Fundamentos de Biologia Molecular

Curso de Licenciatura em Biologia 2º Ano, 1º Semestre Ano Letivo 2018/2019

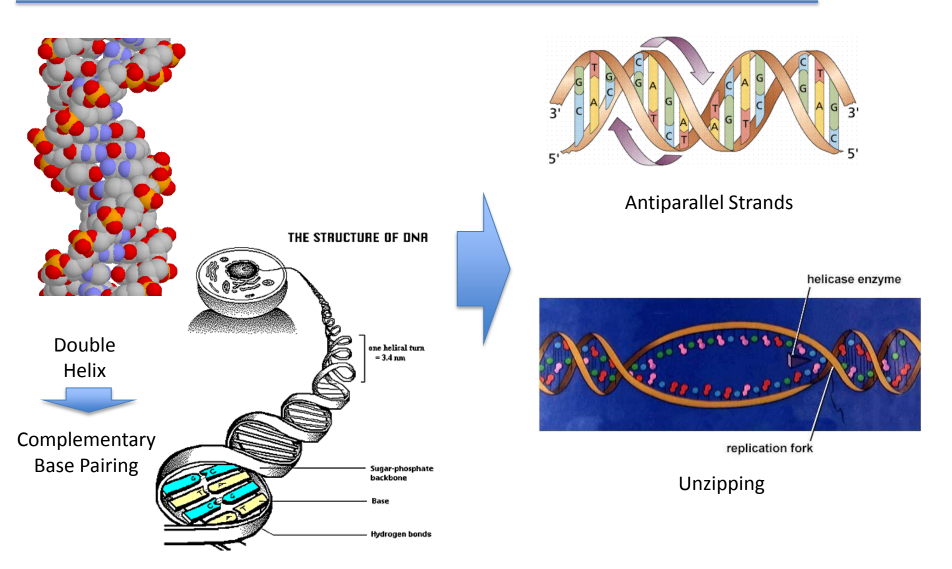
Componente Teórico-Prática



Docente Responsável: **Rita Zilhão** Docente TPs: **Andreia Figueiredo**

- DNA structure
- PCR discovery
- Nobel prizes
- PCR
- PCR components
- Setting up a PCR reaction
- Practical applications

The structure of DNA



The structure of DNA

How do we identify and detect a specific sequence in a genome?

- There are a LOT of other sequences in a genome that we're not interested in detecting. (SPECIFICITY)
- The amount of DNA in samples we're interested in is VERY small. (AMPLIFICATION)

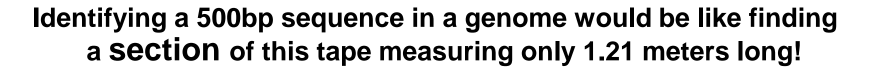


How do we identify and detect a specific sequence in a genome?

- Pine: 68 billion bp
- Corn: 5.0 billion bp
- Soybean: 1.1 billion bp
- Human: 3.4 billion bp
- Housefly: 900 million bp
- Rice: 400 million bp
- E. coli: 4.6 million bp
- HIV: 9.7 thousand bp

Specificity

- The human genome is 3.4 B bp
- If the bases were written in standard 10-point type, on a tape measure...
- ...The tape would stretch for 8635.73 Km!





Detection

How many molecules do we need to be able to see them?

• We need a minimum of 10 ng of fluorescent stained DNA to detect on agarose gel

URI Genomics & Sequencing Center

Calculator for determining the number of copies of a template

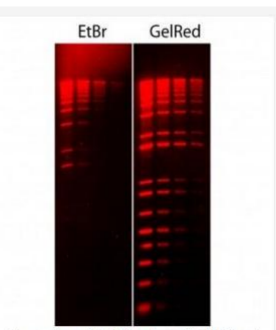
enter amount of DNA (ng): 10

enter length of template (bp): 500

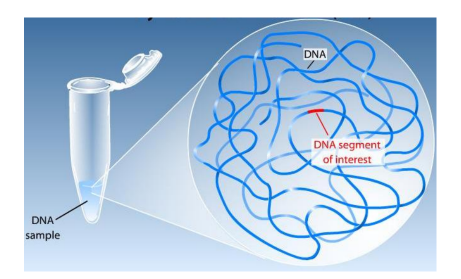
Calculate number of copies!

number of copies : 1.85 x10^ 10

In other words, to "see" a single "gene", the DNA in a sample of 100 cells would have to be multiplied 180 million times!!!!!

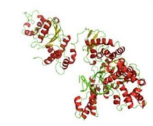


Comparison of ethidium bromide (EtBr) and GelRed in precast gel staining in 1% agarose/TBE gel. Two-fold serial dilutions of 1 kb Plus DNA Ladder (Invitrogen) were loaded in the amounts of 200 ng, 100 ng, 50 ng and 25 ng from left to right. Gels were imaged using 300 nm transilluminator and photographed with an EtBr filter.



PCR is widely used to amplify target DNA regions of known sequences within a heterogeneous collection of DNA sequences

- Taq DNA polymerase isolation from *Thermus* aquaticus by Brock & Freeze - 1969
- Kary Mullis, PCR invention 1988



Habitat of

Thermus aquaticus

Structure of Taq Polymerase



Kary B. Mullis

The inventor of the DNA synthesis process known as the Polymerase Chain Reaction (PCR). The process is an invaluable tool to today's molecular biologists and biotechnology corporations.

Mullis, born in Lenoir, North Carolina, attended the University of Georgia Tech for his undergraduate work in chemistry, and then obtained a Ph. D. in biochemistry from Cal Berkeley.

In 1983, working for Cetus Corporation, Mullis developed the Polymerase Chain Reaction, a technique for the rapid synthesis of a DNA sequence. The simple process involved heating a vial containing the DNA fragment to split the two strands of the DNA molecule, adding oligonucleotide primers to bring about reproduction, and finally using polymerase to replicate the DNA strands. Each cycle doubles the amount of DNA, so multiple cycles increase the amount of DNA exponentially, creating huge numbers of copies of the DNA fragment.

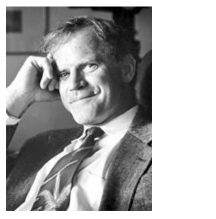


Michael Smith

Smith first conceived of *site-directed mutagenesis* in the early 1970s and devoted several years to working out the details of the technique. The method provided researchers with a new way to study protein function.

•A protein is a compound made up of strings of amino acids that fold into a three-dimensional structure, and the protein's structure determines its function. Instructions for the amino-acid sequence of a protein are contained in its gene, namely, in the sequence of DNA subunits, called nucleotides, that make up that gene. The amino-acid sequence of a protein, and hence its function, can be modified by inducing mutations in the nucleotide sequence of its gene. Once an altered protein has been produced, its structure and function can be compared to those of the natural protein.

Smith developed site-directed mutagenesis, a technique that can be used to modify nucleotide sequences at specific, desired locations within a gene. This has made it possible for researchers to determine the role each amino acid plays in protein structure and function. Aside from its value to basic research, site-directed mutagenesis has many applications in medicine, agriculture, and industry.



