Genetics Practice Test

Modified True/False

Indicate whether the statement is true or false. If false, change the identified word or phrase to make the statement true.

1. A trait is a specific characteristic that varies from one individual to another. _________________________

2. True-breeding plants that produced axial flowers were crossed with true-breeding plants that produced terminal flowers. The resulting offspring all produced terminal flowers because the allele for terminal flowers is recessive. _________________________

3. During the formation of gametes in a hybrid tall plant, the tall allele and the short allele stay together. _________________________

4. If the alleles for a trait did not segregate during gamete formation, offspring would always show the trait of at least one of the parents. _________________________

5. The principles of probability can explain the numerical results of Mendel’s experiments. _________________________

6. The probability that a gamete produced by a pea plant heterozygous for stem height (Tt) will contain the recessive allele is 100%. _________________________

7. If Mendel had found that an F₂ cross of plants that were heterozygous for two traits had made offspring with two phenotypes, this finding would have supported the theory of independent assortment. _________________________

8. A trait in an unidentified plant is controlled by one gene that has two alleles. One allele is dominant over the other. According to Mendel’s principles, one fourth of the offspring made from a cross between two heterozygous plants will show the recessive trait. _________________________

9. If two speckled chickens are mated, according to the principle of codominance, 25% of the offspring are expected to be speckled. _________________________

10. Coat color in rabbits is determined by a single gene that has multiple alleles. _________________________

11. If an organism has 16 chromosomes in each of its egg cells, the organism’s diploid number is 32. _________________________

12. If an organism is heterozygous for a particular gene, the two different alleles will be separated during anaphase II of meiosis, assuming that no crossing-over has occurred. _________________________

13. Mitosis results in two cells, whereas meiosis results in one cell. _________________________

14. If an organism has four linkage groups, it has eight chromosomes. _________________________

15. Genes in the same linkage group are usually inherited separately. _________________________

16. In humans, the mother’s gamete determines the sex of the offspring. _________________________
17. In a human karyotype, 44 of the chromosomes are autosomes. _________________________

18. To make a karyotype, biologists take pictures of cells during mitosis, when they are condensed and easier to view. _________________________

19. A recessive X-linked trait would be more common in males than in females. _________________________

20. If a cat has both orange and black spots, it is homozygous for the alleles on the X chromosome that code for spot color. _________________________

21. If a person has blood type A, he or she cannot safely receive a blood transfusion from a person with blood type O. _________________________

22. Two parents who have Huntington disease may produce an offspring who does not have Huntington disease. _________________________

23. The pedigree chart in Figure 14-10 shows that some people can be carriers of the trait without being afflicted. This means the allele for the trait is dominant. _________________________

24. The pedigree chart in Figure 14-10 shows 4 generations. _________________________

25. A person who has Down syndrome has two copies of chromosome 21. _________________________

26. Down syndrome, Turner’s syndrome, and Klinefelter’s syndrome are all caused by nondisjunction. _________________________

27. When DNA fragments are separated by gel electrophoresis, the longest fragments move fastest. _________________________

28. When sequencing DNA, replication stops when nucleotide bases marked with dye are added to chains of DNA as the DNA is being synthesized. _________________________

29. A haplotype is a group of alleles found on the same chromosome that tend to be inherited all together. _________________________
30. Information from the Human Genome Project can be used to learn more about human diseases.

31. Animal breeders maintain cat and dog breeds by the process of hybridization.

32. Native Americans took teosine and used selective breeding to make corn, a more productive and nutritious plant.

33. Exposing bacteria to radiation or certain chemicals can increase the frequency of mutations that occur within the population.

34. A polyploid plant has more than two copies of each gene.

35. Scientists use genetic markers to determine which cells have been successfully transformed.

36. During a polymerase chain reaction, the number of DNA copies increases exponentially with each cycle.

37. Bacterial cells that have been transformed with a plasmid that carries a genetic marker for resistance to the antibiotic tetracycline will not survive in a culture treated with tetracycline.

38. To transform a plant cell, scientists inject protein into the plant cell.

39. To produce a cloned sheep, Ian Wilmut removed the nucleus from a sheep’s body cell and injected the cell with a nucleus taken from a body cell of another adult sheep.

40. Farmers growing Bt corn can use more insecticide than farmers growing the same amount of non-GM corn.

41. Transgenic plants are often used to simulate humans in medical tests.

42. DNA fingerprinting analyzes sections of DNA that have little or no known function but are similar from person to person.

43. Patenting genetic technology sometimes keeps technology out of the hands of people who need it.

44. Herbicide-resistance in crops can increase yields by allowing farmers to use fewer chemicals to control weeds.

45. Patenting a process, such as PCR, restricts the use of the process to the patent holder and to those who pay a fee.

Completion
Complete each statement.
46. The plants that Gregor Mendel crossed to produce the F₁ generation made up the ________________ generation.

47. Due to the process of segregation, alleles separate during the production of ________________.

48. An organism has 38 chromosomes in a body cell. After mitosis each cell has 38 chromosomes. After meiosis each gamete has ________________ chromosomes.

49. What is the probability of flipping a coin and getting heads 5 times in a row?

\[
\begin{array}{c|c|c}
 & T & t \\
\hline
Tt & T & t \\
TT & TT & Tt \\
T & TT & Tt \\
\end{array}
\]

\[
\begin{array}{c}
T = \text{Tall} \\
t = \text{Short}
\end{array}
\]

Figure 11–1

50. In the Punnett square shown in Figure 11–1, the genotypes of the offspring are ________________.

51. The principle of independent assortment states that ________________ for different traits can segregate independently during the formation of gametes.

52. If pea plants that are homozygous for round, yellow seeds (RRYY) were crossed with pea plants that are heterozygous for round, yellow seeds (RrYy), the expected phenotype(s) of the offspring would be ________________.

53. ________________’s principles can be used to study heredity in dogs, cats and sheep.

54. The reddish-brown pigment that gives color to a fruit-fly’s eye is controlled by three genes, so a fruit fly’s eye color is a ________________.
55. Western white butterflies that hatch in springtime have more pigment in their wings than those that hatch in summer. The darker wings help the butterflies stay warmer by absorbing more ________________ than the lighter-colored wings.

56. The characteristics of an organism are determined by two factors: ________________.

57. In four o’clock plants, flower color is controlled by two alleles that show ________________.

58. An organism’s gametes have ________________ the number of chromosomes found in the organism’s body cells.

59. Crossing-over occurs during the stage of meiosis called ________________.

60. The relative locations of each known gene can be shown on a ________________ map.

61. In humans, sex is determined by the X and ________________ chromosomes.

62. If a couple has five boys, the probability that the next child will be a boy is ________________.

63. A(An) ________________ can be used to determine whether a person has inherited the normal number of chromosomes.

64. A boy who is colorblind inherited the disorder from his ________________.

65. A Barr body is an inactivated ________________ chromosome.

66. The alleles I^A and ________________ for the AB blood group are codominant.

67. A person who has blood type O can safely receive a blood transfusion only from a person who has blood type ________________.

68. A(An) ________________ is a diagram that follows the inheritance of a single gene through several generations of a family.
69. The allele for the trait illustrated in the pedigree chart in Figure 14–3 is dominant. The probability of the couple labeled 2 in the pedigree having a child without the trait is ________________ percent.

70. People who have sickle cell disease inherited ________________ copies of the sickle cell allele.

71. A female with the disorder ________________ inherits only one X chromosome.

72. Nondisjunction can lead to the disorder called ________________, in which a male has an extra X chromosome.

73. An advantage of using a restriction enzyme such as EcoR1 is that it creates ________________ that can match to complimentary base pairs.

74. The new field of Bioinformatics combines the fields of ________________ and ________________.

75. The law that protects people from being discriminated against because of information learned in genetic tests is called the _________________.

76. ________________ is the technique of selective breeding that maintains desirable characteristics in a line of organisms, but increases the risk of genetic defects in certain animal breeds.

77. To produce a fruit that has some characteristics of an orange and some of a grapefruit, you would use the selective breeding technique of _________________.

78. Scientists use ________________ or ________________ to cause mutations in order to increase the genetic variation in bacteria.
79. Bananas planted as crops are____________________ plants which have 2 or 3 times the normal diploid number of chromosomes.

80. Some plasmids have genes that make bacteria resistant to ________________ and allow them serve as genetic markers.

81. __________________ are proteins that cut DNA at specific sequences, as illustrated in Figure 15–1.

82. DNA finger printing has dramatically changed the field of ________________, which is the study of crime scene evidence.

83. Using a ______________________ to locate a gene is like using a magnet to find a needle in a haystack.

84. If DNA fingerprints were made from DNA samples taken from Dolly and the sheep that donated the body cell, they would show ________________ patterns of bands on an electrophoresis gel.

85. The human growth hormone produced by ________ bacteria is identical to the human growth hormone produced by humans because both are coded by the same DNA sequence.

