# Linkage, Recombination, **7** and Eukaryotic Gene Mapping







**(a) (b)**

#### **Pattern baldness is a hereditary trait.**

Recent research demonstrated that a gene for pattern baldness is linked to genetic markers located on the X chromosome, leading to the discovery that pattern baldness is influenced by variation in the androgen-receptor gene. The trait is seen in three generations of the Adams family: (a) John Adams (1735–1826), the second president of the United States, was father to (b) John Quincy Adams (1767–1848), sixth U.S. president, who was father to (c) Charles Francis Adams (1807–1886). [Part a: National Museum of American Art, Washington, D.C./Art Resource, New York. Part b: National Portrait Gallery, Washington, D.C./Art Resource, New York. Part c: Bettmann/Corbis.]

# LINKED GENES AND BALD HEADS

For many, baldness is the curse of manhood. Twenty-five percent of men begin balding by age 30 and almost half are bald to some degree by age 50. In the United States, baldness affects more than 40 million men, who spend hundreds of millions of dollars each year on hair-loss treatment. Baldness is not just a matter of vanity: bald males are more likely to suffer from heart disease, high blood pressure, and prostate cancer.

Baldness can arise for a number of different reasons, including illness, injury, drugs, and heredity. The most-common type of baldness seen in men is pattern baldness—technically known as androgenic alopecia—in which hair is lost prematurely from the front and top of the head. More than 95% of hair loss in men is pattern baldness. Although pattern baldness is also seen in women, it is usually expressed weakly as mild thinning of the hair. The trait is clearly stimulated by male sex hormones (androgens), as evidenced by the observation that males castrated at an early age rarely become bald (but this is not recommended as a preventive treatment for baldness).

A strong hereditary influence on pattern baldness has long been recognized, but the exact mode of inheritance has been controversial. An early study suggested that it was autosomal dominant in males and recessive in females, an example of a sex influenced trait (see Chapter 4). Other evidence and common folklore suggested that a man inherits baldness from his mother's side of the family, exhibiting X-linked inheritance.

In 2005, geneticist Axel Hillmer and his colleagues set out to locate the gene that causes pattern baldness. They suspected that the gene for pattern baldness might be located on the X chromosome, but they had no idea where on the X chromosome the gene might reside. To identify the location of the gene, they conducted a linkage analysis study, in which they

looked for an association between the inheritance of pattern baldness and the inheritance of genetic variants known to be located on the X chromosome. The genetic variants used in the study were single-nucleotide polymorphisms (SNPs, pronounced "snips"), which are positions in the genome where individuals vary in a single nucleotide. The geneticists studied the inheritance of pattern baldness and SNPs in 95 families in which at least two brothers developed pattern baldness at an early age.

What Hillmer and his colleagues found was that pattern baldness and SNPS from the X chromosome were not inherited independently, as predicted by Mendel's principle of independent assortment. Instead, they tended to be inherited together, which occurs when genes are physically linked on the same chromosome and segregate together in meiosis.

As we will learn in this chapter, linkage between genes is broken down by recombination (crossing over), and the amount of recombination between genes is directly related to the distance between them. In 1911, Thomas Hunt Morgan and his student Alfred Sturtevant demonstrated in fruit flies that genes can be mapped by determining the rates of recombination between the genes. Using this method for the families with pattern baldness, Hillmer and his colleagues demonstrated that the gene for pattern baldness is closely linked to SNPs located at position p12–22, a region that includes the androgen-receptor gene, on the X chromosome. The androgen receptor gene encodes a protein that binds male sex hormones. Given the clear involvement of male hormones in the development of pattern baldness, the androgen-receptor gene seemed a likely candidate for causing pattern baldness. Further analysis revealed that certain alleles of the androgen-receptor gene were closely associated with the inheritance of pattern baldness, and the androgen-receptor gene is almost certainly responsible for much of the differences in pattern baldness seen in the families examined. Additional studies conducted in 2008 found that genes on chromosomes 3 and 20 also appear to contribute to the expression of pattern baldness. **TRY PROBLEM 13** 

This chapter explores the inheritance of genes located on the same chromosome. These linked genes do not strictly obey Mendel's principle of independent assortment; rather, they tend to be inherited together. This tendency requires a new approach to understanding their inheritance and predicting the types of offspring produced. A critical piece of information necessary for predicting the results of these crosses is the arrangement of the genes on the chromosomes; thus, it will be necessary to think about the relation between genes and chromosomes. A key to understanding the inheritance of linked genes is to make the conceptual connection between the genotypes in a cross and the behavior of chromosomes in meiosis.

We will begin our exploration of linkage by first comparing the inheritance of two linked genes with the inheritance of two genes that assort independently. We will then examine how crossing over, or recombination, breaks up linked genes. This knowledge of linkage and recombination will be used for predicting the results of genetic crosses in which genes are linked and for mapping genes. Later in the chapter, we will focus on physical methods of determining the chromosomal locations of genes. The final section examines variation in rates of recombination.

# 7.1 Linked Genes Do Not Assort Independently

Chapter 3 introduced Mendel's principles of segregation and independent assortment. Let's take a moment to review these two important concepts. The principle of segregation states that each individual diploid organism possesses two alleles at a locus that separate in meiosis, with one allele going into each gamete. The principle of independent assortment provides additional information about the process of segregation: it tells us that, in the process of separation, the two alleles at a locus act independently of alleles at other loci.

The independent separation of alleles results in *recombination*, the sorting of alleles into new combinations. Consider a cross between individuals homozygous for two different pairs of alleles:  $AA$   $BB \times aa$  bb. The first parent, *AA BB*, produces gametes with the alleles *A B*, and the second parent, *aa bb*, produces gametes with the alleles *a b*, resulting in  $F_1$  progeny with genotype *Aa Bb* (**Figure 7.1**).



**7.1 Recombination is the sorting of alleles into new combinations.**

Recombination means that, when one of the  $F_1$  progeny reproduces, the combination of alleles in its gametes may differ from the combinations in the gametes from its parents. In other words, the  $F_1$  may produce gametes with alleles *A b* or *a B* in addition to gametes with *A B* or *a b*.

Mendel derived his principles of segregation and independent assortment by observing the progeny of genetic crosses, but he had no idea of what biological processes produced these phenomena. In 1903, Walter Sutton proposed a biological basis for Mendel's principles, called the chromosome theory of heredity (see Chapter 3). This theory holds that genes are found on chromosomes. Let's restate Mendel's two principles in relation to the chromosome theory of heredity. The principle of segregation states that a diploid organism possesses two alleles for a trait, each of which is located at the same position, or locus, on each of the two homologous chromosomes. These chromosomes segregate in meiosis, with each gamete receiving one homolog. The principle of independent assortment states that, in meiosis, each pair of homologous chromosomes assorts independently of other homologous pairs. With this new perspective, it is easy to see that the number of chromosomes in most organisms is limited and that there are certain to be more genes than chromosomes; so some genes must be present on the same chromosome and should not assort independently. Genes located close together on the same chromosome are called **linked genes** and belong to the same **linkage group**. Linked genes travel together in meiosis, eventually arriving at the same destination (the same gamete), and are not expected to assort independently.

All of the characteristics examined by Mendel in peas did display independent assortment and, after the rediscovery of Mendel's work, the first genetic characteristics studied in other organisms also seemed to assort independently. How could genes be carried on a limited number of chromosomes and yet assort independently?

The apparent inconsistency between the principle of independent assortment and the chromosome theory of heredity soon disappeared as biologists began finding genetic characteristics that did not assort independently. One of the first cases was reported in sweet peas by William Bateson, Edith Rebecca Saunders, and Reginald C. Punnett in 1905. They crossed a homozygous strain of peas having purple flowers and long pollen grains with a homozygous strain having red flowers and round pollen grains. All the  $F_1$  had purple flowers and long pollen grains, indicating that purple was dominant over red and long was dominant over round. When they intercrossed the  $F_1$ , the resulting  $F_2$  progeny did not appear in the 9 : 3 : 3 : 1 ratio expected with independent assortment (**Figure 7.2**). An excess of  $F_2$  plants had purple flowers and long pollen or red flowers and round pollen (the parental phenotypes). Although Bateson, Saunders, and Punnett were unable to explain these results, we now know that the two loci that they examined lie close together on the same chromosome and therefore do not assort independently.



**7.2 Nonindependent assortment of flower color and pollen shape in sweet peas.**

# 7.2 Linked Genes Segregate Together and Crossing Over Produces Recombination Between Them

Genes that are close together on the same chromosome usually segregate as a unit and are therefore inherited together. However, genes occasionally switch from one homologous chromosome to the other through the process of crossing over (see Chapter 2), as illustrated



**7.3 Crossing over takes place in meiosis and is responsible for recombination.**

in **Figure 7.3**. Crossing over results in recombination; it breaks up the associations of genes that are close together on the same chromosome. Linkage and crossing over can be seen as processes that have opposite effects: linkage keeps particular genes together, and crossing over mixes them up. In Chapter 5, we considered a number of exceptions and extensions to Mendel's principles of heredity. The concept of linked genes adds a further complication to interpretations of the results of genetic crosses. However, with an understanding of how linkage affects heredity, we can analyze crosses for linked genes and successfully predict the types of progeny that will be produced.

#### Notation for Crosses with Linkage

In analyzing crosses with linked genes, we must know not only the genotypes of the individuals crossed, but also the arrangement of the genes on the chromosomes. To keep track of this arrangement, we introduce a new system of notation for presenting crosses with linked genes. Consider a cross between an individual homozygous for dominant alleles at two linked loci and another individual homozygous for recessive alleles at those loci (*AA BB* × *aa bb*). For linked genes, it's necessary to write out the specific alleles as they are arranged on each of the homologous chromosomes:

$$
\frac{A}{A} \qquad \frac{B}{B} \times \frac{a}{a} \qquad \frac{b}{b}
$$

In this notation, each line represents one of the two homologous chromosomes. Inheriting one chromosome from each parent, the  $F_1$  progeny will have the following genotype:

$$
\begin{array}{c|c}\nA & B \\
\hline\na & b\n\end{array}
$$

Here, the importance of designating the alleles on each chromosome is clear. One chromosome has the two dominant alleles *A* and *B*, whereas the homologous chromosome has the two recessive alleles *a* and *b*. The notation can be simplified by drawing only a single line, with the understanding that genes located on the same side of the line lie on the same chromosome:

$$
\begin{array}{cc}\nA & B \\
\hline\na & b\n\end{array}
$$

This notation can be simplified further by separating the alleles on each chromosome with a slash: *AB/ab*.

Remember that the two alleles at a locus are always located on different homologous chromosomes and therefore must lie on opposite sides of the line. Consequently, we would *never* write the genotypes as

$$
\begin{array}{cc}\nA & a \\
\hline\nB & b\n\end{array}
$$

because the alleles *A* and *a* can *never* be on the same chromosome. It is also important to always keep the same order of the genes on both sides of the line; thus, we should *never* write

$$
\begin{array}{cc}\nA & B \\
b & a\n\end{array}
$$

because it would imply that alleles *A* and *b* are allelic (at the same locus).

#### Complete Linkage Compared with Independent Assortment

We will first consider what happens to genes that exhibit complete linkage, meaning that they are located very close together on the same chromosome and do not exhibit crossing over. Genes are rarely completely linked but, by assuming that no crossing over occurs, we can see the effect of linkage more clearly. We will then consider what happens when genes assort independently. Finally, we will consider the results obtained if the genes are linked but exhibit some crossing over.

A testcross reveals the effects of linkage. For example, if a heterozygous individual is test-crossed with a homozygous recessive individual (*Aa Bb*  $\times$  *aa bb*), the alleles that are present in the gametes contributed by the heterozygous parent will be expressed in the phenotype of the offspring,



7.4 A testcross reveals the effects of linkage. Results of a testcross for two loci in tomatoes that determine leaf type and plant height.

because the homozygous parent could not contribute dominant alleles that might mask them. Consequently, traits that appear in the progeny reveal which alleles were transmitted by the heterozygous parent.

Consider a pair of linked genes in tomato plants. One of the genes affects the type of leaf: an allele for mottled leaves (*m*) is recessive to an allele that produces normal leaves (*M*). Nearby on the same chromosome the other gene determines the height of the plant: an allele for dwarf (*d*) is recessive to an allele for tall (*D*).

Testing for linkage can be done with a testcross, which requires a plant heterozygous for both characteristics. A geneticist might produce this heterozygous plant by crossing a variety of tomato that is homozygous for normal leaves and tall height with a variety that is homozygous for mottled leaves and dwarf height:

P 
$$
\begin{array}{c|c}\nM & D \longrightarrow & m & d \\
\hline\nM & D & \longleftarrow & m & d \\
\downarrow & & & \downarrow & \\
F_1 & & & \frac{M & D}{m & d}\n\end{array}
$$

The geneticist would then use these  $F_1$  heterozygotes in a testcross, crossing them with plants homozygous for mottled leaves and dwarf height:

$$
\frac{M}{m} \qquad \frac{D}{d} \times \frac{m}{m} \qquad \frac{d}{d}
$$

The results of this testcross are diagrammed in **Figure 7.4a**. The heterozygote produces two types of gametes: some with the *M D* chromosome and others with the *d* chromosome. Because no crossing over occurs, these gametes are the only types produced by the heterozygote. Notice that these gametes contain only combinations of alleles that were present in the original parents: either the allele for normal leaves together with the allele for tall height (*M* and *D*) or the allele for mottled leaves together with the allele for dwarf height (*m* and *d*). Gametes that contain only original combinations of alleles present in the parents are **nonrecombinant gametes**, or *parental* gametes.

