

Protein Synthesis

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Transcription in Protein Synthesis

Synthesis of RNA

- This process of protein synthesis occurs in **two stages**:
 - Transcription DNA is transcribed and an mRNA molecule is produced
 - mRNA is a single stranded RNA molecule that transfers the information in DNA from the nucleus into the cytoplasm
 - mRNA production requires the enzyme RNA polymerase
 - Translation mRNA (messenger RNA) is translated and an amino acid sequence is produced

The process of transcription

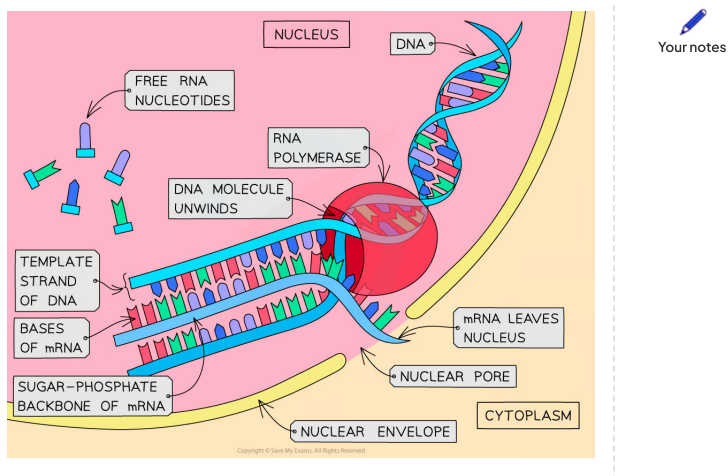
- This stage of protein synthesis occurs **in the nucleus** of the cell
- Part of a DNA molecule **unwinds** (the **hydrogen bonds** between the complementary base pairs **break**)
- This exposes the gene to be transcribed (the gene from which a particular polypeptide will be produced)
- A complementary copy of the code from the gene is made by building a single-stranded nucleic acid molecule known as mRNA (messenger RNA)
- Free RNA nucleotides pair up (via hydrogen bonds) with their complementary (now exposed) bases on one strand (the template strand) of the 'unzipped' DNA molecule
- The sugar-phosphate groups of these RNA nucleotides are then **bonded** together by the enzyme **RNA polymerase** to form the sugar-phosphate backbone of the mRNA molecule
- When the gene has been transcribed (when the mRNA molecule is complete), the hydrogen bonds between the mRNA and DNA strands break and the **double-stranded DNA molecule re-forms**
- The mRNA molecule then leaves the nucleus via a pore in the nuclear envelope
 - This is where the term messenger comes from the mRNA is despatched, carrying a message, to another part of the cell
 - DNA can't make this journey; it's too big to fit through the pores in the nuclear envelope

Transcription in the nucleus diagram



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DNA is transcribed and an mRNA molecule is produced

Examiner Tip

Be careful – DNA polymerase is the enzyme involved in DNA replication; RNA polymerase is the enzyme involved in transcription – don't get these confused.

Hydrogen bonding & Complementary Base Pairing

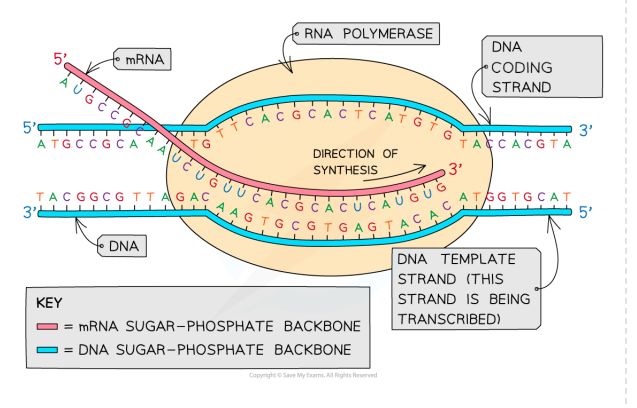
- In the transcription stage of protein synthesis, free RNA nucleotides pair up with the exposed bases on the DNA molecule but only with those bases on one strand of the DNA molecule
- The RNA will have a complementary base sequence to the DNA strand and will bind to the DNA using **hydrogen bonds**
- The adenine of the DNA is complementary to uracil on the new RNA strand, because a thymine RNA nucleotide does not exist