Kary B. Mullis



Michael Smith

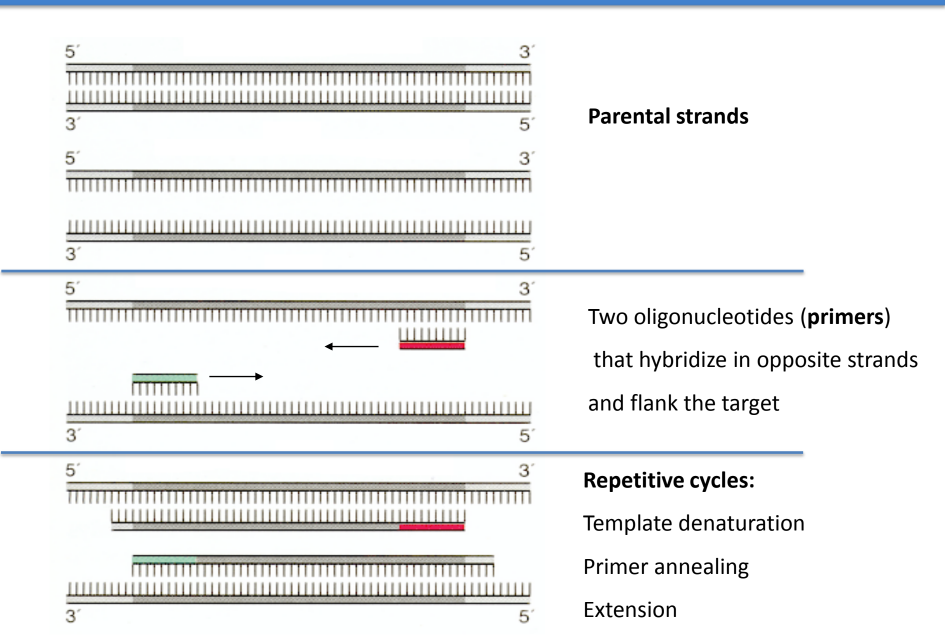
The Nobel Prize in Chemistry 1993 was awarded "*for contributions to the developments of methods within DNA-based chemistry*" jointly with one half to Kary B. Mullis "for his invention of the polymerase chain reaction (PCR) method" and with one half to Michael Smith "*for his fundamental contributions to the establishment of oligonucleotide-based, site-directed mutagenesis and its development for protein studies*".

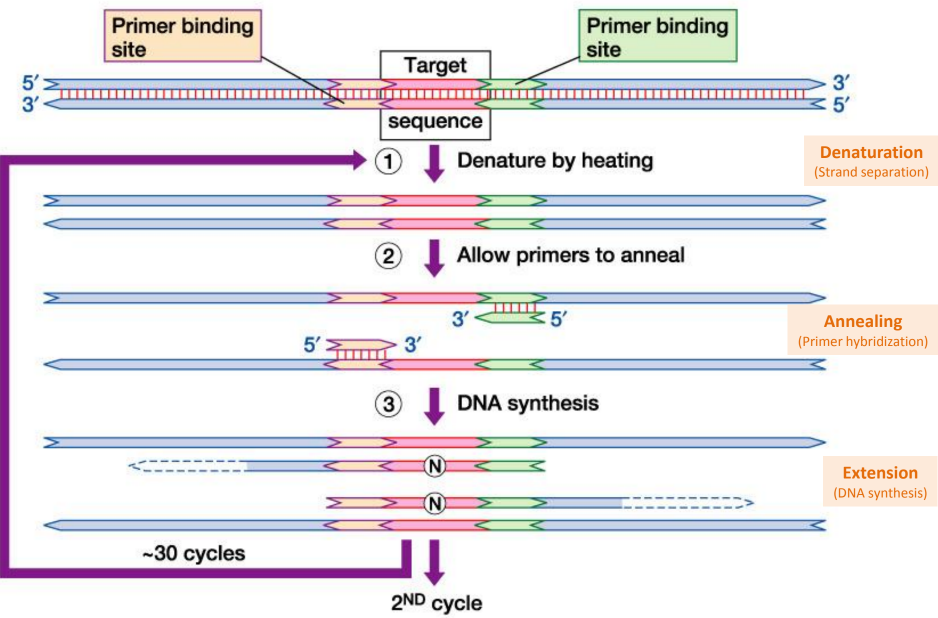
PCR

- Developed by Kary Mullis in 1985,
- PCR results in the selective amplification of a chosen region of a DNA molecule,
- The borders of the amplified region have to be known and two short oligonucleotides have to be synthesized,
- These oligonucleotides, which act as <u>primers</u> for the DNA synthesis, delimit the region to be amplified.



Exponential copies of template DNA





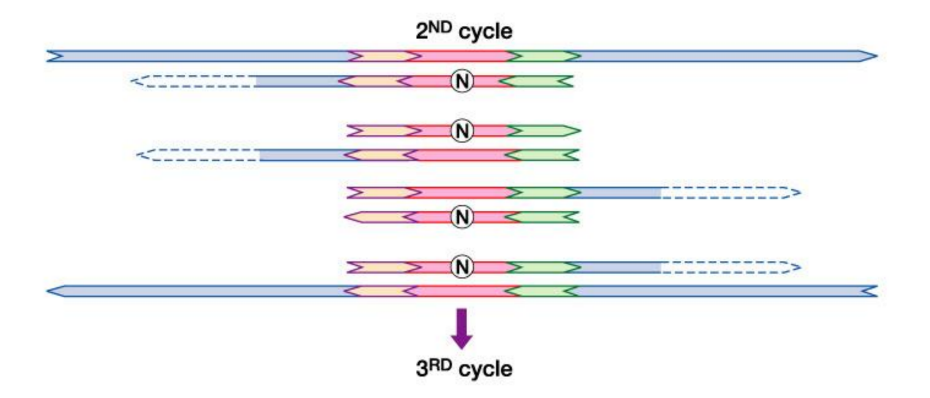


Figure 5-2 part 2 of 3 Human Molecular Genetics, 3/e. (© Garland Science 2004)