The homozygous parent in the testcross produces only one type of gamete; it contains chromosome *m d* and pairs with one of the two gametes generated by the heterozygous parent (see Figure 7.4a). Two types of progeny result: half have normal leaves and are tall:

$$
\begin{array}{cc}\nM & D \\
m & d\n\end{array}
$$

and half have mottled leaves and are dwarf:

$$
\begin{array}{ccc}\nm & d \\
\hline\nm & d\n\end{array}
$$

These progeny display the original combinations of traits present in the P generation and are **nonrecombinant progeny**, or *parental* progeny. No new combinations of the two traits, such as normal leaves with dwarf or mottled leaves with tall, appear in the offspring, because the genes affecting the two traits are completely linked and are inherited together. New combinations of traits could arise only if the physical connection between *M* and *D* or between *m* and *d* were broken.

These results are distinctly different from the results that are expected when genes assort independently (**Figure 7.4b**). If the *M* and *D* loci assorted independently, the heterozygous plant (*Mm Dd*) would produce four types of gametes: two nonrecombinant gametes containing the original combinations of alleles (*M D* and *m d*) and two gametes containing new combinations of alleles (*M d* and *m D*). Gametes with new combinations of alleles are called **recombinant gametes**. With independent assortment, nonrecombinant and recombinant gametes are produced in equal proportions. These four types of gametes join with the single type of gamete produced by the homozygous parent of the testcross to produce four kinds of progeny in equal proportions (see Figure 7.4b). The progeny with new combinations of traits formed from recombinant gametes are termed **recombinant progeny**.

#### CONCEPTS

A testcross in which one of the plants is heterozygous for two completely linked genes yields two types of progeny, each type displaying one of the original combinations of traits present in the P generation. Independent assortment, in contrast, produces four types of progeny in a 1 : 1 : 1 : 1 ratio—two types of recombinant progeny and two types of nonrecombinant progeny in equal proportions.

#### Crossing Over with Linked Genes

Usually, there is some crossing over between genes that lie on the same chromosome, producing new combinations of traits. Genes that exhibit crossing over are incompletely linked. Let's see how it takes place.

**Theory** The effect of crossing over on the inheritance of two linked genes is shown in **Figure 7.5**. Crossing over, which takes place in prophase I of meiosis, is the exchange of genetic material between nonsister chromatids (see Figures 2.16 and 2.18). After a single crossover has taken place, the two chromatids that did not participate in crossing over are unchanged; gametes that receive these chromatids are nonrecombinants. The other two chromatids, which did participate in crossing over, now contain new combinations of alleles; gametes that receive these chromatids are recombinants. For each meiosis in which a single crossover takes place, then, two nonrecombinant gametes and two recombinant gametes will be produced. This result is the same as that produced by independent assortment (see Figure 7.4b); so, when crossing over between two loci takes place in every meiosis, it is impossible to determine whether the genes are on the same chromosome and crossing over took place or whether the genes are on different chromosomes.

For closely linked genes, crossing over does not take place in every meiosis. In meioses in which there is no crossing over, only nonrecombinant gametes are produced. In meioses in which there is a single crossover, half the gametes are recombinants and half are nonrecombinants (because a single crossover affects only two of the four chromatids); so the total percentage of recombinant gametes is always half the percentage of meioses in which crossing over takes place. Even if crossing over between two genes takes place in every meiosis, only 50% of the resulting gametes will be recombinants. Thus, the frequency of recombinant gametes is always half the frequency of crossing over, and the maximum proportion of recombinant gametes is 50%.

#### CONCEPTS

Linkage between genes causes them to be inherited together and reduces recombination; crossing over breaks up the associations of such genes. In a testcross for two linked genes, each crossover produces two recombinant gametes and two nonrecombinants. The frequency of recombinant gametes is half the frequency of crossing over, and the maximum frequency of recombinant gametes is 50%.

#### CONCEPT CHECK 1

For single crossovers, the frequency of recombinant gametes is half the frequency of crossing over because

a. a test cross between a homozygote and heterozygote produces  $\frac{1}{2}$  heterozygous and  $\frac{1}{2}$  homozygous progeny.



**(a) No crossing over**

**7.5 A single crossover produces half nonrecombinant gametes and half recombinant gametes.**

- b. the frequency of recombination is always 50%.
- c. each crossover takes place between only two of the four chromatids of a homologous pair.
- d. crossovers occur in about 50% of meioses.

**Application** Let's apply what we have learned about linkage and recombination to a cross between tomato plants that differ in the genes that encode leaf type and plant height. Assume now that these genes are linked and that some crossing over takes place between them. Suppose a geneticist carried out the testcross described earlier:

$$
\frac{M}{m} \qquad \frac{D}{d} \times \frac{m}{m} \qquad \frac{d}{d}
$$

When crossing over takes place in the genes for leaf type and height, two of the four gametes produced are recombinants. When there is no crossing over, all four resulting gametes are nonrecombinants. Thus, over all meioses, the majority of gametes will be nonrecombinants (**Figure 7.6a**). These gametes then unite with gametes produced by the homozygous recessive parent, which contain only the recessive alleles, resulting in mostly nonrecombinant progeny and a few recombinant progeny (**Figure 7.6b**). In this cross, we see that 55 of the testcross progeny have normal leaves and are tall and 53 have mottled leaves and are dwarf. These plants are the nonrecombinant progeny, containing the original combinations of traits that were present in the parents. Of the 123 progeny, 15 have new combinations of traits that were not seen in the parents: 8 are normal leaved

and dwarf, and 7 are mottle leaved and tall. These plants are the recombinant progeny.

The results of a cross such as the one illustrated in Figure 7.6 reveal several things. A testcross for two independently assorting genes is expected to produce a 1 : 1 : 1 : 1 phenotypic ratio in the progeny. The progeny of this cross clearly do not exhibit such a ratio; so we might suspect that the genes are not assorting independently. When linked genes undergo some crossing over, the result is mostly nonrecombinant progeny and fewer recombinant progeny. This result is what we observe among the progeny of the testcross illustrated in Figure 7.6; so we conclude that the two genes show evidence of linkage with some crossing over.

#### Calculating Recombination Frequency

The percentage of recombinant progeny produced in a cross is called the **recombination frequency**, which is calculated as follows:

recombination<sup>et</sup> = 
$$
\frac{number\ of\ recombination\ program}{total\ number\ of\ progeny} \times 100\%
$$

In the testcross shown in Figure 7.6, 15 progeny exhibit new combinations of traits; so the recombination frequency is:

$$
\frac{8+7}{55+53+8+7} \times 100\% = \frac{15}{123} \times 100\% = 12.2\%
$$

Thus, 12.2% of the progeny exhibit new combinations of traits resulting from crossing over. The recombination frequency can also be expressed as a decimal fraction (0.122). **TRY PROBLEM 15 ->** 



**7.6 Crossing over between linked genes produces nonrecombinant and recombinant offspring.** In this testcross, genes are linked and there is some crossing over.

#### Coupling and Repulsion

In crosses for linked genes, the arrangement of alleles on the homologous chromosomes is critical in determining the outcome of the cross. For example, consider the inheritance of two genes in the Australian blowfly, *Lucilia cuprina*. In this species, one locus determines the color of the thorax: a purple thorax (*p*) is recessive to the normal green thorax  $(p^+)$ . A second locus determines the color of the puparium: a black puparium (*b*) is recessive to the normal brown puparium  $(b^+)$ . The loci for thorax color and puparium color are located close together on the chromosome. Suppose we test-cross a fly that is heterozygous at both loci with a fly that is homozygous recessive at both. Because these genes are linked, there are two possible arrangements on the chromosomes of the heterozygous progeny fly. The dominant alleles for green thorax  $(p^+)$  and brown puparium  $(b<sup>+</sup>)$  might reside on one chromosome of the homologous pair, and the recessive alleles for purple thorax (*p*) and black puparium (*b*) might reside on the other homologous chromosome:

$$
\begin{array}{c|cc}\np^+ & b^+ \\
\hline\np & b\n\end{array}
$$

This arrangement, in which wild-type alleles are found on one chromosome and mutant alleles are found on the other chromosome, is referred to as the **coupling**, or **cis**, **configuration**. Alternatively, one chromosome might bear the alleles for green thorax  $(p^+)$  and black puparium  $(b)$ , and the other chromosome carries the alleles for purple thorax (*p*) and brown puparium  $(b^+)$ :

$$
\begin{array}{c|cc}\np^+ & b \\
\hline\np & b^+\n\end{array}
$$

This arrangement, in which each chromosome contains one wild-type and one mutant allele, is called the **repulsion**, or **trans**, **configuration**. Whether the alleles in the heterozygous parent are in coupling or repulsion determines which phenotypes will be most common among the progeny of a testcross.

When the alleles are in the coupling configuration, the most numerous progeny types are those with a green thorax and brown puparium and those with a purple thorax and black puparium (**Figure 7.7a**); but, when the alleles of the heterozygous parent are in repulsion, the most numerous progeny types are those with a green thorax and black puparium and those with a purple thorax and brown puparium (**Figure 7.7b**). Notice that the genotypes of the parents in Figure 7.7a and b are the same ( $p^+p b^+b \times pp bb$ ) and that the dramatic difference in the phenotypic ratios of the progeny in the two crosses results entirely from the configuration—coupling or repulsion—of the chromosomes. Knowledge of the arrangement of the alleles on the chromosomes is essential to accurately predict the outcome of crosses in which genes are linked.



#### **(a) Alleles in coupling configuration (b) Alleles in repulsion configuration**



#### CONCEPTS

In a cross, the arrangement of linked alleles on the chromosomes is critical for determining the outcome. When two wild-type alleles are on one homologous chromosome and two mutant alleles are on the other, they are in the coupling configuration; when each chromosome contains one wild-type allele and one mutant allele, the alleles are in repulsion.

#### CONCEPT CHECK 2

The following testcross produces the progeny shown: *Aa Bb* × *aa bb*  $\rightarrow$  10 *Aa Bb*, 40 *Aa bb*, 40 *aa Bb*, 10 *aa bb*. Were the genes in the *Aa Bb* parent in coupling or in repulsion?

# CONNECTING CONCEPTS

#### Relating Independent Assortment, Linkage, and Crossing Over

We have now considered three situations concerning genes at different loci. First, the genes may be located on different chromo-

somes; in this case, they exhibit independent assortment and combine randomly when gametes are formed. An individual heterozygous at two loci (*Aa Bb*) produces four types of gametes (*A B, a b, A b,* and *a B*) in equal proportions: two types of nonrecombinants and two types of recombinants, In a testcross, these gametes will result in four types of progeny in equal proportions (**Table 7.1**).

Second, the genes may be completely linked—meaning that they are on the same chromosome and lie so close together that crossing over between them is rare. In this case, the genes do not recombine. An individual heterozygous for two closely linked genes in the coupling configuration

$$
\begin{array}{cc}\nA & B \\
\hline\na & b\n\end{array}
$$

produces only the nonrecombinant gametes containing alleles *A B* or *a b*; the alleles do not assort into new combinations such as *A b* or *a B*. In a testcross, completely linked genes will produce only two types of progeny, both nonrecombinants, in equal proportions (see Table 7.1).

The third situation, incomplete linkage, is intermediate between the two extremes of independent assortment and complete linkage. Here, the genes are physically linked on the same

# **Table 7.1** Results of a testcross  $(Aa \, Bb \times aa \, bb)$  with complete linkage, independent assortment, and linkage with some crossing over



chromosome, which prevents independent assortment. However, occasional crossovers break up the linkage and allow the genes to recombine. With incomplete linkage, an individual heterozygous at two loci produces four types of gametes—two types of recombinants and two types of nonrecombinants—but the nonrecombinants are produced more frequently than the recombinants because crossing over does not take place in every meiosis. In the testcross, these gametes result in four types of progeny, with the nonrecombinants more frequent than the recombinant (see Table 7.1).

Earlier in the chapter, the term recombination was defined as the sorting of alleles into new combinations. We've now considered two types of recombination that differ in the mechanism that generates these new combinations of alleles. **Interchromosomal recombination** takes place between genes located on *different* chromosomes. It arises from independent assortment—the random segregation of chromosomes in anaphase I of meiosis—and is the kind of recombination that Mendel discovered while studying dihybrid crosses. A second type of recombination, **intrachromosomal recombination**, takes place between genes located on the *same* chromosome. This recombination arises from crossing over—the exchange of genetic material in prophase I of meiosis. Both types of recombination produce new allele combinations in the gametes; so they cannot be distinguished by examining the types of gametes produced. Nevertheless, they can often be distinguished by the *frequencies* of types of gametes: interchromosomal recombination produces 50% nonrecombinant gametes and 50% recombinant gametes, whereas intrachromosomal recombination frequently produces less than 50% recombinant gametes. However, when the genes are very far apart on the same chromosome, they assort independently, as if they were on different chromosomes. In this case, intrachromosomal recombination also produces 50% recombinant gametes. Intrachromosomal recombination of genes that lie far apart on the same chromosome and interchromosomal recombination are genetically indistinguishable.