86. The process of _________________________ replaces a faulty gene with a normal, working gene.

87. In DNA fingerprinting, the DNA probe that is used is __________________ to the DNA sequence of the repeats in the sample.

88. All people serving in the United States military are required to give a __________________ for purposes of identification.
89. Critics of GM crops worry that GM plants engineered to be resistant to insects might unintentionally kill ______________ insects.

90. To prevent companies from misusing genetic information, the United States Congress passed the ________________.

Short Answer

91. What attributes of the garden pea plant made it an excellent organism for Gregor Mendel’s genetic studies?

92. Explain how allowing the F\textsubscript{1} generation plants to self-pollinate in producing the F\textsubscript{2} generation allowed Mendel to reach his conclusions about inheritance.

93. What is the difference between segregation and independent assortment?

<table>
<thead>
<tr>
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<th>RY</th>
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<th>rY</th>
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<td>RrYy</td>
<td>rrYy</td>
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</tr>
</tbody>
</table>

**Seed Shape**
- \( R \): Round
- \( r \): Wrinkled

**Seed Color**
- \( Y \): Yellow
- \( y \): Green

94. What is the phenotype ratio of the offspring of the plants in the Punnett square in Figure 11–2?

95. A tall pea plant with yellow seeds is heterozygous for height and seed color (\( TtYy \)). This plant is crossed with a pea plant heterozygous for height but homozygous recessive for seed color (\( Ttyy \)). If 80 offspring are produced, how many are expected to be tall and have yellow seeds?

96. Which kinds of organisms can be used to study Mendel’s principles of heredity?

97. What is the probability that a cross between parents who are both homozygous recessive for trait will have offspring that are homozygous recessive for that trait?

98. Arctic foxes have blue-gray fur in summer and change to white fur in winter. What would be one way to test whether this change is influenced by cooling seasonal temperatures?

99. How many sets of chromosomes are in a diploid cell?
100. Define *homologous chromosomes*.

101. What happens to the number of chromosomes per cell during meiosis?

102. Contrast the cells produced by mitosis with those produced by meiosis.

103. What are the only kinds of cells that undergo meiosis?

104. Why did Gregor Mendel not observe gene linkage during his experiments with pea plants?

105. Figure 11–5, the gene map of a fruit fly’s chromosome 2, shows the relative locations of the star eye, dumpy wing, and black body genes to be 1.3, 13.0, and 48.5, respectively. Between which two genes does crossing-over occur most frequently?

106. If human gametes have just 23 chromosomes, why does a human karyotype show 46?

107. Why are the sex chromosomes considered homologous, even though they vary in size and appearance?

108. Why are all X-linked alleles expressed in males, even if they are recessive?

109. A man who is not colorblind and a woman who is a carrier of the disorder have a son. What is the probability that their son will be colorblind?

110. Why are Barr bodies not found in male cells?

111. A person who has type AB blood is sometimes referred to as a universal recipient. Explain why.

112. What is a pedigree chart?
113. When making a pedigree that shows the inheritance of a recessive allele for a trait within a family, how do you know whether a certain individual should be represented by a shaded symbol?

114. How is the DNA sequence of the allele that causes cystic fibrosis different from that of the normal allele?

115. If malaria were eliminated from a certain area, how do you think the frequency of the sickle cell allele in that area would change? Explain.

116. Why is a person who has Klinefelter’s syndrome (XXY) a male, even though he has two X chromosomes in his cells?

117. How can you tell that the nondisjunction occurred during the formation of the gametes in Figure 14–7?

118. How are DNA fragments separated for sequencing?

119. How do restriction enzymes help make DNA manipulation possible?

120. What is the goal of the Human Genome Project?
121. How are the selective breeding techniques of hybridization and inbreeding opposites?

![Diagram of DNA structure with labeled sections A, B, C, and D]

Figure 15–1

122. What are structures C and D in Figure 15–1, and what is their significance?

123. Briefly explain what plasmids are and how they can be used to create transgenic organisms.

124. What does the polymerase chain reaction enable scientists to do?

125. How did Douglas Prasher find the gene that make the GFP protein?

126. What are two ways in which scientists transform plant cells without using plasmids?

127. Compare the genes in Dolly and the sheep from which she was cloned.

128. Describe how a scientist might make a transgenic animal.

129. How would making crops resistant to herbicides assist farmers?

130. Explain what a DNA probe is and how it could be used to identify a person who carries an allele for a genetic disorder.

131. Why are viruses used in gene therapy?

132. Why do scientists use sections of DNA repeats that have little or no known function to do DNA fingerprinting?
Heterozygous male guinea pigs with black, rough hair ($BbRr$) are crossed with heterozygous female guinea pigs with black, rough hair ($BbRr$). The incomplete Punnett square in Figure 11–6 shows the expected results from the cross.

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<tr>
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<td>$BbRr$</td>
<td>$Bbrr$</td>
<td>$bbRr$</td>
<td>$bbrr$</td>
</tr>
</tbody>
</table>

**Hair Color**
- $B =$ Black
- $b =$ White

**Hair Texture**
- $R =$ Rough
- $r =$ Smooth

**Figure 11–6**

136. **Interpret Tables** Identify the genotype of the offspring that would be represented by the question mark in Figure 11–6.

137. **Interpret Tables** Identify the phenotype of the offspring represented by the question mark in Figure 11–6.

138. **Analyze Data** In Figure 11–6, what are the different phenotypes of the offspring?

139. **Analyze Data** In Figure 11–6, what are the genotypes of the offspring that have black, rough hair?

140. **Calculate** What fraction of the offspring in Figure 11–6 would be expected to have white, smooth hair?
141. Infer What do the letters $R$ and $I$ represent in Figure 11–7?

142. Interpret Visuals In Figure 11–7, what is the genotype of the pink-flowered snapdragons?

143. Infer Explain whether the alleles in Figure 11–7 show dominance, incomplete dominance, or codominance.

144. Infer According to Figure 11–7, if red-flowered snapdragons and ivory-flowered snapdragons are crossed, what percentage of their offspring would be expected to be pink-flowered?

145. Infer Could the red snapdragons shown in the P generation of Figure 11–7 ever have white offspring? Explain your answer.
146. **Interpret Visuals** In Figure 11–8, what is the structure labeled X in stage A?

147. **Interpret Visuals** In Figure 11–8, during which stage might new allele combinations form? Identify the stage.

148. **Infer** If the stages shown in Figure 11–8 were taking place in a female animal, how many eggs would generally result from stage G? Explain your answer.

149. **Interpret Visuals** List the stages in Figure 11–8 in which the cells are 2N and those in which the cells are N.

150. **Infer** In Figure 11–8, in which stage does each cell have a single copy of each gene? Identify the stage.
151. **Infer** In the human karyotype in Figure 14–11, what term is used to describe the pair of chromosomes in each numbered group?

152. **Compare and Contrast** In Figure 14–11, how are the chromosomes that make up each numbered pair similar?

153. **Classify** Which chromosomes in Figure 14–11 are autosomes?

154. **Interpret Visuals** Study the human karyotype in Figure 14–11. Explain whether the person will be male or female, and describe the abnormality that this person has.

155. **Draw Conclusions** Study the abnormality at position 23 in Figure 14–11. What type of abnormality is this, and how might it have occurred?

To determine a person’s blood type, a drop of anti-A serum and a drop of anti-B serum are placed at either end of a microscope slide. Then, a drop of the person’s blood is added to each drop of serum. Clumping in anti-A serum or anti-B serum indicates the presence of antigen A or antigen B in the blood, respectively.
156. **Interpret Visuals** Which numbered slide in Figure 14–12 shows type B blood?

157. **Infer** In Figure 14–12, which slide shows the blood of a person who can safely receive any type of blood in a transfusion? Identify the person’s blood type.

158. **Interpret Visuals** Which slide in Figure 14–12 shows blood that contains no antigens? How do you know?

159. **Infer** In Figure 14–12, what is the genotype or possible genotypes of the person whose blood is shown in slide 1?

160. **Infer** Anti-sera are extracted from blood. Based on what is shown in Figure 14–12, what type of blood does not contain anti-A or anti-B serum? Explain why.

The pedigree shows the inheritance of free earlobes and attached earlobes in five generations of a family. Attached earlobes are caused by a recessive allele (f).
161. **Infer** Is individual 2 in Figure 14–13 homozygous or heterozygous for free earlobes? Explain.

162. **Interpret Visuals** In Figure 14–13, how many children of individuals 4 and 5 have attached earlobes?

163. **Infer** Can you be certain of the genotype of individual 5 in Figure 14–13? Explain.

164. **Predict** Predict the genotype and phenotype of individual 14 in Figure 14–13.

165. **Infer** In Figure 14–13, are any of the descendants of individuals 1 and 2 homozygous for free earlobes?

166. **Interpret Visuals** What process is being illustrated in Figure 15–4?

167. **Interpret Visuals** In Figure 15–4, during which numbered step(s) are bacteria reproducing?

168. **Interpret Visuals** During which numbered steps(s) in Figure 15–4 is a restriction enzyme used?