Complementary base pairing between the DNA and the RNA transcript table

| DNA template strand code | TAC | GGA | AGA | CTT | GGG |
|--------------------------|-----|-----|-----|-----|-----|
| RNA transcript | AUG | CCU | UCU | GAA | CCC |

- The strand of the DNA molecule that carries the genetic code is called the **coding strand**
- The opposite DNA strand is called the **template** strand
- To get an RNA transcript of the coding strand, the template strand is the one that is transcribed to form the mRNA molecule
 - This mRNA molecule will later be translated into an amino acid chain

DNA coding and template strand during transcription diagram







The template strand of the DNA molecule is the one that is transcribed

DNA Templates

- DNA is a very stable molecule due to the hydrogen bonding between the DNA bases of the two strands and the strong phosphodiester bonds between adjacent nucleotides in each strand
 - This means that the genetic code is not prone to spontaneously breaking or changing
- This feature allows single DNA strands to act as **reliable templates for transcription** over several generations of cell replication
- In certain types of somatic cells that do not divide during their lifetimes, such as neurones and some types of muscle cells, the genetic sequence is **conserved** due to this stability and **does not degrade over time**

Transcription & Gene Expression

- There are approximately 20,000 protein-coding genes in the human genome
- Not every protein is needed in every cell
 - For example, the insulin protein is not needed in cardiac muscles of the heart
- As a result, our specialised cells have a way of **switching certain genes off or on** to match the requirements of the cell. This is called **gene expression**
 - Genes that are expressed are 'switched on' and undergo the process of transcription and translation
 - Genes that are not expressed are 'switched off' or silenced, and do not go through the process of transcription and/or translation
- There are various different mechanisms in the cell involved in controlling gene expression
- Transcription is the first stage of gene expression and so this is a key stage at which gene expression can be switched on or off



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Translation in Protein Synthesis

Synthesis of Polypeptides

- Translation involves taking the genetic code from the mRNA and synthesising a polypeptide
 - A polypeptide is a sequence of amino acids covalently bonded together
 - The order of the amino acids is based on the information stored in the genetic code of the mRNA
- This stage of protein synthesis occurs in the cytoplasm of the cell
- The **mRNA template** comes from the process of **transcription**, and so translation always takes place following these events
 - After transcription the mRNA moves out of the nuclear pore and diffuses into the cytoplasm towards the ribosome for translation

Examiner Tip

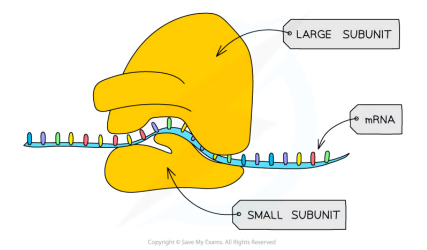
Make sure you learn both stages of protein synthesis fully. Don't forget WHERE these reactions take place – transcription occurs in the nucleus but translation occurs in the cytoplasm!



Roles of RNA & Ribosomes in Translation

- After leaving the nucleus, the mRNA molecule attaches to a ribosome
- A ribosome is a complex structure that is made of a large and small subunit
 - Ribosomes are themselves made of **proteins** and **RNA** (called ribosomal RNA or **rRNA**)
- There are **binding sites on the subunits** for the various other molecules involved in translation
 - The mRNA binds to the small subunit
 - Two tRNA molecules are able to bind to the large subunit simultaneously

