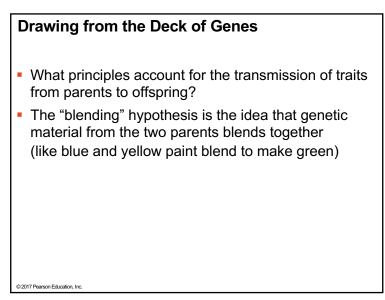


Drawing from the Deck of Genes, Continued

- 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



2

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



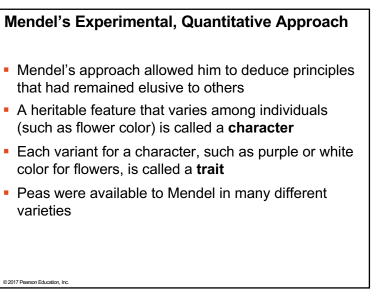
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Figure 14.1a What Principles of Inheritance did Gregor Mendel Discover by Breeding Garden Pea Plants? (Part 1: Mendel with Fellow Monks)



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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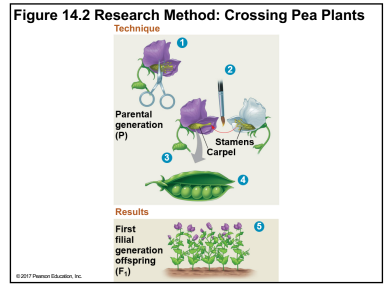
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Mendel's Experimental, Quantitative Approach, Continued

- 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

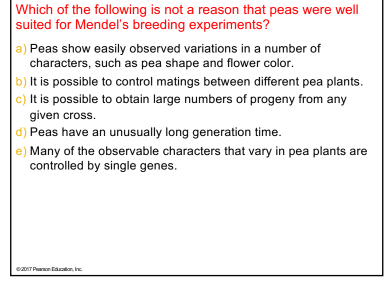
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Mendel's Experimental, Quantitative Approach, Continued-1

- Mendel chose to track only those characters that occurred in two distinct alternative forms
- He also started with varieties that were truebreeding (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 F₁ generation
- When F₁ individuals self-pollinate or cross-pollinate with other F₁ hybrids, the F₂ 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 F₁ hybrids were purple
- When Mendel crossed the F₁ hybrids, many of the F₂ plants had purple flowers, but some had white
- Mendel discovered a ratio of about three purple flowers to one white flower in the F₂ generation

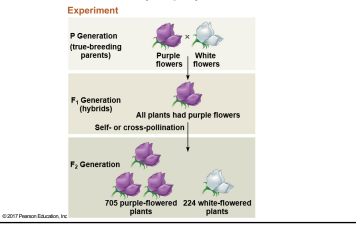
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The Law of Segregation, Continued

- Mendel reasoned that only the purple flower factor was affecting flower color in the F₁ 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 F₂ generation

Figure 14.3_3 Inquiry: When F₁ Hybrid Pea Plants Self- or Cross-pollinate, Which Traits Appear in the F₂ Generation? (Step 3)

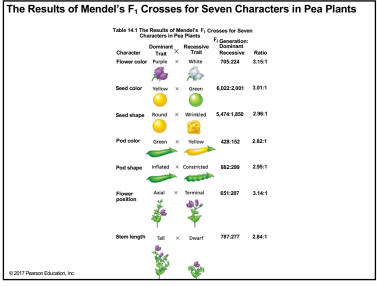


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The Law of Segregation, Continued-1

- 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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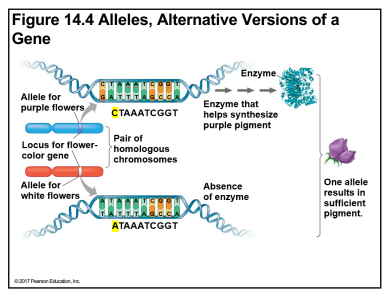


Mendel's Model, Continued

- 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

Mendel's Model Mendel developed a hypothesis to explain the 3:1 inheritance pattern he observed in F₂ 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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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 F₁ hybrids

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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

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 F₁ plants had purple flowers because the allele for that trait is dominant

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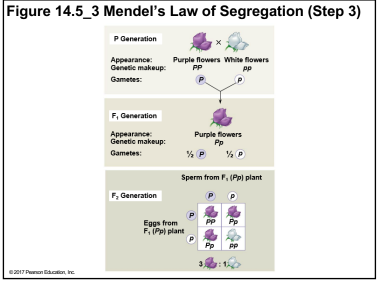
Mendel's Model, Continued-4