169. **Infer** How might the bacterial cell produced in step 6 in Figure 15–4 be useful?

170. **Interpret Visuals** In Figure 15–4, which numbered step produces a recombinant plasmid?
171. **Interpret Visuals** In the process shown in Figure 15–5, which sheep is the source of the nucleus in the fused cell?

172. **Infer** In Figure 15–5, why was the nucleus removed from the egg cell?

173. **Interpret Visuals** Which animal in Figure 15–5 is a clone?

174. **Interpret Visuals** In the cloning shown in Figure 15–5, which sheep provided an egg cell?

175. **Infer** Which two animals in Figure 15–5 are genetically identical?
176. **Interpret Visuals** In Figure 15–6, what do the bands shown in B consist of?

177. **Interpret Visuals** Which group of bands in Figure 15–6 moved faster, C or D? Why?

178. **Infer** What is occurring in A in Figure 15–6?

179. **Infer** In Figure 15–6, why are the bands in shown in B moving toward the positive end of the gel?

180. **Draw Conclusions** In Figure 15–6, were any of the three DNA samples from the same person? Explain your answer.

**Essay**

181. You wish to determine whether a tall pea plant is homozygous or heterozygous for tallness. What cross should you perform to arrive at your answer? Explain your choice of cross.

182. A pea plant with yellow seeds was crossed with a plant with green seeds. The F₁ generation produced plants with yellow seeds. Explain why green seeds reappeared in the F₂ generation.

183. Why are the results of genetic crosses shown in Punnett squares interpreted as probabilities, not certainties? Give some specific reasons.

184. A cross between two organisms heterozygous for two different genes (AaBb) likely results in a 9 : 3 : 3 : 1 phenotype ratio among the offspring. Is the offspring’s genotype ratio the same? Explain your answer.

185. A plant nursery owner wants to guarantee that the seeds she sells produce only pink-flowered four o’clock plants. How should she obtain the seeds?

186. Suppose you have western white butterflies from a population that hatched in spring, and those butterflies then had offspring that hatched in the summer. What would you expect the offspring born in the summer to look like? Explain your answer.

187. Which has more genetic information, a body cell or a gamete? Explain your answer.

188. The stages of meiosis are classified into two phases: meiosis I and meiosis II. Compare and contrast these two phases.

189. Explain why the daughter cells produced by meiosis are genetically different from each other, whereas the daughter cells produced by mitosis are not.

190. Define linkage, and explain how linkage is used to make gene maps.

191. How might karyotypes be useful to doctors?

192. What can you conclude if you observe a male cat that has both black and orange spots? Explain your answer.

193. Why does Huntington disease remain in the human population, even though it is fatal and is caused by a dominant allele?
194. Could the pedigree in Figure 14–10 show the pattern for the transmission of colorblindness? Explain your answer.

195. Explain why the father of a girl who is colorblind must also be colorblind.

196. Compare and contrast the causes of Turner’s syndrome and Down syndrome.


198. What two kinds of enzymes are used in sequencing DNA, and what does each enzyme do in the process?

199. Why might it be incorrect to assume that if one chromosome is larger than another, the larger chromosome has more genes?

200. Shotgun sequencing was one of the techniques used to sequence the human genome. Below are five DNA fragments—labeled A, B, C, D, and E, respectively—that were shotgun sequenced and determined to be part of the same DNA sequence. Notice that the fragments are single stranded. Determine the single-stranded DNA sequence that the fragments below are part of. Explain your reasoning.

Fragment A: GATCTAGGTCATG
Fragment B: ACAG
Fragment C: CAGTCTGATC
Fragment D: AGGTC
Fragment E: CATGCGATC

201. Explain an advantage and a disadvantage of inbreeding.

202. Suppose you want to produce a new animal breed or plant species with certain desirable traits. Write a brief description of the traits you would want the organism to have. Then, explain how you would use selective breeding techniques to produce an organism with those traits.

203. Suppose you are a scientist trying to help people who cannot produce an enzyme needed for proper digestion. How could you use genetic engineering techniques to make transformed bacteria that produce the enzyme?

204. Compare and contrast the techniques used in genetic engineering and in selective breeding to produce organisms with desired traits.
205. How might cloning be useful to animal breeders?

206. What are three general ways that scientists can use recombinant DNA technology to help improve human health?

207. Today, people who have hemophilia can be treated by receiving injections of normal clotting proteins. How would gene therapy be more beneficial to these people? Describe the general procedure that would be used.

208. How does DNA fingerprinting help preserve endangered species, such as elephants?

209. How does the right to patent genetic material and genetic engineering techniques both help and hinder scientific progress in understanding the human genome?

210. What is one potentially controversial use of genetic engineering in people, and what can be done to help manage this issue?
Genetics Practice Test
Answer Section

MODIFIED TRUE/FALSE

1. ANS: T  PTS: 1  DIF: L1
   REF: p. 309  OBJ: 11.1.1 Describe Mendel's studies and conclusions about inheritance.
   STA: UT.BIO.4.1.a | UT.BIO.4.2.a  TOP: Foundation Edition
   BLM: comprehension
2. ANS: F, dominant
   PTS: 1  DIF: L3  REF: p. 310
   OBJ: 11.1.1 Describe Mendel's studies and conclusions about inheritance.
   STA: UT.BIO.4.1.a | UT.BIO.4.2.a  BLM: application
3. ANS: F, separate
   PTS: 1  DIF: L1  REF: p. 325
   OBJ: 11.1.2 Describe what happens during segregation.
   STA: UT.BIO.4.2.a  BLM: knowledge
4. ANS: T  PTS: 1  DIF: L3
   REF: p. 324 | p. 325  OBJ: 11.1.2 Describe what happens during segregation.
   STA: UT.BIO.4.2.a  BLM: evaluation
5. ANS: T  PTS: 1  DIF: L2
   REF: p. 313
   OBJ: 11.2.1 Explain how geneticists use the principles of probability to make Punnett squares.
   STA: UT.BIO.4.2.a | UT.BIO.4.2.b  TOP: Foundation Edition
   BLM: comprehension
6. ANS: F
50%
   PTS: 1  DIF: L2  REF: p. 313 | p. 314
   OBJ: 11.2.1 Explain how geneticists use the principles of probability to make Punnett squares.
   STA: UT.BIO.4.2.a | UT.BIO.4.2.b  TOP: Foundation Edition
   BLM: application
7. ANS: F, contradicted
   PTS: 1  DIF: L3  REF: p. 317
   OBJ: 11.2.2 Explain the principle of independent assortment.
   STA: UT.BIO.4.2.a | UT.BIO.4.2.b  BLM: evaluation
8. ANS: T  PTS: 1  DIF: L2
   REF: p. 316 | p. 318
   OBJ: 11.2.3 Explain how Mendel's principles apply to all organisms.
   STA: UT.BIO.4.2.a  TOP: Foundation Edition
   BLM: analysis
9. ANS: F, 50%
   PTS: 1  DIF: L2  REF: p. 319
11.3.1 Describe the other inheritance patterns.

STA: UT.BIO.4.2.b

BLM: analysis

10. ANS: T

PTS: 1

DIF: L1

REF: p. 320

OBJ: 11.3.1 Describe the other inheritance patterns.

STA: UT.BIO.4.2.b

TOP: Foundation Edition

BLM: knowledge

11. ANS: T

PTS: 1

DIF: L2

REF: p. 323

OBJ: 11.4.1 Contrast the number of chromosomes in body cells and in gametes.

STA: UT.BIO.2.3.a

TOP: Foundation Edition

BLM: analysis

12. ANS: F, anaphase I

PTS: 1

DIF: L3

REF: p. 324 | p. 325

OBJ: 11.4.2 Summarize the events of meiosis.

STA: UT.BIO.2.3.a | UT.BIO.4.1.a

BLM: analysis

13. ANS: F, four cells

PTS: 1

DIF: L1

REF: p. 326 | p. 327

OBJ: 11.4.3 Contrast meiosis and mitosis.

STA: UT.BIO.2.3.a | UT.BIO.4.1.a

TOP: Foundation Edition

BLM: knowledge

14. ANS: F, four

PTS: 1

DIF: L2

REF: p. 328 | p. 329

OBJ: 11.4.4 Describe how alleles from different genes can be inherited together.

STA: UT.BIO.4.2.b

TOP: Foundation Edition

BLM: application

15. ANS: F, together

PTS: 1

DIF: L2

REF: p. 328 | p. 329

OBJ: 11.4.4 Describe how alleles from different genes can be inherited together.

STA: UT.BIO.4.2.b

TOP: Foundation Edition

BLM: application

16. ANS: F, father’s gamete

PTS: 1

DIF: L2

REF: p. 393

OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.

STA: UT.BIO.4.2.b

TOP: Foundation Edition

BLM: application

17. ANS: T

PTS: 1

DIF: L1

REF: p. 393

OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.

STA: UT.BIO.4.2.b

TOP: Foundation Edition

BLM: comprehension

18. ANS: T

PTS: 1

DIF: L2

REF: p. 392

OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.