mRNA in the ribosome diagram



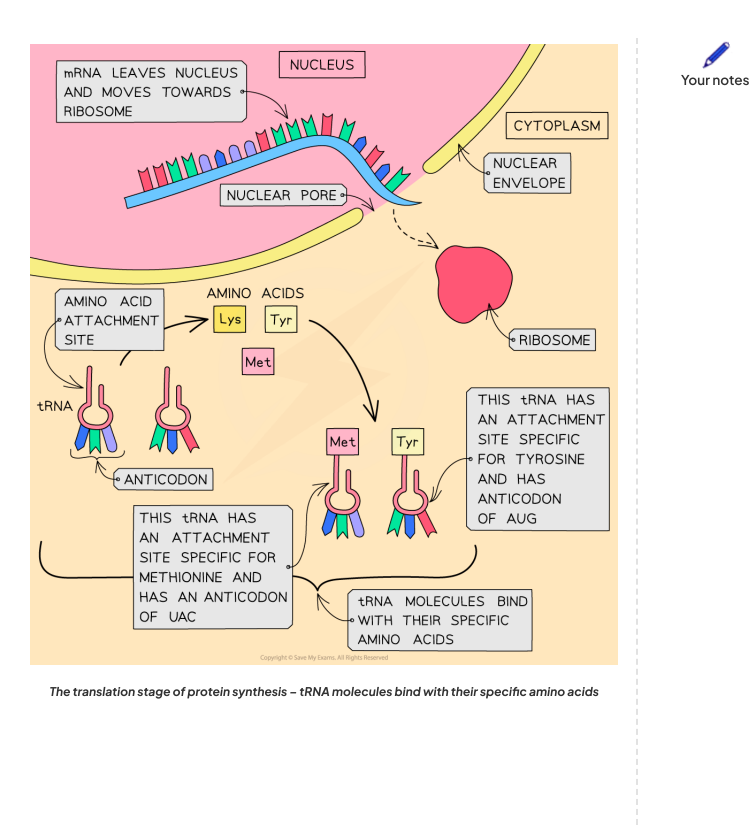
A ribosome is built of large and small subunits, ribosomal RNA and an area on the surface that catalyses the formation of peptide bonds in a newly-synthesised protein

- Translation depends on complementary base pairing between codons on mRNA and anticodons on tRNA
- In the cytoplasm, there are **free molecules of tRNA** (transfer RNA)
- The tRNA molecules bind with their specific amino acids (also in the cytoplasm) and bring them to the mRNA molecule on the ribosome
- The triplet of bases (anticodon) on each tRNA molecule pairs with a complementary triplet (codon) on the mRNA molecule

