

1

### Drawing from the Deck of Genes

- What principles account for the transmission of traits from parents to offspring?
- The “blending” hypothesis is the idea that genetic material from the two parents blends together (like blue and yellow paint blend to make green)

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### Drawing from the Deck of Genes, Continued

- The “particulate” hypothesis is the idea that parents pass on discrete heritable units (genes)
- Mendel documented a particulate mechanism through his experiments with garden peas

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### Figure 14.1 What Principles of Inheritance did Gregor Mendel Discover by Breeding Garden Pea Plants?

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**Figure 14.1a What Principles of Inheritance did Gregor Mendel Discover by Breeding Garden Pea Plants? (Part 1: Mendel with Fellow Monks)**



**Mendel (third from right, holding a sprig of fuchsia) with his fellow monks**

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**Concept 14.1: Mendel used the scientific approach to identify two laws of inheritance**

- Mendel discovered the basic principles of heredity by breeding garden peas in carefully planned experiments

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**Mendel's Experimental, Quantitative Approach**

- Mendel's approach allowed him to deduce principles that had remained elusive to others
- A heritable feature that varies among individuals (such as flower color) is called a **character**
- Each variant for a character, such as purple or white color for flowers, is called a **trait**
- Peas were available to Mendel in many different varieties

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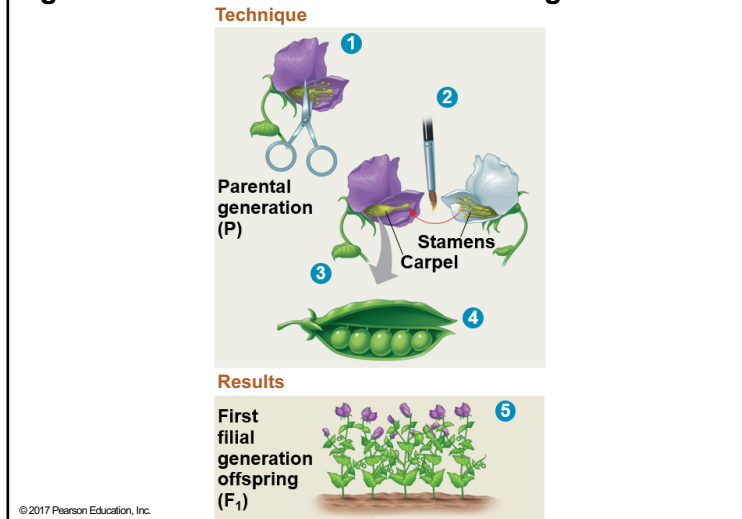
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**Mendel's Experimental, Quantitative Approach, Continued**

- Other advantages of using peas
  - Short generation time
  - Large numbers of offspring
  - Mating could be controlled; plants could be allowed to self-pollinate or could be cross-pollinated

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**Figure 14.2 Research Method: Crossing Pea Plants**

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Which of the following is not a reason that peas were well suited for Mendel's breeding experiments?

- Peas show easily observed variations in a number of characters, such as pea shape and flower color.
- It is possible to control matings between different pea plants.
- It is possible to obtain large numbers of progeny from any given cross.
- Peas have an unusually long generation time.
- Many of the observable characters that vary in pea plants are controlled by single genes.

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### Mendel's Experimental, Quantitative Approach, Continued-1

- Mendel chose to track only those characters that occurred in two distinct alternative forms
- He also started with varieties that were **true-breeding** (plants that produce offspring of the same variety when they self-pollinate)

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### Mendel's Experimental, Quantitative Approach, Continued-2

- In a typical experiment, Mendel mated two contrasting, true-breeding varieties, a process called **hybridization**
- The true-breeding parents are the **P generation**
- The hybrid offspring of the P generation are called the  **$F_1$  generation**
- When  $F_1$  individuals self-pollinate or cross-pollinate with other  $F_1$  hybrids, the  **$F_2$  generation** is produced

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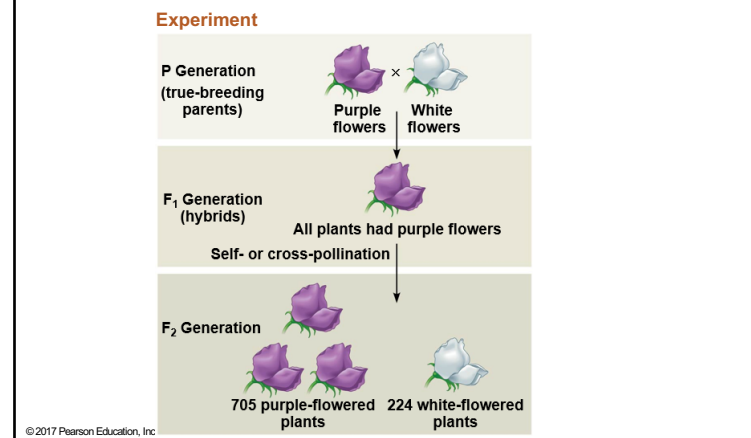
### The Law of Segregation

