

# An Introduction to Polymerase Chain Reaction (PCR)

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# Introduction

- The technique was invented by **Dr. Kary Mullis, 1986**
- for which he received the **Nobel Prize in Chemistry in 1993.**



## PCR Achieves Fame and Fortune

*--becomes standard in molecular biology tool box--*



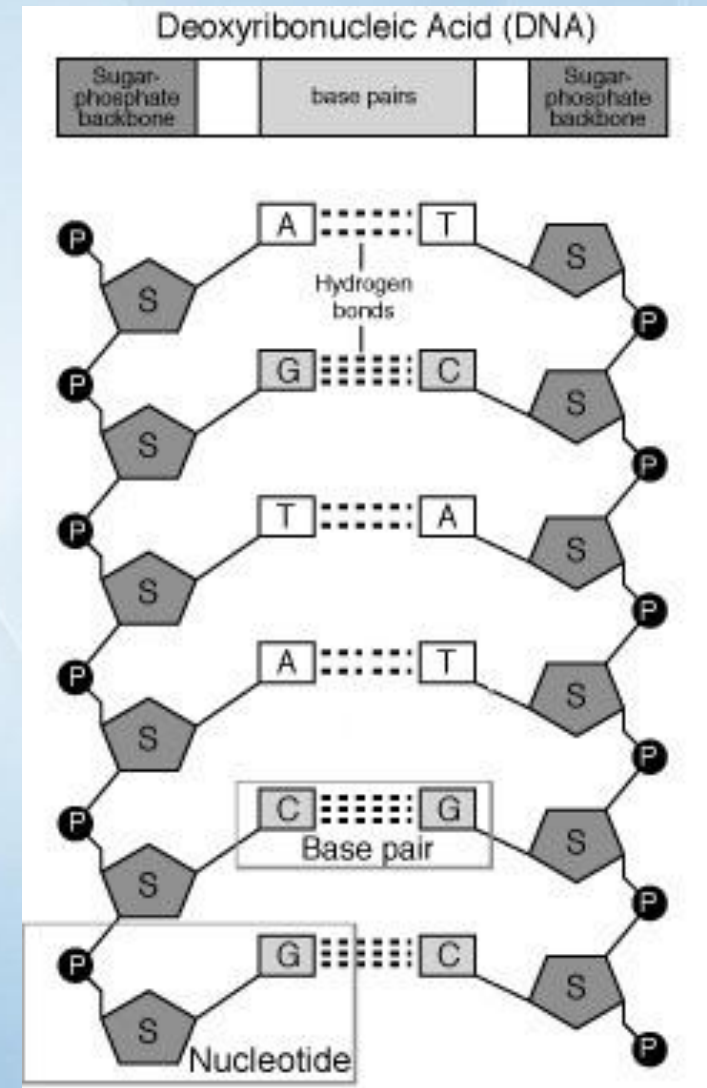
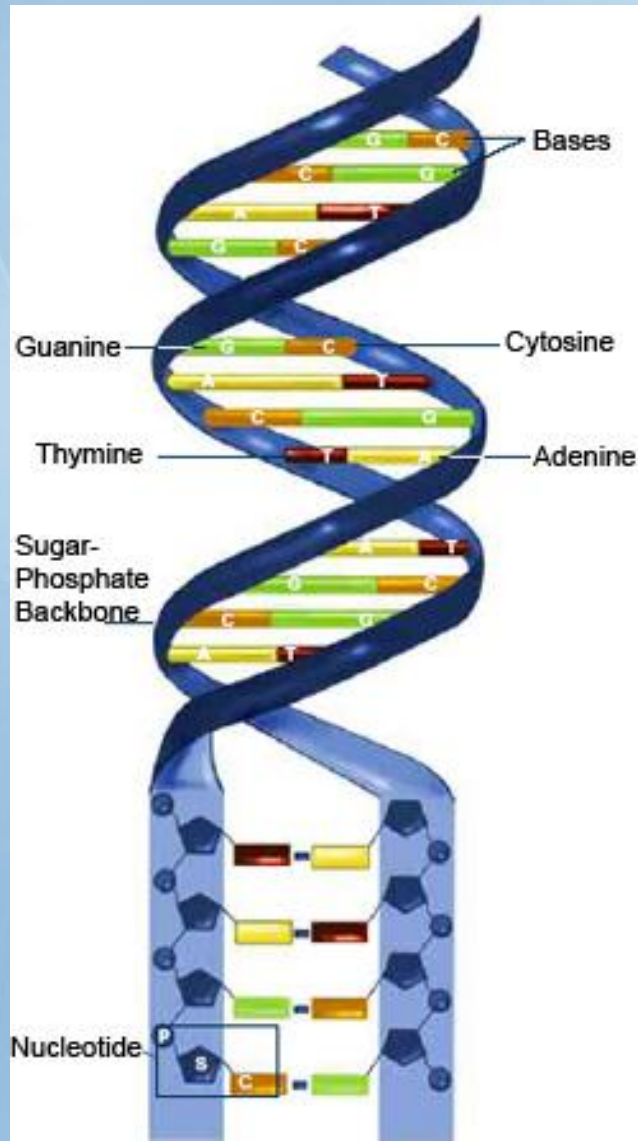
### The Molecule of the Year

RUTH LEVY GUYER AND  
DANIEL E. KOSHLAND, JR.

**S**cience HAS SELECTED THE POLYMERASE CHAIN REACTION AS the major scientific development of 1989 and has chosen for its first "Molecule of the Year" the DNA polymerase molecule that drives the reaction. The list from which the polymerase chain reaction (PCR) was chosen included an impressive array of accomplishments in many areas of science and technology; additional kudos are therefore conferred below to 17 of the other big "stories" that made 1989 an exciting year for scientists and for followers and beneficiaries of science. Although the PCR procedure was introduced several years ago, use of the technique truly burgeoned in 1989; in much the same way, the full potentials of many of the interesting "runner-up" scientific achievements of this year are likely to be realized sometime in the years to come.