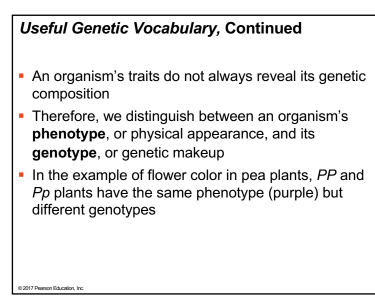
- The model accounts for the 3:1 ratio observed in the F₂ generation of Mendel's crosses
- Possible combinations of sperm and egg can be shown using a **Punnett square**
- A capital letter represents a dominant allele, and a lowercase letter represents a recessive allele

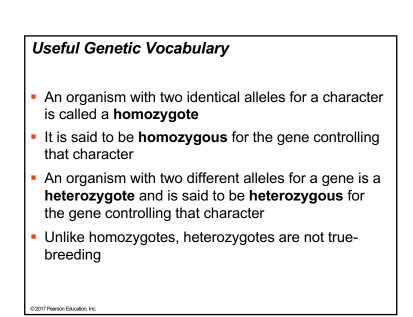
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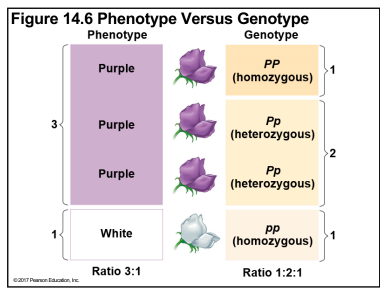
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The Testcross

- 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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The Law of Independent Assortment Mendel derived the law of segregation by following a single character The F₁ offspring produced in this cross were monohybrids, heterozygous for one character A cross between such heterozygotes is called a monohybrid cross

Figure 14.7 Rese	earch Method: 7	The Testcross
Technique	Dominant phenotype unknown genotype: <i>PP</i> or <i>Pp</i> ?	
	Predictions	×
	lf purple-flowered parent is <i>PP</i> :	or If purple-flowered parent is <i>Pp</i> :
	Sperm	Sperm
	(P) (P)	P P
	$\begin{array}{c} P \\ Eggs \\ P \\ \hline P \hline \hline $	$ \begin{array}{c} P \\ Eggs \\ P \end{array} \begin{array}{c} \not \\ P \\ p \end{array} \begin{array}{c} \not \\ P \\ p \\ p \end{array} \begin{array}{c} \not \\ P \\ p \\ p \end{array} \begin{array}{c} \not \\ P \\ p \\ p \end{array} \begin{array}{c} \not \\ P \\ p \\ p \end{array} $
Results		
	All offspring purple	1/2 offspring purple and 1/2 offspring white
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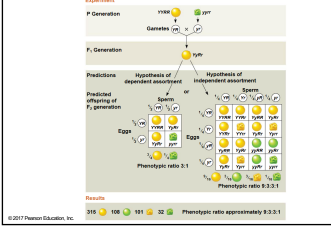
The Law of Independent Assortment, Continued

- 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 F₁ generation, heterozygous for both characters
- A dihybrid cross, a cross between F₁ 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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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

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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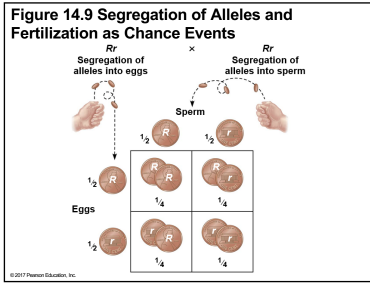
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The Multiplication and Addition Rules Applied to Monohybrid Crosses

- 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 F₁ monohybrid cross can be determined using the multiplication rule
- Segregation in a heterozygous plant is like flipping a coin: Each gamete has a ½ chance of carrying the dominant allele and a ½ chance of carrying the recessive allele

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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

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 F₂ plant from a monohybrid cross will be heterozygous rather than homozygous

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Figure 14.UN01 In-text Figure, Dihybrid Calculations, P. 278 Probability of $YYRR = \frac{1}{4}$ (probability of YY) × $\frac{1}{4}$ (RR) = $\frac{1}{6}$ Probability of $YyRR = \frac{1}{2}$ (Yy) × $\frac{1}{4}$ (RR) = $\frac{1}{8}$

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

Chance of at least two recessive traits	= ⁶ / ₁₆ or ³ / ₈
ppyyrr $1/_4 \times 1/_2 \times 1/_2$	= 1/16
PPyyrr $1/4 \times 1/2 \times 1/2$	= ¹ / ₁₆
Ppyyrr $1/2 \times 1/2 \times 1/2$	$= \frac{2}{16}$
$\mathbf{op} \mathbf{Yyrr} \ 1_{4} \times 1_{2} \times 1_{2}$	= 1/16

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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

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) 1/8 b) 1/16 c) 1/32 d) 1/64 e) That genotype would be impossible.