STA: UT.BIO.4.2.b

TOP: Foundation Edition

BLM: application

19. ANS: F, less

PTS: 1

DIF: L2

REF: p. 395

OBJ: 14.1.2 Describe the patterns of the inheritance of human traits.
20. **ANS:** F, heterozygous  
**PTS:** 1  
**DIF:** L3  
**REF:** p. 396  
**OBJ:** 14.1.2 Describe the patterns of the inheritance of human traits.

21. **ANS:** F, can  
**PTS:** 1  
**DIF:** L1  
**REF:** p. 394  
**OBJ:** 14.1.2 Describe the patterns of the inheritance of human traits.

22. **ANS:** T  
**PTS:** 1  
**DIF:** L3  
**REF:** p. 399  
**OBJ:** 14.1.2 Describe the patterns of the inheritance of human traits.

23. **ANS:** F, recessive  
**PTS:** 1  
**DIF:** L2  
**REF:** p. 396 | p. 397  
**OBJ:** 14.1.3 Explain how pedigrees are used to study human traits.

24. **ANS:** F, 2  
**PTS:** 1  
**DIF:** L1  
**REF:** p. 396 | p. 397  
**OBJ:** 14.1.3 Explain how pedigrees are used to study human traits.

25. **ANS:** F, three  
**PTS:** 1  
**DIF:** L1  
**REF:** p. 401  
**OBJ:** 14.2.2 Summarize the problems caused by nondisjunction.

26. **ANS:** T  
**PTS:** 1  
**DIF:** L3  
**REF:** p. 401  
**OBJ:** 14.2.2 Summarize the problems caused by nondisjunction.

27. **ANS:** F, shortest  
**PTS:** 1  
**DIF:** L1  
**REF:** p. 403 | p. 404  
**OBJ:** 14.3.1 Summarize the methods of DNA analysis.

28. **ANS:** T  
**PTS:** 1  
**DIF:** L2  
**REF:** p. 404 | p. 405  
**OBJ:** 14.3.1 Summarize the methods of DNA analysis.

29. **ANS:** T  
**PTS:** 1  
**DIF:** L3  
**REF:** p. 407  
**OBJ:** 14.3.2 State the goals of the Human Genome Project and explain what we have learned so far.

30. **ANS:** T  
**PTS:** 1  
**DIF:** L1  
**REF:** p. 408  
**OBJ:** 14.3.2 State the goals of the Human Genome Project and explain what we have learned so far.
31. STA: UT.BIO.4.3.a | UT.BIO.4.3.e  BLM: knowledge
   ANS: F, inbreeding
   PTS: 1  DIF: L2  REF: p. 419
   OBJ: 15.1.1 Explain the purpose of selective breeding.
   TOP: Foundation Edition

32. STA: UT.BIO.4.2.c  BLM: knowledge
   ANS: T  PTS: 1  DIF: L2  REF: p. 418 | p. 419
   OBJ: 15.1.1 Explain the purpose of selective breeding.
   TOP: Foundation Edition

33. STA: UT.BIO.4.1.c  BLM: comprehension
   ANS: T  PTS: 1  DIF: L2  REF: p. 418 | p. 420
   OBJ: 15.1.2 Explain how people increase genetic variation.
   TOP: Foundation Edition

34. STA: UT.BIO.4.1.c  BLM: comprehension
   ANS: T  PTS: 1  DIF: L2  REF: p. 425
   OBJ: 15.2.1 Explain how scientists manipulate DNA.
   TOP: Foundation Edition

35. STA: UT.BIO.4.3.a  BLM: knowledge
   ANS: T  PTS: 1  DIF: L3  REF: p. 425
   OBJ: 15.2.1 Explain how scientists manipulate DNA.
   TOP: Foundation Edition

36. STA: UT.BIO.4.3.a  BLM: synthesis
   ANS: F, will
   PTS: 1  DIF: L2  REF: p. 425
   OBJ: 15.2.2 Describe the importance of recombinant DNA.
   TOP: Foundation Edition

37. STA: UT.BIO.4.3.e  BLM: application
   ANS: F, DNA
   PTS: 1  DIF: L2  REF: p. 425
   OBJ: 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.
   TOP: Foundation Edition

38. STA: UT.BIO.4.3.f  BLM: comprehension
   ANS: F, egg
   PTS: 1  DIF: L1  REF: p. 425
   OBJ: 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.
   TOP: Foundation Edition

39. STA: UT.BIO.4.3.f  BLM: knowledge
   ANS: F, less
   PTS: 1  DIF: L3  REF: p. 425
   OBJ: 15.3.1 Describe the benefits of genetic engineering as they relate to agriculture and industry.
   TOP: Foundation Edition

40. STA: UT.BIO.4.3.f | UT.BIO.5.1.d  BLM: analysis
   ANS: F, animals
   PTS: 1  DIF: L2  REF: p. 430
15.3.2 Explain how recombinant DNA technology can improve human health.

ANS: F

PTS: 1  DIF: L2  REF: p. 433 | p. 434

15.3.3 Summarize the process of DNA fingerprinting and explain its uses.

ANS: T

PTS: 1  DIF: L1  REF: p. 436

15.4.1 Describe some of the issues that relate to biotechnology.

ANS: F, more

PTS: 1  DIF: L2  REF: p. 438

15.4.2 Identify some of the pros and cons of genetically modified food.

ANS: T

PTS: 1  DIF: L2  REF: p. 439

15.4.3 Describe some of the ethical issues relating to biotechnology.

PTS: 1  DIF: L2  REF: p. 439

COMPLETION

ANS: P

PTS: 1  DIF: L2  REF: p. 310

11.1.1 Describe Mendel’s studies and conclusions about inheritance.

ANS: gametes, sex cells

PTS: 1  DIF: L2  REF: p. 314

11.1.2 Describe what happens during segregation.

ANS: 19

PTS: 1  DIF: L3  REF: p. 324

11.2.1 Explain how geneticists use the principles of probability to make Punnett squares.

ANS: 1/2 x 1/2 x 1/2 x 1/2 x 1/2 = 1/32

PTS: 1  DIF: L3  REF: p. 313
50. **ANS:** *TT* and *Tt*

**PTS:** 1  
**DIF:** L2  
**REF:** p. 314

**OBJ:** 11.2.1 Explain how geneticists use the principles of probability to make Punnett squares. 

**STA:** UT.BIO.4.2.a | UT.BIO.4.2.b  
**TOP:** Foundation Edition  
**BLM:** analysis

51. **ANS:** genes, chromosomes

**PTS:** 1  
**DIF:** L1  
**REF:** p. 317 | p. 329

**OBJ:** 11.2.2 Explain the principle of independent assortment. 

**STA:** UT.BIO.4.2.a | UT.BIO.4.2.b  
**TOP:** Foundation Edition  
**BLM:** knowledge

52. **ANS:** round yellow seeds only

**PTS:** 1  
**DIF:** L2  
**REF:** p. 317

**OBJ:** 11.2.2 Explain the principle of independent assortment. 

**STA:** UT.BIO.4.2.a | UT.BIO.4.2.b  
**TOP:** Foundation Edition  
**BLM:** analysis

53. **ANS:** Mendel

**PTS:** 1  
**DIF:** L1  
**REF:** p. 318

**OBJ:** 11.2.3 Explain how Mendel's principles apply to all organisms. 

**STA:** UT.BIO.4.2.a  
**TOP:** Foundation Edition  
**BLM:** application

54. **ANS:** polygenic trait

**PTS:** 1  
**DIF:** L2  
**REF:** p. 320

**OBJ:** 11.3.1 Describe the other inheritance patterns. 

**STA:** UT.BIO.4.2.b  
**BLM:** comprehension

55. **ANS:** light energy

**PTS:** 1  
**DIF:** L3  
**REF:** p. 321

**OBJ:** 11.3.2 Explain the relationship between genes and the environment. 

**STA:** UT.BIO.5.1.a  
**TOP:** Foundation Edition  
**BLM:** evaluation

56. **ANS:** genes and environmental conditions

**PTS:** 1  
**DIF:** L1  
**REF:** p. 321

**OBJ:** 11.3.2 Explain the relationship between genes and the environment. 

**STA:** UT.BIO.5.1.a  
**TOP:** Foundation Edition  
**BLM:** knowledge

57. **ANS:** incomplete dominance

**PTS:** 1  
**DIF:** L1  
**REF:** p. 319 | p. 320

**OBJ:** 11.3.2 Explain the relationship between genes and the environment. 

**STA:** UT.BIO.5.1.a  
**TOP:** Foundation Edition  
**BLM:** analysis

58. **ANS:** half

**PTS:** 1  
**DIF:** L2  
**REF:** p. 323

**OBJ:** 11.4.1 Contrast the number of chromosomes in body cells and in gametes. 

**STA:** UT.BIO.2.3.a  
**TOP:** Foundation Edition
59. BLM: comprehension
ANS: prophase I

PTS: 1  DIF: L2  REF: p. 324 | p. 325
OBJ: 11.4.2 Summarize the events of meiosis.
STA: UT.BIO.2.3.a | UT.BIO.4.1.a
BLM: analysis