- When Mendel crossed contrasting, true-breeding white- and purple-flowered pea plants, all of the  $F_1$  hybrids were purple
- When Mendel crossed the  $F_1$  hybrids, many of the  $F_2$  plants had purple flowers, but some had white
- Mendel discovered a ratio of about three purple flowers to one white flower in the  $F_2$  generation

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### Figure 14.3\_3 Inquiry: When $F_1$ Hybrid Pea Plants Self- or Cross-pollinate, Which Traits Appear in the $F_2$ Generation? (Step 3)



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### The Law of Segregation, Continued

- Mendel reasoned that only the purple flower factor was affecting flower color in the  $F_1$  hybrids
- Mendel called the purple flower color a dominant trait and the white flower color a recessive trait
- The factor for white flowers was not diluted or destroyed because it reappeared in the  $F_2$  generation

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### The Law of Segregation, Continued-1

- Mendel observed the same pattern of inheritance in six other pea plant characters, each represented by two traits
- What Mendel called a "heritable factor" is what we now call a gene

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**The Results of Mendel's F<sub>1</sub> Crosses for Seven Characters in Pea Plants**

Table 14.1 The Results of Mendel's F<sub>1</sub> Crosses for Seven Characters in Pea Plants

Character	Dominant Trait	Recessive Trait	F <sub>2</sub> Generation: Dominant/Recessive	Ratio
Flower color	Purple	White	705:224	3.15:1
Seed color	Yellow	Green	6,022:2,001	3.01:1
Seed shape	Round	Wrinkled	5,474:1,850	2.96:1
Pod color	Green	Yellow	428:152	2.82:1
Pod shape	Inflated	Constricted	882:299	2.95:1
Flower position	Axial	Terminal	651:207	3.14:1
Stem length	Tall	Dwarf	787:277	2.84:1

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**Mendel's Model**

- Mendel developed a hypothesis to explain the 3:1 inheritance pattern he observed in F<sub>2</sub> offspring
- Four related concepts make up this model
- These concepts can be related to what we now know about genes and chromosomes

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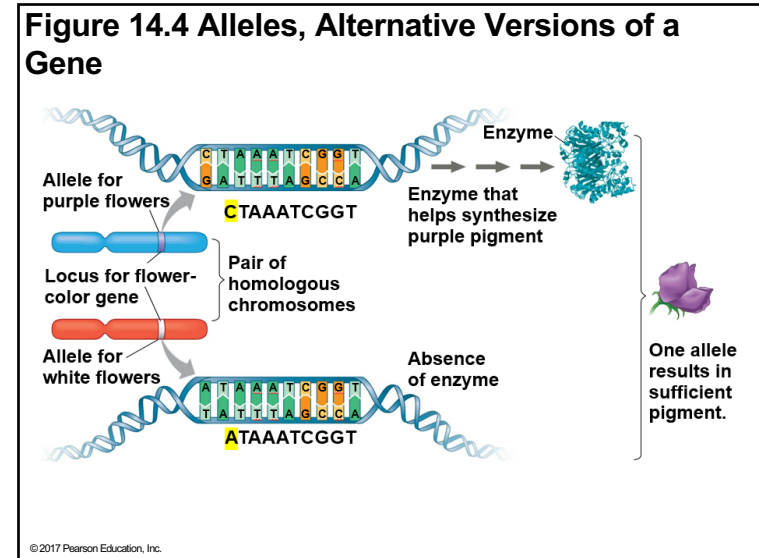
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**Mendel's Model, Continued**

- First: alternative versions of genes account for variations in inherited characters
- For example, the gene for flower color in pea plants exists in two versions, one for purple flowers and the other for white flowers
- These alternative versions of a gene are called **alleles**
- Each gene resides at a specific locus on a specific chromosome

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**Mendel's Model, Continued-1**

- Second: for each character, an organism inherits two alleles, one from each parent
- Mendel made this deduction without knowing about chromosomes
- The two alleles at a particular locus may be identical, as in the true-breeding plants of Mendel's P generation
- Or the two alleles at a locus may differ, as in the F<sub>1</sub> hybrids

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**Mendel's Model, Continued-2**

- Third: if the two alleles at a locus differ, then one (the **dominant allele**) determines the organism's appearance, and the other (the **recessive allele**) has no noticeable effect on appearance
- In the flower-color example, the F<sub>1</sub> plants had purple flowers because the allele for that trait is dominant

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**Mendel's Model, Continued-3**

- Fourth (the **law of segregation**): the two alleles for a heritable character separate (segregate) during gamete formation and end up in different gametes
- Thus, an egg or a sperm gets only one of the two alleles that are present in the organism
- This segregation of alleles corresponds to the distribution of homologous chromosomes to different gametes in meiosis