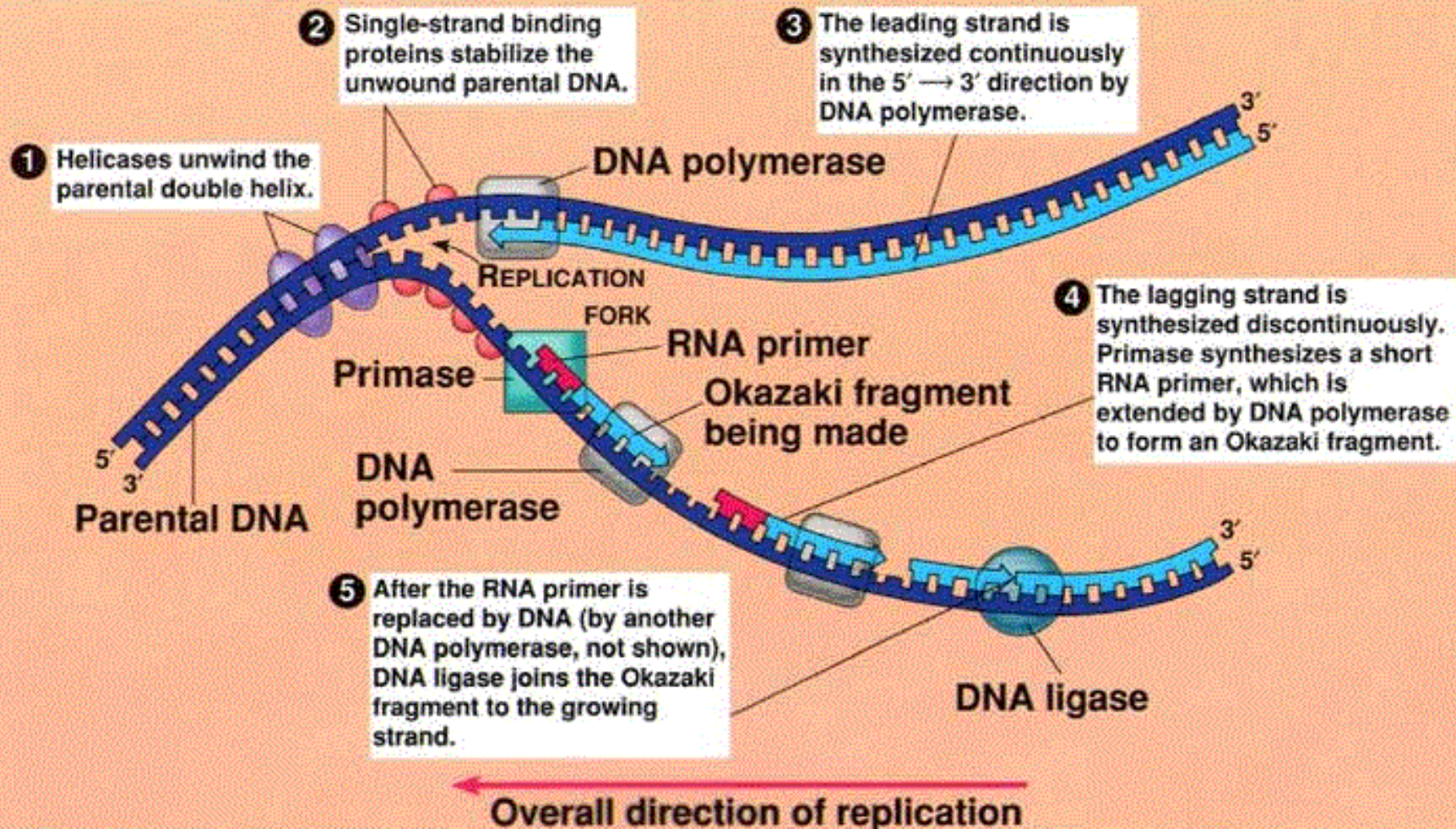


# DNA Structure





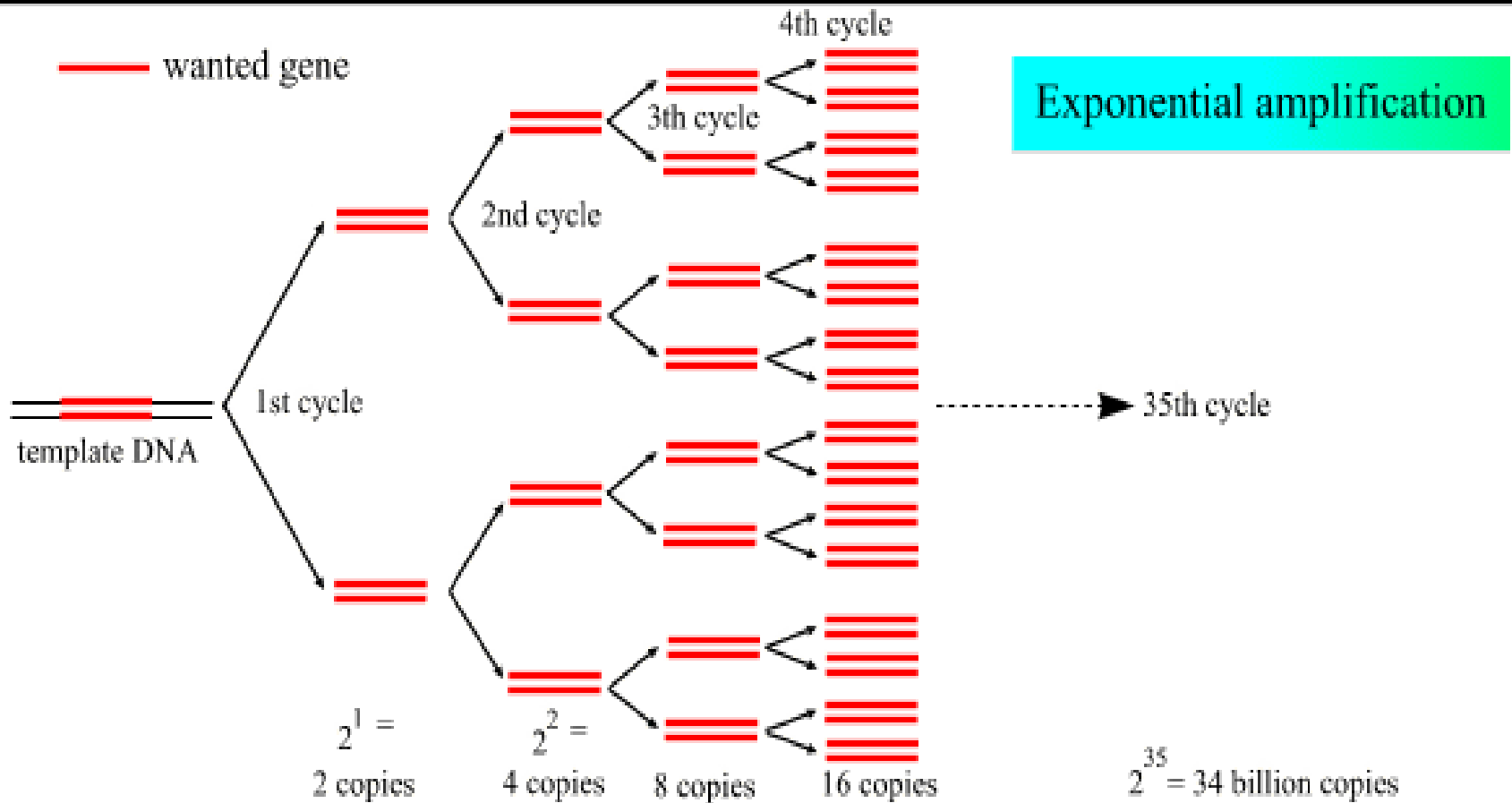
# DNA Replication



# Polymerase Chain Reaction (PCR)

- PCR is a technique which is used to amplify the number of copies of a **specific region of DNA**, (usually fewer than 3000 base pairs) in order to produce enough DNA to be adequately tested.
- **Millions** of copies of a segment of DNA can be made within a few hours
- As a result, it now becomes possible to analyze and characterize the DNA.