tRNA and mRNA before translation diagram

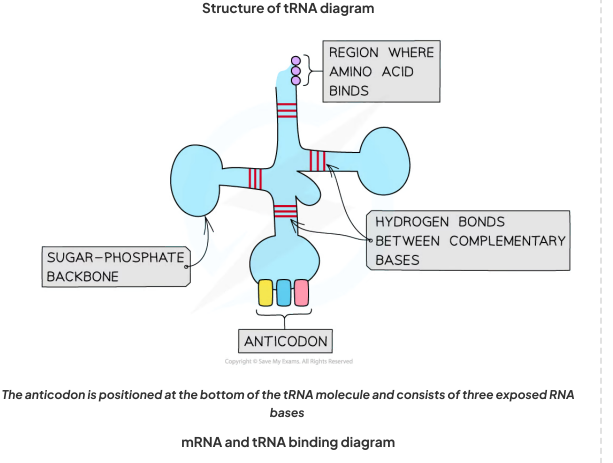


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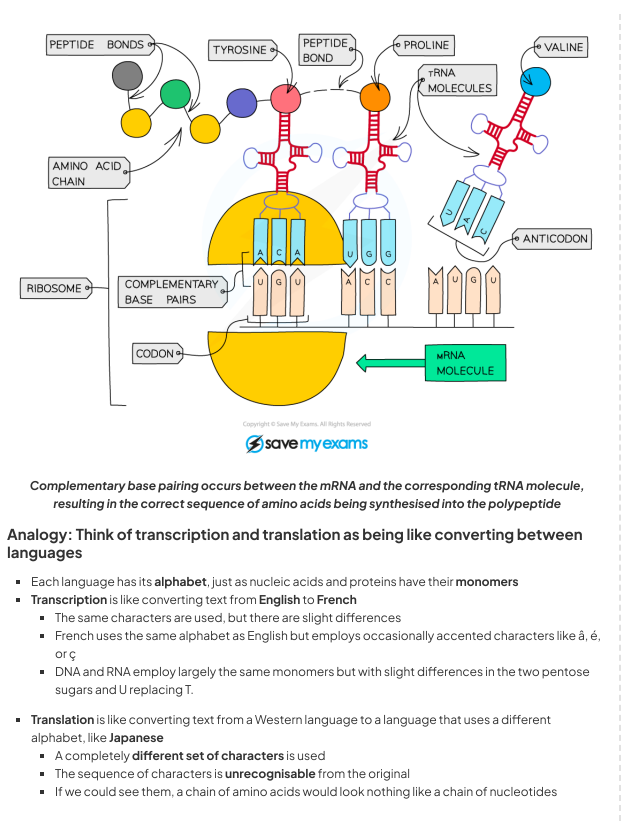
Codons & Anticodons

- Codons of three bases on mRNA correspond to one amino acid in a polypeptide
 - A triplet is a sequence of three DNA bases that codes for a specific amino acid
 - A codon is a sequence of three mRNA bases that codes for a specific amino acid
 - A codon is transcribed from the triplet and is complementary to it
- An anticodon is a sequence of three transfer RNA (tRNA) bases that are complementary to a codon
 - The transfer RNA carries the appropriate amino acid to the ribosome
 - The amino acid can then be condensed **onto the growing polypeptide chain**
- Certain codons carry the command to **stop translation** when the polypeptide chain is complete. These are called **stop codons**



Your notes

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Transcription and Translation Can be Likened to the Conversion Between Languages Table



| Transcription | DNA ightarrow RNA | Similarities | English 	o French | | Similarities |
|---------------|--------------------------|---|---|---|---|
| DNA → RNA | TTACAGCTC → AAUGUCGAG | Both use a similar set of monomers (with a slight difference; U replaces T) | "I received biology lessons at my school" | "J'ai reçu des cours de biologie à mon école" | Both use a similar alphabet (with slight differences: ç,à,é,Ô etc) |

| Translation | $RNA \longrightarrow Protein$ | Differences | French → Japanese | | Differences |
|---------------|-------------------------------|--|---|---------------------|---------------------------------|
| RNA → protein | AAUGUCGAG → Asn-Val-Glu | Both use different monomers (nucleotides & amino acids) | "J'ai reçu des cours de biologie à mon école" | 学校で生物学の授業 を受けました | Both use different alphabets |

O Examiner Tip

Remember that complementary base pairing in RNA means that:

- Adenine (A) will pair up with Uracil (U)
- Cytosine (C) will pair up with Guanine (G)

So if an mRNA codon has a sequence of **CAG**, then its complementary tRNA anticodon will have a sequence of **GUC**.

The Genetic Code

Features of the Genetic Code

- The sequence of DNA nucleotide bases found within a gene is determined by a **triplet** (three-letter) **code**
- Each sequence of three bases (i.e. each triplet of bases) in a gene codes for one amino acid
- These triplets code for different amino acids there are 20 different amino acids that cells use to make up different proteins
- For example:
 - CAG codes for the amino acid valine
 - TTC codes for the amino acid lysine
 - GAC codes for the amino acid leucine
 - CCG codes for the amino acid glycine
- Some of these triplets of bases code for **start** (TAC methionine) **and stop signals**
- These start and stop signals tell the cell where individual genes start and stop
- As a result, the cell reads the DNA correctly and produces the correct sequences of amino acids (and therefore the correct protein molecules) that it requires to function properly
- The genetic code is **non-overlapping**
 - Each base is only read once in which codon it is part of
- There are **four bases**, so there are **64** different codons (triplets) possible (4³ = 64), yet there are only 20 amino acids that commonly occur in biological proteins
 - This is why the code is said to be **degenerate**: multiple codons can code for the same amino acids
 - The degenerate nature of the genetic code can limit the effect of mutations
- The genetic code is also universal, meaning that almost every organism uses the same code (there are a few rare and minor exceptions)
- The same triplet codes code for the same amino acids in all living things (meaning that genetic information is transferable between species)
 - The universal nature of the genetic code is why genetic engineering (the transfer of genes from one species to another) is possible



