



M.Sc Mazin Eidan Hadi & M.Sc Zahraa Aqeel  
[mazin.eidan@uomus.edu.iq](mailto:mazin.eidan@uomus.edu.iq) & [zahraa.aqeel@uomus.edu.iq](mailto:zahraa.aqeel@uomus.edu.iq)

---

## Lab.4

### Polymerase Chain Reaction (PCR)

**Polymerase Chain Reaction (PCR)** is an in-vitro technique for amplification of specific region of DNA whose sequence is known, or which lies between two regions of known sequence, The PCR amplifies only a chosen segment (the target sequence) within the original DNA template, not the whole template DNA molecule.

#### History of PCR

- ❖ Before PCR, DNA of interest could only be amplified by over-expression in cells and this with limited yield
- ❖ 1966, Thomas Brock discovers *Thermus Aquaticus*, a thermostable bacterium in the hot springs of Yellowstone National Park
- ❖ 1983, Kary Mullis postulated the concept of PCR (Nobel Prize in 1993)
- ❖ 1985, Saiki publishes the first application of PCR (beta-Globin)
- ❖ 1985, Cetus Corp. Scientists isolate Thermostable Taq Polymerase (from *T. aquaticus*), which revolutionized PCR.

#### Reaction Components

PCR is used in clinical diagnosis, genetic analysis, genetic engineering, and forensic analysis. In particular, PCR has revolutionized and speeded up the whole area of recombinant DNA technology.

**The components involved in the polymerase chain reaction are as follows:**

1. The original DNA molecule that is to be copied is called the **template** and the segment of it that will actually be amplified is known as the **target sequence**. A trace amount of the DNA template is sufficient.



M.Sc Mazin Eidan Hadi & M.Sc Zahraa Aqeel  
[mazin.eidan@uomus.edu.iq](mailto:mazin.eidan@uomus.edu.iq) & [zahraa.aqeel@uomus.edu.iq](mailto:zahraa.aqeel@uomus.edu.iq)

---

2. Two **PCR primers** are needed to initiate DNA synthesis. These are shortpieces of single-stranded DNA that match the sequences at either end of the target DNA segment.

- Not containing inverted repeat sequences to avoid formation of internal structures
- not complimentary to each other
- 40-60% GC content preferred for better annealing
- $T_m$  of primers can be calculated to determine annealing  $T_0$
- $T_m = .41(\%G+C) + 16.6\log(J+) + 81.5$  where  $J+$  is the concentration of monovalent ions

3. The enzyme **DNA polymerase** is needed to manufacture the DNA copies. The PCR procedure involves several high temperature steps so a heat resistant DNA polymerase is required. This came originally from heat resistant bacterial living in hot springs at temperatures up to 90°C. **Taq polymerase** from **Thermus aquaticus** is most widely used.

4. A supply of nucleotides (**dNTPs**) is needed by the polymerase to make the new DNA. these are supplied as the nucleoside triphosphates.

5. Finally we need a **PCR machine** to keep changing the temperature The PCR process requires cycling through several different temperatures. Because of this, PCR machines are sometimes called thermocyclers.

6- Buffer and  $Mg^{2+}$



M.Sc Mazin Eidan Hadi & M.Sc Zahraa Aqeel  
[mazin.eidan@uomus.edu.iq](mailto:mazin.eidan@uomus.edu.iq) & [zahraa.aqeel@uomus.edu.iq](mailto:zahraa.aqeel@uomus.edu.iq)