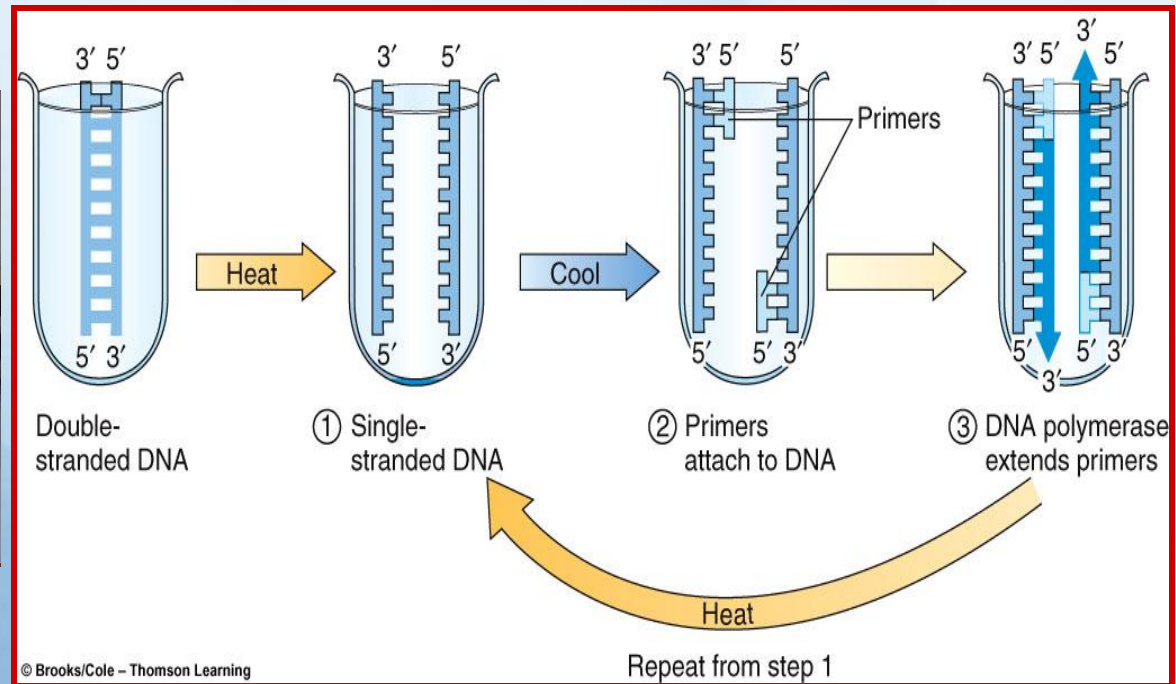
# ● DNA amplification by PCR (overview)



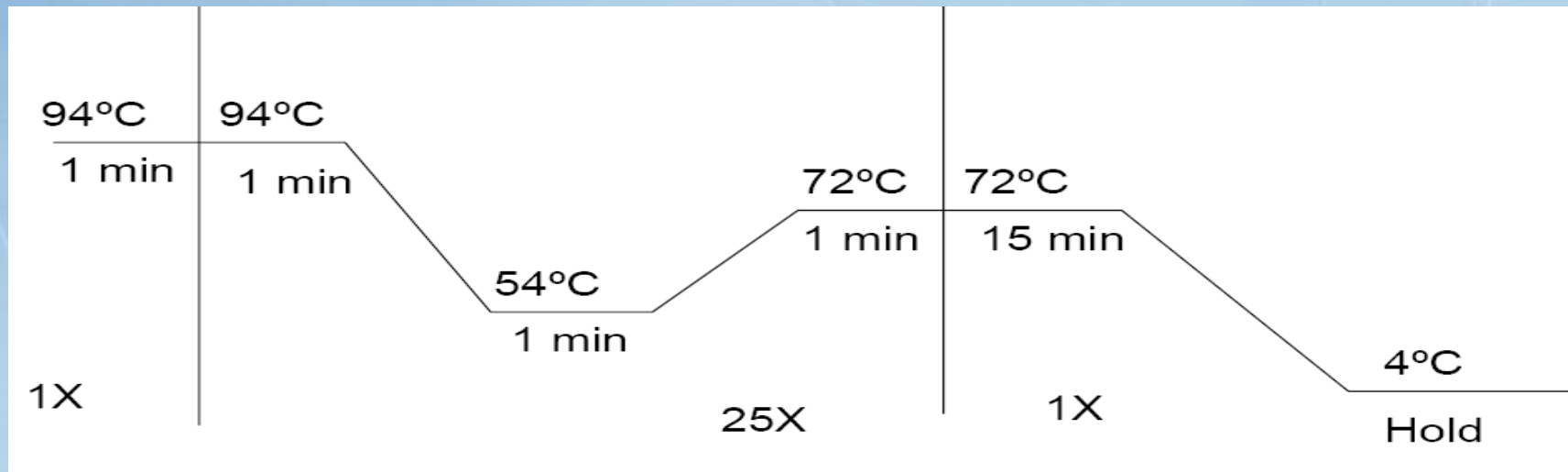


# PCR Cycle

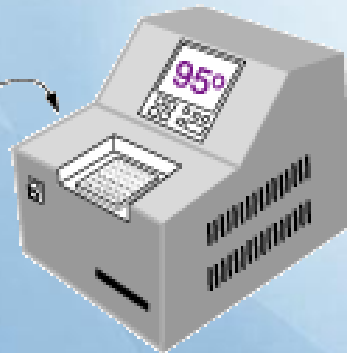
- **Each cycle (Round) of PCR contains 3 steps:**
  - 1- Denaturation**
  - 2- Primer annealing**
  - 3- Primer extension**
- **The cycle usually repeated for 25 – 40 times.**