60. BLM: analysis
ANS: gene

PTS: 1  DIF: L1  REF: p. 328 | p. 329
OBJ: 11.4.4 Describe how alleles from different genes can be inherited together.
STA: UT.BIO.4.2.b  TOP: Foundation Edition
BLM: knowledge

61. BLM: knowledge
ANS: Y

PTS: 1  DIF: L1  REF: p. 393
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b  BLM: knowledge

62. BLM: knowledge
ANS: 50%

PTS: 1  DIF: L2  REF: p. 393
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b  TOP: Foundation Edition
BLM: application

63. BLM: application
ANS: karyotype

PTS: 1  DIF: L2  REF: p. 392
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b  TOP: Foundation Edition
BLM: application

64. BLM: application
ANS: mother

PTS: 1  DIF: L1  REF: p. 395
OBJ: 14.1.2 Describe the patterns of the inheritance of human traits.
STA: UT.BIO.4.2.b  BLM: analysis

65. BLM: analysis
ANS: X

PTS: 1  DIF: L2  REF: p. 396
OBJ: 14.1.2 Describe the patterns of the inheritance of human traits.
STA: UT.BIO.4.2.b  BLM: comprehension

66. BLM: comprehension
ANS: O

PTS: 1  DIF: L3  REF: p. 394
14.1.2 Describe the patterns of the inheritance of human traits.
STA: UT.BIO.4.2.b
BLM: application

ANS: pedigree

PTS: 1   DIF: L1   REF: p. 396 | p. 397

14.1.3 Explain how pedigrees are used to study human traits.
STA: UT.BIO.4.2.b
BLM: knowledge

ANS: 25

PTS: 1   DIF: L3   REF: p. 397

14.2.1 Explain how small changes in DNA cause genetic disorders.
STA: UT.BIO.4.3.d
TOP: Foundation Edition
BLM: application

ANS: Turner’s syndrome

PTS: 1   DIF: L2   REF: p. 401

14.2.2 Summarize the problems caused by nondisjunction.
STA: UT.BIO.4.3.d
TOP: Foundation Edition
BLM: comprehension

ANS: Klinefelter’s syndrome

PTS: 1   DIF: L3   REF: p. 401

14.3.1 Summarize the methods of DNA analysis.
STA: UT.BIO.4.3.a | UT.BIO.4.3.e
BLM: comprehension

ANS: Life Science, Information Science

PTS: 1   DIF: L3   REF: p. 407

14.3.2 State the goals of the Human Genome Project and explain what we have learned so far.
STA: UT.BIO.4.3.a | UT.BIO.4.3.e
BLM: synthesis

ANS: Genetic Information Nondiscrimination Act

PTS: 1   DIF: L2   REF: p. 409

15.1.1 Explain the purpose of selective breeding.
STA: UT.BIO.4.2.c
TOP: Foundation Edition
BLM: knowledge

ANS: Inbreeding

PTS: 1   DIF: L1   REF: p. 419
77. ANS: hybridization

PTS: 1    DIF: L2    REF: p. 418
OBJ: 15.1.1 Explain the purpose of selective breeding.  STA: UT.BIO.4.2.c
TOP: Foundation Edition    BLM: analysis

78. ANS: radiation, chemicals

PTS: 1    DIF: L3    REF: p. 420
OBJ: 15.1.2 Explain how people increase genetic variation.  STA: UT.BIO.4.1.c
BLM: synthesis

79. ANS: polyploid

PTS: 1    DIF: L1    REF: p. 420
OBJ: 15.1.2 Explain how people increase genetic variation.  STA: UT.BIO.4.1.c
TOP: Foundation Edition    BLM: knowledge

80. ANS: antibiotics

PTS: 1    DIF: L2    REF: p. 425
OBJ: 15.2.1 Explain how scientists manipulate DNA.  STA: UT.BIO.4.3.a
TOP: Foundation Edition    BLM: comprehension

81. ANS: Restriction enzymes

PTS: 1    DIF: L2    REF: p. 424
OBJ: 15.2.1 Explain how scientists manipulate DNA.  STA: UT.BIO.4.3.a
TOP: Foundation Edition    BLM: analysis

82. ANS: forensics

PTS: 1    DIF: L2    REF: p. 433 | p. 434
OBJ: 15.2.1 Explain how scientists manipulate DNA.  STA: UT.BIO.4.3.a
TOP: Foundation Edition    BLM: application

83. ANS: DNA probe

PTS: 1    DIF: L2    REF: p. 421
OBJ: 15.2.2 Describe the importance of recombinant DNA.  STA: UT.BIO.4.3.e
TOP: Foundation Edition    BLM: synthesis

84. ANS: identical

identical
the same

PTS: 1    DIF: L3    REF: p. 427
OBJ: 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.  STA: UT.BIO.4.3.f
BLM: synthesis

85. ANS: transgenic

transgenic
genetically engineered

PTS: 1    DIF: L3    REF: p. 425
OBJ: 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.  STA: UT.BIO.4.3.f
BLM: analysis

86. ANS: gene therapy
87. **ANS:** complementary

88. **ANS:** DNA sample

89. **ANS:** beneficial, helpful, needed, good

90. **ANS:** Genetic Information Nondiscrimination Act.

91. **ANS:**
   Garden pea plants produce many offspring, they have traits that come in only two forms, and crosses between the plants can be controlled easily.

92. **ANS:**
   Allowing the F₁ pea plants to self-pollinate caused the recessive phenotype to reappear in the F₂ generation. Self-pollination of the F₁ plants also allowed the 3:1 phenotype ratios to occur, supporting Mendel’s theory. Self-pollination showed that traits controlled by recessive alleles could reappear in the F₂ generation.
Segregation happens when the alleles for each gene separate. Each gamete gets only one allele for each gene. The principle of independent assortment states that the way alleles for one pair of genes segregate does not affect the segregation of other alleles.

The phenotype ratio is 9 round, yellow seeds : 3 round, green seeds : 3 wrinkled, yellow seeds : 1 wrinkled, green seed.

Thirty of the offspring are expected to be tall and have yellow seeds.

Mendel’s principles of heredity apply to all organisms.

A diploid cell has two sets of chromosomes.
Homologous chromosomes are the two sets of chromosomes found in a body cell—one set inherited from the male parent and the other inherited from the female parent.

PTS: 1 DIF: L2 REF: p. 323
OBJ: 11.4.1 Contrast the number of chromosomes in body cells and in gametes.
STA: UT.BIO.2.3.a TOP: Foundation Edition
BLM: comprehension

101. ANS:
The number of chromosomes is cut in half.

PTS: 1 DIF: L1 REF: p. 324
OBJ: 11.4.2 Summarize the events of meiosis.
STA: UT.BIO.2.3.a | UT.BIO.4.1.a TOP: Foundation Edition
BLM: knowledge

102. ANS:
Mitosis produces diploid body cells, whereas meiosis produces haploid gametes.

PTS: 1 DIF: L2 REF: p. 327 OBJ: 11.4.3 Contrast meiosis and mitosis.
STA: UT.BIO.2.3.a | UT.BIO.4.1.a TOP: Foundation Edition
BLM: analysis

103. ANS:
sex cells, gametes

PTS: 1 DIF: L1 REF: p. 323 | p. 325
OBJ: 11.4.3 Contrast meiosis and mitosis.
STA: UT.BIO.2.3.a | UT.BIO.4.1.a TOP: Foundation Edition
BLM: knowledge

104. ANS:
The genes that Mendel studied were located on different chromosomes or were located far apart on the same chromosome.

PTS: 1 DIF: L2 REF: p. 328 | p. 329
OBJ: 11.4.4 Describe how alleles from different genes can be inherited together.
STA: UT.BIO.4.2.b BLM: evaluation

105. ANS:
Crossing-over occurs most frequently between the star eye gene and the black body gene.

PTS: 1 DIF: L3 REF: p. 328 | p. 329
OBJ: 11.4.4 Describe how alleles from different genes can be inherited together.
STA: UT.BIO.4.2.b BLM: synthesis

106. ANS:
A sperm that has 23 chromosomes fertilizes an egg that has 23 chromosomes resulting in a 23 pairs of chromosomes (46 total) in the autosomal cells of the individual.

PTS: 1 DIF: L2 REF: p. 392 | p. 393
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b TOP: Foundation Édition
BLM: comprehension

107. ANS:
The sex chromosomes are homologous because one sex chromosome is inherited from one parent, and the other is inherited from the other parent.
Males have just one X chromosome.

The probability that their son will be colorblind is 50%.

Male cells do not have two X chromosomes.

A person who has type AB blood can safely receive transfusions of all ABO blood types.

A pedigree chart is a drawing that shows the genetic relationships within a family.

The symbol should be shaded if the individual has the trait.