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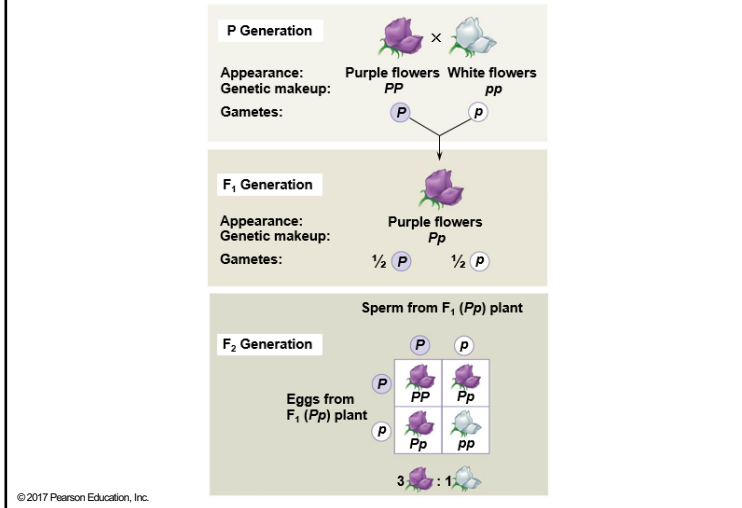
**Mendel's Model, Continued-4**

- The model accounts for the 3:1 ratio observed in the F<sub>2</sub> generation of Mendel's crosses
- Possible combinations of sperm and egg can be shown using a **Punnett square**
- A capital letter represents a dominant allele, and a lowercase letter represents a recessive allele

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Figure 14.5\_3 Mendel's Law of Segregation (Step 3)



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**Useful Genetic Vocabulary**

- An organism with two identical alleles for a character is called a **homozygote**
  - It is said to be **homozygous** for the gene controlling that character
  - An organism with two different alleles for a gene is a **heterozygote** and is said to be **heterozygous** for the gene controlling that character
  - Unlike homozygotes, heterozygotes are not true-breeding
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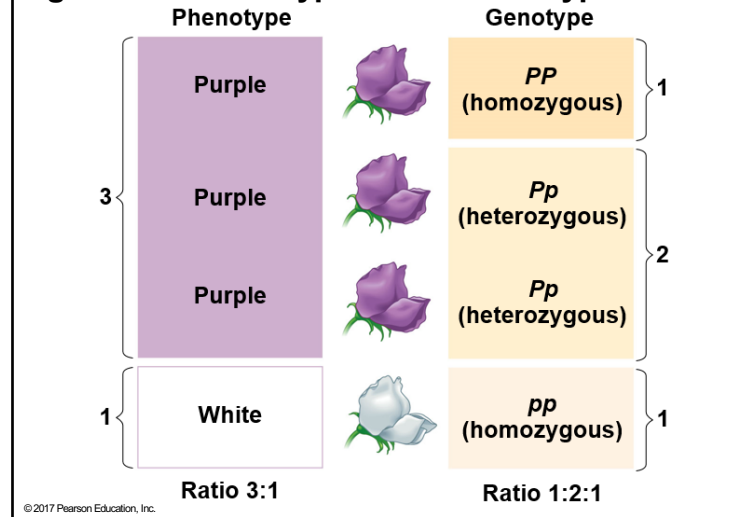
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**Useful Genetic Vocabulary, Continued**

- An organism's traits do not always reveal its genetic composition
  - Therefore, we distinguish between an organism's **phenotype**, or physical appearance, and its **genotype**, or genetic makeup
  - In the example of flower color in pea plants,  $PP$  and  $Pp$  plants have the same phenotype (purple) but different genotypes
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Figure 14.6 Phenotype Versus Genotype



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### The Testcross

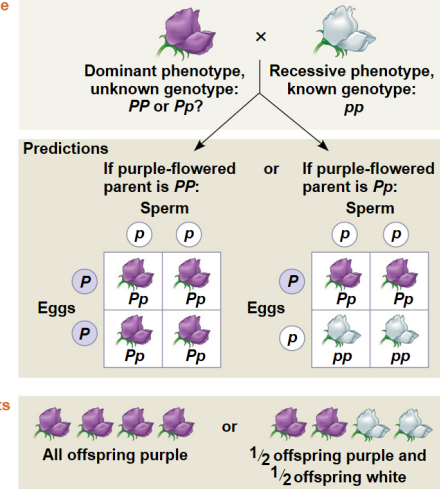
- An individual with the dominant phenotype could be either homozygous dominant or heterozygous
- To determine the genotype we can carry out a **testcross**: breeding the mystery individual with a homozygous recessive individual
- If any offspring display the recessive phenotype, the mystery parent must be heterozygous

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### Figure 14.7 Research Method: The Testcross

Technique



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### The Law of Independent Assortment

- Mendel derived the law of segregation by following a single character
- The  $F_1$  offspring produced in this cross were **monohybrids**, heterozygous for one character
- A cross between such heterozygotes is called a **monohybrid cross**

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### The Law of Independent Assortment, Continued

