairs 398, 403–4, 406
single-strand gap repairs 417–18
Sir2 complex 229
siRNAs (short interfering RNAs) 142F, 259, 260F, 264, 265T, 277
site-directed hydroxyl radical probing 310
site-directed mutagenesis 147–9
site-specific recombination
bacteriophage λ 205, 206F, 342, 419–21
distinguished from homologous recombination 411–12
SIV (simian immunodeficiency viruses) 454–5
6–4 lesions/photoproducts 396F, 397, 399
sliding, as nucleosome remodeling 233
sliding clamps 368, 370
slippage 177
slo gene 284
sncRNAs (short noncoding RNAs) 13–15, 122, 257, 259–60, 264
see also miRNAs
snoRNAs (small nucleolar RNAs) 259, 264, 265T, 444
SNPs (single-nucleotide polymorphisms)
biallelic SNPs 61
as DNA markers 61–3, 152
in genome browsers 131
human genome 107, 152, 456–7
with linkage analysis 68–9
Neanderthal genome 111
plant genomes 83, 114, 458
tiling arrays 128
typing methods 63F
snRNAs (small nuclear RNAs) 259, 264, 265T, 267
snRNPs (small nuclear ribonucleoproteins) 259–60, 280, 281F, 282
SNS (short nascent strand) sequencing 366T, 367
sodium bisulfite deamination 395
software
genome browsers 131
for homology searching 136–7
sequence assemblers 102
solid phase, column chromatography 294
SOLiD (sequencing by oligonucleotide ligation and detection) 96–7
solution hybridization technique 62
somatic, distinguished from sex cells 1
sonication 93, 99, 247
Sorangium cellulosum 189T, 190
sorting sequences 137
SOS protein 334, 335F
SOS response 343, 405
Southern hybridization 35, 36F, 59, 60F, 100, 107
zoo-blotting 125
species
chromosome numbers 155
codon bias in 121–2, 146
concept in prokaryotes 190–4
detectable by metagenomics 194
extent of cytosine methylation 234
gene catalogs compared 169–72
speciation and transposable elements 215
specific features of humans 452
specificity of viruses for 203
speckles 220, 259
spermatozoa, direct genotyping 69
spliceosomes 259, 280, 304, 449
splicing code, possible 284
splicing pathways 169, 279, 282, 449, 452
see also alternative splicing
splicing silencers 281–2
Spm elements 214
SpoOA protein 345–6
sporulation, Bacillus subtilis 343–6
subdomains, protein 316

SSLS (simple sequence length polymorphisms)
  as DNA markers 59–61, 68–9
  in the HGP 109
  mini- and microsatellite 60
  numbers in humans 453
  as STS markers 82, 84

Stahl, Franklin 359

staining, and chromosome banding 79, 158, 159F, 161–2

stalling, cell cycle 229

stalling, replication fork 383, 418

stalling, ribosome 272–3, 311

start codons (initiation codons) 20, 119F, 311

starvation responses 266, 311, 344

STATs (signal transducers and activators of transcription) 333

stem cells 141, 378

stem-loop structures
  foldback RNAs 278
  genes for noncoding RNAs 122, 123F
  miRNAs and 259, 260F
  REP sequences and 187
  ribosome binding 313
  selenocysteine codon 21, 22F
  transcription termination 271–2, 273F, 276
  translation prevention 313

steroid hormones 274, 331, 332F

steroid receptor protein 331–2

sterols, as specific to eukaryotes 434

sticky ends 32, 33F, 34, 37, 38F, 41

stop codons (termination codons) 20–1, 119F, 121F, 276

strand break repair systems 403–5

streptavidin 93, 94F

Streptococcus spp.
  S. agalactiae 191–2
  S. pneumoniae 3, 4F
  S. pyogenes 144

Streptomyces coelicolor 183–4, 185T

stress
  B. subtilis sporulation 345
  ribosome inactivation response 311–12
  stress response in plants 287–8

stringent response 311–12

STRs (short tandem repeats) 60–1, 63, 151
  see also microsatellite DNA

structural activity, GO 171

STS (sequence-tagged sites) 81–4
  STS content mapping 81–4, 106F, 107, 109, 113
  STS markers 81–4, 109, 131F

stuffer fragments 45

Sturtevant, Arthur 68, 74

subdomains, DNA 365–6

subdomains, protein 316

substitution mutations 456

sugar pucker 10

suicide enzymes 399

SUMO (small ubiquitin-related modifier) 230, 314

supercoiling
  and buoyant density 162
  in DNA replication 358, 363, 365, 367
  DnaA attachment and 365
  in the E. coli genome 182–3
  negative supercoiling 182, 365, 367
  reverse gyrase and 363

surveillance mechanism 276, 402

Sutton, W S 3

SWI/SNF complex 230, 233

symbiosis
  endosymbiosis 190, 195–6
  satellite RNAs and virusoids 209

synctia 348–9

synteny 124, 225, 441

systems biology 2, 322–4, 341

T

T-cell receptor diversity 339–41

T-DNAs 10, 49, 50F

t-loops 377, 378F

T4 polynucleotide kinase 38

TADs (topologically associated domains) 225–7

TAFs (TBP-associated factors) 229, 273

TAF- and initiator-dependent cofactors (TICs) 273

tandem gene duplications 438, 439F

tandem mass spectrometry 298–300

tandem repeats
  in mice and humans 442–3
  as repetitive DNA sequences 176–7
  see also satellite DNA; STRs