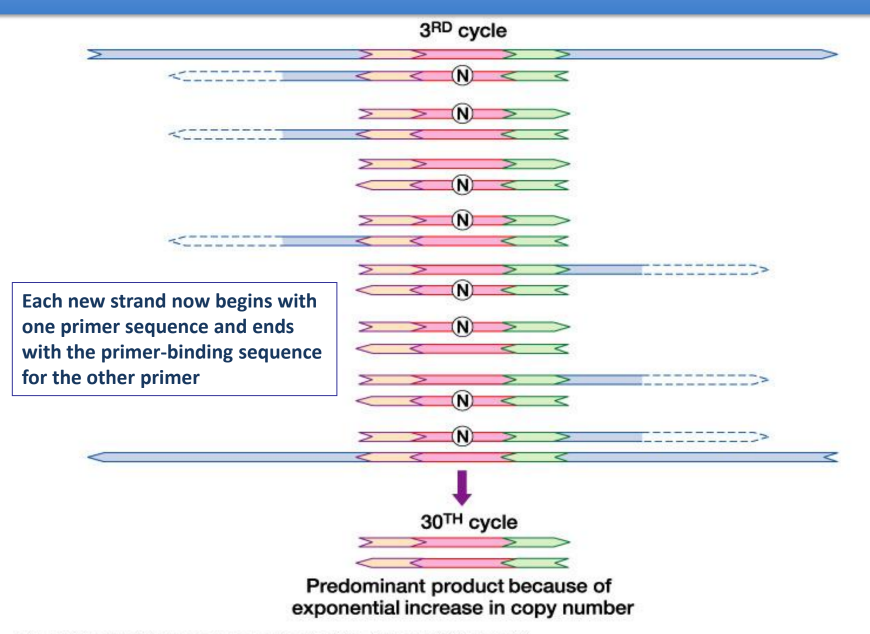
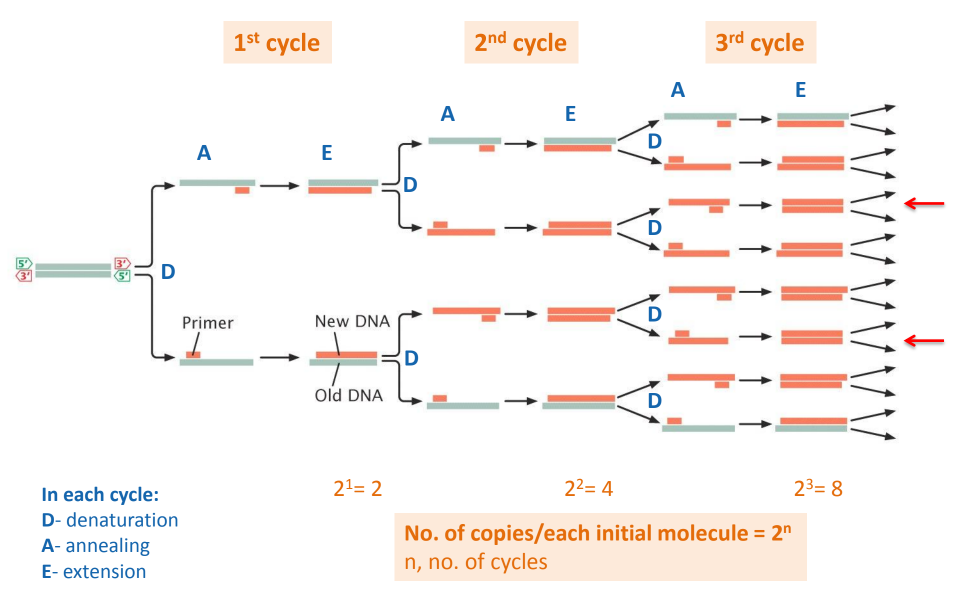


Figure 5-2 part 3 of 3 Human Molecular Genetics 3/e (@ Garland Science 2004)



http://openwetware.org/wiki/BME100_f2013:W900_Group17_L4

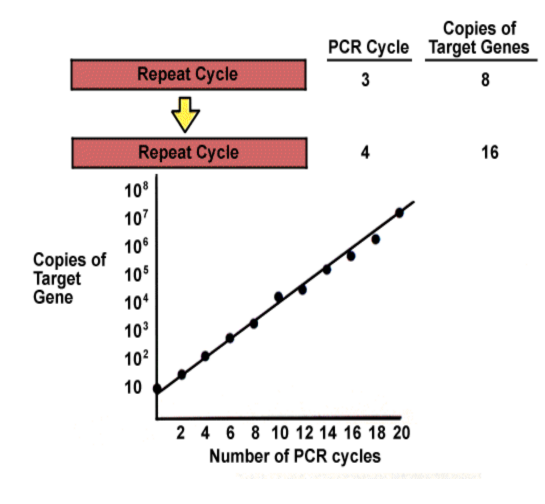


Table 18.5	Number of copies of DNA fragment in PCR amplification
Number of PCR Cycles (<i>n</i>	Number of Double-Stranded () Copies of Original DNA (2")
0	1
1	2
2	4
3	8
4	16
5	32
6	64
7	128
8	256
9	512
10	1,024
20	1,048,576
30	1,073,741,824

n, no. of cycles

Brock, Molecular Biology of Microorganisms

Pierce, Genetics, a conceptual approach

PCR

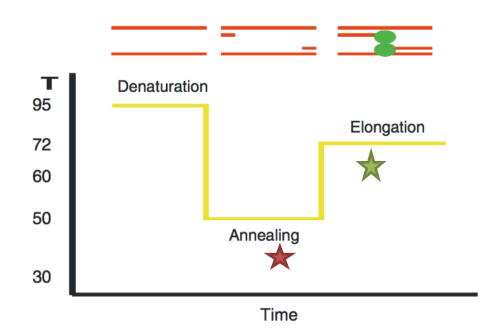


Fig. 1. The PCR temperature cycle: (1) the temperature is raised to about 95 °C to melt the double stranded DNA, (2) the temperature is lowered to let primers anneal, (3) the temperature is set to 72 °C to let the polymerase extend the primers.

melting temperature (T_m)- dependent

Size- and polymerase- dependent

Kubista et al. 2006, Molecular Aspects of Medicine 27, 95–125

PCR reaction

- Water
- Buffer •
- DNA template •
- Primers •
- **Nucleotides** ٠
- Mg²⁺ions •

5'

5'

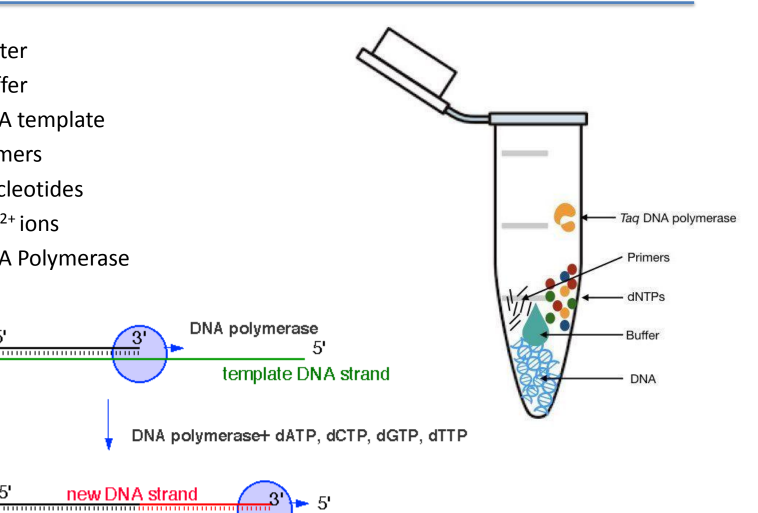
3'

3'

DNA Polymerase •

3'

new DNA strand

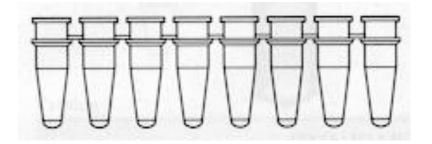


PCR reaction components

- Water
- Buffer
- DNA template
- Primers
- Nucleotides
- Mg²⁺ ions
- DNA Polymerase

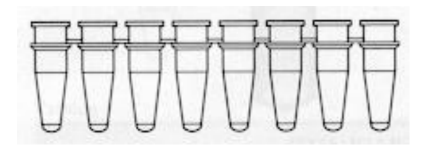
• Water

The medium for all other components.