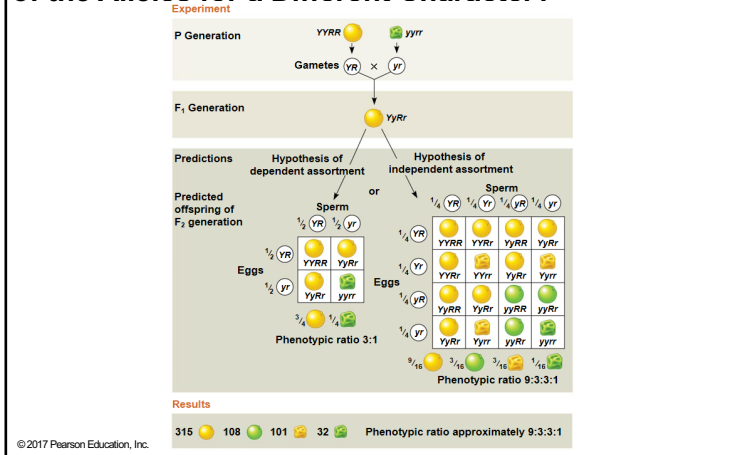
- Mendel identified his second law of inheritance by following two characters at the same time
- Crossing two true-breeding parents differing in two characters produces **dihybrids** in the  $F_1$  generation, heterozygous for both characters
- A **dihybrid cross**, a cross between  $F_1$  dihybrids, can determine whether two characters are transmitted to offspring as a package or independently

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**Figure 14.8 Inquiry: Do the Alleles for One Character Assort into Gametes Dependently or Independently of the Alleles for a Different Character?**



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**The Law of Independent Assortment, Continued-1**

- Using a dihybrid cross, Mendel developed the **law of independent assortment**
- It states that each pair of alleles segregates independently of any other pair of alleles during gamete formation
- This law applies only to genes on different, nonhomologous chromosomes or those far apart on the same chromosome
- Genes located near each other on the same chromosome tend to be inherited together

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**Concept 14.2: Probability laws govern Mendelian inheritance**

- Mendel's laws of segregation and independent assortment reflect the rules of probability
- When tossing a coin, the outcome of one toss has no impact on the outcome of the next toss
- In the same way, the alleles of one gene segregate into gametes independently of another gene's alleles

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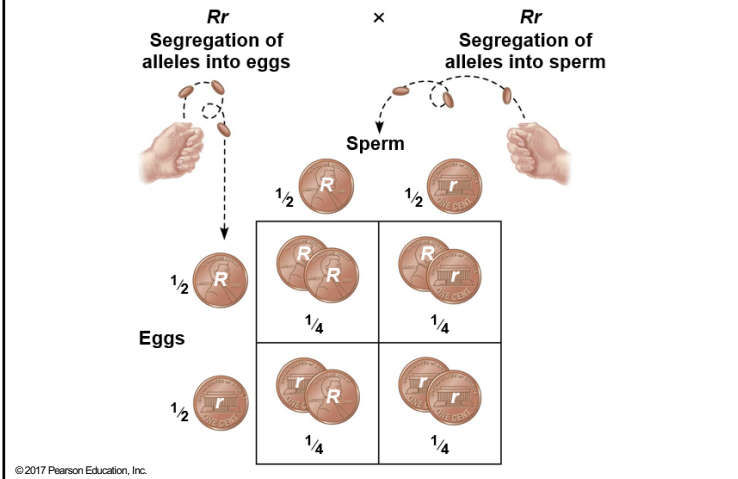
**The Multiplication and Addition Rules Applied to Monohybrid Crosses**

- The **multiplication rule** states that the probability that two or more independent events will occur together is the product of their individual probabilities
- Probability in an F<sub>1</sub> monohybrid cross can be determined using the multiplication rule
- Segregation in a heterozygous plant is like flipping a coin: Each gamete has a 1/2 chance of carrying the dominant allele and a 1/2 chance of carrying the recessive allele

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**Figure 14.9 Segregation of Alleles and Fertilization as Chance Events**



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**The Multiplication and Addition Rules Applied to Monohybrid Crosses, Continued**

- The **addition rule** states that the probability that any one of two or more mutually exclusive events will occur is calculated by adding together their individual probabilities
- The rule of addition can be used to figure out the probability that an  $F_2$  plant from a monohybrid cross will be heterozygous rather than homozygous

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### Solving Complex Genetics Problems with the Rules of Probability

- We can apply the rules of probability to predict the outcome of crosses involving multiple characters
- A multicharacter cross is equivalent to two or more independent monohybrid crosses occurring simultaneously
- In calculating the chances for various genotypes, each character is considered separately, and then the individual probabilities are multiplied

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**Figure 14.UN01 In-text Figure, Dihybrid Calculations, P. 278**

$$\begin{aligned} \text{Probability of } YYRR &= 1/4 (\text{probability of } YY) \times 1/4 (RR) = 1/16 \\ \text{Probability of } YyRR &= 1/2 (Yy) \times 1/4 (RR) = 1/8 \end{aligned}$$

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**Figure 14.UN02 In-text Figure, Trihybrid Probabilities, P. 278**

$ppyyRr$	$\frac{1}{4}$ (probability of $pp$ ) $\times$ $\frac{1}{2}$ ( $yy$ ) $\times$ $\frac{1}{2}$ ( $Rr$ )	$= \frac{1}{16}$
$ppYyrr$	$\frac{1}{4} \times \frac{1}{2} \times \frac{1}{2}$	$= \frac{1}{16}$
$Ppyyrr$	$\frac{1}{2} \times \frac{1}{2} \times \frac{1}{2}$	$= \frac{2}{16}$
$PPyyrr$	$\frac{1}{4} \times \frac{1}{2} \times \frac{1}{2}$	$= \frac{1}{16}$
$ppyyrr$	$\frac{1}{4} \times \frac{1}{2} \times \frac{1}{2}$	$= \frac{1}{16}$
<b>Chance of at least two recessive traits</b>		$= \frac{6}{16}$ or $\frac{3}{8}$

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An individual with the genotype  $AaBbEeHH$  is crossed with an individual who is  $aaBbEehh$ . What is the likelihood of having offspring with the genotype  $AabbEEHh$ ?