# Evidence for the Physical Basis of Recombination

Walter Sutton's chromosome theory of inheritance, which stated that genes are physically located on chromosomes, was supported by Nettie Stevens and Edmund Wilson's discovery that sex was associated with a specific chromosome in insects (p. 75 in Chapter 4) and Calvin Bridges's demonstration that nondisjunction of X chromosomes was related to the inheritance of eye color in *Drosophila* (pp. 82–83). Further evidence for the chromosome theory of heredity came in 1931, when Harriet Creighton and Barbara McClintock (**Figure 7.8**) obtained evidence that intrachromosomal recombination was the result of physical exchange between chromosomes. Creighton and McClintock discovered a strain of corn that had an abnormal chromosome 9, containing a densely staining knob at one end and a small piece of another chromosome attached to the other end. This aberrant chromosome allowed them to visually distinguish the two members of a homologous pair.

They studied the inheritance of two traits in corn determined by genes on chromosome 9. At one locus, a dominant allele (*C*) produced colored kernels, whereas a recessive allele (*c*) produced colorless kernels. At a second, linked locus, a dominant allele (*Wx*) produced starchy kernels, whereas a recessive allele (*wx*) produced waxy kernels. Creighton and McClintock obtained a plant that was heterozygous at both loci in repulsion, with the alleles for colored and waxy on



**7.8 Barbara McClintock (left) and Harriet Creighton (right) provided evidence that genes are located on chromosomes.**  [Karl Maramorosch/Courtesy of Cold Spring Harbor Laboratory Archives.]

the aberrant chromosome and the alleles for colorless and starchy on the normal chromosome:



They crossed this heterozygous plant with a plant that was homozygous for colorless and heterozygous for waxy (with both chromosomes normal):

$$
\frac{C}{c} \qquad \frac{wx}{Wx} \times \frac{c}{c} \qquad \frac{Wx}{wx}
$$

This cross will produce different combinations of traits in the progeny, but the only way that colorless and waxy progeny can arise is through crossing over in the doubly heterozygous parent:



Note: Not all progeny genotypes are shown.

Notice that, if crossing over entails physical exchange between the chromosomes, then the colorless, waxy progeny resulting from recombination should have a chromosome with an extra piece but not a knob. Furthermore, some of the colored, starchy progeny should possess a knob but not the extra piece. This outcome is precisely what Creighton and McClintock observed, confirming the chromosomal theory of inheritance. Curt Stern provided a similar demonstration by using chromosomal markers in *Drosophila* at about the same time. We will examine the molecular basis of recombination in more detail in Chapter 12.

### Predicting the Outcomes of Crosses with Linked Genes

Knowing the arrangement of alleles on a chromosome allows us to predict the types of progeny that will result from a cross entailing linked genes and to determine which of these types will be the most numerous. Determining the *proportions* of the types of offspring requires an additional piece of information—the recombination frequency. The recombination frequency provides us with information about how often the alleles in the gametes appear in new combinations and therefore allows us to predict the proportions of offspring phenotypes that will result from a specific cross with linked genes.

In cucumbers, smooth fruit (*t*) is recessive to warty fruit (*T*) and glossy fruit (*d*) is recessive to dull fruit (*D*). Geneticists have determined that these two genes exhibit a recombination frequency of 16%. Suppose we cross a plant homozygous for warty and dull fruit with a plant homozygous for smooth and glossy fruit and then carry out a testcross by using the  $F_1$ :

$$
\frac{T}{t} \qquad \frac{D}{d} \times \frac{t}{t} \qquad \frac{d}{d}
$$

What types and proportions of progeny will result from this testcross?

Four types of gametes will be produced by the heterozygous parent, as shown in **Figure 7.9**: two types of nonrecombinant gametes  $(T \t D$  and  $t \t d$  ) and two types of recombinant gametes ( *T d* and *t D* ). The recombination frequency tells us that 16% of the gametes produced by the heterozygous parent will be recombinants. Because there are two types of recombinant gametes, each should arise with a frequency of  $16\%/2 = 8\%$ . This frequency can also be represented as a probability of 0.08. All the other gametes will be nonrecombinants; so they should arise with a frequency of  $100\% - 16\% = 84\%$ . Because there are two types of nonrecombinant gametes, each should arise with a frequency of  $84\%/2 = 42\%$  (or 0.42). The other parent in the testcross is homozygous and therefore produces only a single type of gamete ( *t d* ) with a frequency of 100% (or 1.00).

Four types of progeny result from the testcross (see Figure 7.9). The expected proportion of each type can be determined by using the multiplication rule, multiplying together the probability of each gamete. Testcross progeny with warty and dull fruit

$$
\begin{array}{cc}\nT & D \\
\hline\nt & d\n\end{array}
$$

appear with a frequency of 0.42 (the probability of inheriting a gamete with chromosome *T D* from the heterozygous parent)  $\times$  1.00 (the probability of inheriting a gamete



**7.9 The recombination frequency allows a prediction of the proportions of offspring expected for a cross entailing linked genes.**

with chromosome  $t \underline{d}$  from the recessive parent)  $= 0.42$ . The proportions of the other types of  $F_2$  progeny can be calculated in a similar manner (see Figure 7.9). This method can be used for predicting the outcome of any cross with linked genes for which the recombination frequency is known. TRY PROBLEM 17 >

#### Testing for Independent Assortment

In some crosses, the genes are obviously linked because there are clearly more nonrecombinant progeny than recombinant progeny. In other crosses, the difference between independent assortment and linkage is not so obvious. For example, suppose we did a testcross for two pairs of genes, such as  $Aa$   $Bb \times aa$   $bb$ , and observed the following numbers of progeny: 54 *Aa Bb*, 56 *aa bb*, 42 *Aa bb*, and 48 *aa Bb*. Is this outcome the 1 : 1 : 1 : 1 ratio we would expect if *A* and *B* assorted independently? Not exactly, but it's pretty close. Perhaps these genes assorted independently and chance produced the slight deviations between the observed numbers and the expected  $1:1:1:1$  ratio. Alternatively, the genes might be linked, with considerable crossing over taking place between them, and so the number of nonrecombinants is only slightly greater than the number of recombinants. How do we distinguish between the role of chance and the role of linkage in producing deviations from the results expected with independent assortment?

We encountered a similar problem in crosses in which genes were unlinked—the problem of distinguishing between deviations due to chance and those due to other factors. We addressed this problem (in Chapter 3) with the goodnessof-fit chi-square test, which helps us evaluate the likelihood that chance alone is responsible for deviations between the numbers of progeny that we observed and the numbers that we expected by applying the principles of inheritance. Here, we are interested in a different question: Is the inheritance of alleles at one locus independent of the inheritance of alleles at a second locus? If the answer to this question is yes, then the genes are assorting independently; if the answer is no, then the genes are probably linked.

A possible way to test for independent assortment is to calculate the expected probability of each progeny type, assuming independent assortment, and then use the goodness-of-fit chi-square test to evaluate whether the observed numbers deviate significantly from the expected numbers. With independent assortment, we expect  $\frac{1}{4}$  of each phenotype:  $\frac{1}{4}$  *Aa Bb*,  $\frac{1}{4}$  *aa bb*,  $\frac{1}{4}$  *Aa bb*, and  $\frac{1}{4}$  *aa Bb*. This expected probability of each genotype is based on the multiplication rule of probability, which we considered in Chapter 3. For example, if the probability of  $Aa$  is  $\frac{1}{2}$  and the probability of *Bb* is  $\frac{1}{2}$ , then the probability of *Aa Bb* is  $\frac{1}{2} \times \frac{1}{2} = \frac{1}{4}$ . In this calculation, we are making two assumptions: (1) the probability of each single-locus genotype is  $\frac{1}{2}$ , and (2) genotypes at the two loci are inherited independently  $\binom{1}{2} \times \binom{1}{2} = \binom{1}{4}$ .

One problem with this approach is that a significant chi-square value can result from a violation of either assumption. If the genes are linked, then the inheritance of genotypes at the two loci are not independent (assumption 2), and we will get a significant deviation between observed and expected numbers. But we can also get a significant deviation if the probability of each single-locus genotype is not  $\frac{1}{2}$  (assumption 1), even when the genotypes are assorting independently. We may obtain a significant deviation, for example, if individuals with one genotype have a lower probability of surviving or the penetrance of a genotype is not 100%. We could test both assumptions by conducting a series of chi-square tests, first testing the inheritance of genotypes at each locus separately (assumption 1) and then testing for independent assortment (assumption 2). However, a faster method is to test for independence in genotypes with a *chi-square test of independence*.

**The chi-square test of independence** The chisquare test of independence allows us to evaluate whether the segregation of alleles at one locus is independent of the segregation of alleles at another locus, without making any assumption about the probability of single-locus genotypes. To illustrate this analysis, we will examine results from a cross between German cockroaches, in which yellow body (*y*) is recessive to brown body  $(y^+)$  and curved wings  $(cv)$  are recessive to straight wings  $(cv^+)$ . A testcross  $(y^+y c v^+ c v \times yy c v c v)$ produced the progeny shown in **Figure 7.10a**.

To carry out the chi-square test of independence, we first construct a table of the observed numbers of progeny, somewhat like a Punnett square, except, here, we put the genotypes that result from the segregation of alleles at one locus along the top and the genotypes that result from the segregation of alleles at the other locus along the side (**Figure 7.10b**). Next, we compute the total for each row, the total for each column, and the grand total (the sum of all row totals or the sum of all column totals, which should be the same). These totals will be used to compute the expected values for the chi-square test of independence.

Now, we compute the values expected if the segregation of alleles at the *y* locus is independent of the segregation of alleles at the *cv* locus. If the segregation of alleles at each locus is independent, then the proportion of progeny with *y*<sup>+</sup>*y* and *yy* genotypes should be the same for cockroaches with genotype  $cv^+cv$  and for cockroaches with genotype *cvcv*. The converse is also true; the proportions of progeny with  $c\nu^+c\nu$  and

**7.10 A chi-square test of independence can be used to determine if genes at two loci are assorting independently.**



*cvcv* genotypes should be the same for cockroaches with genotype  $y^+y$  and for cockroaches with genotype  $yy$ . With the assumption that the alleles at the two loci segregate independently, the expected number for each cell of the table can be computed by using the following formula:

$$
expected number = \frac{row total \times column total}{grand total}
$$

For the cell of the table corresponding to genotype  $y^+$ *y*  $cv^+$ *cv* (the upper-left-hand cell of the table in Figure 7.10b) the expected number is:

$$
\frac{96 \text{ (row total)} \times 91 \text{ (column total)}}{201 \text{ (grand total)}} = \frac{8736}{201} = 43.46
$$

With the use of this method, the expected numbers for each cell are given in **Figure 7.10c**.

We now calculate a chi-square value by using the same formula that we used for the goodness-of-fit chi-square test in Chapter 3:

$$
x^2 = \sum \frac{(observed - expected)^2}{expected}
$$

Recall that  $\Sigma$  means "sum," and that we are adding together the (observed  $-$  expected)<sup>2</sup>/expected value for each type of progeny. With the observed and expected numbers of cockroaches from the testcross, the calculated chi-square value is 30.73 (**Figure 7.10d**).

To determine the probability associated with this chisquare value, we need the degrees of freedom. Recall from Chapter 3 that the degrees of freedom are the number of ways in which the observed classes are free to vary from the expected values. In general, for the chi-square test of independence, the degrees of freedom equal the number of rows in the table minus 1 multiplied by the number of columns in the table minus 1 (**Figure 7.10e**), or

$$
df =
$$
(number of rows – 1) × (number of columns – 1)

In our example, there were two rows and two columns, and so the degrees of freedom are:

$$
df = (2 - 1) \times (2 - 1) = 1 \times 1 = 1
$$

Therefore, our calculated chi-square value is 30.73, with 1 degree of freedom. We can use Table 3.5 to find the associated probability. Looking at Table 3.5, we find that our calculated chi-square value is larger than the largest chi-square value given for 1 degree of freedom, which has a probability of 0.005. Thus, our calculated chi-square value has a probability less than 0.005. This very small probability indicates that the genotypes are not in the proportions that we would expect if independent assortment were taking place. Our conclusion, then, is that these genes are not assorting independently and must be linked. As is the case for the goodness-of-fit chi-square test, geneticists generally consider that any chi-square value for the test of independence with a

probability less than 0.05 is significantly different from the expected values and is therefore evidence that the genes are not assorting independently. **TRY PROBLEM 16** 

#### Gene Mapping with Recombination **Frequencies**

Thomas Hunt Morgan and his students developed the idea that physical distances between genes on a chromosome are related to the rates of recombination. They hypothesized that crossover events occur more or less at random up and down the chromosome and that two genes that lie far apart are more likely to undergo a crossover than are two genes that lie close together. They proposed that recombination frequencies could provide a convenient way to determine the order of genes along a chromosome and would give estimates of the relative distances between the genes. Chromosome maps calculated by using the genetic phenomenon of recombination are called **genetic maps**. In contrast, chromosome maps calculated by using physical distances along the chromosome (often expressed as numbers of base pairs) are called **physical maps**.