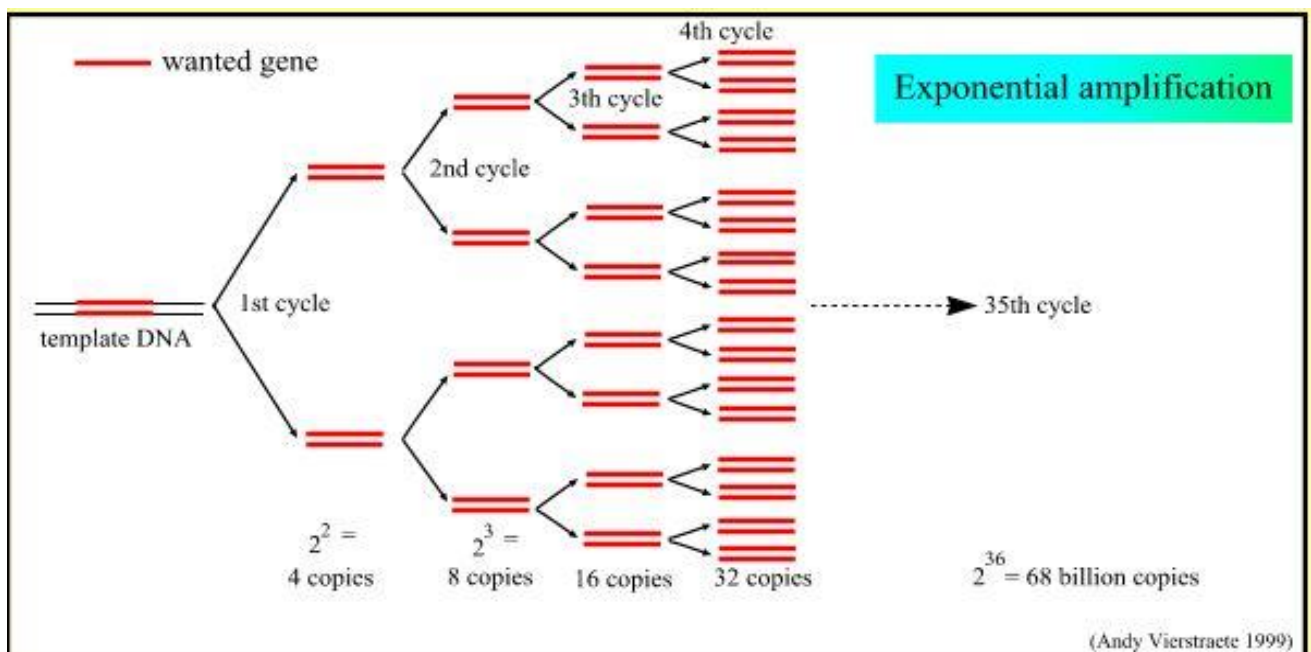
## The PCR Cycle

Consist of 3 steps

1 - **Denaturation** of DNA at 95°C

2-**Primer hybridization**(annealing)at40-50°C 3- **DNA**

**synthesis** (Primer extension) at 72°C.

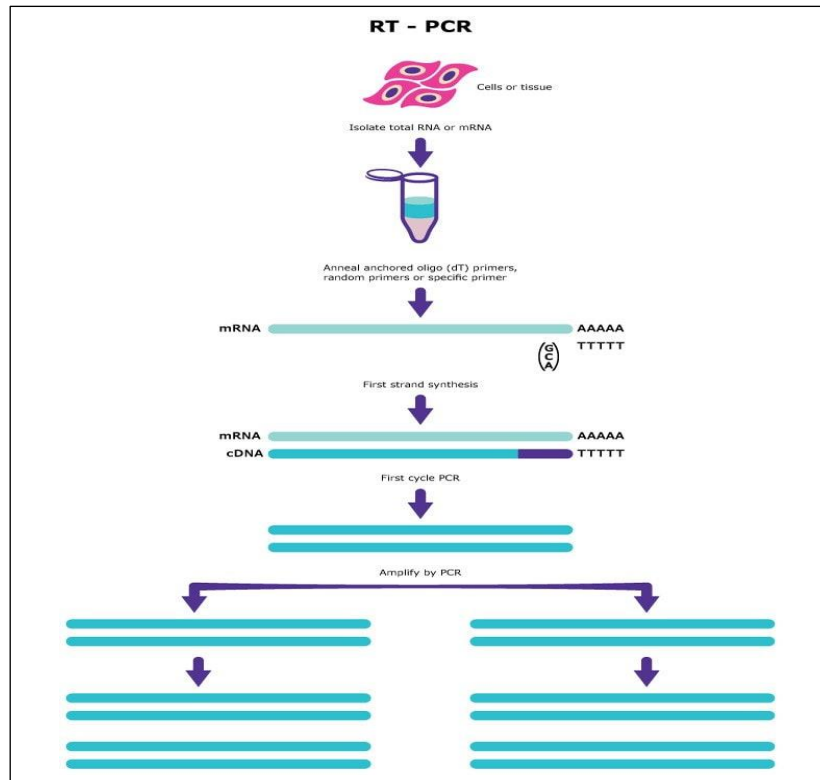


## Reverse Transcription PCR (RT-PCR)

Reverse Transcription PCR Uses RNA as the initial template, yields ds cDNA, Real time RT-PCR is a nuclear-derived method for detecting the presence of specific genetic material in any pathogen, including a virus. Originally, the method used radioactive isotope markers to detect targeted genetic materials, this technique allows scientists to see the results almost immediately while the process is still ongoing, whereas conventional RT-PCR only provides results at the end of the process. Real time RT-PCR is one of the most widely used laboratory methods for detecting the COVID-19 virus, Ebola virus and Zika virus.



M.Sc Mazin Eidan Hadi & M.Sc Zahraa Aqeel  
[mazin.eidan@uomus.edu.iq](mailto:mazin.eidan@uomus.edu.iq) & [zahraa.aqeel@uomus.edu.iq](mailto:zahraa.aqeel@uomus.edu.iq)



## Detection of amplification products

- 1- Gel electrophoresis
- 2- Sequencing of amplified fragment
- 3- Southern blot .... ect

## Advantage of PCR

- ❖ Automated, fast, reliable (reproducible) results
- ❖ Contained :(less chances of contamination)
- ❖ High output
- ❖ Sensitive
- ❖ Broad uses
- ❖ Defined, easy to follow protocols.



M.Sc Mazin Eidan Hadi & M.Sc Zahraa Aqeel  
[mazin.eidan@uomus.edu.iq](mailto:mazin.eidan@uomus.edu.iq) & [zahraa.aqeel@uomus.edu.iq](mailto:zahraa.aqeel@uomus.edu.iq)

---

## Application of PCR

- ❖ Genome mapping and gene function determination
- ❖ Biodiversity studies ( e.g. evolution studies)
- ❖ Diagnostics (prenatal testing of genetic diseases, early detection of cancer, viral infections...)
- ❖ Detection of drug resistance genes
- ❖ Forensic (DNA fingerprinting)