- a)  $\frac{1}{8}$
- b)  $\frac{1}{16}$
- c)  $\frac{1}{32}$
- d)  $\frac{1}{64}$
- e) That genotype would be impossible.

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**Concept 14.3: Inheritance patterns are often more complex than predicted by simple Mendelian genetics**

- The relationship between genotype and phenotype is rarely as simple as in the pea plant characters Mendel studied
- Many heritable characters are not determined by only one gene with two alleles
- However, the basic principles of segregation and independent assortment apply even to more complex patterns of inheritance

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**Extending Mendelian Genetics for a Single Gene**

- Inheritance of characters by a single gene may deviate from simple Mendelian patterns in the following situations:
  - When alleles are not completely dominant or recessive
  - When a gene has more than two alleles
  - When a gene produces multiple phenotypes

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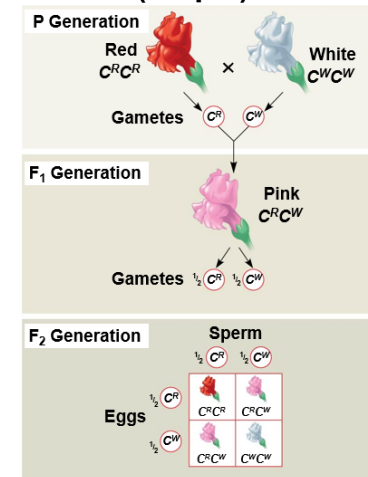
### Degrees of Dominance

- **Complete dominance** occurs when phenotypes of the heterozygote and dominant homozygote are identical
- In **incomplete dominance**, the phenotype of  $F_1$  hybrids is somewhere between the phenotypes of the two parental varieties
- In **codominance**, two dominant alleles affect the phenotype in separate, distinguishable ways

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### Figure 14.10\_3 Incomplete Dominance in Snapdragon Color (Step 3)



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### Degrees of Dominance, Continued

#### The Relationship Between Dominance and Phenotype

- In the case of pea shape, the dominant allele codes for an enzyme that converts an unbranched form of starch in the seed to a branched form
- The recessive allele codes for a defective form of the enzyme, which leads to an accumulation of unbranched starch
- This causes water to enter the seed, which then wrinkles as it dries

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### Degrees of Dominance, Continued-1

- **Tay-Sachs disease** is fatal; a dysfunctional enzyme causes an accumulation of lipids in the brain
  - At the organismal level, the allele is recessive
  - At the biochemical level, the phenotype (i.e., the enzyme activity level) is incompletely dominant
  - At the molecular level, the alleles are codominant

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**Degrees of Dominance, Continued-2**

**Frequency of Dominant Alleles**

- Dominant alleles are not necessarily more common in populations than recessive alleles
- One baby out of 400 in the United States is born with extra fingers or toes
- This condition, polydactyly, is caused by a dominant allele, found much less frequently in the population than the recessive allele

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**Multiple Alleles**



- Most genes exist in populations in more than two allelic forms
- For example, the four phenotypes of the ABO blood group in humans are determined by three alleles for the enzyme that attaches A or B carbohydrates to red blood cells:  $I^A$ ,  $I^B$ , and  $i$
- The enzyme encoded by the  $I^A$  allele adds the A carbohydrate, whereas the enzyme encoded by the  $I^B$  allele adds the B carbohydrate; the enzyme encoded by the  $i$  allele adds neither

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



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**Figure 14.11 Multiple Alleles for the ABO Blood Groups**

a) The three alleles for the ABO blood groups and their carbohydrates

Allele	$I^A$	$I^B$	$i$
Carbohydrate	A 	B 	none

b) Blood group genotypes and phenotypes

Genotype	$I^A I^A$ or $I^A i$	$I^B I^B$ or $I^B i$	$I^A I^B$	$ii$
Red blood cell with surface carbohydrates				
Phenotype (blood group)	A	B	AB	O

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**Pleiotropy**

- Most genes have multiple phenotypic effects, a property called **pleiotropy**
- For example, pleiotropic alleles are responsible for the multiple symptoms of certain hereditary diseases, such as cystic fibrosis and sickle-cell disease

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### Extending Mendelian Genetics for Two or More Genes

- Some traits may be determined by two or more genes
- In epistasis, one gene affects the phenotype of another due to interaction of their gene products
- In polygenic inheritance, multiple genes independently affect a single trait