Distances on genetic maps are measured in **map units** (abbreviated m.u.); one map unit equals 1% recombination. Map units are also called **centiMorgans** (cM), in honor of Thomas Hunt Morgan; 100 centiMorgans equals 1 **Morgan**. Genetic distances measured with recombination rates are approximately additive: if the distance from gene *A* to gene *B* is 5 m.u., the distance from gene *B* to gene *C* is 10 m.u., and the distance from gene *A* to gene *C* is 15 m.u., then gene *B* must be located between genes *A* and *C*. On the basis of the map distances just given, we can draw a simple genetic map for genes *A, B,* and *C*, as shown here:



We could just as plausibly draw this map with *C* on the left and *A* on the right:



Both maps are correct and equivalent because, with information about the relative positions of only three genes, the most that we can determine is which gene lies in the middle. If we obtained distances to an additional gene, then we could position *A* and *C* relative to that gene. An additional gene *D*, examined through genetic crosses, might yield the following recombination frequencies:





**7.11 A two-strand double crossover between two linked genes produces only nonrecombinant gametes.**

Notice that *C* and *D* exhibit the greatest amount of recombination; therefore, *C* and *D* must be farthest apart, with genes *A* and *B* between them. Using the recombination frequencies and remembering that  $1 \text{ m.u.} = 1\%$  recombination, we can now add *D* to our map:



By doing a series of crosses between pairs of genes, we can construct genetic maps showing the linkage arrangements of a number of genes.

Two points should be emphasized about constructing chromosome maps from recombination frequencies. First, recall that we cannot distinguish between genes on different chromosomes and genes located far apart on the same chromosome. If genes exhibit 50% recombination, the most that can be said about them is that they belong to different groups of linked genes (different linkage groups), either on different chromosomes or far apart on the same chromosome.

The second point is that a testcross for two genes that are far apart on the same chromosome tends to underestimate the true physical distance, because the cross does not reveal double crossovers that might take place between the two genes (**Figure 7.11**). A double crossover arises when two separate crossover events take place between two loci. (For now, we will consider only double crossovers that occur between two of the four chromatids of a homologus pair—a two-strand double crossover. Double crossovers that take place among three and four chromatids will be considered later.) Whereas a single crossover produces combinations of alleles that were not present on the original parental chromosomes, a second crossover

between the same two genes reverses the effects of the first, thus restoring the original parental combination of alleles (see Figure 7.11). We therefore cannot distinguish between the progeny produced by two-strand double crossovers and the progeny produced when there is no crossing over at all. As we shall see in the next section, we can detect double crossovers if we examine a third gene that lies between the two crossovers. Because double crossovers between two genes go undetected, map distances will be underestimated whenever double crossovers take place. Double crossovers are more frequent between genes that are far apart; therefore genetic maps based on short distances are usually more accurate than those based on longer distances.

#### **CONCEPTS**

A genetic map provides the order of the genes on a chromosome and the approximate distances from one gene to another based on recombination frequencies. In genetic maps, 1% recombination equals 1 map unit, or 1 centiMorgan. Double crossovers between two genes go undetected; so map distances between distant genes tend to underestimate genetic distances.

#### CONCEPT CHECK 3

How does a genetic map differ from a physical map?

### Constructing a Genetic Map with the Use of Two-Point Testcrosses

Genetic maps can be constructed by conducting a series of testcrosses. In each testcross, one of the parents is heterozygous for a different pair of genes, and recombination frequencies are calculated between pairs of genes. A testcross between two genes is called a **two-point testcross** or a twopoint cross for short. Suppose that we carried out a series of two-point crosses for four genes, *a*, *b*, *c,* and *d,* and obtained the following recombination frequencies:



We can begin constructing a genetic map for these genes by considering the recombination frequencies for each pair of genes. The recombination frequency between *a* and *b* is 50%, which is the recombination frequency expected with independent assortment. Therefore, genes *a* and *b* may either be on different chromosomes or be very far apart on the same chromosome; so we will place them in different linkage groups with the understanding that they may or may not be on the same chromosome:

Linkage group 1



The recombination frequency between *a* and *c* is 50%, indicating that they, too, are in different linkage groups. The recombination frequency between *b* and *c* is 20%; so these genes are linked and separated by 20 map units:

Linkage group 1



The recombination frequency between *a* and *d* is 50%, indicating that these genes belong to different linkage groups, whereas genes *b* and *d* are linked, with a recombination frequency of 10%. To decide whether gene *d* is 10 m.u. to the left or to the right of gene *b*, we must consult the *c*-to*d* distance. If gene *d* is 10 m.u. to the left of gene *b,* then the distance between *d* and *c* should be approximately the sum of the distance between b and c and between c and d: 20 m.u. + 10 m.u. = 30 m.u. If, on the other hand, gene *d* lies to the right of gene *b,* then the distance between gene *d* and gene *c* will be much shorter, approximately 20 m.u.  $- 10$  m.u.  $=$ 

10 m.u. The summed distances will be only approximate because any double crossovers between the two genes will be missed and the map distance will be underestimated.

By examining the recombination frequency between *c* and *d*, we can distinguish between these two possibilities. The recombination frequency between *c* and *d* is 28%; so gene *d* must lie to the left of gene *b*. Notice that the sum of the recombination between *d* and *b* (10%) and between *b* and *c* (20%) is greater than the recombination between *d* and *c* (28%). As already discussed, this discrepancy arises because double crossovers between the two outer genes go undetected, causing an underestimation of the true map distance. The genetic map of these genes is now complete:





# 7.3 A Three-Point Testcross Can Be Used to Map Three Linked Genes

Genetic maps can be constructed from a series of testcrosses for pairs of genes, but this approach is not particularly efficient, because numerous two-point crosses must be carried out to establish the order of the genes and because double crossovers are missed. A more efficient mapping technique is a testcross for three genes—a **three-point testcross**, or three-point cross. With a three-point cross, the order of the three genes can be established in a single set of progeny and some double crossovers can usually be detected, providing more-accurate map distances.

Consider what happens when crossing over takes place among three hypothetical linked genes. **Figure 7.12** illustrates a pair of homologous chromosomes of an individual that is heterozygous at three loci (*Aa Bb Cc*). Notice that the genes are in the coupling configuration; that is, all the dominant alleles are on one chromosome ( *A B C* ) and all the recessive alleles are on the other chromosome a b c). Three types of crossover events can take place between these three genes: two types of single crossovers (see Figure 7.12a and b) and a double crossover (see Figure 7.12c). In each type of crossover, two of the resulting chromosomes are recombinants and two are nonrecombinants.

Notice that, in the recombinant chromosomes resulting from the double crossover, the outer two alleles are the



**among three linked loci.**

same as in the nonrecombinants, but the middle allele is different. This result provides us with an important clue about the order of the genes. In progeny that result from a double crossover, only the middle allele should differ from the alleles present in the nonrecombinant progeny.

### Constructing a Genetic Map with the Three-Point Testcross

To examine gene mapping with a three-point testcross, we will consider three recessive mutations in the fruit fly *Drosophila melanogaster*. In this species, scarlet eyes (*st*) are recessive to wild-type red eyes  $(st^+)$ , ebony body color  $(e)$ is recessive to wild-type gray body color  $(e^+)$ , and spineless (*ss*)—that is, the presence of small bristles—is recessive to wild-type normal bristles  $(s<sup>+</sup>)$ . The loci encoding these three characteristics are linked and located on chromosome 3.

We will refer to these three loci as *st*, *e*, and *ss*, but keep in mind that either the recessive alleles (*st*, *e*, and *ss*) or the dominant alleles (*st*<sup>+</sup>, *e* <sup>+</sup>, and *ss*<sup>+</sup>) may be present at each locus. So, when we say that there are 10 m.u. between *st* and *ss*, we mean that there are 10 m.u. between the loci at which mutations *st* and *ss* occur; we could just as easily say that there are 10 m.u. between  $st^+$  and  $ss^+$ .

To map these genes, we need to determine their order on the chromosome and the genetic distances between them. First, we must set up a three-point testcross, a cross between a fly heterozygous at all three loci and a fly homozygous for recessive alleles at all three loci. To produce flies heterozygous for all three loci, we might cross a stock of flies that are homozygous for wild-type alleles at all three

loci with flies that are homozygous for recessive alleles at all three loci:

P  
\n
$$
\begin{array}{c}\n \text{St}^+ e^+ \text{St}^+ \\
 \text{St}^+ e^+ \text{St}^+ \\
 \text{St}^+ e^+ \text{St}^+ \\
 \downarrow \\
 \text{St}^+ e^+ \text{St}^+ \\
 \text{St}^+ e^+ \text{St}^+ \\
 \end{array}
$$

The order of the genes has been arbitrarily assigned because, at this point, we do not know which is the middle gene. Additionally, the alleles in these heterozygotes are in coupling configuration (because all the wild-type dominant alleles were inherited from one parent and all the recessive mutations from the other parent), although the testcross can also be done with alleles in repulsion.

In the three-point testcross, we cross the  $F_1$  heterozygotes with flies that are homozygous for all three recessive mutations. In many organisms, it makes no difference whether the heterozygous parent in the testcross is male or female (provided that the genes are autosomal) but, in *Drosophila*, no crossing over takes place in males. Because crossing over in the heterozygous parent is essential for determining recombination frequencies, the heterozygous flies in our testcross must be female. So we mate female  $F_1$ flies that are heterozygous for all three traits with male flies that are homozygous for all the recessive traits:

$$
\frac{st^+}{st} \quad \frac{e^+}{es} \quad \text{Female} \quad \times \quad \frac{st}{st} \quad \frac{e}{es} \quad \text{Male}
$$







The progeny produced from this cross are listed in **Figure 7.13**. For each locus, two classes of progeny are produced: progeny that are heterozygous, displaying the dominant trait, and progeny that are homozygous, displaying the recessive trait. With two classes of progeny possible for each of the three loci, there will be  $2^3 = 8$  classes of phenotypes possible in the progeny. In this example, all eight phenotypic classes are present but, in some three-point crosses, one or more of the phenotypes may be missing if the number of progeny is limited. Nevertheless, the absence of a particular class can provide important information about which combination of traits is least frequent and, ultimately, about the order of the genes, as we will see.

To map the genes, we need information about where and how often crossing over has taken place. In the homozygous recessive parent, the two alleles at each locus are the same, and so crossing over will have no effect on the types of gametes produced; with or without crossing over, all gametes from this parent have a chromosome with three recessive alleles ( *st e ss* ). In contrast, the heterozygous parent has different alleles on its two chromosomes, and so crossing over can be detected. The information that we need for mapping, therefore, comes entirely from the gametes produced by the heterozygous parent. Because chromosomes contributed by the homozygous parent carry only recessive alleles, whatever alleles are present on the chromosome contributed by the heterozygous parent will be expressed in the progeny.

As a shortcut, we often do not write out the complete genotypes of the testcross progeny, listing instead only the alleles expressed in the phenotype, which are the alleles inherited from the heterozygous parent. This convention is used in the discussion that follows.

#### **CONCEPTS**

To map genes, information about the location and number of crossovers in the gametes that produced the progeny of a cross is needed. An efficient way to obtain this information is to use a three-point testcross, in which an individual heterozygous at three linked loci is crossed with an individual that is homozygous recessive at the three loci.

#### CONCEPT CHECK 4

Write the genotypes of all recombinant and nonrecombinant progeny expected from the following three-point cross:



**Determining the gene order** The first task in mapping the genes is to determine their order on the chromosome. In Figure 7.13, we arbitrarily listed the loci in the order *st*, *e*, *ss*, but we had no way of knowing which of the three loci was between the other two. We can now identify the middle locus by examining the double-crossover progeny.

First, determine which progeny are the nonrecombinants; they will be the two most-numerous classes of progeny. (Even if crossing over takes place in every meiosis, the nonrecombinants will constitute at least 50% of the progeny.) Among the progeny of the testcross in Figure 7.13,

the most numerous are those with all three dominant traits<br>( $st^+$   $e^+$   $ss^+$ ) and those with all three recessive  $ss^{+}$  ) and those with all three recessive traits ( *st e ss* ).

Next, identify the double-crossover progeny. These progeny should always have the two least-numerous phenotypes, because the probability of a double crossover is always less than the probability of a single crossover. The least-common progeny among those listed in Figure 7.13 are progeny with spineless<br>bristles ( $st^+$   $e^+$   $ss$ ) and progeny with scarlet eves bristles  $(s t^+ e^+ - s s)$  and progeny with scarlet eyes and ebony body ( $\frac{st}{e}$  e  $\frac{ss^+}{ss^+}$ ); so they are the doublecrossover progeny.

