

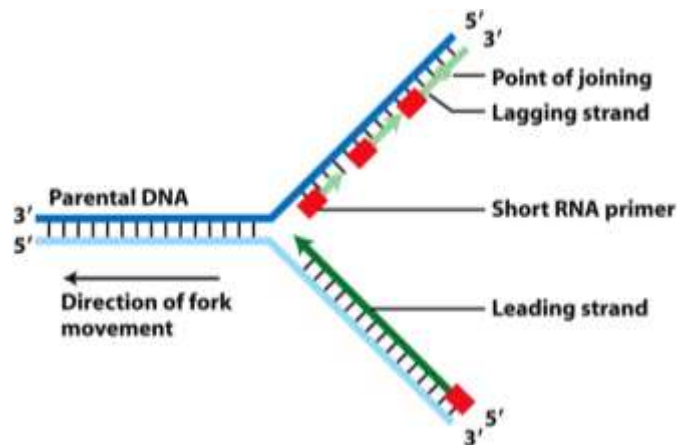
1.2. Replication of DNA

DNA has the capability of _____ itself, to make an identical copy of its DNA. This process is performed at the beginning of every cell division so that when the cell divides, each daughter cell will inherit an _____ copy of the DNA.

There are five requirements for the replication of DNA:

1 -	each strand can be used as a template to create a new DNA strand
2 -	start process since DNA polymerase can only add nucleotides to an existing strand of DNA
3 -	DNA Polymerase - adds nucleotides to a growing strand of DNA Ligase - joins the fragments together
4 -	needed to form the new strands
5 -	a chemical which supplies energy for the process to occur

Stages of DNA replication



Stage 1 - The DNA double helix molecule _____ and weak _____ bonds between base pairs _____, allowing the two strands to separate ('unzip'). These template strands become stabilised and expose their bases at a Y-shaped **replication fork**.

Stage 2 - _____ adds complementary free DNA nucleotides (found in the nucleus) to the now exposed bases on both strands (A-T and C-G) in a ___' to ___' direction. A **primer** is needed to start replication, since DNA polymerase can only add nucleotides to a pre-existing chain.

1. **Leading strand** is synthesised _____. DNA polymerase adds nucleotides to the deoxyribose (3') ended strand in a 5' to 3' direction.
2. **Lagging strand** is synthesised in _____. Nucleotides cannot be added to the phosphate (5') end because DNA polymerase can only add DNA nucleotides in a 5' to 3' direction. The lagging strand is therefore synthesised in fragments. The fragments are then sealed together by an enzyme called _____.

Stage 3 - The two new strands twist to form a _____. Each is identical to the original strand. Each DNA molecules is known as being _____.

Polymerase chain reaction (PCR)

The Polymerase Chain Reaction (PCR) is a technique used to increase or amplify DNA *in vitro* (process happens outside the body of an organism) so that many copies of it can be made from a very small amount.

The PCR process

1. The DNA to be amplified, is **heated** to ____°C to **break** the _____ bonds between the bases, therefore **separating the two strands**.
2. The DNA is then **cooled** to approximately ____°C to allow the complementary _____ to **bind** to its **target sequence** at the 3' end of the original DNA strands.
3. It is then **heated again** to ____°C.
4. **Heat-tolerant** _____ is added, which **adds nucleotides to the primers** at the 3' end of the original DNA strands. **Two strands are formed**.
5. Repeated cycles of heating and cooling are then carried out using the original and the new copies of the DNA to produce millions of copies, within about 3 hours.

Positive and negative controls

It is essential to set up control experiments to allow you to verify that you are measuring what you intended.

control	contains a template of DNA with a known sequence to which the primers are complementary. This means that if it is unsuccessful, you know that there is something wrong with the set-up, e.g. problems with the primers or the conditions.
control	would have a template which is not complementary to the primers or might have no DNA template. This means that if there is amplification, there must be some contamination.

Practical applications of PCR

This technique allows scientists to easily and cheaply turn a single strand of DNA into millions of copies which can then be used for analysis.

The analysis of DNA is used in:

- the Human Genome Project,
- phylogenetics,
- research,
- _____ testing,
- prenatal diagnosis of inherited _____ disorders
- and the detection of infection.