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### Epistasis

- In **epistasis**, expression of a gene at one locus alters the phenotypic expression of a gene at a second locus
- For example, in Labrador retrievers and many other mammals, coat color depends on two genes
- One gene determines the pigment color (with alleles *B* for black and *b* for brown)
- The other gene (with alleles *E* for color and *e* for no color) determines whether the pigment will be deposited in the hair

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### Epistasis, Continued

- If heterozygous black labs (genotype *BbEe*) are mated, we might expect the dihybrid  $F_2$  ratio of 9:3:3:1
- However, a Punnett square shows that the phenotypic ratio will be 9 black to 3 chocolate to 4 yellow labs
- Epistatic interactions produce a variety of ratios, all of which are modified versions of 9:3:3:1

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### Figure 14.12 An Example of Epistasis

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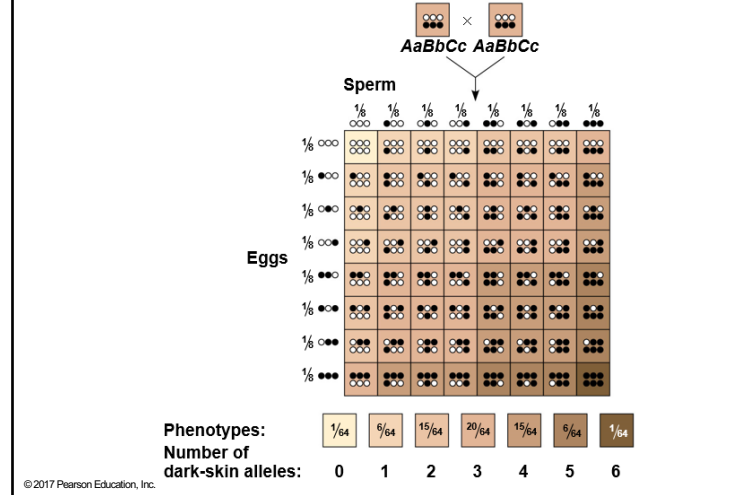
**Polygenic Inheritance**

- **Quantitative characters** are those that vary in the population along a continuum
- Quantitative variation usually indicates **polygenic inheritance**, an additive effect of two or more genes on a single phenotype
- Height is a good example of polygenic inheritance: Over 180 genes affect height
- Skin color in humans is also controlled by many separately inherited genes

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**Figure 14.13 A Simplified Model for Polygenic Inheritance of Skin Color**



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**Nature and Nurture: The Environmental Impact on Phenotype**

- Another departure from Mendelian genetics arises when the phenotype for a character depends on environment as well as genotype
- The phenotypic range is broadest for polygenic characters
- Traits that depend on multiple genes combined with environmental influences are called **multifactorial**

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**Figure 14.14 The Effect of Environment on Phenotype**



(a) Hydrangeas grown in basic soil

(b) Hydrangeas of the same genetic variety grown in acidic soil with free aluminum

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### A Mendelian View of Heredity and Variation

- An organism's phenotype includes its physical appearance, internal anatomy, physiology, and behavior
- An organism's phenotype reflects its overall genotype and unique environmental history

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### Concept 14.4: Many Human Traits follow Mendelian Patterns of Inheritance

- Humans are not good subjects for genetic research
  - Generation time is too long
  - Parents produce relatively few offspring
  - Breeding experiments are unacceptable
- However, basic Mendelian genetics endures as the foundation of human genetics

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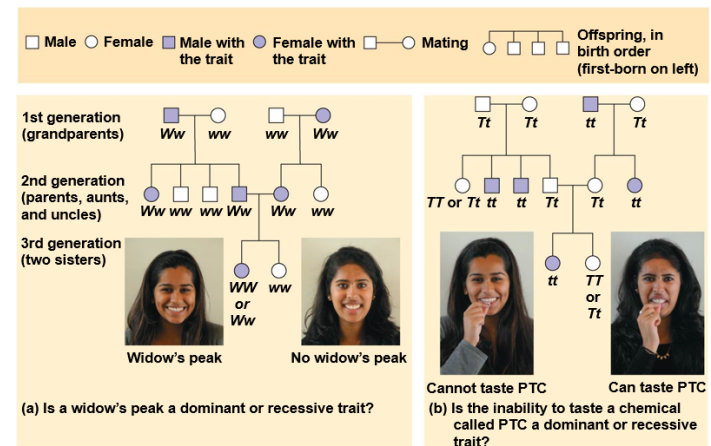
### Pedigree Analysis

- A **pedigree** is a family tree that describes the interrelationships of parents and children across generations
- Inheritance patterns of particular traits can be traced and described using pedigrees

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### Figure 14.15 Pedigree Analysis



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### Pedigree Analysis, Continued

- Pedigrees can also be used to make predictions about future offspring
- We can use the multiplication and addition rules to predict the probability of specific phenotypes

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### Recessively Inherited Disorders

- Many genetic disorders are inherited in a recessive manner
- These range from relatively mild to life-threatening