PCR reaction components

- Water
- Buffer
- DNA template
- Primers
- Nucleotides
- Mg²⁺ ions
- DNA Polymerase

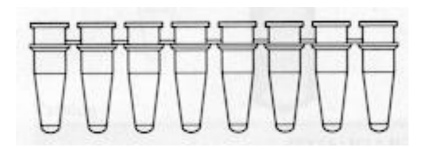


• Buffer

- Stabilizes the
 DNA polymerase,
 DNA, and
 nucleotides
- 500 mM KCl
- 100 mM Tris-HCl, pH 8.3
- Triton X-100 or Tween

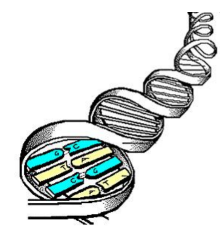
PCR reaction components

- Water
- Buffer
- DNA template
- Primers
- Nucleotides
- Mg²⁺ions
- DNA Polymerase



DNA template

- Contains region to be amplified
- Any DNA desired
- Purity not required
- Should be free
 of polymerase
 inhibitors



PCR reaction components

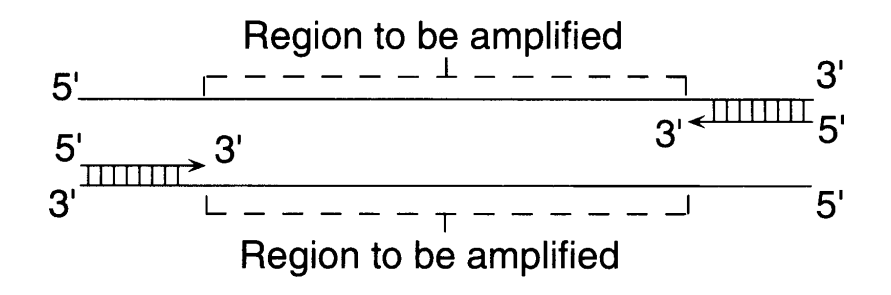
- Water
- Buffer
- DNA template
- Primers
- Nucleotides
- Mg²⁺ions
- DNA Polymerase



Primers

- Specific for ends of amplified region
- Forward and Reverse
- Annealing temps should be known
 - Depends on primer length, GC content, etc.
- Length 15-30 nt
- Conc 0.1 1.0 μM (pMol/μl)

Choosing primer sequences

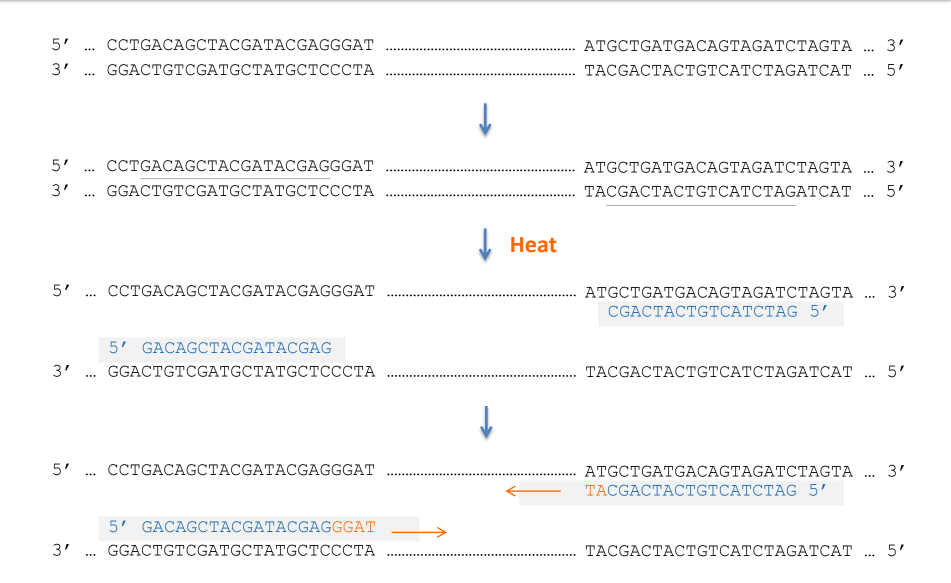


Primer design tips

- Typical PCR primers are anything between 18-28 nucleotides in length
- The G+C composition should ideally be similar to that of the desired amplicon and should in general be between 40-60%
- The calculated Tm for a primer pair should be balanced
- Rule of thumb: Tm = 2(A+T) + 4(G+C) and -1.5 °C for every mismatch
- A Tm 55°C -72°C is desired (62-65°C is best)
- Check for complementarity in 3' ends of primer pairs this leads to primer dimer artifacts
- Avoid any significant secondary structure within primers i.e. internal palindromic sequences
- Runs of 3 or more C's and G's at 3' ends promote mispriming in G/C rich regions
- Avoid an A and especially a T at the 3' end of a primer
- Avoid any potential mismatches in the 3'end of primers (pay attention when designing degenerate primers)

Primer design using Primer3 disclaimer Primer3 Home Checks for mispriming in template Primer3 (v. 0.4.0) Pick primers from a DNA sequence FAQ/WIKI Website: http://bioinfo.ut.ee/primer3-0.4.0/ Primer3plus interface <u>cautions</u> There is a newer version of Primer3 available at http://primer3.ut.ee Paste source sequence below (5'->3', string of ACGTNacgtn -- other letters treated as N -- numbers and blanks ignored). FASTA format ok. Please N-out undesirable sequence (vector, ALUs, LINEs, etc.) or use a Mispriming Library (repeat library): NONE Place here query sequence for which primers will be designed Pick left primer, or use left primer below: Pick hybridization probe (internal oligo), or use oligo below: Pick right primer, or use right primer below (5' to 3' on opposite strand): Pick Primers Reset Form Sequence Id: A string to identify your output Min-Max size of generated amplicon E.g. 50,2 requires primers to surround the 2 bases at positions 50 and 51. Or mark the source sequence with [and]: e.g. ...ATCT[CCCC]TCAT.. means that Targets: primers must flank the central CCCC. (also counting with intermediate Excluded E.g. 401,7 68,3 forbids selection of primers in the 7 bases starting at 401 and the 3 bases at 68. Or mark the source sequence with < and >: e.g. values) Product Size Ranges 150-250 100-300 301-400 401-500 501-600 601-700 701-850 851-1000 Max 3' Stability 9.0 Number To Return e.g. - 200-500 Max Repeat Mispriming 12.00 Pair Max Repeat Mispriming 24.00 Max Template Mispriming 12.00 Pair Max Template Mispriming 24.00 Amplicons with size ranging from 200 Pick Primers Reset Form bp to 500 bp **General Primer Picking Conditions** Primer Size Min: 18 Max: 27 Opt: 20 Primer Tm Min: 57.0 Opt: 60.0 Max: 63.0 Max Tm Difference: 100.0 Table of thermodynamic parameters: Breslauer et al. 1986 • Product Tm Min: Opt Max: Primer GC% Min: 20.0 Opt: Max: 80.0 Min, Max and Opt size/melting Max Self Complementarity: Max 3' Self Complementarity: 3.00 8.00 temperature and GC content of Max #N's: Max Poly-X: primers Inside Target Penalty: Outside Target Penalty: Note: you can set Inside Target Penalty to allow primers inside a target. 0 First Base Index: CG Clamp: 0 Schildkraut and Lifson 1965 V Concentration of monovalent cations: 50.0 Salt correction formula: Concentration of divalent cations 0.0 Concentration of dNTPs 0.0 Annealing Oligo Concentration: 50.0 (Not the concentration of oligos in the reaction mix but of those annealing to template.) 🖉 Liberal Base 🔲 Show Debuging Info 🖉 Do not treat ambiguity codes in libraries as consensus 🗏 Lowercase masking