Deducing Amino Acid Sequences

• By observing the **genetic code in the mRNA** it is possible to determine the **sequence of amino acids** that are coded for in the **polypeptide**

mRNA codons and amino acids table

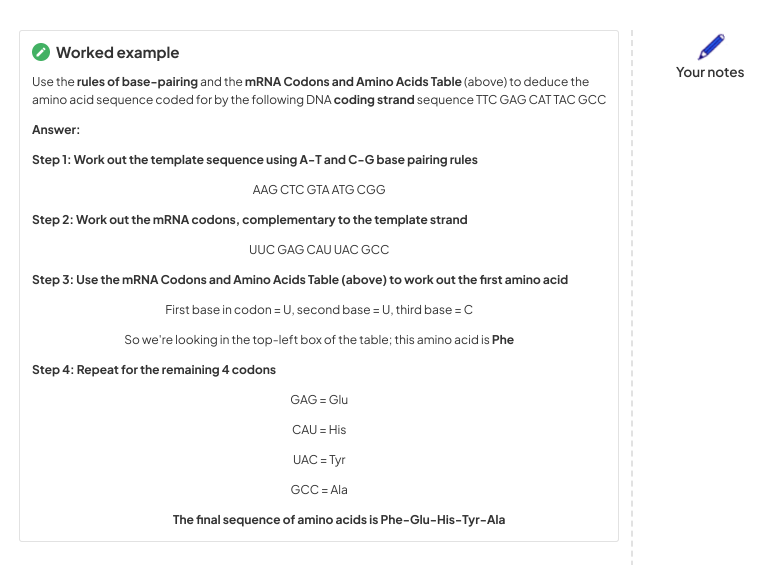
| | | U | С | А | G | | |
|--------|---|--|---------------------------------|--|---|---------|--------|
| | U | $\left. egin{array}{c} UUU \\ UUC \end{array} ight\} \operatorname{Phe} \\ UUA \\ UUG \end{array} ight\} \operatorname{Leu}$ | UCU UCC UCA UCG | UAU UAC } Tyr UAA Stop UAG Stop | UGU UGC UGA Stop UGG Trp | U C < G | |
| LETTER | С | CUU CUC CUA CUG | CCU CCC CCA CCG Pro | CAU His CAC CAA CAG Gln | CGU CGC CGA CGG Arg | ン い < の | THIRD |
| FIRST | A | AUU AUC AUA AUG Met | ACU ACC ACA ACG | $\left. \begin{array}{c} AAU \\ AAC \end{array} \right\} \left. \begin{array}{c} Asn \\ Asn \\ AAA \\ AAG \end{array} \right\} Lys \\ \end{array}$ | $\left. \begin{matrix} AGU \\ AGC \end{matrix} \right\} \begin{matrix} Ser \\ AGA \\ AGG \end{matrix} \right\} \begin{matrix} Arg \end{matrix}$ | してくの | LETTER |
| | G | GUU GUC GUA GUG | GCU GCC GCA GCG | GAU GAC GAA GAG GLu | GGU GGC GGA GGG | U C < G | |