The DNA sequence of the allele that causes cystic fibrosis has a deletion of three bases.
115. **ANS:**
The frequency of the sickle cell allele would probably decrease because the allele would no longer be beneficial in heterozygous individuals.

**PTS:** 1  
**DIF:** L3  
**REF:** p. 400  
**OBJ:** 14.2.1 Explain how small changes in DNA cause genetic disorders.  
**STA:** UT.BIO.4.3.d  
**BLM:** evaluation

116. **ANS:**
If human cells have a Y chromosome, the person is a male regardless of how many X chromosomes are in the cells.

**PTS:** 1  
**DIF:** L2  
**REF:** p. 401  
**OBJ:** 14.2.2 Summarize the problems caused by nondisjunction.  
**STA:** UT.BIO.4.3.d  
**TOP:** Foundation Edition  
**BLM:** analysis

117. **ANS:**
You can tell that nondisjunction occurred because the gametes have an abnormal number of chromosomes.

**PTS:** 1  
**DIF:** L1  
**REF:** p. 401  
**OBJ:** 14.2.2 Summarize the problems caused by nondisjunction.  
**STA:** UT.BIO.4.3.d  
**BLM:** analysis

118. **ANS:**
The fragments are separated using gel electrophoresis.

**PTS:** 1  
**DIF:** L1  
**REF:** p. 403 | p. 404  
**OBJ:** 14.3.1 Summarize the methods of DNA analysis.  
**STA:** UT.BIO.4.3.a | UT.BIO.4.3.e  
**BLM:** comprehension

119. **ANS:**
Restriction enzymes cut large DNA molecules into smaller, more manageable pieces.

**PTS:** 1  
**DIF:** L2  
**REF:** p. 403  
**OBJ:** 14.3.1 Summarize the methods of DNA analysis.  
**STA:** UT.BIO.4.3.a | UT.BIO.4.3.e  
**TOP:** Foundation Edition  
**BLM:** synthesis

120. **ANS:**
The goal of the Human Genome Project is to attempt to sequence all human DNA.

**PTS:** 1  
**DIF:** L1  
**REF:** p. 406  
**OBJ:** 14.3.2 State the goals of the Human Genome Project and explain what we have learned so far.  
**STA:** UT.BIO.4.3.a | UT.BIO.4.3.e  
**BLM:** knowledge

121. **ANS:**
In hybridization, organisms with dissimilar traits are crossed. In inbreeding, organisms with similar traits are crossed.

**PTS:** 1  
**DIF:** L3  
**REF:** p. 419 | p. 420  
**OBJ:** 15.1.1 Explain the purpose of selective breeding.  
**STA:** UT.BIO.4.2.c  
**BLM:** analysis

122. **ANS:**
Structures C and D are the sticky ends of a DNA fragment, which allow the fragment to be inserted into a piece of DNA that has complementary sticky ends.
A plasmid is a circular DNA molecule that is naturally found in bacteria. Scientists use them to transform bacteria and plants.

The polymerase chain reaction enables scientists to make many copies of a gene.

He used the amino acid sequence of GFP and the genetic code to find the mRNA sequences that code GFP. Once he had the mRNA, he used it as a probe to find its DNA complement, which was the GFP gene.

Scientists inject DNA directly into plant cells. They also remove the cell walls of plant cells and allow the cells to take up DNA on their own.

Dolly and the sheep from which she was cloned have identical genes.

Sample answer: Scientists can inject DNA directly into the nucleus of an egg cell, where it may be inserted into the chromosomes.

Farmers can spray their crops with herbicides that will kill the weeds and leave the crop plants unharmed.
A DNA probe is short piece of DNA designed to detect a certain gene. A probe can be made to be complementary to part of the sequence of a disease-causing allele, and it will only bind to that specific allele. This enables scientists to see who has the allele and who does not.

Viruses are used in gene therapy because they can transfer genes into human cells.

These sections of DNA vary widely from person to person.

Because the DNA in the Y chromosome and mitochondria do not undergo crossing over and are passed from parents to children unchanged. If two Y chromosomes or mtDNA samples from two people match, there is a good chance they are related.

Using herbicide resistant plants might encourage farmers to spray much more herbicide into the environment.

No. The Genetic Information Nondiscrimination Act protects against discrimination based on genetic information.
The genotype of the offspring is \( bbRR \).

137. \textbf{ANS:} The phenotype of the offspring is white, rough hair.

138. \textbf{ANS:} The phenotypes of the offspring are black, rough hair; black, smooth hair; white, rough hair; and white, smooth hair.

139. \textbf{ANS:} Offspring with black, rough hair have the genotypes \( BBRR, BBRr, BbRR, \) and \( BbRr \).

140. \textbf{ANS:} One sixteenth of the offspring would be expected to have white, smooth hair.

141. \textbf{ANS:} \( R \) represents the allele for red flowers. \( I \) represents the allele for ivory flowers.

142. \textbf{ANS:} The genotype of the pink-flowered snapdragons is \( RI \).
One hundred percent of the offspring would be expected to be pink-flowered.

PTS: 1  DIF: L2  REF: p. 319
OBJ: 11.3.1 Describe the other inheritance patterns.  STA: UT.BIO.4.2.b
TOP: Foundation Edition  BLM: application

145.  ANS:
It is not possible for the red snapdragon to have white offspring. They are homozygous (RR) and therefore do not carry any white alleles. The red snapdragons in Figure 11–7 will always pass the R allele on to offspring and therefore the offspring can only be red or pink.

PTS: 1  DIF: L3  REF: p. 319 | p. 320
OBJ: 11.3.1 Describe the other inheritance patterns.  STA: UT.BIO.4.2.b
BLM: evaluation

146.  ANS:
The structure is a tetrad.

PTS: 1  DIF: L1  REF: p. 324
OBJ: 11.4.2 Summarize the events of meiosis.  STA: UT.BIO.2.3.a | UT.BIO.4.1.a
TOP: Foundation Edition  BLM: analysis

147.  ANS:
New allele combinations might form during stage A, which is prophase I.

PTS: 1  DIF: L3  REF: p. 324
OBJ: 11.4.2 Summarize the events of meiosis.  STA: UT.BIO.2.3.a | UT.BIO.4.1.a
BLM: synthesis

148.  ANS:
Generally, one egg would result. One of the four haploid cells would form an egg.

PTS: 1  DIF: L2  REF: p. 324 | p. 325
OBJ: 11.4.2 Summarize the events of meiosis.  STA: UT.BIO.2.3.a | UT.BIO.4.1.a
TOP: Foundation Edition  BLM: comprehension

149.  ANS:
The cells in stages A, B, and C are 2N. The cells in stages D, E, F, and G are N.

PTS: 1  DIF: L3  REF: p. 324 | p. 325
OBJ: 11.4.2 Summarize the events of meiosis.  STA: UT.BIO.2.3.a | UT.BIO.4.1.a
BLM: synthesis

150.  ANS:
Each cell in stage G, telophase II, has a single copy of each gene.

PTS: 1  DIF: L2  REF: p. 324 | p. 325
OBJ: 11.4.2 Summarize the events of meiosis.  STA: UT.BIO.2.3.a | UT.BIO.4.1.a
BLM: analysis

151.  ANS:
The chromosomes in each group are called homologous chromosomes.

PTS: 1  DIF: L1  REF: p. 392 | p. 393
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.  STA: UT.BIO.4.2.b
BLM: knowledge

152.  ANS:
The chromosomes in each pair are similar in shape and size. They also contain the same genes, though not necessarily the same alleles of those genes.

PTS: 1 DIF: L1 REF: p. 392 | p. 393
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b BLM: knowledge

153. ANS:
The chromosomes in groups 1 through 22 are autosomes.

PTS: 1 DIF: L2 REF: p. 392 | p. 393
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b TOP: Foundation Edition BLM: knowledge

154. ANS:
This person will be male, due to the presence of a Y chromosome. This person has Klinefelter’s syndrome and is likely to be infertile.

PTS: 1 DIF: L3 REF: p. 392 | p. 393
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b BLM: synthesis

155. ANS:
The karyotype shows a chromosomal abnormality. Position 23 has 3 chromosomes instead of the usual two. This abnormality was caused by nondisjunction of sex chromosomes during meiosis while a sperm or egg cell was developing in a parent.

PTS: 1 DIF: L3 REF: p. 392 | p. 393
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b BLM: evaluation

156. ANS:
Slide 2 shows type B blood.

PTS: 1 DIF: L2 REF: p. 394
OBJ: 14.1.2 Describe the patterns of the inheritance of human traits.
STA: UT.BIO.4.2.b BLM: application

157. ANS:
Slide 3 shows the blood of a person who can safely receive any type of blood. The person’s blood type is AB.

PTS: 1 DIF: L3 REF: p. 394
OBJ: 14.1.2 Describe the patterns of the inheritance of human traits.
STA: UT.BIO.4.2.b BLM: synthesis

158. ANS:
The blood on slide 4 contains no antigens. There is no clumping in either drop of blood.

PTS: 1 DIF: L3 REF: p. 394
OBJ: 14.1.2 Describe the patterns of the inheritance of human traits.
STA: UT.BIO.4.2.b BLM: synthesis

159. ANS:
The person’s genotype is \(I^A I^A\) or \(I^A i\).
Type AB blood does not contain anti-A or anti-B serum because it contains both antigen A and antigen B. If it did contain either serum, the serum would cause the blood to clump in the person’s body.