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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 F₁ 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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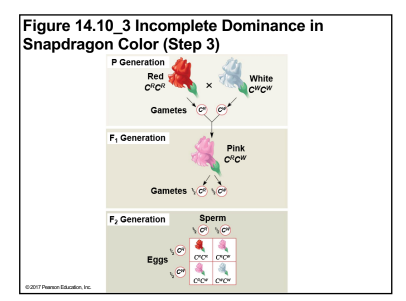
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Degrees of Dominance, Continued

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

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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Figure 14.11 Multiple Alleles for the ABO Blood Groups a) The three alleles for the ABO blood groups and their carbohydrates Allele ŢΑ ŢВ i Carbohydrate Α 🛆 в 🔾 none b) Blood group genotypes and phenotypes I^AI^A or I^Ai I^BI^B or I^Bi Genotype IAIB ii Red blood cell with surface carbohydrates Phenotype в AB 0 Α (blood group) © 2017 Pearson Education. Inc.

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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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, Continued

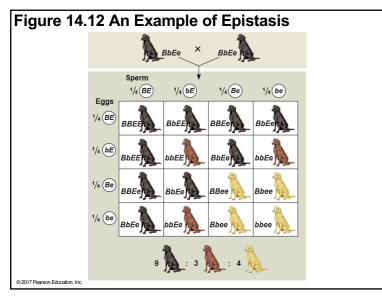
- If heterozygous black labs (genotype *BbEe*) are mated, we might expect the dihybrid F₂ ratio of 9:3:3:1
- However, a Punnett square shows that the phenotypic ratio will be 9 black to 3 chocolate to 4 yellow labs
- Epistatic interactions produce a variety of ratios, all of which are modified versions of 9:3:3:1

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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Polygenic Inheritance

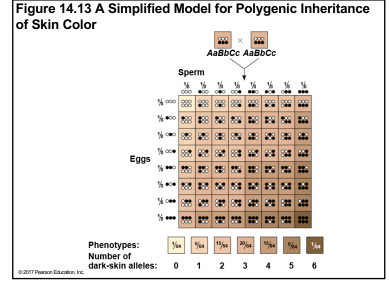
- Quantitative characters are those that vary in the population along a continuum
- Quantitative variation usually indicates polygenic inheritance, an additive effect of two or more genes on a single phenotype
- Height is a good example of polygenic inheritance: Over 180 genes affect height
- Skin color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

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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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Pedigree Analysis

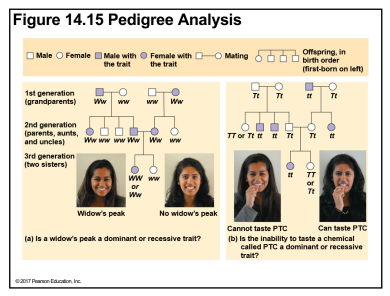
- A pedigree is a family tree that describes the interrelationships of parents and children across generations
- Inheritance patterns of particular traits can be traced and described using pedigrees

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, Continued

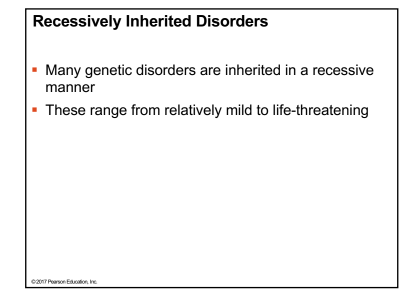
- Pedigrees can also be used to make predictions about future offspring
- We can use the multiplication and addition rules to predict the probability of specific phenotypes

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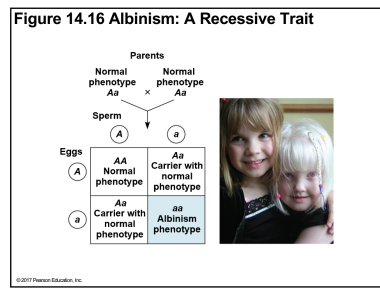
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The Behavior of Recessive Alleles

- 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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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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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