SECOND LETTER

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ST LETTER



Elongation of the Polypeptide Chain

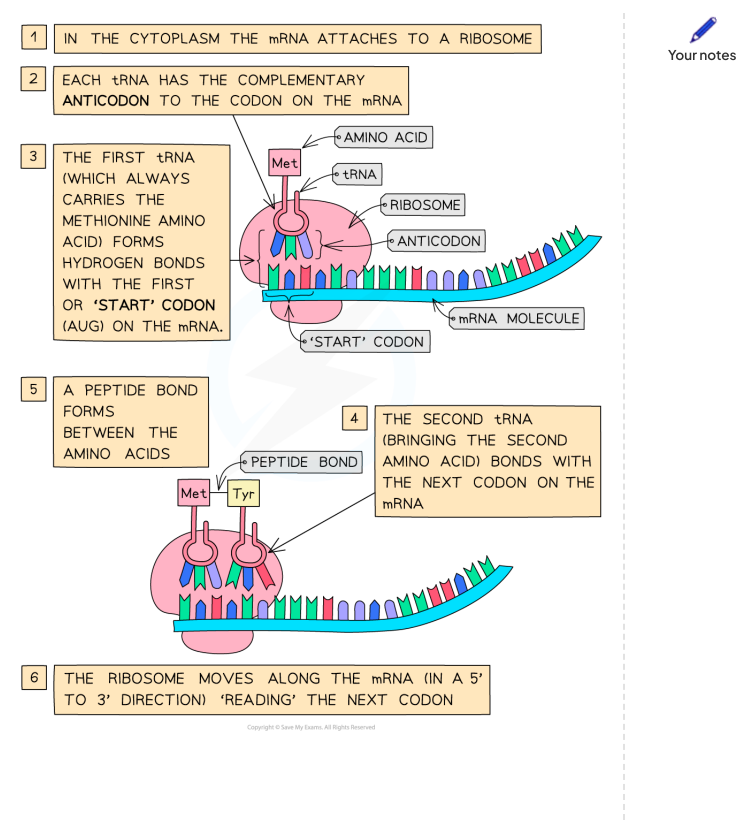
- During translation **two tRNA molecules fit onto the ribosome at any one time**, bringing the amino acid they are each carrying side by side
 - The ribosome will move along the mRNA molecule, one codon at a time
- A **peptide bond** is then formed (by condensation) between the two amino acids
 - The formation of a peptide bond between amino acids is an anabolic reaction
 - It requires energy, in the form of ATP
 - The ATP needed for translation is provided by the **mitochondria** within the cell
- This process continues until a '**stop' codon** on the mRNA molecule is reached this acts as a signal for translation to stop and at this point the amino acid chain coded for by the mRNA molecule is complete
- This amino acid chain is then released from the ribosome and forms the final polypeptide

The process of translation diagram



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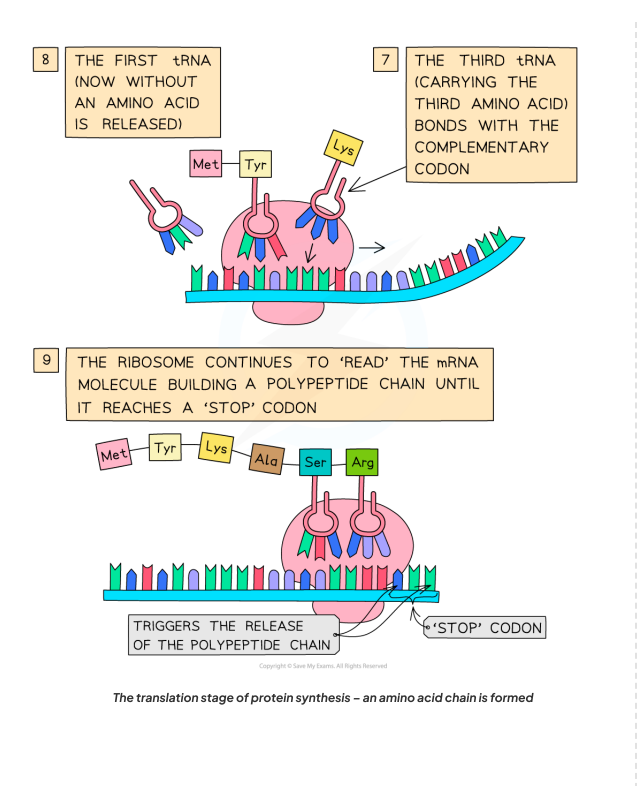
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Your notes