Three orders of genes on the chromosome are possible: the eye-color locus could be in the middle ( *e st ss* ), the body-color locus could be in the middle ( *st e ss* ), or the bristle locus could be in the middle ( *st ss e* ). To determine which gene is in the middle, we can draw the chromosomes of the heterozygous parent with all three possible gene orders and then see if a double crossover produces the combination of genes observed in the double-crossover progeny. The three possible gene orders and the types of progeny produced by their double crossovers are:



The only gene order that produces chromosomes with the set of alleles observed in the least-numerous progeny or double crossovers ( $st^+$  e<sup>+</sup> ss</sup> and  $\frac{st}{s}$  e  $\frac{ss^+}{s}$  in Figure 7.13) is the one in which the *ss* locus for bristles lies in the middle (gene-order 3). Therefore, this order ( *st ss e* ) must be the correct sequence of genes on the chromosome.

With a little practice, we can quickly determine which locus is in the middle without writing out all the gene orders. The phenotypes of the progeny are expressions of the alleles inherited from the heterozygous parent. Recall that, when we looked at the results of double crossovers (see Figure 7.12), only the alleles at the middle locus differed from the nonrecombinants. If we compare the nonrecombinant progeny with double-crossover progeny, they should differ only in alleles of the middle locus (**Table 7.2**).

Let's compare the alleles in the double-crossover prog-<br>eny  $\frac{st^+}{s^+}$   $\frac{e^+}{s^+}$  with those in the nonrecombinant eny  $\frac{st^+}{s^+}$   $\frac{e^+}{e^+}$  *ss* with those in the nonrecombinant progeny  $\frac{st^+}{e^+}$   $\frac{es^+}{s^+}$ . We see that both have an allele  $e^+$  *ss*<sup>+</sup> . We see that both have an allele

#### **Table 7.2** Steps in determining gene order in a three-point cross

- 1. Identify the nonrecombinant progeny (two mostnumerous phenotypes).
- 2. Indentify the double-crossover progeny (two leastnumerous phenotypes).
- 3. Compare the phenotype of double-crossover progeny with the phenotype of nonrecombinant progeny. They should be alike in two characteristics and differ in one.
- 4. The characteristic that differs between the double crossover and the nonrecombinant progeny is encoded by the middle gene.

for red eyes  $(st^+)$  and both have an allele for gray body  $(e^+)$ , but the nonrecombinants have an allele for normal bristles  $(s<sup>+</sup>)$ , whereas the double crossovers have an allele for spineless bristles (*ss*). Because the bristle locus is the only one that differs, it must lie in the middle. We would obtain the same results if we compared the other class of double-crossover progeny  $(st \, e \, s\bar{s}^+)$  with other nonrecombinant progeny ( *st e ss* ). Again, the only locus that differs is the one for bristles. Don't forget that the nonrecombinants and the double crossovers should differ at only one locus; if they differ at two loci, the wrong classes of progeny are being compared.

#### CONCEPTS

To determine the middle locus in a three-point cross, compare the double-crossover progeny with the nonrecombinant progeny. The double crossovers will be the two least-common classes of phenotypes; the nonrecombinants will be the two most-common classes of phenotypes. The double-crossover progeny should have the same alleles as the nonrecombinant types at two loci and different alleles at the locus in the middle.

#### CONCEPT CHECK 5

A three-point test cross is carried out between three linked genes. The resulting nonrecombinant progeny are  $s^+$   $r^+$   $c^+$  and  $s$   $r$   $c$  and the double-crossover progeny are *s*  $r c^{+}$  and  $s^{+} r^{+}$  c. Which is the middle locus?

**Determining the locations of crossovers** When we know the correct order of the loci on the chromosome, we should rewrite the phenotypes of the testcross progeny in Figure 7.13 with the alleles in the correct order so that we can determine where crossovers have taken place (**Figure 7.14**).

Among the eight classes of progeny, we have already identified two classes as nonrecombinants ( $St^+$  ss<sup>+</sup> e<sup>+</sup> and *st ss e* ) and two classes as double crossovers  $\left(\begin{array}{ccc} st^+ & ss & e^+ \end{array}\right)$  and  $\left(\begin{array}{ccc} st^- & ss & e^- \end{array}\right)$ . The other



**7.14 Writing the results of a three-point testcross with the loci in the correct order allows the locations of crossovers to be determined.** These results are from the testcross illustrated in Figure 7.13, with the loci shown in the correct order. The location of a crossover is indicated by a slash (/). The sex of the progeny flies has been designated arbitrarily.

four classes include progeny that resulted from a chromosome that underwent a single crossover: two underwent single crossovers between *st* and *ss*, and two underwent single crossovers between *ss* and *e*.

To determine where the crossovers took place in these progeny, compare the alleles found in the single-crossover progeny with those found in the nonrecombinants, just as we did for the double crossovers. For example, consider progeny with chromosome  $st^+$  ss e. The first allele (st<sup>+</sup>) came from the nonrecombinant chromosome  $st^+$  *ss<sup>+</sup>*  $e^+$  and the other two alleles (*ss* and *e*) must have come from the other nonrecombinant chromosome *st ss e* through crossing over:



This same crossover also produces the  $st$   $ss^+$ progeny.

This method can also be used to determine the location of crossing over in the other two types of single-crossover progeny. Crossing over between *ss* and *e* produces  $st^+$   $ss^+$  e and  $st$   $ss$   $e^+$  chromosomes:



We now know the locations of all the crossovers; their locations are marked with a slash in Figure 7.14.

**Calculating the recombination frequencies** Next, we can determine the map distances, which are based on the frequencies of recombination. We calculate recombination frequency by adding up all of the recombinant progeny, dividing this number by the total number of progeny from the cross, and multiplying the number obtained by 100%. To determine the map distances accurately, we must include all crossovers (both single and double) that take place between two genes.

Recombinant progeny that possess a chromosome that underwent crossing over between the eye-color locus (*st*) and the bristle locus (*ss*) include the single crossovers  $(\underline{st}^+$  / ss e and  $\underline{st}$  / ss<sup>+</sup> e<sup>+</sup> ) and the two double crossovers  $(st^+$  / ss /  $e^+$  and *st* /  $ss^+$  / e<sub>)</sub>; see Figure 7.14. There are a total of 755 progeny; so the recombination frequency between *ss* and *st* is:

*st*−ss recombination frequency =

$$
\frac{(50+52+5+3)}{755} \times 100\% = 14.6\%
$$

The distance between the *st* and *ss* loci can be expressed as 14.6 m.u.



**7.15** *Drosophila melanogaster* **has four linkage groups corresponding to its four pairs of chromosomes.** Distances between genes within a linkage group are in map units.

The map distance between the bristle locus (*ss*) and the body locus (*e*) is determined in the same manner. The recombinant progeny that possess a crossover between *ss* and *e* are the single crossovers  $st^+$  ss<sup>+</sup> / e<sub> and st ss / e<sup>+</sup></sub> and the double crossovers <u>st / ss / e<sup>+</sup></u> *st* / *ss*<sup>+</sup> / *e* . The recombination frequency is:

*ss*−*e* recombination frequency =

$$
\frac{(43+41+5+3)}{755} \times 100\% = 12.2\%
$$

Thus, the map distance between *ss* and *e* is 12.2 m.u.

Finally, calculate the map distance between the outer two loci, *st* and *e*. This map distance can be obtained by summing the map distances between *st* and *ss* and between *ss* and *e* (14.6 m.u. + 12.2 m.u. = 26.8 m.u.). We can now use the map distances to draw a map of the three genes on the chromosome:



A genetic map of *D. melanogaster* is illustrated in **Figure 7.15**.

**Interference and the coefficient of coincidence** Map distances give us information not only about the distances

that separate genes, but also about the proportions of recombinant and nonrecombinant gametes that will be produced in a cross. For example, knowing that genes *st* and *ss* on the third chromosome of *D. melanogaster* are separated by a distance of 14.6 m.u. tells us that 14.6% of the gametes produced by a fly heterozygous at these two loci will be recombinants. Similarly, 12.2% of the gametes from a fly heterozygous for *ss* and *e* will be recombinants.

Theoretically, we should be able to calculate the proportion of double-recombinant gametes by using the multiplication rule of probability (see Chapter 3), which states that the probability of two independent events occurring together is calculated by multiplying the probabilities of the independent events. Applying this principle, we should find that the proportion (probability) of gametes with double crossovers between *st* and *e* is equal to the probability of recombination between *st* and *ss* multiplied by the probability of recombination between *ss* and  $e$ , or 0.146  $\times$  0.122  $= 0.0178$ . Multiplying this probability by the total number of progeny gives us the *expected* number of double-crossover progeny from the cross:  $0.0178 \times 755 = 13.4$ . Only 8 double crossovers—considerably fewer than the 13 expected—were observed in the progeny of the cross (see Figure 7.14).

This phenomenon is common in eukaryotic organisms. The calculation assumes that each crossover event is independent and that the occurrence of one crossover does not influence the occurrence of another. But crossovers are frequently *not* independent events: the occurrence of one crossover tends to inhibit additional crossovers in the same region of the chromosome, and so double crossovers are less frequent than expected.

The degree to which one crossover interferes with additional crossovers in the same region is termed the **interference**. To calculate the interference, we first determine the **coefficient of coincidence**, which is the ratio of observed double crossovers to expected double crossovers:

coefficient of coincidence =

number of observed double crossovers number of expected double crossovers

For the loci that we mapped on the third chromosome of *D. melanogaster* (see Figure 7.14), we find that the

coefficient of coincidence =

$$
\frac{5+3}{0.146 \times 0.122 \times 755} = \frac{8}{13.4} = 0.6
$$

which indicates that we are actually observing only 60% of the double crossovers that we expected on the basis of the single-crossover frequencies. The interference is calculated as

 $interference = 1 - coefficient of coincidence$ 

So the interference for our three-point cross is:

interference  $= 1 - 0.6 = 0.4$ 

This value of interference tells us that 40% of the double-crossover progeny expected will not be observed, because of interference. When interference is complete and no double-crossover progeny are observed, the coefficient of coincidence is 0 and the interference is 1.

Sometimes a crossover *increases* the probability of another crossover taking place nearby and we see *more* double-crossover progeny than expected. In this case, the coefficient of coincidence is greater than 1 and the interference is negative. **TRY PROBLEM 28**

#### CONCEPTS

The coefficient of coincidence equals the number of double crossovers observed divided by the number of double crossovers expected on the basis of the single-crossover frequencies. The interference equals 1 − the coefficient of coincidence; it indicates the degree to which one crossover interferes with additional crossovers.

#### CONCEPT CHECK 6

In analyzing the results of a three-point testcross, a student determines that the interference is −0.23. What does this negative interference value indicate?

- a. Fewer double crossovers took place than expected on the basis of single-crossover frequencies.
- b. More double crossovers took place than expected on the basis of single-crossover frequencies.
- c. Fewer single crossovers took place than expected.
- d. A crossover in one region interferes with additional crossovers in the same region.

#### CONNECTING CONCEPTS

#### Stepping Through the Three-Point Cross

We have now examined the three-point cross in considerable detail and have seen how the information derived from the cross can be used to map a series of three linked genes. Let's briefly review the steps required to map genes from a three-point cross.

- **1. Write out the phenotypes and numbers of progeny produced in the three-point cross.** The progeny phenotypes will be easier to interpret if you use allelic symbols for the traits (such as *st*<sup>+</sup> *e*<sup>+</sup> *ss*).
- **2. Write out the genotypes of the original parents used to produce the triply heterozygous individual** in the testcross and, if known, the arrangement (coupling or repulsion) of the alleles on their chromosomes.
- **3. Determine which phenotypic classes among the progeny are the nonrecombinants and which are the double crossovers.** The nonrecombinants will be the two most-

common phenotypes; double crossovers will be the two least-common phenotypes.

- **4. Determine which locus lies in the middle.** Compare the alleles present in the double crossovers with those present in the nonrecombinants; each class of double crossovers should be like one of the nonrecombinants for two loci and should differ for one locus. The locus that differs is the middle one.
- **5. Rewrite the phenotypes with the genes in correct order.**
- **6. Determine where crossovers must have taken place to give rise to the progeny phenotypes.** To do so, compare each phenotype with the phenotype of the nonrecombinant progeny.
- **7. Determine the recombination frequencies.** Add the numbers of the progeny that possess a chromosome with a crossover between a pair of loci. Add the double crossovers to this number. Divide this sum by the total number of progeny from the cross, and multiply by 100%; the result is the recombination frequency between the loci, which is the same as the map distance.
- **8. Draw a map of the three loci.** Indicate which locus lies in the middle, and indicate the distances between them.
- **9. Determine the coefficient of coincidence and the interference.** The coefficient of coincidence is the number of observed double-crossover progeny divided by the number of expected double-crossover progeny. The expected number can be obtained by multiplying the product of the two singlerecombination probabilities by the total number of progeny in the cross. **TRY PROBLEM 31 ->**

# Worked Problem

In *D. melanogaster*, cherub wings (*ch*), black body (*b*), and cinnabar eyes (*cn*) result from recessive alleles that are all located on chromosome 2. A homozygous wild-type fly was mated with a cherub, black, and cinnabar fly, and the resulting  $F_1$  females were test-crossed with cherub, black, and cinnabar males. The following progeny were produced from the testcross:



**a.** Determine the linear order of the genes on the chromosome (which gene is in the middle?).

**b.** Calculate the recombinant distances between the three loci.

**c.** Determine the coefficient of coincidence and the interference for these three loci.

#### • Solution

**a.** We can represent the crosses in this problem as follows:

P 
$$
\frac{ch^{+} b^{+} cn^{+}}{ch^{+} b^{+} cn^{+}} \times \frac{ch}{ch} \frac{b}{b} cn
$$
  
\nF<sub>1</sub> 
$$
\frac{ch^{+} b^{+} cn^{+}}{ch} \frac{ch}{b} cn
$$
  
\nTestcross 
$$
\frac{ch^{+} b^{+} cn^{+}}{ch} \times \frac{ch}{ch} \frac{b}{b} cn
$$

Note that we do not know, at this point, the order of the genes; we have arbitrarily put *b* in the middle.