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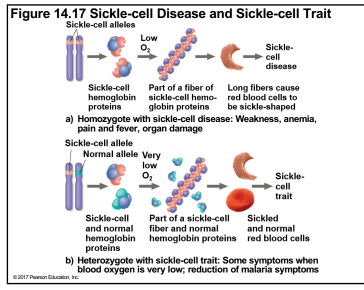
Sickle-Cell Disease: A Genetic Disorder with Evolutionary Implications, Continued

- Heterozygotes (said to have sickle-cell trait) are usually healthy but may suffer some symptoms
- About one out of ten African Americans has sicklecell 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

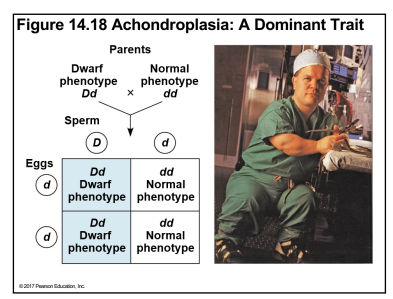
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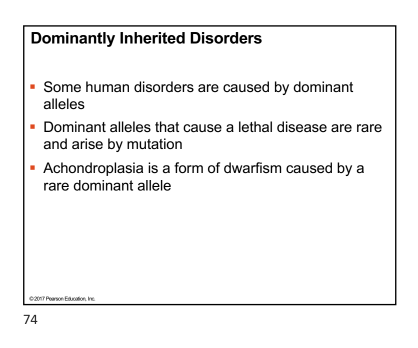
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Dominantly Inherited Disorders, Continued

- 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

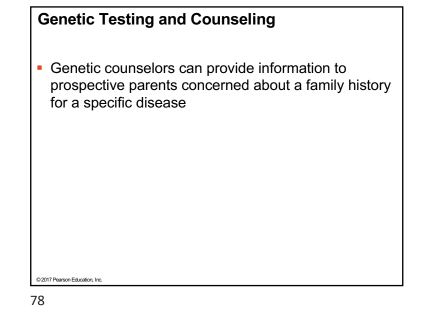
- Many diseases, such as heart disease, cancer, alcoholism, and mental illnesses, have both genetic and environmental components
- No matter what our genotype, our lifestyle has a tremendous effect on phenotype

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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



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 ²⁄₃ chance of being a carrier
- If both are carriers, there is a ¼ chance of each child having the recessive illness
- The overall probability of them having a child with the illness is ²/₃ × ²/₃ × ¹/₄ = 1/9

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Tests for Identifying Carriers

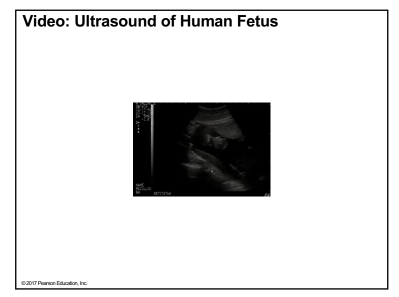
- For a growing number of diseases, tests are available that identify carriers and help define the odds more accurately
- The tests enable people to make more informed decisions about having children
- However, they raise other issues, such as whether affected individuals fully understand their genetic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

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Figure 14.19 Testing a Fetus for Genetic Disorders (a) Amniocentesis (b) Chorionic villus sampling (CVS) Ultra-Ultra- Amniotic fluid Suction tube inserted sound withdrawn sound through cervix monitor monitor Fetus etus Placenta Placenta Centrifugation Chorionic Uterus Cervix villi Cervix Uterus Fluid Biochemical Several hours and genetic Fetal cells Fetal 2 tests cells Several weeks Several hours © 2017 Pearson Education, Inc.

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*



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.UN06 Sum	mary of Key Concepts	s: 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 × BbEe BE bE Be bE A A Be A A Be A A A A A Be A A
Polygenic inheritance	A single phenotypic character is affected by two or more genes	AaBbCc x x x AaBbCc
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Relationship among alleles of a single gene	Description	Example
Complete dominance of one allele	Heterozygous phenotype same as that of homo- zygous dominant	РР 🍌 Рр 焼
Incomplete dominance of either allele	Heterozygous phenotype intermediate between the two homozygous phenotypes	🥰 ዺ 🔏 crcr crcw cwcw
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 , İ
Pleiotropy	One gene affects multiple phenotypic characters	Sickle-cell disease

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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?

- <mark>a)</mark> 0%
- <mark>b)</mark> 25%
- <mark>c)</mark> 50%
- <mark>d)</mark> 75%
- <mark>e)</mark> 100%

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