Pick Primers Reset Form



Primer storage

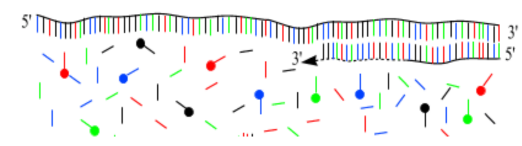
- Oligonucleotides are chemically stable. Left dry, they should be good for years.
 Once hydrated, they are susceptible to degradation by nucleases. If handled correctly, they should be stable for years
- Any DNA oligonucleotide can be degraded by microbial or fingertip nucleases
- Primers should be stored in deionized water or buffered solutions containing EDTA and kept frozen when not in use
- Make a concentrated stock solution in water or TE (10 mM Tris pH 8.0, 1 mM EDTA). A convenient stock could be 100 μM, stored at -20^oC
- Stock solutions should be distributed into several tubes for long term storage so that accidental contamination of a tube will not lead to loss of the entire synthesis

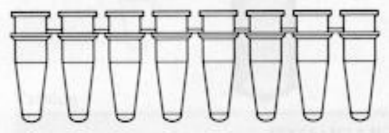
PCR reaction components

- Water
- Buffer
- DNA template
- Primers
- Nucleotides (dNTPs)
- Mg²⁺ ions
- DNA Polymerase

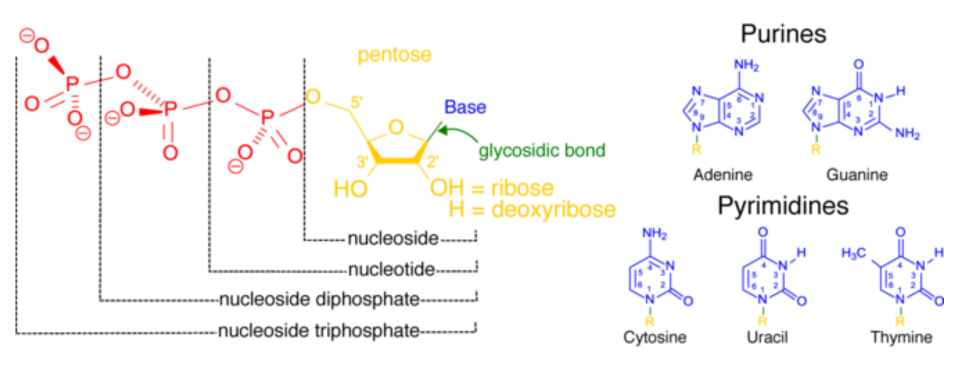


- Added to the growing chain
- Activated NTP's
- datp, dgtp, dctp, dttp
- Stored at 10mM, pH 7.0
- Add to 20-200 μM in assay





PCR reaction components



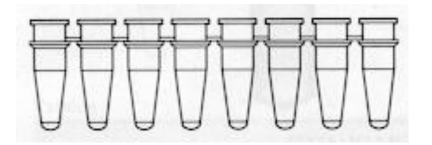
dNTP, deoxyribonucleotide triphosphate. Each dNTP is made up of a phosphate group, a deoxyribose sugar and a nitrogenous base. There are four different dNTPs and can be split into two groups: the purines (**A**,**G**) and the pyrimidines (**T**,**C**) that connect through hydrogen bridges.

This figure also shows the five common nitrogenous bases found in DNA and RNA on the right.

PCR reaction components



- Water
- Buffer
- DNA template
- Primers
- Nucleotides
- Mg²⁺ ions
- DNA Polymerase



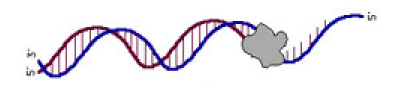
- Mg²⁺ ions
 - Essential co-factor of DNA polymerase
 - Insufficient concentration reduces yield
 - Stabilizes the DNA doublehelix
 - Too much: DNA extra stable, non-specific priming, band smearing
 - Used at 0.5 to 3.5 μM in the assay

PCR reaction components

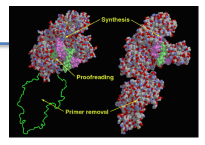
- Water
- Buffer
- DNA template
- Primers
- Nucleotides
- Mg²⁺ions
- DNA Polymerase

DNA Polymerase

- The enzyme that does the extension
- TAQ or similar
- Heat-stable
- Approx 1 U / rxn
 - 5' TACGCGGTAACGGTATGTTCGACCGTTTAGCTACCGA



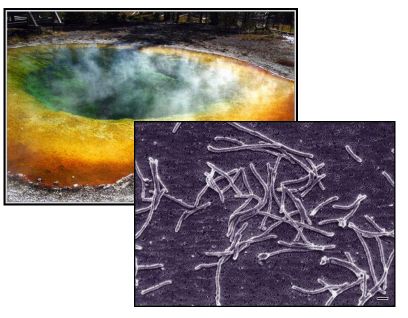




Thermostable DNA polymerase (taq DNA polymerase)

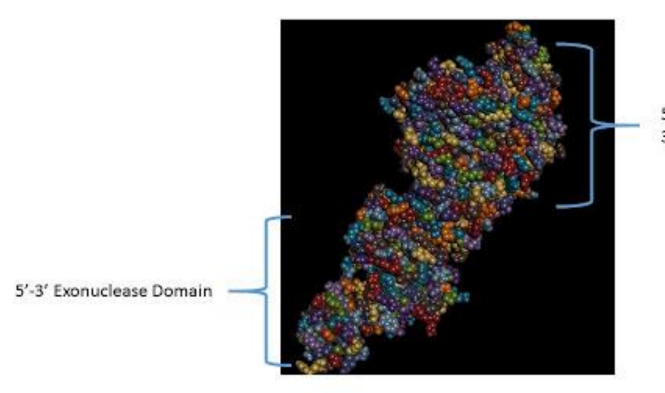
When studies of biological organisms in hot springs began in the 1960s, scientists thought that the life of thermopile bacteria could not be sustained in temperatures above about 55 °C.

Thomas D. Brock and Hudson Freeze (Indiana University) reported a new species of thermopile bacterium which they named *Thermus aquaticus*. The bacterium was first discovered in the Lower Geyser Basin of Yellowstone National Park, near the major geysers and has since been found in similar thermal habitats around the world.