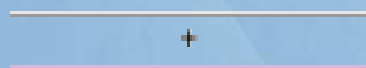
# Programming the Thermocycler



Reaction buffer  
DNA template  
PCR primers  
Taq DNA pol



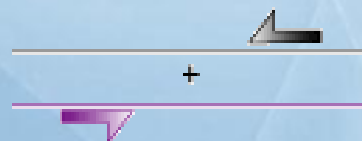
Denature Template



1



Anneal Primers



2



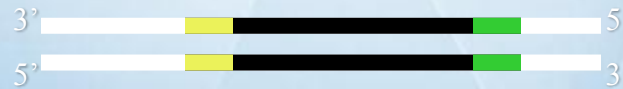
DNA Synthesis



3



# PCR



# PCR

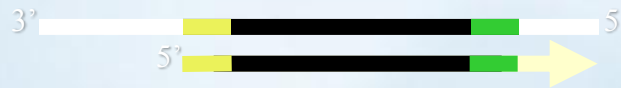
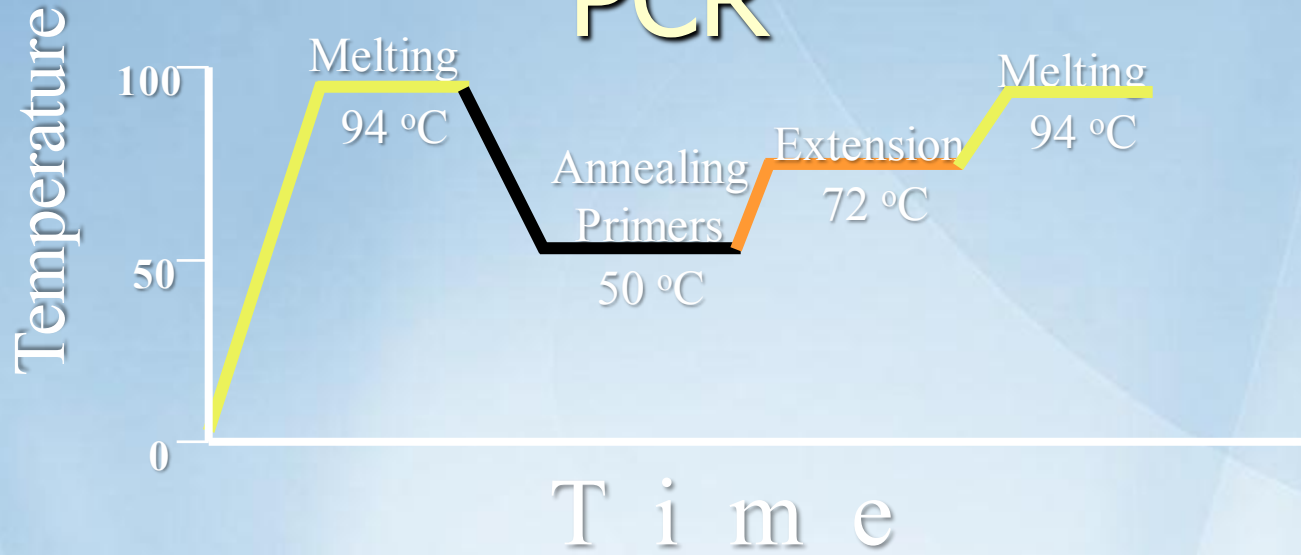


3' 5'

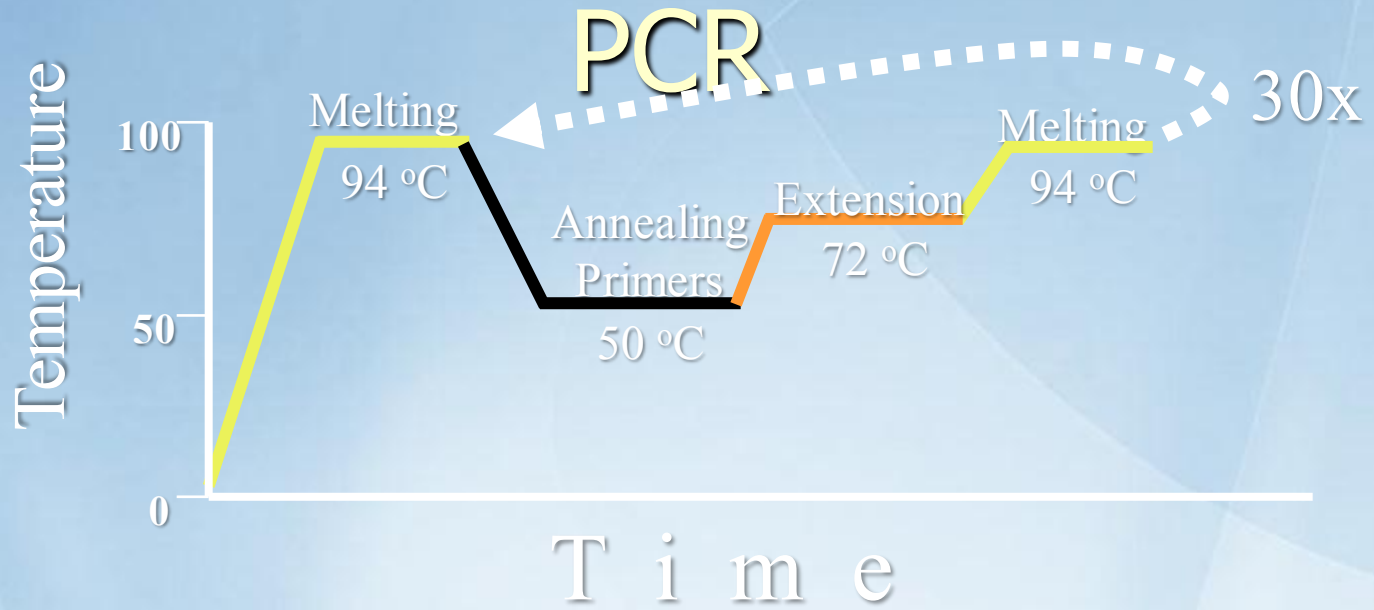


5' 3'

# PCR







3' ——— 5'

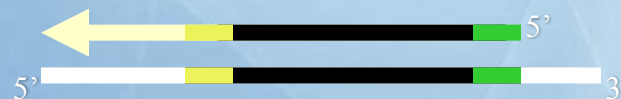
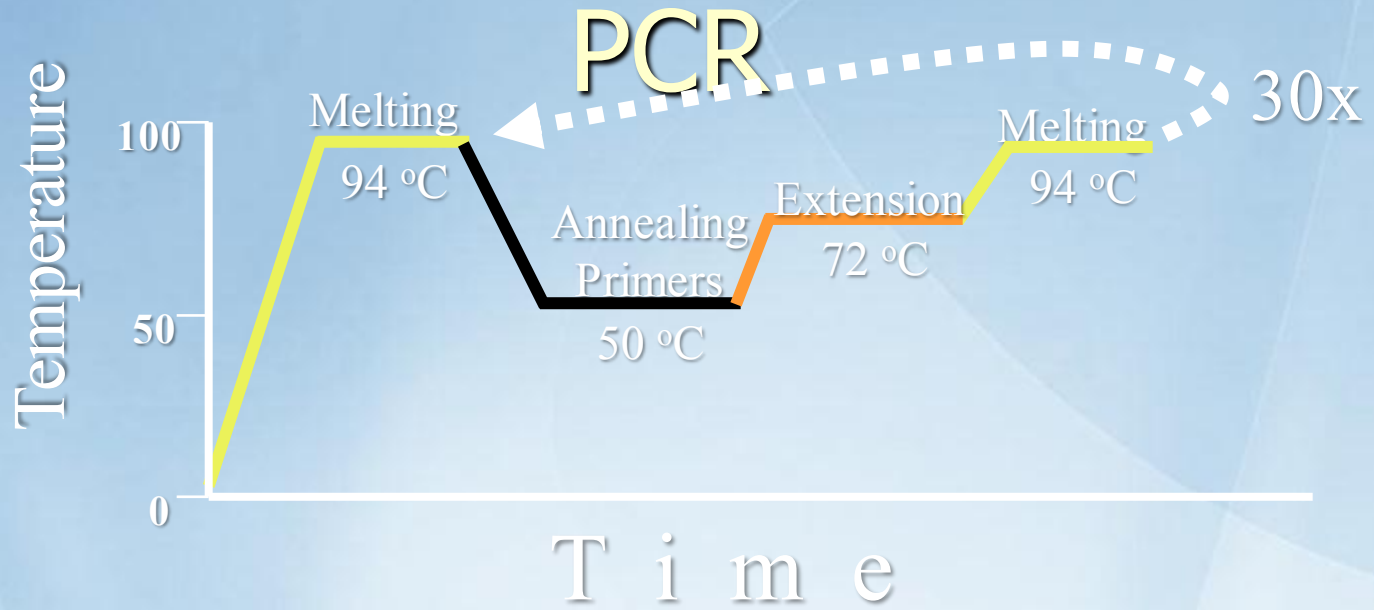


