Prokaryotic DNA Replication and Telomere Replication

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- DNA replication is essential before dividing of a cell so each daughter cell has a complete copy of genetic information.
- The enzyme that copy DNA, polymerize nucleotides in the 5'>3' directions although the two polynucleotide strands of DNA run in opposite directions, yet both strands appear to grow in the same directions.
- Replication mechanisms requires lots of enzymes and proteins.



Fig: Nucleotides in DNA backbone are bonded from phosphate to sugar between 3' & 5' carbons.

DNA Replication is Semi-conservative

Semi-conservative model of Watson and Crick predicts that when a double helix replicates, each daughter molecule will have one old strand and one newly synthesized strand.
Competing models were the conservative model (the two parent strands rejoin) and the dispersive model (each strand is a mix of old and new).

Experiments by Matthew Meselson and Franklin Stahl supported the semi-conservative model of DNA replication.
They labeled the nucleotides of the old strands with a (¹⁵N) heavy isotope of nitrogen, while any new nucleotides were labeled with a lighter isotope (¹⁴N).

Comparison of three proposed Models





CONCLUSION OF THE EXPERIMENT:

• The first replication produced a band of hybrid DNA, eliminating the conservative model.

 A second replication produced both light and hybrid DNA, eliminating the dispersive model and supporting the semi-conservative model.

Prokaryotic Replication- A closer look

I. Origin of Replication- The Ori C and Prepriming Complex

- Replication begins at particular sites called origins of replication, where the two DNA strands are separated, opening up and form 'replication bubble'.
- Ori C consists of 245bp and contains 3 repeats of a13bp sequence and 4 repeats of a 9bp sequence called 13mers and 9mers respectively.
- A eukaryotic chromosome may have hundreds or even thousands of origins of replication.
- Replication proceeds in both directions from each origin, until the entire molecule is copied.



➤ At first 20 Dna-A protein molecules bind to the four 9bp repeates, bindind needs ATP, which help to denature 13bp repeats rich in A=T.

➢Now Dna −B which serves as a helicase along with Dna −C proteins bound to unbound region in two hexamers that unwinds DNA bidirectionally to create the replication fork. All these processes need ATP.

- At the end of each replication bubble there is a a Y-shaped region where new DNA strands are elongating, called replication fork.
- Helicase enzymes that unwind the double helix at the replication forks.
- Topoisomerase I and II relax super coiled DNA strands ahead of replication forks by breaking, swiveling, and rejoining DNA strands.
- Single-strand binding proteins (SSBP) bind to and stabilize DNA single-strand.





- The initial nucleotide strand is a short RNA primer (5–10 nucleotides long), and the 3' end serves as the starting point for the new DNA strand.
- An enzyme called primase adds RNA nucleotides one at a time using the parental DNA as a template.

Structure of DNA Polymerase III

> DNA polymerase III is a holoenzyme, which has two core enzymes (Pol III), each consisting of three subunits (α , ϵ and θ), a sliding clamp that has two beta subunits, and a clamp-loading complex which has multiple subunits (δ , τ , γ , ψ , and χ).

➢Core enzyme catalyzes DNA polymerization activity.

➢ S subunit binds DNA at the junction between template and primer as a dimeric ring. The ring holds the catalytic core polymerase and functions like a clamp.



Leading & Lagging strands

- DNA polymerases add nucleotides only to the free 3' end of a growing strand; therefore, a new DNA strand can elongate only in the 5' to 3' direction.
- Along one template strand of DNA, the DNA polymerase III synthesizes a leading strand continuously, moving toward the replication fork.
- The lagging strand is synthesized as a series of segments called Okazaki fragments, which are joined together by DNA ligase.





Fig: Lagging strand synthesis

III.<u>The Termination</u>

E. coli chromosome and several other plasmids carry specific sequence, called 'Ter sites', diametrically opposite from OriC; where TBP (ter binding protein) binds.
There are three Ter sites (ter A, ter D and ter E) for counter clockwise fork and three Ter sites (ter B, ter C and ter F) for clockwise fork.

These six Ter sites stall the fork, by inhibiting DNA helicase.

A SUMMARY OF DNA REPLICATION



Proofreading and Repairing DNA

- DNA polymerase I having 5'→3' and 3'→5' exonuclease activity and 5'→3' polymerization activity.
- DNA polymerase I proofread newly synthesized DNA, replacing any incorrect nucleotides and increasing accuracy by a hundred to a thousand fold.
- mismatch repair enzymes help to correct errors in base pairing in DNA.
- DNA can be damaged by exposure to harmful chemical or physical agents, which causes spontaneous changes.
- nucleotide excision repair, a nuclease help to cuts out and replaces damaged part of DNA.

Replication of Telomeres

- Eukaryotic chromosomal DNA molecules have special nucleotide sequences (5'TTGGGG3') at their ends called telomeres.
- Limitations of DNA polymerase create problems for the linear DNA of eukaryotic chromosomes. This is not a problem for prokaryotes, as most of which have circular chromosomes.
- The usual replication machinery unable to complete the 5' ends, so repeated rounds of replication produce shorter DNA molecules with uneven ends in the chain.

- An enzyme called telomerase, a ribonuclease protein catalyzes the lengthening of telomeres and bind to G rich sequence and add 'TTGGGG' to end of telomere sequence of template lagging strand by built in RNA strand template.
- This 'TTGGGG' repeats form hairpin loop .
- This loop provide3'OH end to DNA polymerase-I to fill the gap after the removal of RNA primer.
- After this hairpin is removed and thus shortening of DNA is prevented.



Significance

- Telomeres do not prevent the shortening of DNA molecules, but they do postpone the erosion of genes near the ends of DNA molecules in germ cells.
- Telomere prevent DNAse from degrading the end of linear DNA molecules.
- Facilitates replication of the end of linear DNA molecule without loss.
- It gives stability and integrity to chromosome.
- It has been proposed that the shortening of telomeres is connected to aging
- There is evidence of enhanced telomerase activity in cancer cells, which may allow cancer cells to persist.