The next step is to determine which of the testcross progeny are nonrecombinants and which are double crossovers. The nonrecombinants should be the most-frequent phenotype; so they must be the progeny with phenotypes encoded by  $c h^+ b^+ c n^+$  and  $c h b c n$ . These genotypes are consistent with the genotypes of the parents, given earlier. The double crossovers are the leastfrequent phenotypes and are encoded by *ch*<sup>+</sup> *b*<sup>+</sup> *cn* and *ch b cn*<sup>+</sup> .

We can determine the gene order by comparing the alleles present in the double crossovers with those present in the nonrecombinants. The double-crossover progeny should be like one of the nonrecombinants at two loci and unlike it at one locus; the allele that differs should be in the middle. Compare the double-crossover progeny  $ch\ b\ cn^+$  with the nonrecombinant *ch b cn*. Both have cherub wings (*ch*) and black body (*b*), but the double-crossover progeny have wildtype eyes  $(cn^+)$ , whereas the nonrecombinants have cinnabar eyes (*cn*). The locus that determines cinnabar eyes must be in the middle.

**b.** To calculate the recombination frequencies among the genes, we first write the phenotypes of the progeny with the genes encoding them in the correct order. We have already identified the nonrecombinant and double-crossover progeny; so the other four progeny types must have resulted from single crossovers. To determine *where* single crossovers took place, we compare the alleles found in the single-crossover progeny with those in the nonrecombinants. Crossing over must have taken place where the alleles switch from those found in one nonrecombinant to those found in the other nonrecombinant. The locations of the crossovers are indicated with a slash:



Next, we determine the recombination frequencies and draw a genetic map:

*ch-cn* recombination frequency =

$$
\frac{40+4+41+5}{1800} \times 100\% = 5\%
$$

*cn-b* recombination frequency =

$$
\frac{105+4+102+5}{1800} \times 100\% = 12\%
$$

*ch-b* map distance = 
$$
5\% + 12\% = 17\%
$$



**c.** The coefficient of coincidence is the number of observed double crossovers divided by the number of expected double crossovers. The number of expected double crossovers is obtained by multiplying the probability of a crossover between *ch* and *cn* (0.05)  $\times$  the probability of a crossover between *cn* and  $b(0.12) \times$  the total number of progeny in the cross (1800):

coefficient of coincidence 
$$
=
$$
  $\frac{4+5}{0.05 \times 0.12 \times 1800} = 0.83$ 

Finally, the interference is equal to  $1 -$  the coefficient of coincidence:

$$
interference = 1 - 0.83 = 0.17
$$

To increase your skill with three-point crosses, try working Problem 29 at the end of this chapter.

#### Effect of Multiple Crossovers

So far, we have examined the effects of double crossovers taking place between only two of the four chromatids (strands) of a homologous pair. These crossovers are called two-strand crossovers. Double crossovers including three and even four of the chromatids of a homologous pair also may take place (**Figure 7.16**). If we examine only the alleles at loci on either side of both crossover events, two-strand double crossovers result in no new combinations of alleles, and no recombinant gametes are produced (see Figure 7.16). Three-strand double crossovers result in two of the four gametes being recombinant, and four-strand double crossovers result in all four gametes being recombinant. Thus, two-strand double crossovers produce 0% recombination, three-strand double crossovers produce 50% recombination, and four-strand double crossovers produce 100% recombination. The overall result is that all types of double crossovers, taken together, produce an average of 50% recombinant progeny.

As we have seen, two-strand double crossover cause alleles on either side of the crossovers to remain the same and produce no recombinant progeny. Three-strand and four-strand crossovers produce recombinant progeny, but these progeny are the same types produced by single crossovers. Consequently, some multiple crossovers go unde-



**7.16 Results of two-, three-, and four-strand double crossovers on recombination between two genes.**

detectable recombinants



**7.17 Percent recombination underestimates the true physical distance between genes at higher map distances.**

tected when the progeny of a genetic cross are observed. Therefore, map distances based on recombination rates will underestimate the true physical distances between genes, because some multiple crossovers are not detected among the progeny of a cross. When genes are very close together, multiple crossovers are unlikely, and the distances based on recombination rates accurately correspond to the physical distances on the chromosome. But, as the distance between genes increases, more multiple crossovers are likely, and the discrepancy between genetic distances (based on recombination rates) and physical distances increases. To correct for this discrepancy, geneticists have developed mathematical **mapping functions**, which relate recombination frequencies to actual physical distances between genes (**Figure 7.17**). Most of these functions are based on the Poisson distribution, which predicts the probability of multiple rare events. With the use of such mapping functions, map distances based on recombination rates can be more accurately estimated.

#### Mapping Human Genes

Efforts in mapping human genes are hampered by the inability to perform desired crosses and the small number of progeny in most human families. Geneticists are restricted to analyses of pedigrees, which are often incomplete and provide limited information. Nevertheless, a large number of human traits have been successfully mapped with the use of pedigree data to analyze linkage. Because the number of progeny from any one mating is usually small, data from several families and pedigrees are usually combined to test for independent assortment. The methods used in these types of analysis are beyond the scope of this book, but an example will illustrate how linkage can be detected from pedigree data.

One of the first documented demonstrations of linkage in humans was between the locus for nail–patella syndrome and the locus that determines the ABO blood types. Nail–patella syndrome is an autosomal dominant disorder characterized by abnormal fingernails and absent or rudimentary kneecaps. The ABO blood types are determined by an autosomal locus with multiple alleles (see Chapter 5). Linkage between the genes encoding these traits was established in families in which both traits segregate. Part of one such family is illustrated in **Figure 7.18**.

Nail–patella syndrome is rare, and so we can assume that people having this trait are heterozygous (*Nn*); unaffected people are homozygous (*nn*). The ABO genotypes can be inferred from the phenotypes and the types of offspring produced. Person I-2 in Figure 7.18, for example, has blood-type B, which has two possible genotypes:  $I^{\text{B}}I^{\text{B}}$ or  $I^{\text{B}}i$  (see Figure 5.6). Because some of her offspring are blood-type O (genotype *ii*) and must have therefore inherited an *i* allele from each parent, female I-2 must have genotype *I* B *i*. Similarly, the presence of blood-type O offspring



**7.18 Linkage between ABO blood types and nail–patella syndrome was established by examining families in whom both traits segregate.** The pedigree shown here is for one such family. The ABO blood type is indicated in each circle or square. The genotype, inferred from phenotype, is given below each circle or square.

in generation II indicates that male I-1, with blood-type A, also must carry an *i* allele and therefore has genotype *I* <sup>A</sup>*i*. The parents of this family are:

#### $I^{\text{A}}$ *i Nn*  $\times$   $I^{\text{B}}$ *i nn*

From generation II, we can see that the genes for nail–patella syndrome and the blood types do not appear to assort independently. All children in generation II with nail–patella syndrome have either blood-type B or bloodtype O; all those with blood-type A have normal nails and kneecaps. If the genes encoding nail–patella syndrome and the ABO blood types assorted independently, we would expect that some children in generation II would have blood-type A and nail–patella syndrome, inheriting both the *I* A and *N* alleles from their father. This outcome indicates that the arrangements of the alleles on the chromosomes of the crossed parents are:

$$
\frac{I^{\mathcal{A}}}{i} \qquad \frac{n}{N} \times \frac{I^{\mathcal{B}}}{i} \qquad \frac{n}{n}
$$

The pedigree indicates that there is no recombination among the offspring (generation II) of these parents, but there are two instances of recombination among the persons in generation III. Persons II-1 and II-2 have the following genotypes:

$$
\frac{i^B}{i} \qquad \frac{n}{N} \times \frac{I^A}{i} \qquad \frac{n}{n}
$$

Their child III-2 has blood-type A and does not have nail–patella syndrome; so he must have genotype

$$
\frac{I^{\mathrm{A}}}{i} \qquad n
$$

and must have inherited both the *i* and the *n* alleles from his father. These alleles are on different chromosomes in the father; so crossing over must have taken place. Crossing over also must have taken place to produce child III-3.

In the pedigree of Figure 7.18, 13 children are from matings in which the genes encoding nail–patella syndrome and ABO blood types segregate; 2 of them are recombinants. On this basis, we might assume that the loci for nail–patella syndrome and ABO blood types are linked, with a recombination frequency of  $\frac{2}{13} = 0.154$ . However, it is possible that the genes *are* assorting independently and that the small number of children just makes it seem as though the genes are linked. To determine the probability that genes are actually linked, geneticists often calculate **lod** (logarithm of odds) scores.

To obtain a lod score, we calculate both the probability of obtaining the observed results with the assumption that the genes are linked with a specified degree of recombination and the probability of obtaining the observed results with the assumption of independent assortment. We then determine the ratio of these two probabilities, and the logarithm of this ratio is the lod score. Suppose that the probability of obtaining a particular set of observations with the assumption of linkage and a certain recombination frequency is 0.1 and that the probability of obtaining the same observations with the assumption of independent assortment is 0.0001. The ratio of these two probabilities is  $^{0.1}/_{0.0001}$  = 1000, the logarithm of which (the lod score) is 3. Thus, linkage with the specified recombination is 1000 times as likely as independent assortment to produce what was observed. A lod score of 3 or higher is usually considered convincing evidence for linkage.  $TRY PROBLEM 33 \rightarrow$ 

#### Mapping with Molecular Markers

For many years, gene mapping was limited in most organisms by the availability of **genetic markers**—that is, variable genes with easily observable phenotypes for which inheritance could be studied. Traditional genetic markers include genes that encode easily observable characteristics such as flower color, seed shape, blood types, and biochemical differences. The paucity of these types of characteristics in many organisms limited mapping efforts.

In the 1980s, new molecular techniques made it possible to examine variations in DNA itself, providing an almost unlimited number of genetic markers that can be used for creating genetic maps and studying linkage relations. The earliest of these molecular markers consisted of restriction fragment length polymorphisms (RFLPs), which are variations in DNA sequence detected by cutting the DNA with restriction enzymes (see Chapter 19). Later, methods were developed for detecting variable numbers of short DNA sequences repeated in tandem, called microsatellites. More recently, DNA sequencing allows the direct detection of individual variations in the DNA nucleotides. All of these methods have expanded the availability of genetic markers and greatly facilitated the creation of genetic maps.

Gene mapping with molecular markers is done essentially in the same manner as mapping performed with traditional phenotypic markers: the cosegregation of two or more markers is studied, and map distances are based on the rates of recombination between markers. These methods and their use in mapping are presented in more detail in Chapters 19 and 20.

### Locating Genes with Genomewide Association Studies

The traditional approach to mapping genes, which we have learned in this chapter, is to examine progeny phenotypes in genetic crosses or among individuals in a pedigree, looking for associations between the inheritance of particular phenotype and the inheritance of alleles at other loci. This type of gene mapping is called **linkage analysis**, because it is based on the detection of physical linkage between genes, as measured by the rate of recombination, in progeny from a cross. Linkage analysis has been a powerful tool in the genetic analysis of many different types of organisms.

Another alternative approach to mapping genes is to conduct **genomewide association studies**, looking for nonrandom associations between the presence of a trait and alleles at many different loci scattered across the genome. Unlike linkage analysis, this approach does not trace the inheritance of genetic markers and a trait in a genetic cross or family. Rather, it looks for associations between traits and particular suites of alleles in a *population*.

Imagine that we are interested in finding genes that contribute to bipolar disease, a psychiatric illness characterized by severe depression and mania. When a mutation that predisposes a person to bipolar disease first arises in a population, it will occur on a particular chromosome and be associated with a specific set of alleles on that chromosome (**Figure 7.19**). A specific set of linked alleles, such as this, is called a **haplotype**, and the nonrandom association between alleles in a haplotype is called **linkage disequilibrium**. Because of the physical linkage between the bipolar mutation and the other alleles of the haplotype, bipolar illness and the haplotype will tend to be inherited together. Crossing over, however, breaks up the association between the alleles of the haplotype (see Figure 7.19), reducing the linkage disequilibrium between them. How long the linkage disequilibrium persists depends on the amount of recombination between alleles at different loci. When the loci are far apart, linkage disequilibrium breaks down quickly; when the loci are close together, crossing over is less common and linkage disequilibrium will persist longer. The important point is that linkage disequilibrium—the nonrandom association between alleles—provides information about the distance between genes. A strong association between a trait such as bipolar illness and a set of linked genetic markers indicates that one or more genes contributing to bipolar illness are likely to be near the genetic markers.