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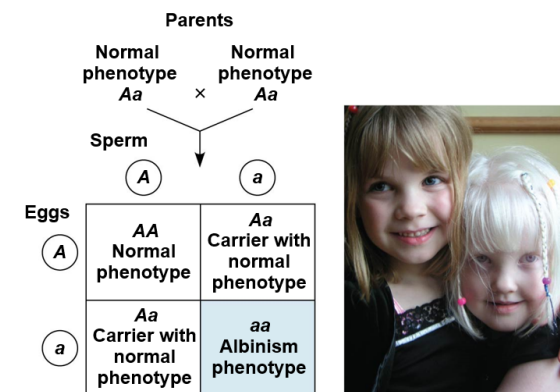
### The Behavior of Recessive Alleles

- Recessively inherited disorders show up only in individuals homozygous for the allele
- **Carriers** are heterozygous individuals who carry the recessive allele but are phenotypically normal
- Most individuals with recessive disorders are born to carrier parents
- Albinism is a recessive condition characterized by a lack of pigmentation in skin and hair

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### Figure 14.16 Albinism: A Recessive Trait



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### The Behavior of Recessive Alleles, Continued

- If a recessive allele that causes a disease is rare, it is unlikely that two carriers will meet and mate
- Consanguineous matings (i.e., between close relatives) increase the chance that both parents of a child carry the same rare allele
- Most societies and cultures have laws or taboos against marriages between close relatives

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### Cystic Fibrosis

- **Cystic fibrosis** is the most common lethal genetic disease in the United States, striking one out of every 2,500 people of European descent
- The cystic fibrosis allele results in defective or absent chloride transport channels in plasma membranes, leading to a buildup of chloride ions outside the cell
- Symptoms include mucus buildup in some internal organs and abnormal absorption of nutrients in the small intestine

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### Sickle-Cell Disease: A Genetic Disorder with Evolutionary Implications

- **Sickle-cell disease** affects one out of 400 African-Americans
- It is caused by the substitution of a single amino acid in the hemoglobin protein in red blood cells
- In homozygous individuals, all hemoglobin is abnormal (sickle-cell)
- Symptoms include physical weakness, pain, organ damage, and even paralysis

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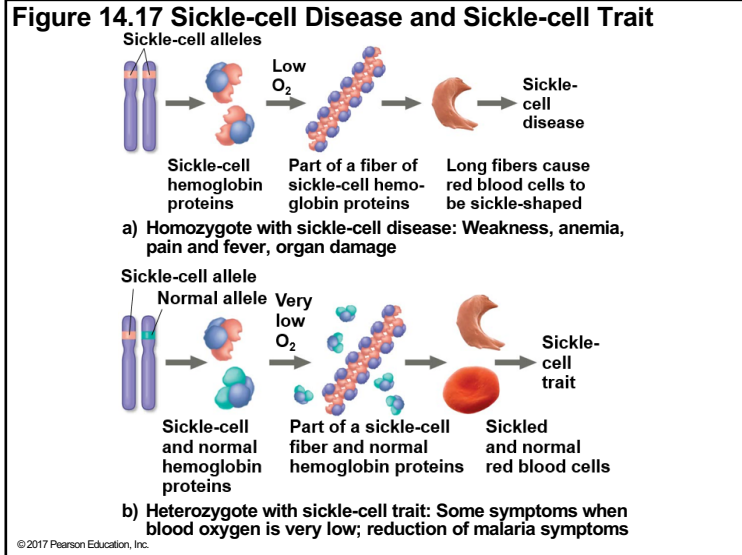
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### Sickle-Cell Disease: A Genetic Disorder with Evolutionary Implications, Continued

- Heterozygotes (said to have sickle-cell trait) are usually healthy but may suffer some symptoms
- About one out of ten African Americans has sickle-cell trait, an unusually high frequency
- Heterozygotes are less susceptible to the malaria parasite, so there is an advantage to being heterozygous in regions where malaria is common

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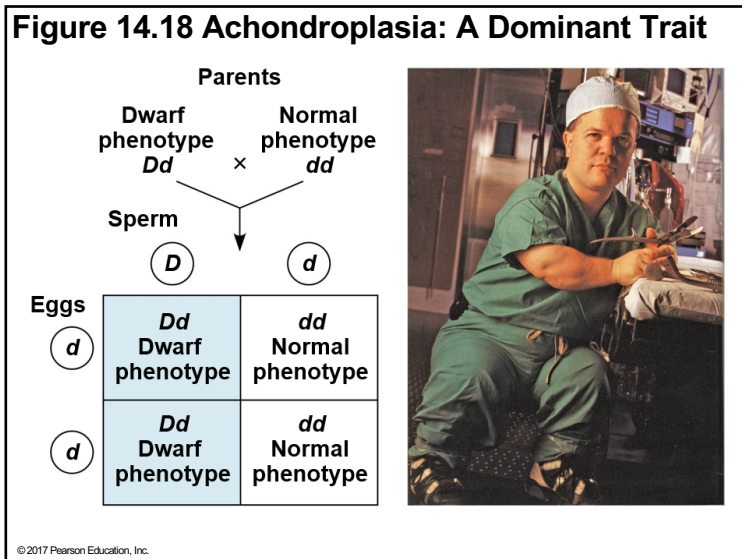
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**Dominantly Inherited Disorders**