TAP (tandem affinity purification) 304F, 305

Taq DNA polymerase 31T, 39, 63, 90–1, 126

target enrichment 108

TATA box 267, 273, 424

TATA-binding protein (TBP) 229, 249T, 273

Tatum, Edward 323

tautomers 391, 394–5

TBP (TATA-binding protein) 229, 249T, 273

telomeras 260, 375–9

telomeres/telomeric regions
  in cell senescence and cancer 378–9
  characteristic repeat sequence 161
  chromosome recognition 158
  DNA-protein interactions 159–61
  gene density and 162
  in metaphase chromosomes 158–61
  and retroposons 379
  transfer of repeat sequences 379

telomeric DNA as a minisatellite 60, 177

Temin, Howard 207
temperate infection cycles see lysogenic
template-dependent DNA polymerases 8–9, 28–30, 31T, 89
template-dependent RNA polymerases 12
template-independent DNA polymerases 38, 367
template-independent RNA polymerases 14
template switching 424f, 425
terminal deoxyxynucleotidyl transferase 29f, 37–8, 367, 368T
termination codons 20–1, 119f, 121f, 276
terminator sequences 271, 276, 373–4
terminology, gene function annotation 138–8
tertiary protein structure 16–17, 137, 315
test crosses 69–71
tetracycline resistance 49f, 213
Tetrahymena 228, 377T
TFIIA transcription factor 273–4
TFIIB transcription factor 267, 273–4
TFIID transcription factor 267, 273–4
TFIIE transcription factor 229, 274
TFIIF transcription factor 229, 273–4, 402
TFIIF transcription factor 273–4, 402
TFIIIA transcription factor 249T
TGFβ (transforming growth factor β) 332
thalassemia 41
thermal cycle sequencing 90–1
Thermosipho spp. 194
Thermotoga maritima 193
Thermus aquaticus (Taq) 31
Taq DNA polymerase 31T, 39, 63, 90–1, 126
thio-substitution of RNA 15
4-thiouracil 15f, 275
third- and fourth-generation sequencing 97–8
30 nm fiber 157, 221, 223, 228–9
3’ or 3’-OH terminus 6
3’-transduction 447
3C (chromosome conformation capture) method 224–5, 247
thymine
  relation to uracil 432
  structure 5F
thymine dimers 396
Ti plasmid 49, 50F
Tibet 453
tiling arrays
  genomewide RNA mapping 128–9
  mRNA copy numbers 258
  transcriptome studies 262
see also microarray analysis
time-of-flight spectrometry 297–8
tissues, extent of gene expression 258
TK (thymidine kinase) 82–3
Tn3-type or unit transposons 213, 214f, 422
TOCSY (total correlation spectroscopy) 243
top-down approaches, protein profiling 293–4, 297–8, 300
topoisomerases
  breakage-and-reunion model 358–9
  DNA gyrase 182, 363
  evolution 362–3
  integrase 342, 420–1, 425
  linking number 361–2
  nuclear matrix 222
  overcoming stresses 370
  Top 2 and topoisoamerase IV 374
totipotency 141
TPA (tissue plasminogen activator) 446–7
trans-displacement, nucleosomes 233
transcribed sequences, hybridization tests 125
transcript-specific regulation 313–14
transcription
  defined 2
  mapping transcript ends 126
  multiple transcripts 125
  pausing replication 372, 383
  synthesis of RNA 263–275
  termination, bacterial RNA 271–2
transcription-coupled repair 402, 406
transcription factors
  EPAS1 453
  FOXP2 451–2
  homeotic genes as 350, 352
  as proteins 22
  regulatory role 274–5
  sequence-specific protein binding 244
  in the two-hybrid system 302
see also general transcription factor; TFIIA etc.
transcription start points 223, 231, 258, 274
transcriptional modes, plant stress 287–8
transcriptome
  and cancer research 286–7
  components of 257–63
  degradation of components 275–8
  different definitions 15–16, 257
  and genome annotation 284–6
  genome expression and 2
  influence of RNA processing 278–84
  link to proteome 19–20
  overview 11–16
  and plant stress 287–8
  and protein-protein interactions 306
  synthesis of components 263–75
transduction, by bacteriophages 73, 74f, 192
transfection 45
transfer, as nucleosome remodeling 233
transfer RNA see tRNAs
transformation
  in DNA cloning 43, 50
  in genetic mapping 73
transforming principle 3, 4F
transgenic mice 144
translation
defined 2
regulating for individual mRNAs 313–14
synthesis of protein 308–314
translational efficiency 314
transporter activity, GO 170–1
transposable elements
insertion sequences as 187
moving gene segments 447
three categories 210, 422
transposases 213–15
transposition
interspersed repeats from 177
replicative and conservative transposition 210, 421–5
retrotransposition 210, 423–4, 438
Shapiro model 422
use of recombination 412
transposon tagging 142
transposons
control using RNAi 277
DNA transposons 163, 166, 213–15, 447
as interspersed repeats 163, 166
as mobile genetic elements 210–15
P element transposon 142
retrotransposons 250
RNA transposons 210–13
yeast Ty1 transposon 142
treble clef fingers 249T, 251, 332
TRiC chaperonins 316, 318
Trichomonas vaginalis 448
trichothiodystrophy 406
trimethylpsoralen 182
triplex structures 80, 416
trisomies 439
Triticum aestivum 166F, 445
Triticum turgidum 445
tRNAs (transfer RNAs)
as adaptors in protein synthesis 19
locating genes for 122
pre-tRNAs 14
as retrotransposition primers 425
role 13–15
truncated genes 123F, 175
truncated LTR sequences 165
trxG (trithorax group) proteins 337
trypsin 293F, 294, 437
tryptophan biosynthesis 73, 323
tryptophan operon 188, 272, 273F, 323
Tsix gene 236, 261
Tudor protein domain 137–8
tumor suppressor genes 151
tumor-suppressor proteins 383
Tus (terminator utilization substance) 373–4
two-dimensional gel electrophoresis 294–5, 297, 299
two-hybrid system 301–4, 306
Ty1/copia retroelement 211, 424–5
Ty3/gypsy retroelement 211, 424
tyrosine, and topoisomerase action 362
tyrosine kinase-associated receptors 332–3
tyrosine kinases 347