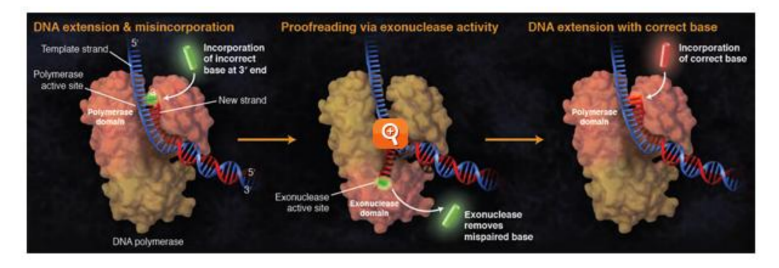
Taq DNA polimerase was isolated from *T. aquaticus* in 1969 by Brock & Freeze 1969

Structure of DNA Polymerase



5'-3' Polymerase activity domain and 3'-5' Proof reading activity domain

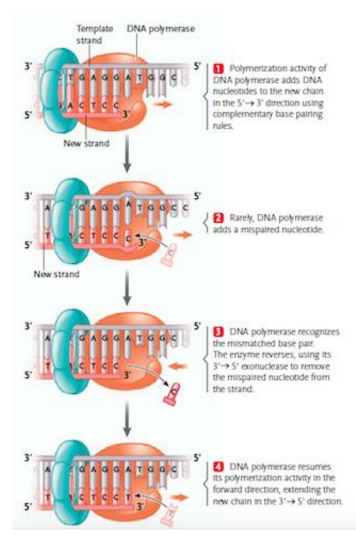
Proofreading



Extension proceeds along the template strand at the 3' end of the newly synthesized strand. When the polymerase recognizes an error, the mismatched base is transferred to the exonuclease active site and the base is excised. The extended strand returns to the polymerase domain, re-anneals to the template strand, and replication continues.

https://international.neb.com/tools-and-resources/feature-articles/polymerase-fidelity-what-is-it-and-what-does-it-mean-for-your-pcr

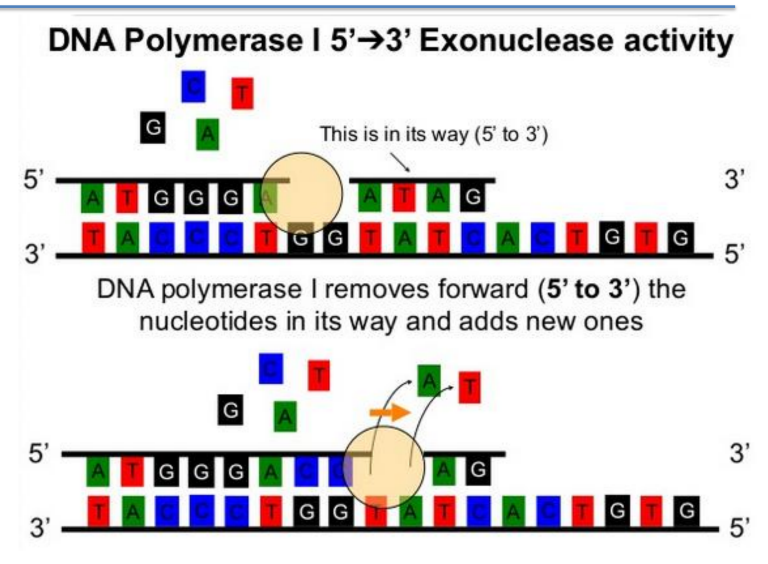
Proofreading



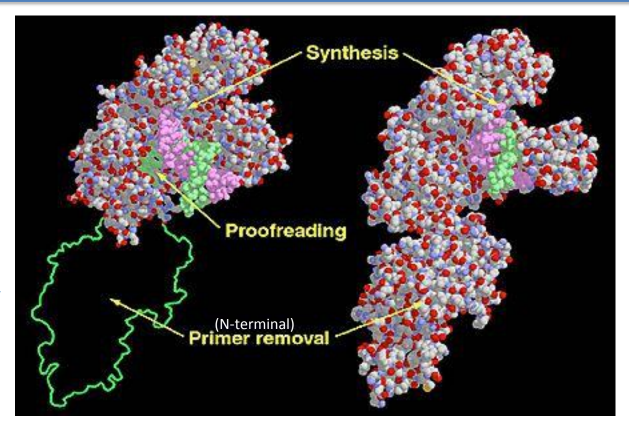
	KOD (%)	Phusion HF (%)	Pt <i>Taq</i> (%)	Expand HF (%)	FastStart HF (%)	Sequal Prep Long (%)
Overall error rate ^a	0.21	0.11	0.34	0.25	0.23	0.29
Insertions	0.10	0.07	0.14	0.11	0.11	0.11
Deletions	0.06	0.02	0.08	0.07	0.05	0.06
Substitutions	0.01	0.01	0.07	0.04	0.03	0.07
Dots or Dot ^b	0.04	0.01	0.05	0.04	0.04	0.05

https://www.thermofisher.com/pt/en/home/brands/thermo-scientific/molecular-biology/phusion.html

5' – 3' exonuclease activity



Klenow fragment



Small fragment (35 kDa) 5'-3' exonuclease activity

- Klenow fragment (68 kDa)
- 5'-3' polymerase activity
- 3'-5' exonuclease activity

- DNA polymerase I (103 kDa)
- 5'-3' polymerase activity
- 5'-3' exonuclease activity
- 3'-5' exonuclease activity

Example of a PCR reaction

Sterile Water	38.0	μl
10X PCR Buffer	5.0	μl
MgCl2 (50mM)	2.5	μl
dNTP's (10mM each)	1.0	μl
PrimerFWD (25 pmol/ul)	1.0	μl
PrimerREV	1.0	μl
DNA Polymerase	0.5	μl
DNA Template	1.0	μl
Total Volume	50.0	μl

Example of a PCR reaction

Sterile Water 10X PCR Buffer MgCl2 (50mM) dNTP's (10mM each)	•	Mixing Common Rea	igents Saves	Time – master	mix concept
PrimerFWD (25 pmol/ul PrimerREV DNA Polymerase DNA Template Total Volume	1) 1.0 μ1 1.0 μ1 0.5 μ1 1.0 μ1	Component Sterile Water 10X PCR Buffer		20X (μl) 760 100 50 20 20 20 20 10 	
		Total Volume	50.0	980	
		FREE	用日	Aliquot 49 ul	

0

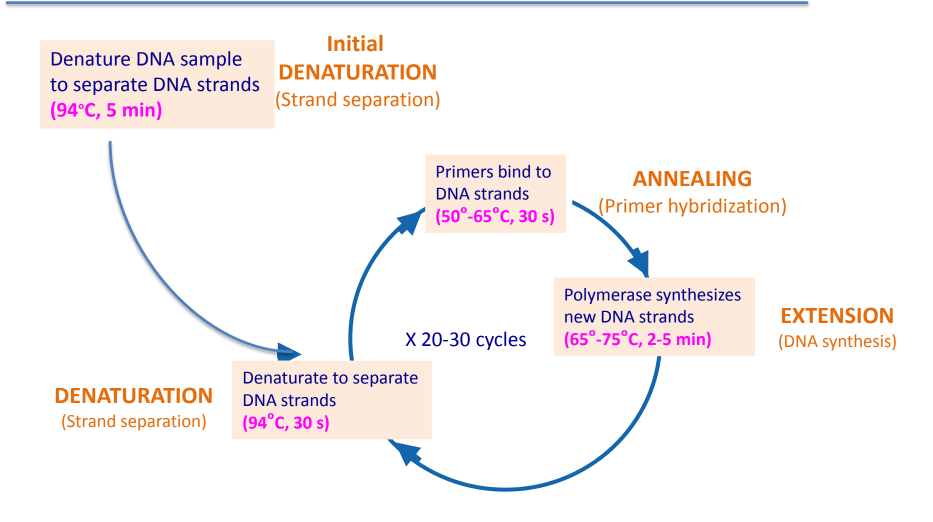
9

Thermal cycling conditions

Step	Time	Temperature	Comments
1. Denaturation	4 min	94°C	Denaturation of template and primer-dimers
2. Denaturation cycle	0.5 min	94°C	
3. Annealing cycle	0.5 min	55°C	Or 5°C below the lowest primer melting temperature
4. Extension cycle	1 min/kb	72°C	
5. Repeat steps 2–4			Repeat 30 times
6. Final extension	10 min	72°C	
7. End	Infinite	4°C	



