Name	e:			Date:		Honors Biology 2020
	<u>Mu</u>	<u>tations Rev</u>	<u>iew - An A</u>	<u>pplication</u>	of Protein Sy	<u>nthesis</u>
		A mutation	is any error o	or change in	the DNA sequen	ce.
Part   Direc co sp	<b>I: Learning type</b> tions: For each odon chart to tra bace in the DNA	<mark>s of Point Mut</mark> DNA/mRNA st anslate the lar strand where	<u>tations</u> rand provide nguage from the mutation	d break the nitrogen bas n has occurre	mRNA strand int es to amino acid ed.	o codons, then use the s. Also, highlight the
1.	Master: mRNA:	T ACCGTO AUGGCAO	G	AA CAT A UUGUAU	A C T I G A	
	Amino Acids:		,,		,, 	
2.	Master: mRNA:	T ACCGTO AUGGCAO	GCATGG A CGUACCU	AA CAT A UUGUAL	A C T J G A	
	Amino Acids:	,	,,	·	،،	
	Comp	are this DNA/r	mRNA sequer	nce to Questi	on 1. Mark any c	difference you find.
	What	kind of mutati	ion is this?			
3.	Master: mRNA:	T ACCGTO AUGGCAO	G A T G G A A C U A C C U U	A CA TAC UGUAUG	CT GA	
	Amino Acids:	,	,,	·	,, 	
	Comp	are this DNA/r	mRNA sequer	ice to Questi	on 1. Mark any c	difference you find.
	What	kind of mutati	ion is this?			
4.	Master: mRNA:	T ACCAG AUGGUC	T G G A T G G A C C U A C C	A A A C A T U U U G U A	T A C T A U G A	
	Amino Acids:		<i>,</i>		رر	
			,	·		
	Comp	are this DNA/r	mRNA sequer	ice to Questi	on 1. Mark any d	difference you find.
	What	kind of mutati	ion is this?			
5.	Master: mRNA:	T A C C G T ( A U G G C A (	G	A G C A T A U C G U A U	C T G A	
	Amino Acids:		,		,,	
	Comp	are this DNA/r	, mRNA sequer	, nce to Questi	 on 1. Mark any c	difference you find.
	What	kind of mutati	ion is this?			

6.	Master:	TAC CC T ACC TGCCA TAGTATT						
	mRNA:	A U G G G A U G G A C G G U A U C A U A A						
	Amino Acids:							
	/	,,						
7.	Master:	Т АССС Т АСТ ТGС САТАGТ АТТ						
	mRNA:	A U G G G A U G A A C G G U A U C A U A A						
	Amino Acids:							
	,	,,						
	Compare this DNA/mRNA sequence to Question 6. Mark any difference you find.							
	What kind of mutation is this?							

#### *Identify the type of mutations in the examples above:*

First decide whether the mutation is a **substitution**, **deletion** or **insertion** and then determine whether they are a **missense** mutation, a **nonsense** mutation or a **silent** mutation. Which mutation(s) result in a **frameshift** mutation?

**Please note**: The mutations shown above involving one nucleotide are called **point mutations**. There is another class of mutations called **chromosomal mutations**. Chromosomal mutations affect segments of chromosomes and involve multiple genes and long segments of nucleotides. A **Translocation** is an example of a chromosomal mutation (when a piece of one chromosome attaches to another, non-homologous chromosome). When entire segments of chromosomes are lost, that is referred to as a **deletion**. There could also be an **inversion** where a segment of a chromosome undergoes breakage and rearrangement within itself.

## <u>Part II: An Application of Mutations in Humans: Sickle Cell Anemia</u> How does a mutation in the gene for hemoglobin result in Sickle Cell Anemia?