- Some human disorders are caused by dominant alleles
- Dominant alleles that cause a lethal disease are rare and arise by mutation
- Achondroplasia is a form of dwarfism caused by a rare dominant allele

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**Dominantly Inherited Disorders, Continued**

- The timing of onset of a disease significantly affects its inheritance
- Huntington's disease** is a degenerative disease of the nervous system
- The disease has no obvious phenotypic effects until the individual is about 35 to 40 years of age
- Once the deterioration of the nervous system begins, the condition is irreversible and fatal

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### Multifactorial Disorders

- Many diseases, such as heart disease, cancer, alcoholism, and mental illnesses, have both genetic and environmental components
- No matter what our genotype, our lifestyle has a tremendous effect on phenotype

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### Genetic Testing and Counseling

- Genetic counselors can provide information to prospective parents concerned about a family history for a specific disease

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### *Counseling Based on Mendelian Genetics and Probability Rules*

- Suppose a couple both have a brother who died from the same recessively inherited disease
- A genetic counselor can help determine the risk that a couple will have a child with a particular disease
- It is important to remember that each child represents an independent event in the sense that its genotype is unaffected by the genotypes of older siblings

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### *Counseling Based on Mendelian Genetics and Probability Rules, Continued*

- If both members of the couple had a sibling with the recessively inherited illness, both of their parents were carriers
- Thus each has a  $\frac{2}{3}$  chance of being a carrier
- If both are carriers, there is a  $\frac{1}{4}$  chance of each child having the recessive illness
- The overall probability of them having a child with the illness is  $\frac{2}{3} \times \frac{2}{3} \times \frac{1}{4} = 1/9$

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### Tests for Identifying Carriers

- For a growing number of diseases, tests are available that identify carriers and help define the odds more accurately
- The tests enable people to make more informed decisions about having children
- However, they raise other issues, such as whether affected individuals fully understand their genetic test results, and how the test results are used

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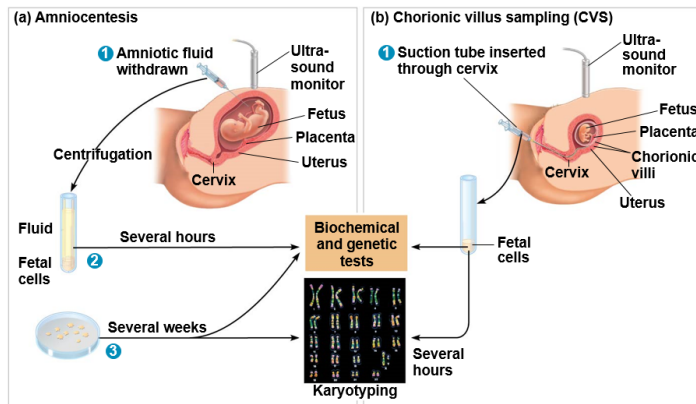
### Fetal Testing

- In **amniocentesis**, the liquid that bathes the fetus is removed and tested
- In **chorionic villus sampling (CVS)**, a sample of the placenta is removed and tested
- Other techniques, such as ultrasound, allow fetal health to be assessed visually *in utero*

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### Figure 14.19 Testing a Fetus for Genetic Disorders



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### Video: Ultrasound of Human Fetus



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### Newborn Screening

- Some genetic disorders can be detected at birth by simple tests that are now routinely performed in most hospitals in the United States
- One common test is for phenylketonuria (PKU), a recessively inherited disorder that occurs in one of every 10,000-15,000 births in the United States

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### Figure 14.UN05 Summary of Key Concepts: Single-gene Mendelian Extensions

Relationship among alleles of a single gene	Description	Example
Complete dominance of one allele	Heterozygous phenotype same as that of homozygous dominant	$PP$ $Pp$
Incomplete dominance of either allele	Heterozygous phenotype intermediate between the two homozygous phenotypes	$C^R C^R$ $C^R C^W$ $C^W C^W$
Codominance	Both phenotypes expressed in heterozygotes	$I^A I^B$
Multiple alleles	In the population, some genes have more than two alleles	ABO blood group alleles $I^A, I^B, i$
Pleiotropy	One gene affects multiple phenotypic characters	Sickle-cell disease

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### Figure 14.UN06 Summary of Key Concepts: Multi-gene Mendelian Extensions

Relationship among two or more genes	Description	Example
Epistasis	The phenotypic expression of one gene affects the expression of another gene	$BbEe$ $\times$ $BbEe$  9  : 3  : 4
Polygenic inheritance	A single phenotypic character is affected by two or more genes	$AaBbCc$ $\times$ $AaBbCc$ 

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John, age 47, has just been diagnosed with Huntington's disease, which is caused by a rare dominant allele. His daughter, age 25, has a 2-year-old son. No one else in the family has the disease. What is the probability that the daughter will develop the disease?

- a) 0%
- b) 25%
- c) 50%
- d) 75%
- e) 100%

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