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**ANATOMY**

**BCH 204 (MEDICAL BIOCHEMISTRY)**

1.) DNA Replication, also known as **Semi-Conservative Replication**, is the process by which DNA is essentially “doubled”. It is an important process that takes place within the dividing cell.

 **Stages of DNA replication**

DNA replication can be thought of in three stages; Initiation, Elongation, Termination

a. **Initiation**

DNA synthesis is initiated at particular points within the DNA strand known as ‘origins’, which are specific coding regions. These origins are targeted by initiator proteins, which go on to recruit more proteins that help aid the replication process, forming a replication complex around the DNA origin. There are multiple origin sites, and when replication of DNA begins, these sites are referred to as Replication Forks.

Within the replication complex is the enzyme DNA Helicase, which unwinds the double helix and exposes each of the two strands, so that they can be used as a template for replication. It does this by hydrolysing the ATP used to form the bonds between the nucleobases, therefore breaking the bond between the two strands.

DNA can only be extended via the addition of a free nucleotide triphosphate to the 3’- end of a chain. As the double helix runs antiparallel, but DNA replication only occurs in one direction, it means growth of the two new strands is very different (and will be covered in Elongation).

DNA Primase is another enzyme that is important in DNA replication. It synthesizes a small RNA primer, which acts as a ‘kick-starter’ for DNA Polymerase. DNA Polymerase is the enzyme that is ultimately responsible for the creation and expansion of the new strands of DNA.

b. **Elongation**

Once the DNA Polymerase has attached to the original, unzipped two strands of DNA (i.e. the template strands), it is able to start synthesizing the new DNA to match the templates. This enzyme is only able to extend the primer by adding free nucleotides to the 3’-end of the strand, causing difficulty as one of the template strands has a 5’-end from which it needs to extend from.

One of the templates is read in a 3’ to 5’ direction, which means that the new strand will be formed in a 5’ to 3’ direction (as the two strands are antiparallel to each other)*.* This newly formed strand is referred to as the Leading Strand. Along this strand, DNA Primase only needs to synthesize an RNA primer once, at the beginning, to help initiate DNA Polymerase to continue extending the new DNA strand. This is because DNA Polymerase is able to extend the new DNA strand normally, by adding new nucleotides to the 3’ end of the new strand (how DNA Polymerase usually works).

However, the other template strand is antiparallel, and is therefore read in a 5’ to 3’ direction, meaning the new DNA strand being formed will run in a 3’ to 5’ direction. This is an issue as DNA Polymerase doesn’t extend in this direction. To counteract this, DNA Primase synthesizes a new RNA primer approximately every 200 nucleotides, to prime DNA synthesis to continue extending from the 5’ end of the new strand. To allow for the continued creation of RNA primers, the new synthesis is delayed and is such called the Lagging Strand.

The leading strand is one complete strand, while the lagging strand is not. It is instead made out of multiple ‘mini-strands’, known of Okazaki fragments. These fragments occur due to the fact that new primers are having to be synthesized, therefore causing multiple strands to be created, as opposed to the one initial primer that is used with the leading strand.

c. **Termination**

The process of expanding the new DNA strands continues until there is either no more DNA template left to replicate (i.e. at the end of the chromosome), or two replication forks meet and subsequently terminate. The meeting of two replication forks is not regulated and happens randomly along the course of the chromosome.

Once DNA synthesis has finished, it is important that the newly synthesized strands are bound and stabilized.  With regards to the lagging strand, two enzymes are needed to achieve this; RNAase H removes the RNA primer that is at the beginning of each Okazaki fragment, and DNA Ligase joins two fragments together creating one complete strand.

Now with two new strands being finally finished, the DNA has been successfully replicated, and will just need other intrinsic cell systems to ‘proof-read’ the new DNA to check for any errors in replication, and for the new single strands to be stabilized.

2.) **DNA ENZYMES**

 At the replication fork, many replication enzymes assemble on the DNA into a complex molecular machine called the [replisome](https://en.wikipedia.org/wiki/Replisome%22%20%5Co%20%22Replisome). The following is a list of major DNA replication enzymes that participate in the replisome:

a. [**DNA helicase**](https://en.wikipedia.org/wiki/DNA_helicase): Also known as helix destabilizing enzyme. Helicase separates the two strands of DNA at the [Replication Fork](https://en.wikipedia.org/wiki/Replication_Fork) behind the topoisomerase.

b. [**DNA polymerase**](https://en.wikipedia.org/wiki/DNA_polymerase): The enzyme responsible for catalyzing the addition of nucleotide substrates to DNA in the 5′ to 3′ direction during DNA replication. Also performs proof-reading and error correction. There exist many different types of DNA Polymerase, each of which perform different functions in different types of cells.

c. [**DNA clamp**](https://en.wikipedia.org/wiki/DNA_clamp): A protein which prevents elongating DNA polymerases from dissociating from the DNA parent strand.

d. [**Single-strand DNA-binding protein**](https://en.wikipedia.org/wiki/Single-strand_DNA-binding_protein): Bind to ssDNA and prevent the DNA double helix from re-annealing after DNA helicase unwinds it, thus maintaining the strand separation, and facilitating the synthesis of the nascent strand.

e. [**Topoisomerase**](https://en.wikipedia.org/wiki/Topoisomerase): Relaxes the DNA from its super-coiled nature.

f. [**DNA gyrase**](https://en.wikipedia.org/wiki/DNA_gyrase): Relieves strain of unwinding by DNA helicase; this is a specific type of topoisomerase.

g. [**DNA ligase**](https://en.wikipedia.org/wiki/DNA_ligase): Re-anneals the semi-conservative strands and joins [Okazaki Fragments](https://en.wikipedia.org/wiki/Okazaki_Fragments) of the lagging strand.

h. [**Primase**](https://en.wikipedia.org/wiki/Primase): Provides a starting point of RNA (or DNA) for DNA polymerase to begin synthesis of the new DNA strand.

i. [**Telomerase**](https://en.wikipedia.org/wiki/Telomerase): Lengthens telomeric DNA by adding repetitive nucleotide sequences to the ends of [eukaryotic chromosomes](https://en.wikipedia.org/wiki/Eukaryotic_chromosome_fine_structure). This allows germ cells and stem cells to avoid the Hayflick limit on cell division.