Thermal cycling steps



Visualization of PCR products

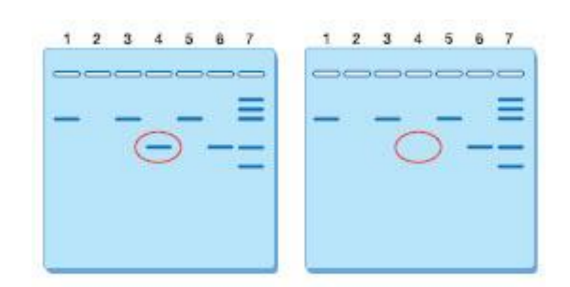


•After thermal cycling, tubes are taken out of the PCR machine.

•Contents of tubes are loaded onto an agarose gel.

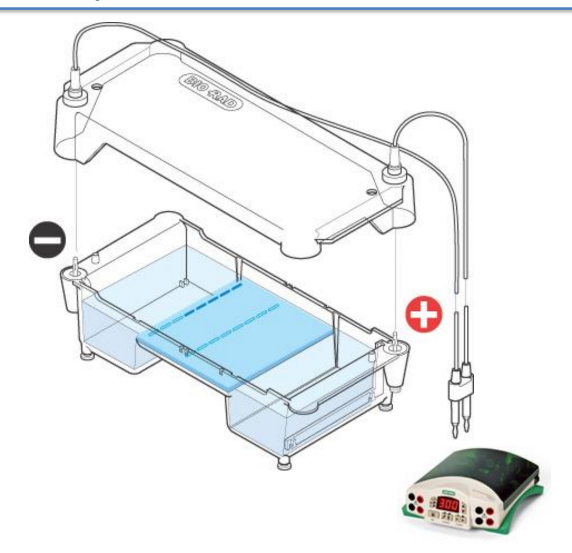
- •DNA is separated by size using an electric field.
- •DNA is then stained.

•PCR products are visible as different "bands".

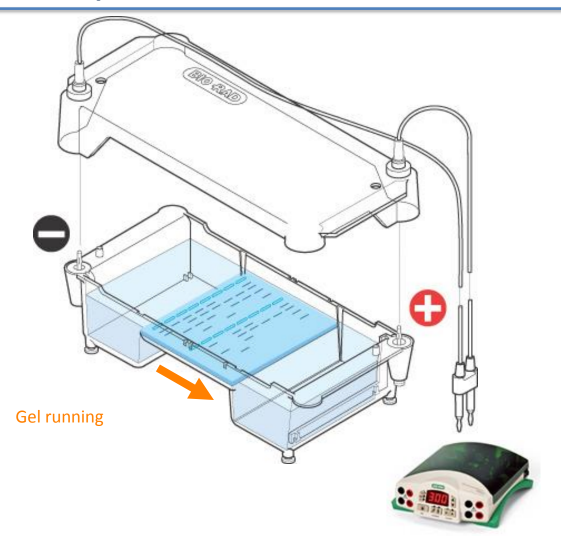




Visualization of PCR products

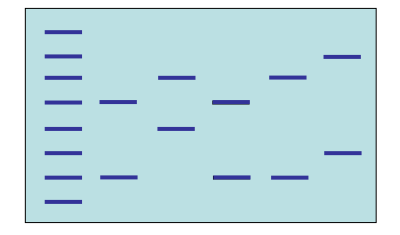


Visualization of PCR products



Visualization of PCR products

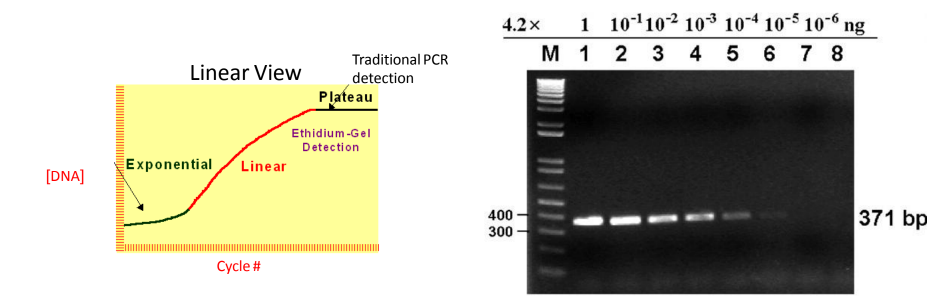
The final result of the traditional PCR procedure is a gel with a series of bands:



Bands can be compared against each other, and to known size-standards, to determine the presence or absence of a specific amplification product.