5' ———→

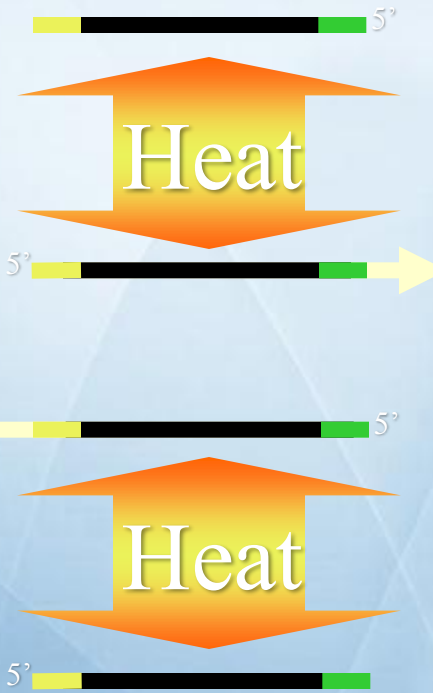
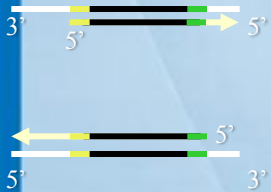
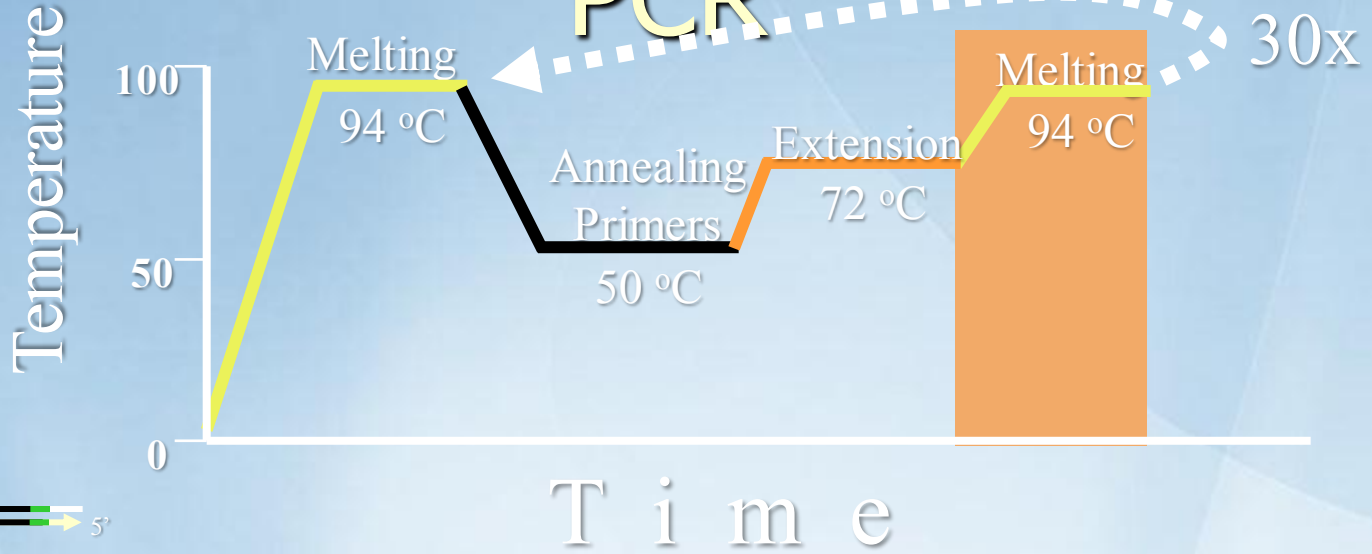
←—— 5'



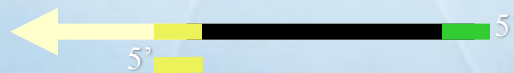
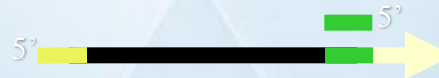
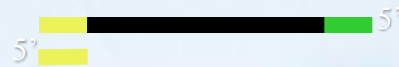
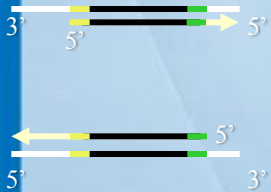
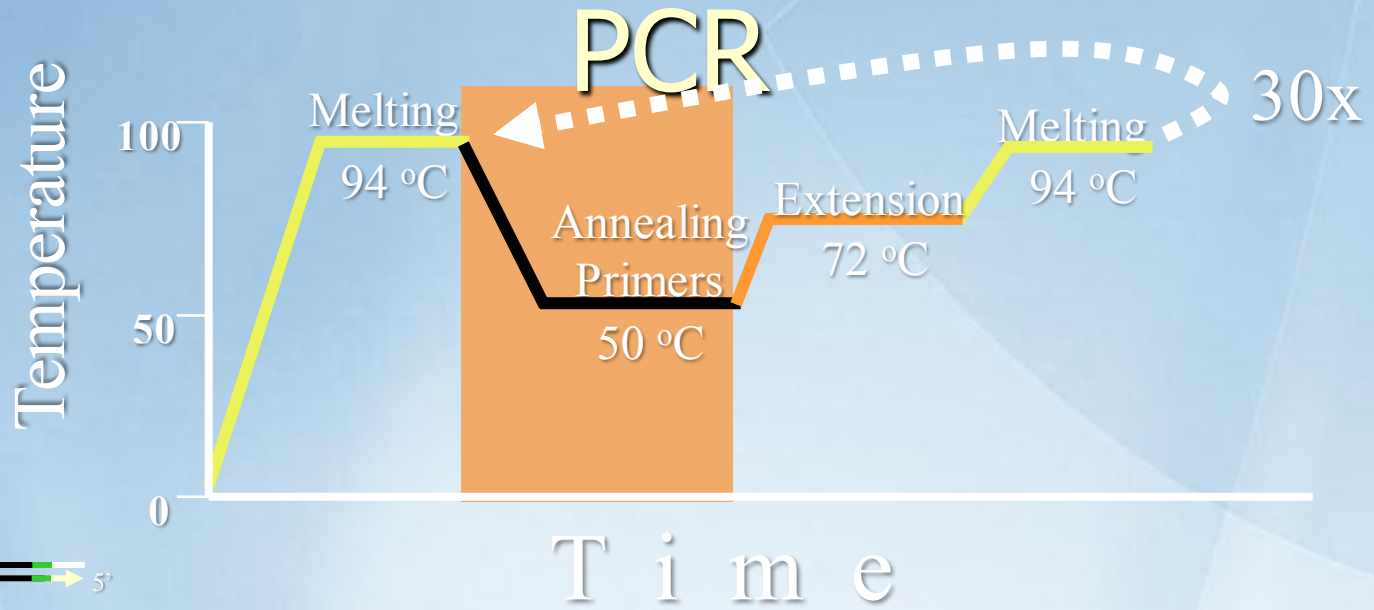
5' ——— 5' 3'