U
U-RNAs see snRNAs
ubiquitin ligases 314, 381
ubiquitin-receptor protein 314
ubiquitination 163, 230–3, 314, 337
UCE (upstream control elements) 266, 267F
UCSC genome browser
use of recombination 412
untranslated regions (UTRs)
in transcript analysis 124, 128
3’-UTR 21, 22F, 278, 393
5’-UTR 313, 393
upstream control elements (UCE) 266, 267F
upstream promoter elements 266–7, 274–5
upstream regulatory sequences 121, 175
uracil
4-thio- 15F, 275
5-bromo- (5-bU) 394–5
dihydro- 15F
relative stability of thymine 432
urea 245, 315–16
Urey, Harold 430F
UvrABC endonucleases

V
v- onc oncogenes 208F, 209
vaccinia virus 206
vault RNAs 259–60
Venter, Craig 110
vertebrate genome evolution 441, 448F
vertical resolution of Holliday structures 413
Vibrio cholerae
chromosome and chromid count 186
O1 El Tor strain 184, 185T, 189T
retroelements 213
viral DNA, detection by PCR 41
viral retroelements 207, 210–11
virulence and the lytic infection cycle 205
viruses
genomes of eukaryotic 206–9
picornaviruses 312
polyprotein use 319
and RNA silencing 276
whether living 203, 209
see also bacteriophages
virusoids 209, 431
Vitamin C (ascorbic acid) 174
VLPs (virus-like particles) 211
VNTRs (variable number of tandem repeats) 60
   see also minisatellite DNA

W
water content
   DNA structure and 6
   in protein-DNA interactions 254
Watson, James 4–6, 7F, 8–9, 357–8
Watson-Crick base pairs 9, 80, 358, 433
Werner’s syndrome 406
wheat (Triticum spp.)
   allopolyploidization 445
   genome characteristics 57-8, 166T, 442F
   gliadins 22, 258
   hierarchical shotgun sequencing 104
   landraces 457
   radiation hybrids 83
whole-genome duplication 439–42
wild type, phenotypes 73
winged helix-turn-helix motif 249F, 250

X
X-gal 44
X inactivation 235–6, 261
X-ray crystallography
   DNA binding proteins 241–3
   double helical DNA structure 358
   ribosomes 310–11
   RuvA, RuvB and RecC proteins 416
X-ray diffraction principles 242
X-ray diffraction studies 6–7, 310, 358
X-rays, radiation hybrids 82
xeroderma pigmentosum 406
Xist gene 236, 261

Y
yeasts
   genome characteristics 164–5
   yeast two-hybrid system 301–4, 306
   see also Saccharomyces cerevisiae;
      Schizosaccharomyces pombe
YIp5 cloning vector 49

Z
Zea mays (maize) 21F, 166–7, 197T, 211, 214–15
zinc finger domains
   Cys2H2 172F
   Cys3His2 zinc fingers 137, 250
   GATA 172F, 250
   versions 250–1, 332
zoo-blotting 125