The individual is heterozygous \((Ff)\), since her daughter has attached earlobes. The daughter inherited one allele for attached earlobes from individual 2 and another from individual 1.

No, the genotype of individual 5 is uncertain because his children have free earlobes. Thus, individual 5 could be homozygous \((FF)\) or heterozygous \((Ff)\) for free earlobes. If, however, one of his children had attached earlobes, it would be certain that individual 5 was heterozygous.

Individual 14 will be homozygous for the recessive allele \((ff)\) and will have attached earlobes.

Transformation
167. **ANS:**
Bacteria are reproducing during step 7.

**PTS:** 1  
**DIF:** L2  
**REF:** p. 424 | p. 425  
**OBJ:** 15.2.1 Explain how scientists manipulate DNA.  
**STA:** UT.BIO.4.3.a  
**TOP:** Foundation Edition  
**BLM:** analysis

168. **ANS:**
A restriction enzyme is used between steps 2 and 3, and in step 4.

**PTS:** 1  
**DIF:** L3  
**REF:** p. 424 | p. 425  
**OBJ:** 15.2.2 Describe the importance of recombinant DNA.  
**STA:** UT.BIO.4.3.e  
**BLM:** evaluation

169. **ANS:**
The bacterial cell produced in step 6 contains the gene for human growth hormone. It can produce human growth hormone identical to the hormone produced in human cells in great quantities.

**PTS:** 1  
**DIF:** L3  
**REF:** p. 424 | p. 425  
**OBJ:** 15.2.2 Describe the importance of recombinant DNA.  
**STA:** UT.BIO.4.3.e  
**BLM:** synthesis

170. **ANS:**
Step 5 produces a recombinant plasmid.

**PTS:** 1  
**DIF:** L2  
**REF:** p. 424 | p. 425  
**OBJ:** 15.2.1 Explain how scientists manipulate DNA.  
**STA:** UT.BIO.4.3.a  
**TOP:** Foundation Edition  
**BLM:** analysis

171. **ANS:**
Sheep A

**PTS:** 1  
**DIF:** L2  
**REF:** p. 426 | p. 427  
**OBJ:** 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.  
**STA:** UT.BIO.4.3.f  
**TOP:** Foundation Edition  
**BLM:** application

172. **ANS:**
The nucleus was removed from the egg cell to make sure that all of the DNA in the clone was from a single sheep.

**PTS:** 1  
**DIF:** L2  
**REF:** p. 426 | p. 427  
**OBJ:** 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.  
**STA:** UT.BIO.4.3.f  
**TOP:** Foundation Edition  
**BLM:** analysis

173. **ANS:**
The lamb is a clone.

**PTS:** 1  
**DIF:** L1  
**REF:** p. 426 | p. 427  
**OBJ:** 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.  
**STA:** UT.BIO.4.3.f  
**TOP:** Foundation Edition  
**BLM:** application

174. **ANS:**
Sheep B
PTS: 1 DIF: L1 REF: p. 426 | p. 427
OBJ: 15.2.1 Explain how scientists manipulate DNA. STA: UT.BIO.4.3.a
TOP: Foundation Edition BLM: application
175. ANS:
Sheep A and the lamb are genetically identical.

PTS: 1 DIF: L2 REF: p. 426 | p. 427
OBJ: 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.
STA: UT.BIO.4.3.f TOP: Foundation Edition BLM: analysis
176. ANS:
The bands consist of DNA fragments.

PTS: 1 DIF: L2 REF: p. 433 | p. 434
OBJ: 15.3.3 Summarize the process of DNA fingerprinting and explain its uses.
STA: UT.BIO.4.3.f TOP: Foundation Edition BLM: comprehension
177. ANS:
The bands in group D moved faster because they consist of smaller DNA fragments.

PTS: 1 DIF: L2 REF: p. 433 | p. 434
OBJ: 15.3.3 Summarize the process of DNA fingerprinting and explain its uses.
STA: UT.BIO.4.3.f TOP: Foundation Edition BLM: application
178. ANS:
The restriction enzyme is cutting the DNA into fragments.

PTS: 1 DIF: L3 REF: p. 433
OBJ: 15.3.3 Summarize the process of DNA fingerprinting and explain its uses.
STA: UT.BIO.4.3.f BLM: comprehension
179. ANS:
The bands consist of DNA, which is negatively charged.

PTS: 1 DIF: L2 REF: p. 433
OBJ: 15.3.3 Summarize the process of DNA fingerprinting and explain its uses.
STA: UT.BIO.4.3.f BLM: application
180. ANS:
No, none of the DNA samples were from the same person because they produced different patterns of bands on the gel.

ESSAY
181. ANS:
The tall pea plant should be crossed with a short pea plant. If the tall pea plant is homozygous, all of the offspring will be tall. If the tall pea plant is heterozygous, it is likely that about half of the offspring will be tall and half will be short.

PTS: 1  DIF: L2  REF: p. 312
OBJ: 11.1.1 Describe Mendel's studies and conclusions about inheritance.
STA: UT.BIO.4.1.a | UT.BIO.4.2.a  TOP: Foundation Edition
BLM: analysis

182. ANS:
When the heterozygous yellow-seed F1 plants produced gametes, their dominant allele for yellow seeds segregated from their recessive allele for green seeds. As a result, some of their gametes had the dominant allele, and others had the recessive allele. When the F1 plants self-pollinated, some male gametes with the recessive allele fused with female gametes with the recessive allele during fertilization. Some of the offspring that resulted had two alleles for green seeds and therefore had green seeds.

PTS: 1  DIF: L2  REF: p. 314
OBJ: 11.1.2 Describe what happens during segregation.  STA: UT.BIO.4.2.a
BLM: application

183. ANS:
The kinds of offspring produced by genetic crosses are the results of chance. For example, the number of gametes produced that contain particular alleles is not certain. Likewise, the fusion of two gametes with particular alleles is not certain. Thus, the results of genetic crosses shown in Punnett squares are just probable results.

PTS: 1  DIF: L3  REF: p. 314
OBJ: 11.2.1 Explain how geneticists use the principles of probability to make Punnett squares.  STA: UT.BIO.4.2.a | UT.BIO.4.2.b
BLM: evaluation

184. ANS:
The genotype ratio of the offspring is not the same as their phenotype ratio. The same phenotype can be produced by several different genotypes. For example, offspring that are heterozygous for both traits (AaBb) will have the same phenotype as offspring that are homozygous for both traits (AABB).

PTS: 1  DIF: L3  REF: p. 317
OBJ: 11.2.2 Explain the principle of independent assortment.  STA: UT.BIO.4.2.a | UT.BIO.4.2.b
BLM: synthesis

185. ANS:
The alleles that determine flower color in four o’clock plants show incomplete dominance. She should use pollen from white-flowered four o’clock plants to pollinate red-flowered four o’clock plants, or vice versa. She should then collect seeds from the offspring. All of these hybrid seeds will produce only pink-flowered four o’clock plants.

PTS: 1  DIF: L3  REF: p. 319
OBJ: 11.3.1 Describe the other inheritance patterns.  STA: UT.BIO.4.2.b
BLM: synthesis

186. ANS:
I would expect the offspring to be lighter-colored than their parents. The phenotype for color in these butterflies is determined by two components: genes and environmental conditions. The offspring would have their parents’ genetic material, but they would have a different environmental influence. The butterflies born in the summer would have lower levels of pigment and so would be lighter.
A body cell has more genetic information than a gamete. A gamete is haploid, so it has just one allele for each gene. A body cell is diploid. It has two alleles for each gene. So, a body cell has twice as much genetic information as a sex cell.

Both meiosis I and meiosis II contain a prophase, a metaphase, and an anaphase. However, chromosomes replicate prior to meiosis I but not prior to meiosis II. Also, during meiosis I, tetrads form and align along the center of the cell. Then, the homologous chromosomes are separated and two haploid daughter cells form. During meiosis II, sister chromatids align along the center of the cell and are then separated. Four haploid daughter cells form.

During meiosis, the pairs of homologous chromosomes in the parent cell form tetrads and then separate. As a result, each daughter cell receives only one chromosome from each homologous pair, and the particular chromosomes that it receives are random. Thus, each daughter cell has a different combination of chromosomes. Also, crossing-over may occur during meiosis and may result in new combinations of alleles on the chromosomes in the daughter cells. In contrast, during mitosis, homologous chromosomes usually do not form tetrads and separate, and therefore crossing-over usually does not occur.