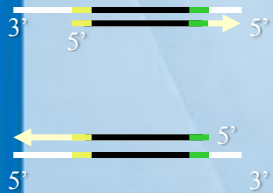
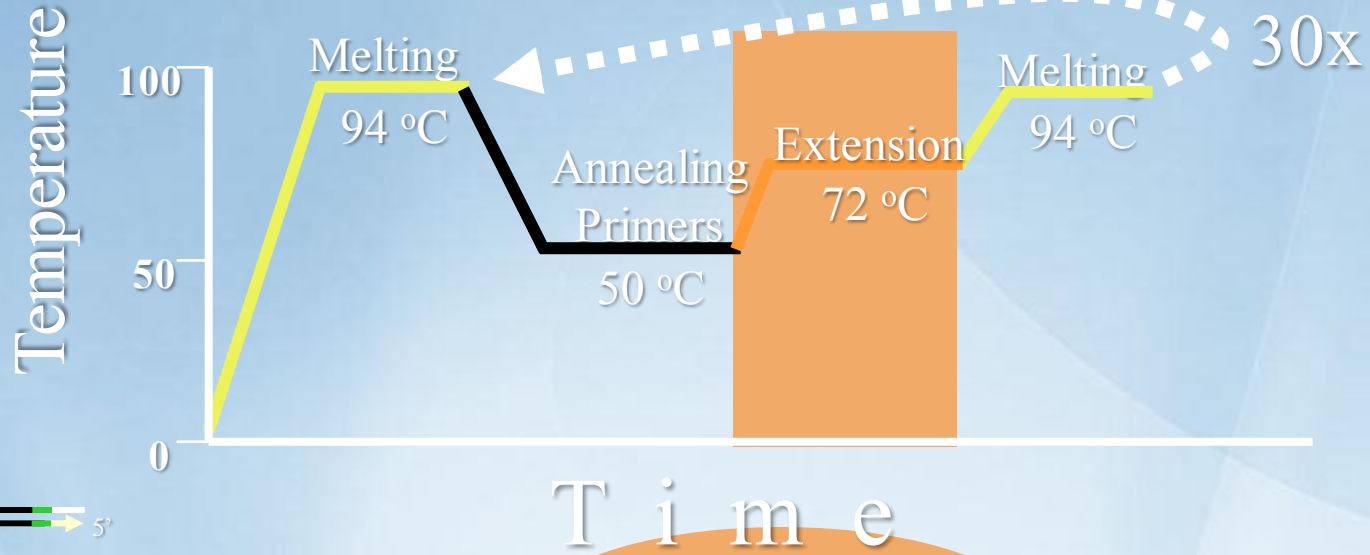
# PCR



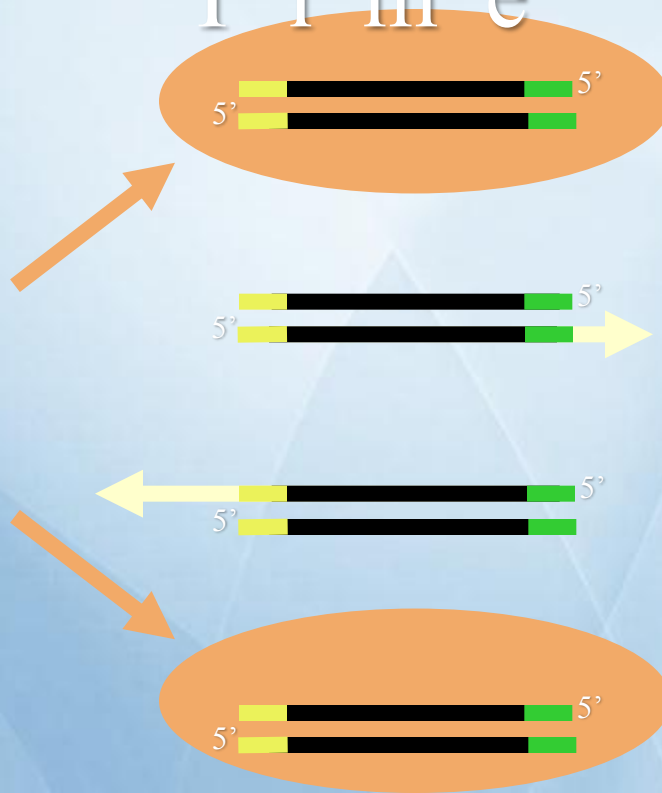




# PCR



Fragments of defined length



# Movie



# What do we need for PCR?



**PCR tube**

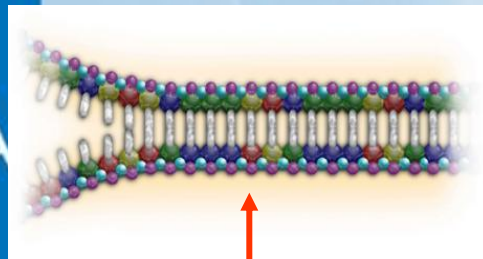
**THERMOCYCLER**

# What do we need for PCR?

- **Target DNA (Template).**
- **Two primers: (forward and reverse)**
- **Nucleotides: (the 4 dNTP'S: A, T, C, G)**
- **Heat-stable DNA polymerase:**  
(like **Taq DNA Polymerase**)
- **Buffer and Cofactor  $\text{MgCl}_2$  ( $\text{Mg}^{++}$ ,  $\text{K}^+$ ).**
- **Thermal cycler.**

