

Name: _____

Period__

Date:_____

Pedigree Worksheet

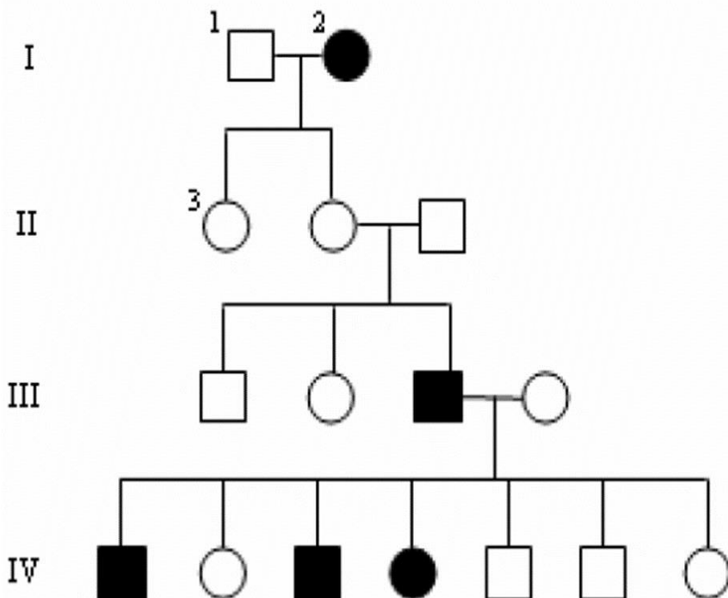
A sex-linked recessive characteristic is determined by an allele that is carried only on the X chromosomes. The shorter Y chromosome does not carry an allele for a sex-linked trait. Since there is only one X in his genotype, XY, a male who carries a particular recessive allele on the X chromosome will have the sex-linked condition. A female who carries a recessive allele in one X chromosome will not show the condition if there is a dominant allele on her other X chromosome. She will express the recessive condition only if she inherits two recessive alleles – one from each parent. Her chances of inheriting the condition are thus greatly reduced. One sex linked trait is hemophilia, a condition in which the blood does not clot properly. Most people who have hemophilia are men.

A. Interpreting a human pedigree. (use the pedigree below)

1. Number all individuals on the pedigree at the top of each shape.
2. How many males are there? _____
3. How many males have hemophilia? _____
4. How many female are there? _____
5. How many females have hemophilia? _____
6. How many marriages are there? _____
7. How many children did the first couple (couple in row I) have? _____
8. How many children did the third couple (couple in row III) have? _____
9. How many generations are there? _____
10. How many members are there in the fourth generation? _____

The genotypes of the males in a pedigree for sex-linked inheritance are easy to determine since normal blood clotting (N) is dominant and hemophilia is recessive (n). Since these alleles are on the X chromosome only, a male represented by a clear square will have the genotype $X^N Y$. A darkened square will be $X^n Y$.

Label under each individual on the pedigree. Females with hemophilia have an easy genotype to identify. They are all $X^n X^n$. Both recessive alleles must be present for a female to have hemophilia. If one dominant allele is present (X^N), the individual would be normal for clotting. Females who do not show the trait for hemophilia may be homozygous dominant ($X^N X^N$) or heterozygous ($X^N X^n$). A heterozygous female is called a carrier. Examination of offspring can often determine which genotype the parents have. **If any child (son or daughter) has hemophilia, then the female must be heterozygous.** If her son has hemophilia, he has genotype $X^n Y$. He inherited the Y from the father, so the other allele is his genotype (X^n) had to come from the mother. If a daughter has hemophilia ($X^n X^n$), she inherited an X^n from each parent, thus making the genotype for the normal mother $X^N X^n$.



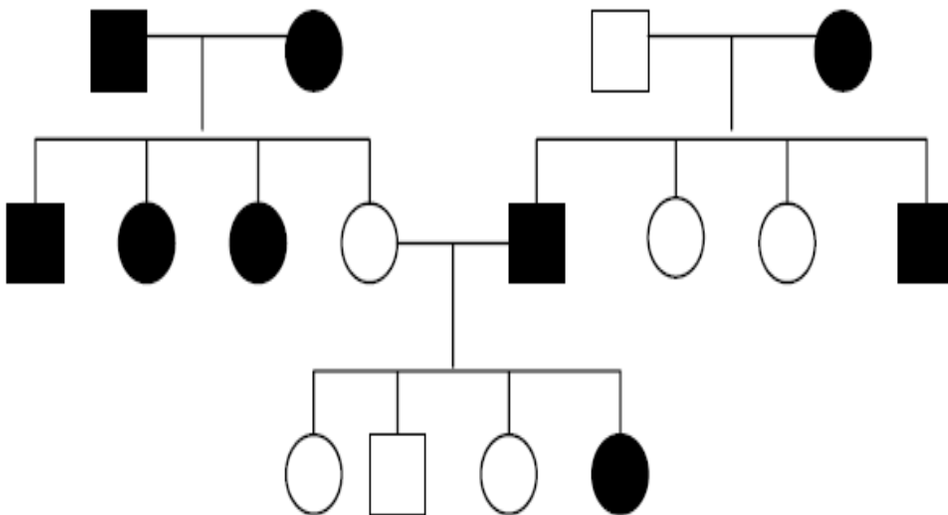
11. What is the genotype for Individual #1? _____
Phenotype? _____
12. What is the genotype for Individual #10? _____
Phenotype? _____
13. What is the genotype for the first born male in generation III? _____
Phenotype? _____
14. How many females have the genotype $X^n X^n$?
15. What would be the genotype for the female who marries into the family in generation III?

B. Determining the pattern of inheritance and assigning genotypes.

When working through a pedigree, the first thing you need to do is figure out which characteristic is dominant—the shaded one or the open one. Then you need to choose a letter (let's use B) and begin assigning genotypes. Remember that recessive individuals are **always** homozygous, so assign their genotypes first. Then go back and look at all of the dominant individuals. For some, you will only be able to determine one allele of the genotype, so just write a “?” underneath and write the possible genotypes.

Fur color is a autosomal trait. (Not sex-linked)

Fur Color in Mice



1. Which characteristic is **dominant**? White or Black?

2. Which characteristic is **recessive**? White or Black?

3. Determine the genotypes of all individuals.

You will have three inconclusive genotypes.
Write all possible genotypes beneath each individual.

C. Create Your Own Pedigree

In the space below, use colored pencils to create a pedigree with the following information:

Ray and Elaine were married in 1970. They both had normal vision. They had 2 daughters and then a son. Both daughters, Alicia and Candace, had normal vision and never had any children of their own. The son, Mike, was colorblind. The son married Beth who also had normal vision and they had 2 children of their own, first Greg then Victoria. Victoria was colorblind, but Greg was not. **Colorblindness is a sex-linked recessive trait.**

Do not forget what shapes are male and female. Place the names and genotypes of the people under their shape. Shade in carriers as necessary.

Outline- in color your individuals who fit the following:

Red- for colorblindness

White- for regular vision

Blue- for individuals with regular vision but are carriers

Green- unknown genotype