In recent years, geneticists have mapped millions of genetic variants called **single-nucleotide polymorphisms** (SNPs), which are positions in the genome where people vary in a single nucleotide base (see Chapter 20). It is now possible to quickly and inexpensively genotype people for hundreds of thousands or millions of SNPs. This genotyping has provided genetic markers needed for conducting genomewide association studies, in which SNP haplotypes of people with a particular disease, such as bipolar illness, are compared with the haplotypes of healthy persons. Nonrandom associations between SNPs and the disease suggest that one or more genes that contribute to the disease are closely linked to the SNPs. Genomewide association studies do not usually locate specific genes, but rather associate the inheritance of a trait or disease with a specific chromosomal region. After such an association has been established, geneticists can examine the chromosomal region for genes that might be responsible for the trait. Genomewide association studies have been instrumental in the discovery of genes or chromosomal regions that affect a number of genetic diseases and important human traits,



**7.19 Genomewide association studies are based on the nonrandom association of a mutation (***D***<sup>−</sup>) that produces a trait and closely linked genes that constitute a haplotype.**

including bipolar disease, height, skin pigmentation, eye color, body weight, coronary artery disease, blood-lipid concentrations, diabetes, heart attacks, bone density, and glaucoma, among others.

#### **CONCEPTS**

The development of molecular techniques for examining variation in DNA sequences has provided a large number of genetic markers that can be used to create genetic maps and study linkage relations. Genomewide association studies examine the nonrandom association of genetic markers and phenotypes to locate genes that contribute to the expression of traits.

# 7.4 Physical-Mapping Methods Are Used to Determine the Physical Positions of Genes on Particular Chromosomes

Genetic maps reveal the relative positions of genes on a chromosome on the basis of frequencies of recombination, but they do not provide information that allow us to

place groups of linked genes on particular chromosomes. Furthermore, the units of a genetic map do not always precisely correspond to physical distances on the chromosome, because a number of factors other than physical distances between genes (such as the type and sex of the organism) can influence recombination. Because of these limitations, physical-mapping methods that do not rely on recombination frequencies have been developed.

#### Deletion Mapping

One method for determining the chromosomal location of a gene is **deletion mapping**. Special staining methods have been developed that reveal characteristic banding patterns on the chromosomes (see Chapter 9). The absence of one or more of the bands that are normally on a chromosome reveals the presence of a chromosome deletion, a mutation in which a part of a chromosome is missing. Genes can be assigned to regions of chromosomes by studying the association between a gene's phenotype or product and particular chromosome deletions.

In deletion mapping, an individual that is homozygous for a recessive mutation in the gene of interest is crossed with an individual that is heterozygous for a deletion (**Figure 7.20**). If the gene of interest is in the region of the chromosome represented by the deletion (the red part of the chromosomes in Figure 7.20), approximately half of the progeny will display the mutant phenotype (see Figure 7.20a). If the gene is not within the deleted region, all of the progeny will be wild type (see Figure 7.20b).

Deletion mapping has been used to reveal the chromosomal locations of a number of human genes. For example, Duchenne muscular dystrophy is a disease that causes progressive weakening and degeneration of the muscles. From its X-linked pattern of inheritance, the mutated allele causing this disorder was known to be on the X chromosome, but its precise location was uncertain. Examination of a number of patients having Duchenne muscular dystrophy, who also possessed small deletions, allowed researchers to position the gene on a small segment of the short arm of the X chromosome. **TRY PROBLEM 34**

#### Somatic-Cell Hybridization

Another method used for positioning genes on chromosomes is **somatic-cell hybridization**, which requires the fusion of different types of cells. Most mature somatic (nonsex) cells can undergo only a limited number of divisions and therefore cannot be grown continuously. However, cells that have been altered by viruses or derived from tumors that have lost the normal constraints on cell division will divide indefinitely; this type of cell can be cultured in the laboratory to produce a **cell line**.

Cells from two different cell lines can be fused by treating them with polyethylene glycol or other agents that alter their plasma membranes. After fusion, the cell possesses



**7.20 Deletion mapping can be used to determine the chromosomal location of a gene.**  An individual homozygous for a recessive mutation in the gene of interest (*aa*) is crossed with an individual heterozygous for a deletion.



**7.21 Somatic-cell hybridization can be used to determine which chromosome contains a gene of interest.**

two nuclei and is called a **heterokaryon**. The two nuclei of a heterokaryon eventually also fuse, generating a hybrid cell that contains chromosomes from both cell lines. If human and mouse cells are mixed in the presence of polyethylene glycol, fusion results in human–mouse somatic-cell hybrids (**Figure 7.21**). The hybrid cells tend to lose chromosomes as they divide and, for reasons that are not understood, chromosomes from one of the species are lost preferentially. In human–mouse somatic-cell hybrids, the human chromosomes tend to be lost, whereas the mouse chromosomes are retained. Eventually, the chromosome number stabilizes when all but a few of the human chromosomes have been lost. Chromosome loss is random and differs among cell lines. The presence of these "extra" human chromosomes in the mouse genome makes it possible to assign human genes to specific chromosomes.

To map genes by using somatic-cell hybridization requires a panel of different hybrid cell lines. Each cell line is examined microscopically and the human chromosomes that it contains are identified. The cell lines of the panel are chosen so that they differ in the human chromosomes that they have retained. For example, one cell line might possess human chromosomes 2, 4, 7, and 8, whereas another might possess chromosomes 4, 19, and 20. Each cell line in the panel is examined for evidence of a particular human gene. The human gene can be detected by looking either for the for the gene itself (discussed in Chapter 19) or for the protein that it produces. Correlation of the presence of the gene with the presence of specific human chromosomes often allows the gene to be assigned to the correct chromosome. For example, if a gene is detected in both of the aforementioned cell lines, the gene must be on chromosome 4, because it is the only human chromosome common to both cell lines (**Figure 7.22**).

Sometimes somatic-cell hybridization can be used to position a gene on a specific part of a chromosome. Some



**Human chromosomes present**

**7.22 Somatic-cell hybridization is used to assign a gene to a particular human chromosome.** A panel of six cell lines, each line containing a different subset of human chromosomes, is examined for the presence of the gene product (such as an enzyme). Four of the cell lines (A, B, D, and F) have the gene product. The only chromosome common to all four of these cell lines is chromosome 4, indicating that the gene is located on this chromosome.



**7.23 Genes can be localized to a specific part of a chromosome by using somatic-cell hybridization.**

hybrid cell lines carry a human chromosome with a chromosome mutation such as a deletion or a translocation. If the gene is present in a cell line with the intact chromosome but missing from a line with a chromosome deletion, the gene must be located in the deleted region (**Figure 7.23**). Similarly, if a gene is usually absent from a chromosome but consistently appears whenever a translocation (a piece of another chromosome that has broken off and attached itself to the chromosome in question) is present, it must be present on the translocated part of the chromosome.

# Worked Problem

A panel of cell lines was created from human–mouse somaticcell fusions. Each line was examined for the presence of human chromosomes and for the production of human haptoglobin (a protein). The following results were obtained:



On the basis of these results, which human chromosome carries the gene for haptoglobin?

#### • Solution

First, identify the cell lines that are positive for the protein (human haptoglobin) and determine the chromosomes that they have in common. Lines B and C produce human haptoglobin; the only chromosomes that they have in common are chromosomes 1 and 16. Next, examine all the cell lines that possess either chromosomes 1 and 16 and determine whether they produce haptoglobin. Chromosome 1 is found in cell lines A, B, C, and D. If the gene for human haptoglobin were found on chromosome 1, human haptoglobin would be present in all of these cell lines. However, lines A and D do not produce human haptoglobin; so the gene cannot be on chromosome 1. Chromosome 16 is found only in cell lines B and C, and only these lines produce human haptoglobin; so the gene for human haptoglobin lies on chromosome 16.

For more practice with somatic-cell hybridizations, work Problem 35 at the end of this chapter.

# Physical Chromosome Mapping Through Molecular Analysis

So far, we have explored methods to indirectly determine the chromosomal location of a gene by deletion mapping or by looking for gene products. Researchers now have the information and technology to actually see where a gene lies. Described in more detail in Chapter 19, in situ hybridization is a method for determining the chromosomal location of a particular gene through molecular analysis. This method requires the creation of a probe for the gene, which is a singlestranded DNA complement to the gene of interest. The probe is radioactive or fluoresces under ultraviolet light so that it can be visualized. The probe binds to the DNA sequence of the gene on the chromosome. The presence of radioactivity or fluorescence from the bound probe reveals the location of the gene on a particular chromosome (**Figure 7.24**).

In addition to allowing us to see where a gene is located on a chromosome, modern laboratory techniques now allow researchers to identify the precise location of a gene at the nucleotide level. For example, with DNA sequencing (described fully in Chapter 19), physical distances between genes can be determined in numbers of base pairs.



**7.24 In situ hybridization is another technique for determining the chromosomal location of a gene.** The red fluorescence is produced by a probe for sequences on chromosome 9; the green fluorescence is produced by a probe for sequences on chromosome 22. [Courtesy of Applied Imaging Corp.]

#### CONCEPTS

Physical-mapping methods determine the physical locations of genes on chromosomes and include deletion mapping, somatic-cell hybridization, in situ hybridization, and direct DNA sequencing.

# 7.5 Recombination Rates Exhibit Extensive Variation

In recent years, geneticists have studied variation in rates of recombination and found that levels of recombination vary widely—among species, among and along chromosomes of a single species, and even between males and females of the same species. For example, about twice as much recombination takes place in humans as in the mouse and the rat. Within the human genome, recombination varies among chromosomes, with chromosomes 21 and 22 having the highest rates and chromosomes 2 and 4 having the lowest rates. Researchers have also detected differences between male and female humans in rates of recombination: the autosomal chromosomes of females undergo about 50% more recombination than do the autosomal chromosomes of males.

Geneticists have found numerous recombination *hotspots*, where recombination is at least 10 times as high as the average elsewhere in the genome. The human genome may contain an estimated 25,000 to 50,000 such recombination hotspots. Approximately 60% of all crossovers take place in hotspots. For humans, recombination hotspots tend to be found near but not within active genes. Recombination hotspots have been detected in the genomes of other organisms as well. In comparing recombination hotspots in the genomes of humans and chimpanzees (our closest living relative), geneticists have determined that, although the DNA sequences of humans and chimpanzees are extremely similar, their recombination hotspots are at entirely different places.

#### **CONCEPTS**

Rates of recombination vary among species, among and along chromosomes, and even between males and females.

#### CONCEPTS SUMMARY

- Linked genes do not assort independently. In a testcross for two completely linked genes (no crossing over), only nonrecombinant progeny are produced. When two genes assort independently, recombinant progeny and nonrecombinant progeny are produced in equal proportions. When two genes are linked with some crossing over between them, more nonrecombinant progeny than recombinant progeny are produced. •
- Recombination frequency is calculated by summing the number of recombinant progeny, dividing by the total number of progeny produced in the cross, and multiplying by 100%. The recombination frequency is half the frequency of crossing over, and the maximum frequency of recombinant gametes is 50%. •
- Coupling and repulsion refer to the arrangement of alleles on a chromosome. Whether genes are in coupling configuration or in repulsion determines which combination of phenotypes will be most frequent in the progeny of a testcross. •
- Interchromosomal recombination takes place among genes located on different chromosomes through the random segregation of chromosomes in meiosis. Intrachromosomal recombination takes place among genes located on the same chromosome through crossing over. •
- A chi-square test of independence can be used to determine if genes are linked. •
- Recombination rates can be used to determine the relative order of genes and distances between them on a chromosome. One percent recombination equals one map

unit. Maps based on recombination rates are called genetic maps; maps based on physical distances are called physical maps.