# PCR Procedure

**All the required components are inserted into an Eppendorf tube**



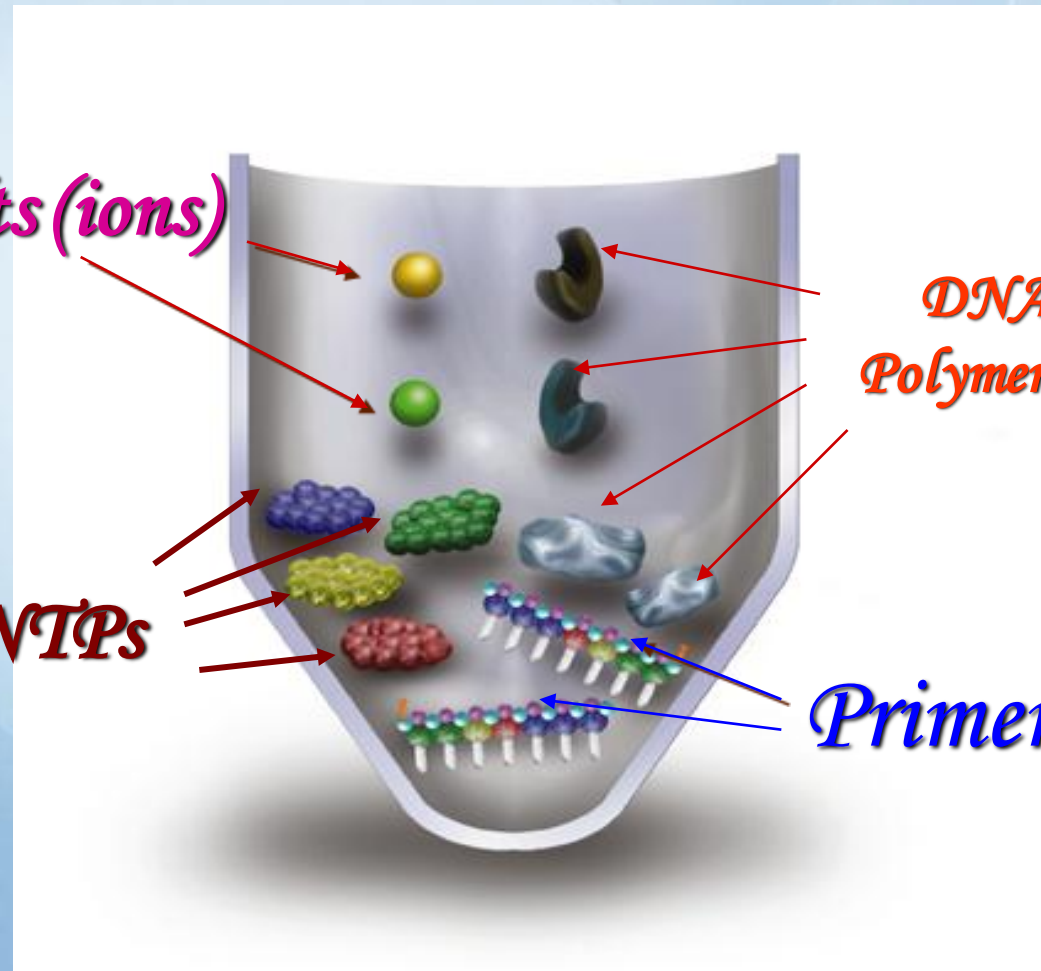
*Template DNA*

*salts (ions)*

*dNTPs*

*DNA  
Polymerase*

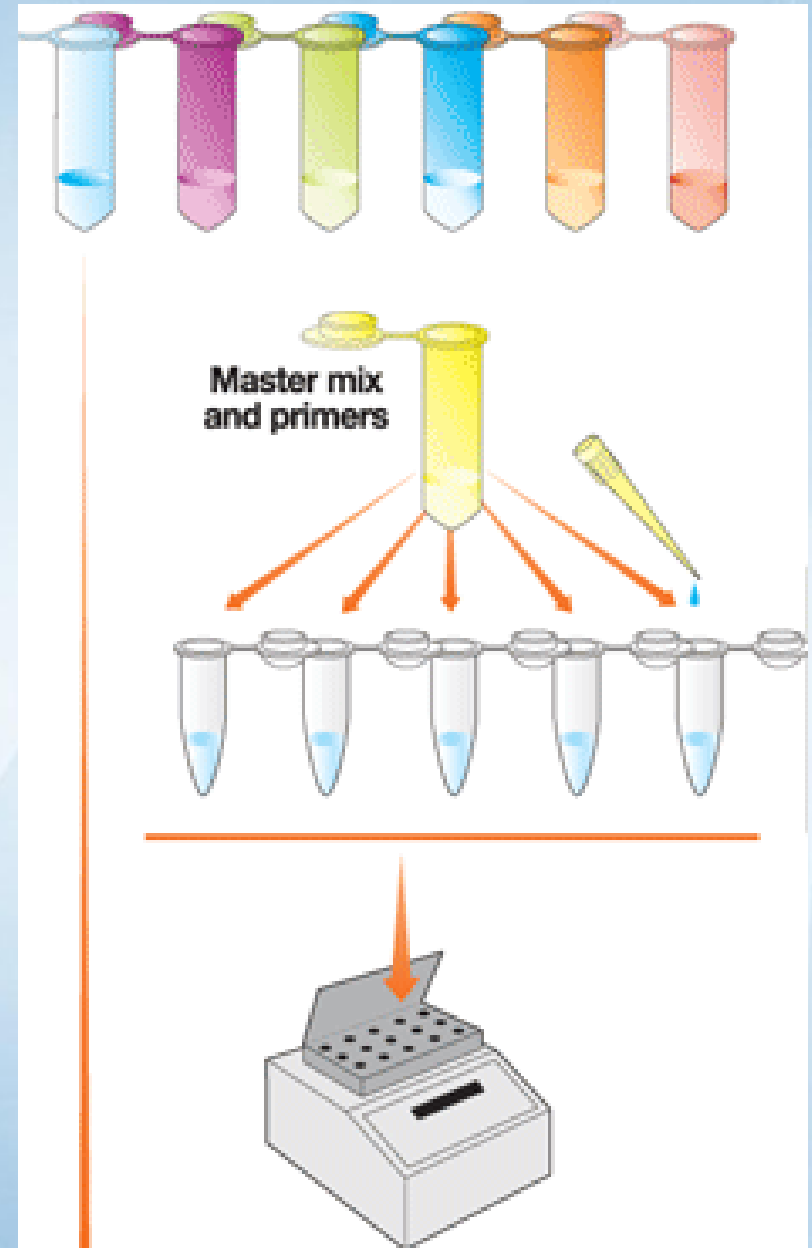
*Primers*





# PROCEDURE .....

- DNA (Template).
- Forward primer
- Reverse primer
- dNTP's
- *Taq* DNA Polymerase
- Buffer
- H<sub>2</sub>O