Protein Structure & Mutations

Protein Structure & Mutations

- A gene mutation is a change in the sequence of bases in a DNA molecule; this may result in a new allele
 - Mutations occur all the time and occur randomly
 - Mutations are **copying errors** that take place when DNA is replicated during **S phase** of interphase
- As the DNA base sequence determines the sequence of amino acids that make up a polypeptide, mutations in a gene can sometimes lead to a change in the polypeptide for which the gene codes
- Most mutations are **harmful** or **neutral** (have no effect) but some can be **beneficial**
- Inheritance of mutations:
 - Mutations present in normal body cells are not inherited; they are eliminated once the affected cells die
 - Mutations within gametes are inherited by offspring, so can lead to **heritable** genetic conditions
- Point mutations are mutations where one base in the DNA sequence is altered; this can result in a changed amino acid at this location

Example of a point mutation: sickle cell disease

- A small change to a gene can have **serious consequences** for an organism
- Sickle cell disease is a genetic disorder caused by a single point mutation within the gene that codes for the alpha-globin polypeptide in haemoglobin (Hb)
 - Most humans have the allele Hb^A
 - The mutation results in a new allele **Hb**^S

The sickle cell mutation

- Within the haemoglobin gene a point mutation changes the DNA triplet GAG to GTG on the coding strand
- The resulting DNA triplet (CAC) on the template strand is transcribed into the **mRNA codon GUG**, instead of GAG
- During translation the amino acid valine (Val) replaces the original amino acid glutamic acid (Glu)
 - This occurs at the **sixth position** of the polypeptide

Sickle cell anaemia point mutation diagram

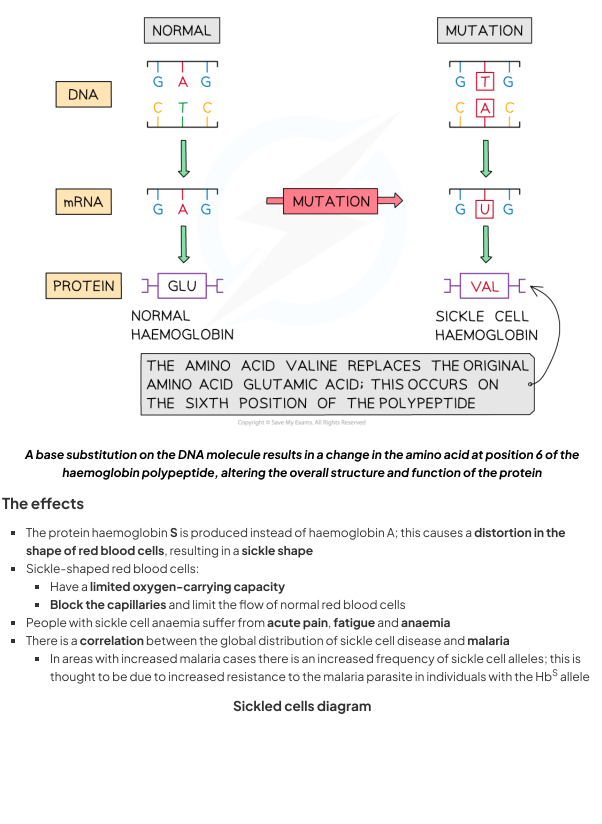


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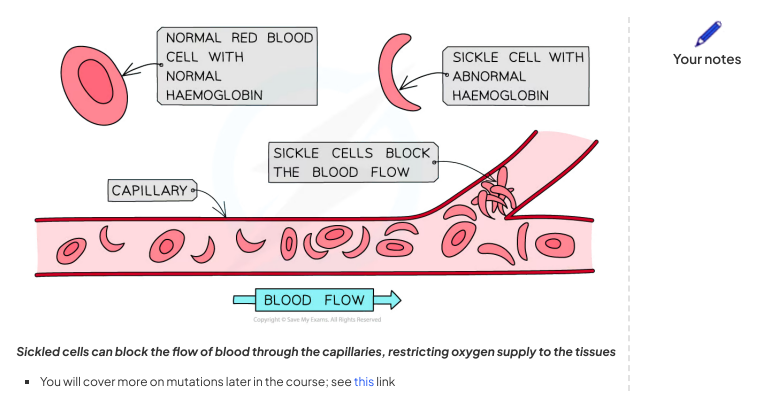
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