**Background:** Different versions of the same gene are called **alleles**. These different alleles share the same general sequence of nucleotides, but they differ in at least one nucleotide in the sequence. Different alleles can result in different characteristics by the following sequence:

Differences in the nucleotide sequence in the gene

- → lead to differences in the nucleotide sequence in mRNA
- ightarrow lead to differences in the amino acid sequence in the protein
- → lead to differences in the structure & function of the protein
- → lead to differences in a person's characteristics

Hemoglobin is a protein inside red blood cells that binds to oxygen and allows for its transport throughout the body. If someone has an allele that codes for sickle cell hemoglobin, they will produce mis-shaped hemoglobin instead of typical-forming hemoglobin and their capacity for oxygen transport will be reduced. Additionally, the altered hemoglobin then can affect the overall shape of the red blood cell and the cells appear sickled in shape. The sickled red blood cells can then clump together and ultimately, the person with this condition suffers from anemia (the lack of enough red blood cells to adequately transport oxygen leading to fatigue, shortness of breath and other serious complications).

The following exercise will help you to understand how differences between the varieties of the hemoglobin gene can be expressed (either causing sickle cell or not). Then you will learn how the differences between the typical and sickle cell hemoglobin proteins can result in good health or sickle cell anemia.

- 1. In the table below, compare the DNA for the <u>Beginning of the **Typical** Hemoglobin Gene</u> to the <u>Beginning of the **Sickle Cell** Hemoglobin Gene</u>. Identify the difference.
- 2. Complete the table below. (Use the codon translator sheet)

## Typical Hemoglobin Gene (more common in the population)

Beginning of a <b>typical</b> Hemoglobin <b>Gene</b>	CAC GTA GAC TGA GGA CTC						
Transcription produces:	codon 1	codon 2	codon 3	codon 4	codon 5	codon 6	
Beginning of a <b>typical</b> Hemoglobin <b>mRNA</b>							
Translation produces:	amino acid 1	amino acid 2	amino acid 3	amino acid 4	amino acid 5	amino acid 6	
Beginning of a <b>typical</b> Hemoglobin <i>Protein</i>							

## Sickle Cell Hemoglobin Gene

Beginning of <b>Sickle Cell</b> Hemoglobin <b>Gene</b>	CAC GTA GAC TGA GGA CAC							
Transcription produces:	codon 1	codon 2	codon 3	codon 4	codon 5	codon 6		
Beginning of <b>Sickle Cell</b> Hemoglobin <i>mRNA</i>								
Translation produces:	amino acid 1	amino acid 2	amino acid 3	amino acid 4	amino acid 5	amino acid 6		
Beginning of <b>Sickle Cell</b> Hemoglobin <i>Protein</i>								

- 3. What is the difference in the amino acid sequence of the hemoglobin molecules synthesized by translating the sickle cell vs. typical hemoglobin mRNA molecules?
- 4. What kind of mutation is this?

# Part III (Extension): The effect of Sickle Cell Gene:

Each complete hemoglobin protein has more than 100 amino acids. Sickle cell hemoglobin and typical hemoglobin differ in only a single amino acid. This difference in a single amino acid results in the very different properties of sickle cell hemoglobin, compared to typical hemoglobin.

If a person inherits two copies of the sickle cell hemoglobin gene and produces only sickle cell hemoglobin, then the sickle cell hemoglobin will tend to clump together in long rods. These long rods of clumped-together sickle cell hemoglobin change the shape of the red blood cells from their typical disk shape to a sickle shape (see chart). The sickle-shaped red blood cells can block the blood flow in the tiny capillaries, causing pain and damage to body organs. In addition, the sickle-shaped red blood cells do not last nearly as long as normal red blood cells, so the person does not have enough red blood cells, resulting in anemia.

Genotype → Protein		<b>→</b>	Phenotype	
HH (2 alleles for typical hemoglobin)	<b>→</b>	Typical hemoglobin in red blood cells	→	Disk-shaped red blood cells → healthy individual
hh (2 alleles for sickle cell hemoglobin)	<b>→</b>	Sickle cell hemoglobin in red blood cells CLUMPED HEMOGLOBIN	<b>→</b>	Sickle-shaped red blood cells → pain, damage to body organs, anemia

In summary, the sickle cell allele results in production of the sickle cell hemoglobin protein, which results in the health problems observed in sickle cell anemia. This is a dramatic example of the importance of the nucleotide sequence in a gene, which determines the amino acid sequence in a protein, which in turn influences the characteristics of an individual.

#### In summary:

How does someone's DNA determine whether they develop sickle cell anemia?