# PROCEDURE .....

PCR

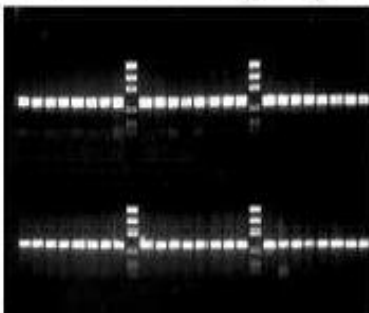


Agarose gel electrophoresis

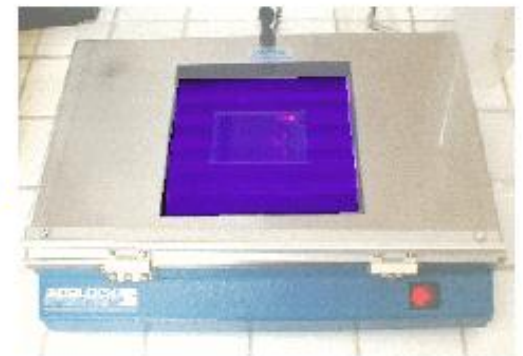


**3-4 hours**

Reliable PCR from Every Sample



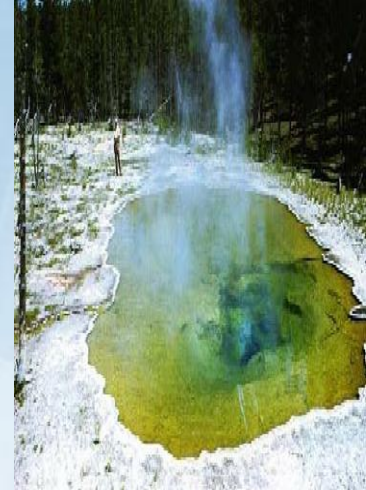
The final product



UV visualisation

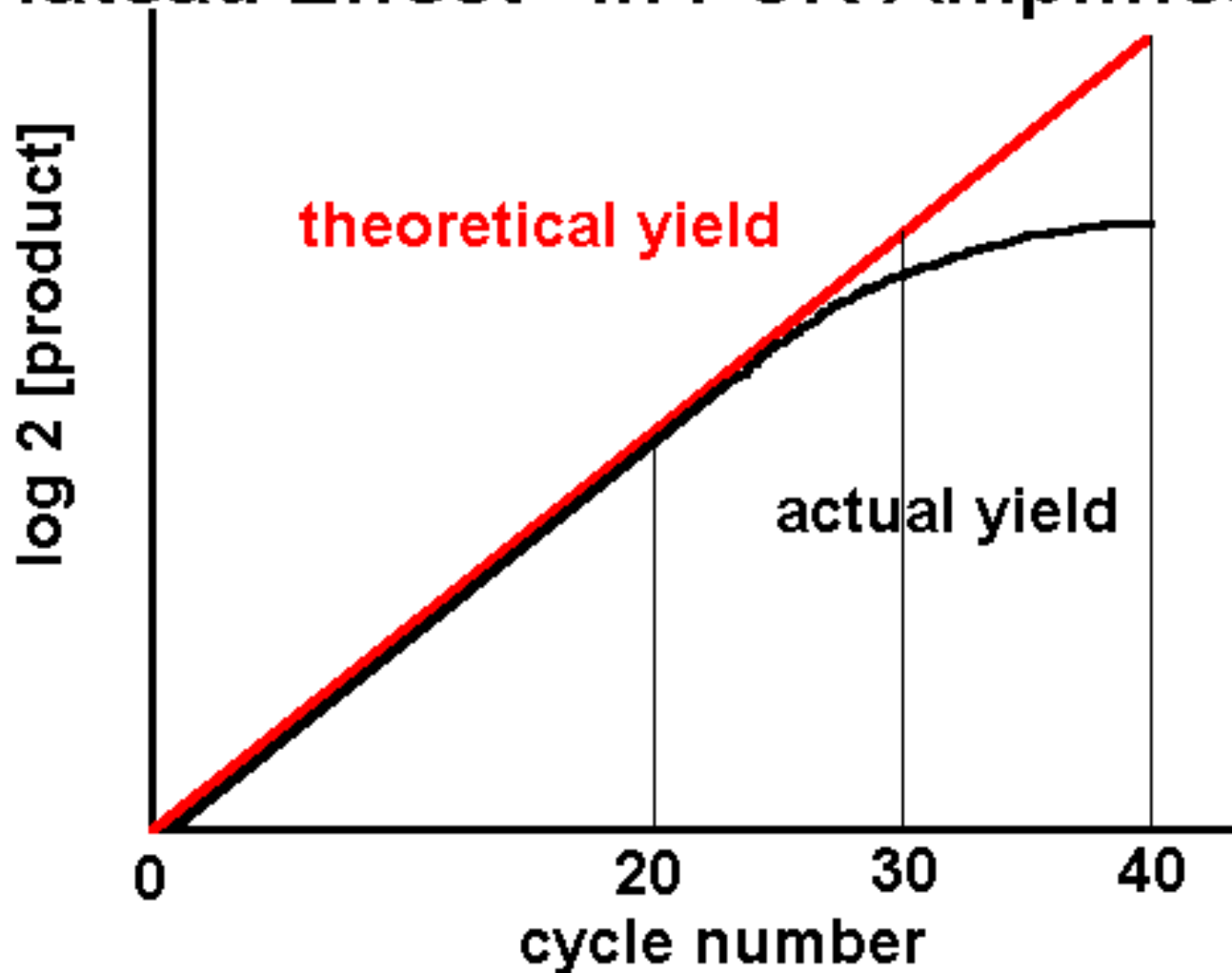
# Development....

- PCR work was first published (1985) using Klenow polymerase - unstable with heat
- First reports using DNA polymerase from *Thermus aquaticus* (1988)
- Developed automatic "thermocycler" programmable heat block



# Cycle Number

## "Plateau Effect" in PCR Amplification






# Types of PCR

- **Standard PCR (conventional )**
- **RT-PCR (Reverse Transcriptase PCR)**
- **Real Time PCR (qRT-PCR)**

# Applications of PCR

- 
- Classification of organisms
  - Genotyping
  - Mutagenesis
  - Mutation detection
  - Sequencing
  - Detection of pathogens
  - DNA fingerprinting
  - Genetic engineering
  - Research

# Variations of the PCR

- Colony PCR
- Nested PCR
- Multiplex PCR
- AFLP PCR
- Hot Start PCR
- Inverse PCR
- Long PCR
- Long Accurate PCR
- Reverse Transcriptase PCR
- Allele specific PCR
- Real time PCR