- Genetic maps can be constructed by examining recombination rates from a series of two-point crosses or by examining the progeny of a three-point testcross.
- Some multiple crossovers go undetected; thus, genetic maps based on recombination rates underestimate the true physical distances between genes. •
- Human genes can be mapped by examining the cosegregation of traits in pedigrees. •
- A lod score is the logarithm of the ratio of the probability of obtaining the observed progeny with the assumption of linkage to the probability of obtaining the observed progeny with the assumption of independent assortment. A lod score of 3 or higher is usually considered evidence for linkage.
- Molecular techniques that allow the detection of variable differences in DNA sequence have greatly facilitated gene mapping.
- Genomewide association studies locate genes that affect particular traits by examining the nonrandom association of a trait with genetic markers from across the genome.
- Nucleotide sequencing is another method of physically mapping genes. •
- Rates of recombination vary widely, differing among species, among and along chromosomes within a single species, and even between males and females of the same species. •

#### IMPORTANT TERMS

linked genes (p. 163) linkage group (p. 163) nonrecombinant (parental) gamete (p. 166) nonrecombinant (parental) progeny (p. 166) recombinant gamete (p. 166) recombinant progeny (p. 166) recombination frequency (p. 167) coupling (cis) configuration (p. 168) repulsion (trans) configuration (p. 168) interchromosomal recombination (p. 170)

intrachromosomal recombination (p. 170) genetic map (p. 174) physical map (p. 174) map unit (m.u.) (p. 174) centiMorgan (p. 174) Morgan (p. 174) two-point testcross (p. 175) three-point testcross (p. 176) interference (p. 182) coefficient of coincidence (p. 182) mapping function (p. 185) lod (logarithm of odds) score (p. 186) genetic marker (p. 186) linkage analysis (p. 186) genomewide association studies (p. 187) haplotype (p. 187) linkage disequilibrium (p. 187) single-nucleotide polymorphism (SNP) (p. 187) deletion mapping (p. 188) somatic-cell hybridization (p. 188) cell line (p. 188) heterokaryon (p. 189)

#### ANSWERS TO CONCEPT CHECKS



**4.**  $\frac{m^+}{m} \frac{p^+}{p} s$  $m^{+}$  ps *m ps mp s m*  $+$  +  $+$  +  $+$  +  $+$  +  $+$  +  $+$  +  $+$ *p s*  $m^+$   $p^+s$ *m ps mps m p*  $^+$  +  $^+$  +  $^+$ *s*  $m^+$  p s *m ps*  $m$   $p^+$  s *mps mps mps*  $+$  +  $+$  +  $+$  +  $+$ 

#### WORKED PROBLEMS

**1.** In guinea pigs, white coat (*w*) is recessive to black coat (*W*) and wavy hair (*v*) is recessive to straight hair (*V*). A breeder crosses a guinea pig that is homozygous for white coat and wavy hair with a guinea pig that is black with straight hair. The  $F_1$  are then crossed with guinea pigs having white coats and wavy hair in a series of testcrosses. The following progeny are produced from these testcrosses:



**a.** Are the genes that determine coat color and hair type assorting independently? Carry out chi-square tests to test your hypothesis.

**b.** If the genes are not assorting independently, what is the recombination frequency between them?

#### • Solution

**a.** Assuming independent assortment, outline the crosses conducted by the breeder:

P *ww vv WW VV* A F1 *Ww Vv* A Testcross *Ww Vv ww vv* A *Ww Vv* ₁  $\frac{1}{4}$  black, straight *Ww vv* ₁  $\frac{1}{4}$  black, wavy *ww Vv* ₁  $\frac{1}{4}$  white, straight  $ww$   $vv$  $\frac{1}{4}$  white, wavy

Because a total of 83 progeny were produced in the testcrosses, we expect  $\frac{1}{4} \times 83 = 20.75$  of each. The observed numbers of progeny from the testcross (30, 10, 12, 31) do not appear to fit the expected numbers (20.75, 20.75, 20.75, 20.75) well; so independent assortment may not have taken place.

To test the hypothesis, carry out a chi-square test of independence. Construct a table, with the genotypes of the first locus along the top and the genotypes of the second locus along the side. Compute the totals for the rows and columns and the grand total.



The expected value for each cell of the table is calculated with the following formula:

> expected number  $=$   $\frac{\text{row total} \times \text{column total}}{\text{grand total}}$ grand total

Using this formula, we find the expected values (given in parentheses) to be:



Using these observed and expected numbers, we find the calculated chi-square value to be:

$$
\chi^{2} = \sum \frac{(\text{observed} - \text{expected})^{2}}{\text{expected}}
$$
  
=  $\frac{(30 - 20.24)^{2}}{20.24} + \frac{(10 - 19.76)^{2}}{19.76} + \frac{(12 - 21.76)^{2}}{21.76} + \frac{(31 - 21.24)^{2}}{21.24}$   
= 4.71 + 4.82 + 4.38 + 4.48 = 18.39

The degrees of freedom for the chi-square test of independence are  $df =$  (number of rows  $-1$ ) × (number of columns − 1). There are two rows and two columns, and so the degrees of freedom are:

$$
df = (2 - 1) \times (2 - 1) = 1 \times 1 = 1
$$

In Table 3.5, the probability associated with a chi-square value of 18.39 and 1 degree of freedom is less than 0.005, indicating that chance is very unlikely to be responsible for the differences between the observed numbers and the numbers expected with independent assortment. The genes for coat color and hair type have therefore not assorted independently.

**b.** To determine the recombination frequencies, identify the recombinant progeny. Using the notation for linked genes, write the crosses:

> *W V w v*

$$
\begin{array}{ccccccc}\nP & & & & \frac{W & V}{W & V} & \times & \frac{W & V}{W & V} \\
& & & & & \downarrow & & & \downarrow\n\end{array}
$$

$$
\mathbf{F}_1 =
$$



The recombination frequency is:

$$
\frac{\text{number of recombinant progeny}}{\text{total number progeny}} \times 100\%
$$

or

recombinant frequency =

$$
\frac{10+12}{30+31+10+12} \times 100\% = \frac{22}{83} \times 100\% = 26.5
$$

**2.** A series of two-point crosses were carried out among seven loci (*a*, *b*, *c*, *d*, *e*, *f*, and *g*), producing the following recombination frequencies. Using these recombination frequencies, map the seven loci, showing their linkage groups, the order of the loci in each linkage group, and the distances between the loci of each group:



#### • Solution

To work this problem, remember that 1% recombination equals 1 map unit and a recombination frequency of 50% means that genes at the two loci are assorting independently (located in different linkage groups).

The recombination frequency between *a* and *b* is 10%; so these two loci are in the same linkage group, approximately 10 m.u. apart.



The recombination frequency between *a* and *c* is 50%; so *c* must lie in a second linkage group.



The recombination frequency between *a* and *d* is 14%; so *d* is located in linkage group 1. Is locus *d* 14 m.u. to the right or to the left of gene *a?* If *d* is 14 m.u. to the left of *a*, then the *b*-to-*d* distance should be 10 m.u.  $+$  14 m.u.  $=$  24 m.u. On the other hand, if *d* is to the right of *a*, then the distance between *b* and *d* should be 14 m.u.  $-10$  m.u.  $= 4$  m.u. The *b–d* recombination frequency is 4%; so *d* is 14 m.u. to the right of *a*. The updated map is:



Linkage group 2

The recombination frequencies between each of loci *a*, *b*, and *d*, and locus *e* are all 50%; so *e* is not in linkage group 1 with *a*, *b*, and *d*. The recombination frequency between *e* and *c* is 8 m.u.; so *e* is in linkage group 2:

*c*



Linkage group 2



There is 50% recombination between *f* and all the other genes; so *f* must belong to a third linkage group:



Finally, position locus *g* with respect to the other genes. The recombination frequencies between *g* and loci *a*, *b*, and *d* are all 50%; so *g* is not in linkage group 1. The recombination frequency between *g* and *c* is 12 m.u.; so *g* is a part of linkage group 2. To determine whether *g* is 12 m.u. to the right or left of *c*, consult the *g*–*e* recombination frequency. Because this recombination frequency is 18%, *g* must lie to the left of *c:*



Note that the *g*-to-*e* distance (18 m.u.) is shorter than the sum of the *g*-to-*c* (12 m.u.) and *c*-to-*e* distances (8 m.u.) because of undetectable double crossovers between *g* and *e*.

**3.** Ebony body color (*e*), rough eyes (*ro*), and brevis bristles (*bv*) are three recessive mutations that occur in fruit flies. The loci for these mutations have been mapped and are separated by the following map distances:



The interference between these genes is 0.4.

A fly with ebony body, rough eyes, and brevis bristles is crossed with a fly that is homozygous for the wild-type traits. The resulting  $F_1$  females are test-crossed with males that have

ebony body, rough eyes, and brevis bristles; 1800 progeny are produced. Give the expected numbers of phenotypes in the progeny of the testcross.

#### • Solution

The crosses are:

P A F1 A Testcross *e ro bv <sup>e</sup> ro bv <sup>e</sup> ro bv e ro bv e ro bv e ro bv e ro bv <sup>e</sup> ro bv <sup>e</sup> ro bv e ro bv*

In this case, we know that *ro* is the middle locus because the genes have been mapped. Eight classes of progeny will be produced from this cross:



To determine the numbers of each type, use the map distances, starting with the double crossovers. The expected number of double crossovers is equal to the product of the single-crossover probabilities:

expected number of double crossovers = 0.20  $\times$  0.12  $\times$  1800  $= 43.2$ 

However, there is some interference; so the observed number of double crossovers will be less than the expected. The interference is 1 − coefficient of coincidence; so the coefficient of coincidence is:

 $coefficient of coincidence = 1 - interface$ 

The interference is given as 0.4; so the coefficient of coincidence equals  $1 - 0.4 = 0.6$ . Recall that the coefficient of coincidence is:

$$
coefficient of coincidence =
$$

number of observed double crossovers number of expected double crossovers

Rearranging this equation, we obtain:

number of observed double crossovers =

coefficient of coincidence  $\times$  number of expected double crossovers

number of observed double crossovers  $= 0.6 \times 43.2 = 26$ 

A total of 26 double crossovers should be observed. Because there are two classes of double crossovers  $(e^+$  / ro / bv<sup>+</sup> and  $\frac{e}{r}$  /  $\frac{f}{r}$  /  $\frac{b\nu}{c}$ , we expect to observe 13 of each class.

Next, we determine the number of single-crossover progeny. The genetic map indicates that the distance between *e* and *ro* is 20 m.u.; so 360 progeny (20% of 1800) are expected to have resulted from recombination between these two loci. Some of them will be single-crossover progeny and some will be double-crossover progeny. We have already determined that the number of doublecrossover progeny is 26; so the number of progeny resulting from a single crossover between *e* and *ro* is 360 − 26 = 334, which will be divided equally between the two single-crossover phenotypes  $(e / ro^{+} / bv^{+}$  and  $e^{+} / ro$  by).

The distance between *ro* and *bv* is 12 m.u.; so the number of progeny resulting from recombination between these two genes is  $0.12 \times 1800 = 216$ . Again, some of these recombinants will be single-crossover progeny and some will be double-crossover progeny. To determine the number of progeny resulting from a single crossover, subtract the double crossovers:  $216 - 26 = 190$ . These single-crossover progeny will be divided between the two singlecrossover phenotypes ( $e^+$  *ro*<sup>+</sup> / *bv* and  $e$  *ro* / *bv*<sup>+</sup>); so there will be  $190/2 = 95$  of each of these phenotypes. The remaining progeny will be nonrecombinants, and they can be obtained by subtraction:  $1800 - 26 - 334 - 190 = 1250$ ; there are two nonrecombinants  $(\frac{e^+}{e^+} \cdot \frac{f_0}{e^+} + \frac{f_0}{e^+} + \frac{f_0}{e^+} + \frac{f_0}{e^+} + \frac{f_0}{e^+})$ ; so there will be  $\frac{1250}{2} = 625$  of each. The numbers of the various phenotypes are listed here:



**4.** The locations of six deletions have been mapped to the *Drosophila* chromosome as shown in the following diagram.



Recessive mutations *a*, *b*, *c*, *d*, *e*, *f*, and *g* are known to be located in the same regions as the deletions, but the order of the mutations on the chromosome is not known. When flies homozygous for the recessive mutations are crossed with flies homozygous for the deletions, the following results are obtained, where the letter "m" represents a mutant phenotype and a plus

sign  $(+)$  represents the wild type. On the basis of these data, determine the relative order of the seven mutant genes on the chromosome:



#### • Solution

The offspring of the cross will be heterozygous, possessing one chromosome with the deletion and wild-type alleles and its homolog without the deletion and recessive mutant alleles. For loci within the deleted region, only the recessive mutations will be present in the offspring, which will exhibit the mutant phenotype. The presence of a mutant trait in the offspring therefore indicates that the locus for that trait is within the region covered by the deletion. We can map the genes by examining the expression of the recessive mutations in the flies with different deletions.

Mutation *a* is expressed in flies with deletions 4, 5, and 6 but not in flies with other deletions; so *a* must be in the area that is unique to deletions 4, 5, and 6:



#### COMPREHENSION QUESTIONS

#### Section 7.1

**\*1.** What does the term recombination mean? What are two causes of recombination?

#### Section 7.2

- **\*2.** In a testcross for two genes, what types of gametes are produced with (a) complete linkage, (b) independent assortment, and (c) incomplete linkage?
- **3.** What effect does crossing over have on linkage?
- **4.** Why is the frequency of recombinant gametes always half the frequency of crossing over?

Mutation *b* is expressed only when deletion 1 is present; so *b* must be located in the region of the chromosome covered by deletion 1 and none of the other deletions:



Using this procedure, we can map the remaining mutations. For each mutation, we look for the areas of overlap among deletions that express the mutations and exclude any areas of overlap that are covered by other deletions that do not express the mutation:



- **\*5.** What is the difference between genes in coupling configuration and genes in repulsion? What effect does the arrangement of linked genes (whether they are in coupling configuration or in repulsion) have on the results of a cross?
- **6.** How would you test to see if two genes are linked?
- **7.** What is the difference between a genetic map and a physical map?
- **\*8.** Why do calculated recombination frequencies between pairs of loci that are located far apart underestimate the true genetic distances between loci?