Linkage is the condition in which two genes are located on the same chromosome. As a result, the genes’ alleles are usually inherited together. However, if crossing-over occurs between the alleles of the linked genes, those alleles no longer are inherited together. The frequency of crossing-over between linked genes (or the frequency in which linked genes are inherited separately) is used to determine the relative locations of genes on the same chromosome, resulting in a gene map. The greater the frequency of separate inheritance, the farther apart the genes on the chromosome.

Doctors can use karyotypes to determine the sex of an individual. They can also use karyotypes to determine whether an individual has an abnormal number of chromosomes or noticeable chromosomal mutations, such as large deletions, additions, or translocations.
OBJ: 14.1.1 Identify the types of human chromosomes in a karyotype.
STA: UT.BIO.4.2.b     TOP: Foundation Edition
BLM: application

192. ANS:
The male cat inherited an extra X chromosome. One of the X chromosomes carries the allele for black spots, while the other carries the allele for orange spots. The two X chromosomes are randomly inactivated in the cat’s cells, producing the orange and black spots.

PTS: 1     DIF: L3     REF: p. 396
OBJ: 14.1.2 Describe the patterns of the inheritance of human traits.
STA: UT.BIO.4.2.b     BLM: evaluation

193. ANS:
The symptoms of Huntington disease usually do not appear until a person is middle-aged. Thus, although the allele is dominant, people who have Huntington disease may have one or more children before becoming aware that they are carrying the allele for Huntington disease. The allele is passed on to the children and therefore remains in the population.

PTS: 1     DIF: L2     REF: p. 399
OBJ: 14.1.2 Describe the patterns of the inheritance of human traits.
STA: UT.BIO.4.2.b     TOP: Foundation Edition
BLM: analysis

194. ANS:
The figure could not show the transmission of colorblindness. If it did, female number 5 would have to be either colorblind or a carrier of the allele for colorblindness. According to the pedigree in the figure, the mother (1) would have one recessive allele for colorblindness and one normal allele. The father (2), having just one X chromosome, would have one recessive allele for colorblindness and no normal alleles. So, the mother would be a carrier, and the father would be colorblind. All of the females born from these parents would inherit the father’s abnormal allele on one of their X chromosomes. So, all the females in this pedigree would either be colorblind (if they also got an abnormal allele from their mother), or carriers of the trait (if they inherited their mother’s X chromosome with the normal allele). Since female number 5 is neither colorblind nor a carrier, this cannot be a pedigree for a sex-linked, recessive trait, such as colorblindness.

PTS: 1     DIF: L3     REF: p. 396 | p. 397
OBJ: 14.1.3 Explain how pedigrees are used to study human traits.
STA: UT.BIO.4.2.b     BLM: evaluation

195. ANS:
Because the allele for colorblindness is recessive and X-linked, the girl must have inherited the alleles for colorblindness on the X chromosomes from both her mother and father. Since the father has a single X chromosome, it must carry the allele for colorblindness, and he must be colorblind.

PTS: 1     DIF: L2     REF: p. 395
OBJ: 14.2.1 Explain how small changes in DNA cause genetic disorders.
STA: UT.BIO.4.3.d     TOP: Foundation Edition
BLM: analysis

196. ANS:
The two syndromes are similar because both Turner’s syndrome and Down syndrome are the result of nondisjunction, the failure of chromosomes to separate in meiosis. So, in both cases, a fertilized egg has an abnormal number of chromosomes. The chromosomal problem in each case is different, however. In the case of Turner’s syndrome the fertilized egg has just one sex chromosome. In the case of Down syndrome, the fertilized egg has three copies of chromosome 21.
197. **ANS:**
Both Turner’s syndrome and Klinefelter’s syndrome result from nondisjunction of the sex chromosomes. A female with Turner’s syndrome inherits only one X chromosome. A male with Klinefelter’s syndrome has at least one extra X chromosome.

198. **ANS:**
Sequencing DNA requires restriction enzymes and DNA polymerase. Restriction enzymes cut up the huge DNA molecule into smaller, more manageable fragments. DNA polymerase replicates those pieces using normal bases and some bases that are tagged with dye. Each base can be tagged with a different color dye. The dye-tagged bases stop replication, and after running the bases on a gel, scientists can determine the sequence of the DNA by reading the order of the colored bands on the gel.

199. **ANS:**
The larger chromosome might be larger because it has longer stretches of repetitive DNA than does the smaller chromosome. Because repetitive DNA does not code for proteins, it does not contain genes. Thus, in this case, the larger chromosome does not have more genes than the smaller chromosome.

200. **ANS:**
The DNA sequence is ACAGTCTGATCTAGGTCATGCGATC. The sequence is determined by finding overlapping regions among the five fragments as shown below.

<table>
<thead>
<tr>
<th>Fragment</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>ACAG</td>
</tr>
<tr>
<td>C</td>
<td>CAGTCTGATC</td>
</tr>
<tr>
<td>A</td>
<td>GATCTAGGTCATG</td>
</tr>
<tr>
<td>E</td>
<td>CATGCGATC</td>
</tr>
<tr>
<td>D</td>
<td>AGGTC</td>
</tr>
</tbody>
</table>

201. **ANS:**
Examples will vary. The advantage of inbreeding is that it maintains desired traits within a breed, such as curly hair in poodles. The disadvantage is that it can lead to genetic defects within a breed, because of the likelihood that an individual could inherit two defective, recessive alleles.
Answers will vary. Students should include in their answers the traits that they will select for and the methods that they will use to select for those traits, such as hybridization and inducing mutations.

PTS: 1  DIF: L3  REF:  p. 419 | p. 420  
OBJ: 15.1.2 Explain how people increase genetic variation.  STA:  UT.BIO.4.1.c  
BLM:  synthesis

203. ANS:
Extract DNA from the cells of people who can make the digestion enzyme. Cut the DNA with restriction enzyme, then use gel electrophoresis and a DNA probe to locate the gene. Use the polymerase chain reaction to copy the gene. Choose a plasmid that has an antibiotic-resistance genetic marker, and cut the plasmid with the same restriction enzyme used to cut out the human gene. Insert the copies of the human gene into the plasmids. Allow bacterial cells to take in the plasmids. Select for transformed bacteria by growing them in a culture containing the antibiotic. These bacteria will make the digestion enzyme.

OBJ: 15.2.1 Explain how scientists manipulate DNA.  STA:  UT.BIO.4.3.a  
BLM:  synthesis

204. ANS:
In genetic engineering, organisms with desired traits are produced by directly changing the DNA of the organisms. This is done by cutting out desirable genes from the DNA of certain organisms and inserting them into the DNA of other organisms. In selective breeding, organisms with desired traits are produced by selecting organisms for their traits and then mating, or crossing, them. Selective breeding does not directly change the DNA of living organisms.

OBJ: 15.2.2 Describe the importance of recombinant DNA.  STA:  UT.BIO.4.3.e  
BLM:  analysis

205. ANS:
Animal breeders might first produce a particular desirable animal by using the technique of hybridization or by inducing mutations. Then, instead of using inbreeding to maintain the animal’s desirable traits, they might produce clones of that animal. The clones would be genetically identical to the original animal and thus would have all of its desirable traits.

PTS: 1  DIF: L2  REF:  p. 427  
OBJ: 15.2.3 Define transgenic and describe the usefulness of some transgenic organisms to humans.  STA:  UT.BIO.4.3.f  
TOP:  Foundation Edition  
BLM:  analysis

206. ANS:
Sample answer: Genetic engineering can help improve human health in many ways.  
• First, scientists can use genetic engineering to make more nutritional crops, such as golden rice. When people have better nutrition, they are less likely to get certain diseases.  
• Second, scientists can use transgenic animals in medical research. Animals with modified genomes are used as models in medical experiments.  
• Third, scientists can treat diseases using genetic engineering. Some diseases can be treated with drugs made through genetic engineering, and some diseases can be treated directly through gene therapy.

PTS: 1  DIF: L2  REF:  p. 429  
OBJ: 15.3.2 Explain how recombinant DNA technology can improve human health.
Gene therapy would cure the disorder rather than just treat it. The normal gene for clotting protein would be inserted into a virus’s genetic material. Then, appropriate cells from the patient would be infected with the virus. The normal clotting-protein gene, along with the viral genetic material, would be incorporated into the cells’ DNA. As a result, the cells would begin to transcribe the gene and produce the clotting protein.

Poachers kill elephants for their tusks, but officials can use genetic information to identify the herds from which the poached elephants came from, and better police those areas.

On one hand, the ability to patent genetic material helps progress by encouraging scientists to make new discoveries. With a patent on a gene or a new technique, the scientist may make money. On the other hand, if other scientists want to use the new technique or gene, they have to pay the patent holder. The need to pay may prohibit some scientists from using those tools in research, perhaps hindering progress in understanding the genome.

Sample answer: One potentially controversial issue is whether or not parents should be able to “design” their children. That is to say, should parents be able to choose their children’s height and features, such as hair color and eye color? All citizens have a duty to ensure that the tools of science are used properly, and should work toward developing a consensus on how the tools of genetic engineering should be used.