Visualization of PCR products

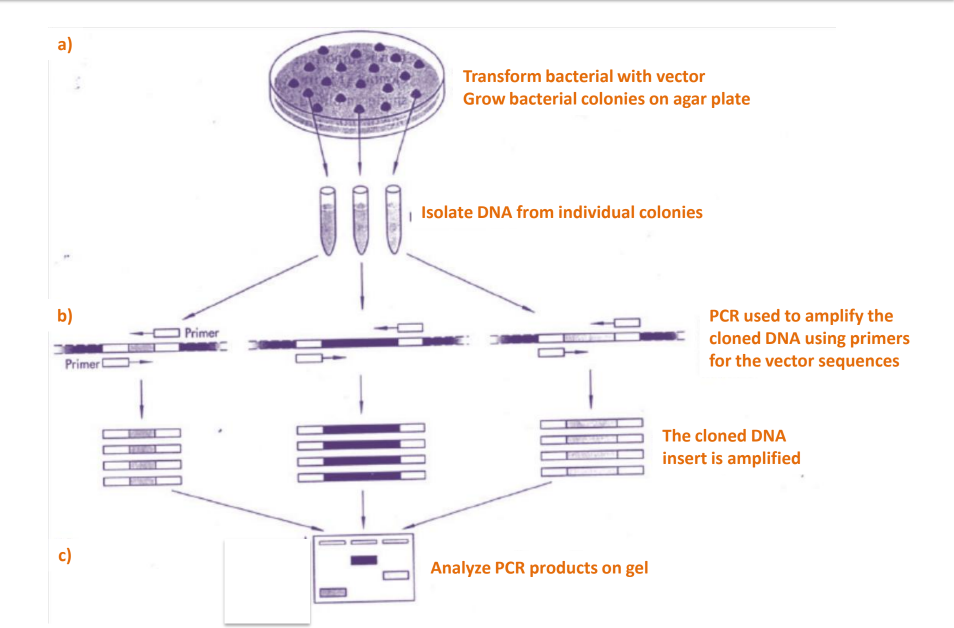
- End point detection
- Low resolution

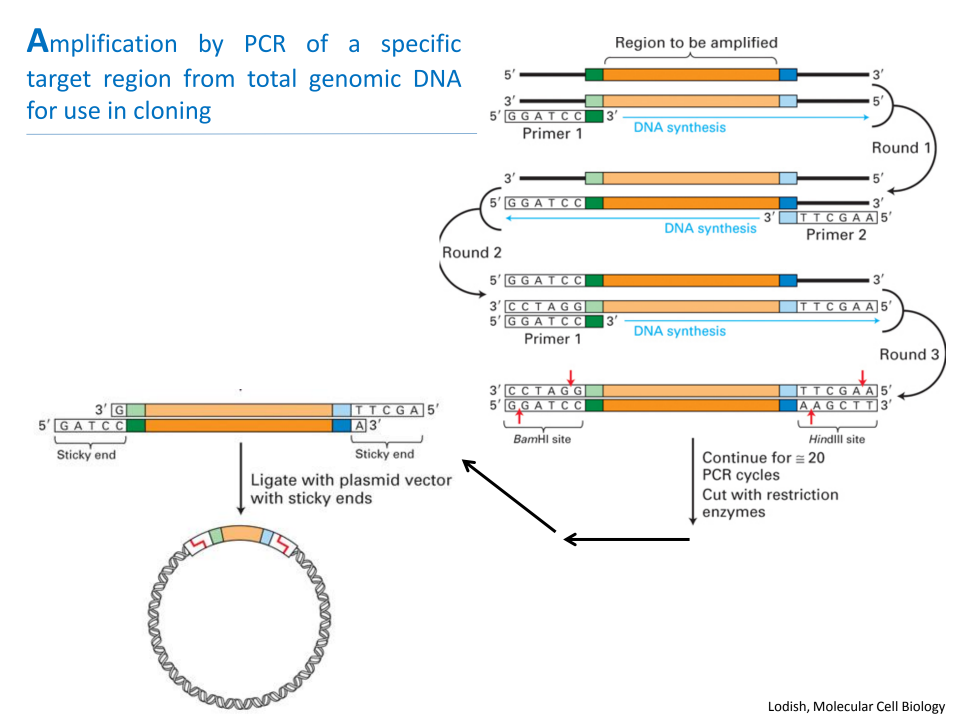


Applications

1. Diagnosis of pathogens	 a) PCR b) Nested PCR c) Quantitative PCR d) Multiplex PCR e) Differential on-line and real time PCR
2. Typing genetic markers	a) RFLPsb) AFLPsc) Short tandem repeat polymorphisms
3. DNA template for	a) Genomic mutation screening intron-specific primers flanking exonsb) RT-PCR cDNA used as templates for pairs of exon-specific primers to generate overlapping fragments
4. Detecting point mutations	a) Restriction site polymorphismsb) Allele specific amplification
5. cDNA cloning	a) DOP-PCR b) RACE
6. Genomic DNA cloning	New members of a DNA family a) DOP-PCR. Whole genome or subgenomic amplification b) DOP-PCR c) Linker-primed PCR
7. Genome walking	a) Inverse PCRb) Bubble linker (vectored) PCRc) IRE-PCR
8. DNA templates for DNA sequencing	a) ssDNA by asymmetric PCRb) dsDNA for direct sequencing or for cloning followed by sequencing
9. <i>In vitro</i> mutagenesis	a) 5' add-on mutagenesis to create a recombinant PCR productb) Mispaired primers to change a single predetermined nucleotide

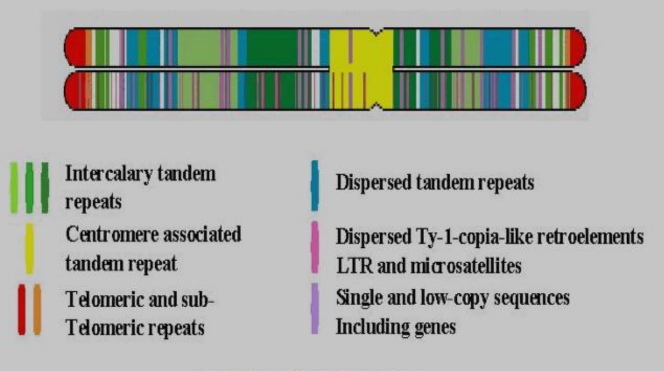
- Amplification of specific DNA fragments to clone, or to isolate a specific clone (gene discovery)
- Distinction between alleles
- Detection of polymorphisms, Molecular Typing, Taxonomy, Forensics, Molecular Diagnosis
- Phylogenetic studies
- Direct isolation of a specific segment of DNA of different origins (genomic, plasmidic etc)
- Preparation of probes
- Tagging of genes by insertion mutations, site-directed mutagenesis
- Cloned DNA molecules are sequenced rapidly by new methods based on PCR





-Tandem repeats - copies which lie adjacent to each other, either directly or inverted: Satellite DNA, Minisatellite, Microsatellite





A model of a plant chromosome.

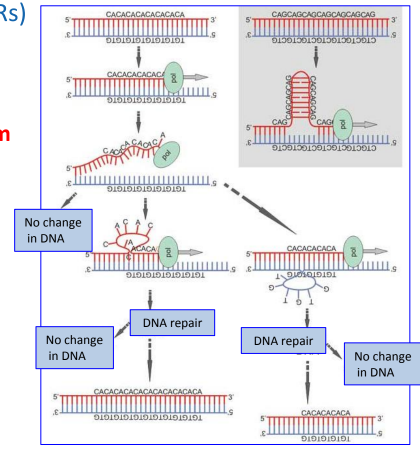
(1) Satellite DNA-typically in centromeres and heterochromatin (2)Minisatellite- repeat units of more than 10 base pairs, in many places in the genome, inc. centromeres (3) Microsatellite-repeat units of less than 10 base pairs; incl. telomeres

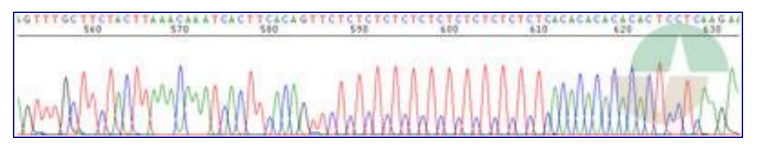
VNTR (Variable Number of Tandem Repeats) Minisatellites or Microsatellites (STRs or SSRs)

Polymorphism in microsatellites can arise from polymerase slippage

In plants, the most common plant repeat motif is AA/TT, followed by AT/TA and CT/GA.

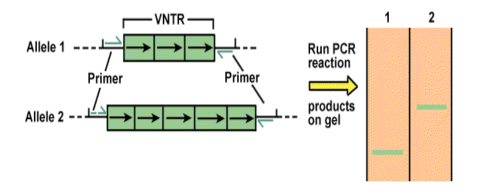
In mammals the GT/CA motif, being the most abundant dinucleotide repeat, was found to be considerably less frequent in plants.





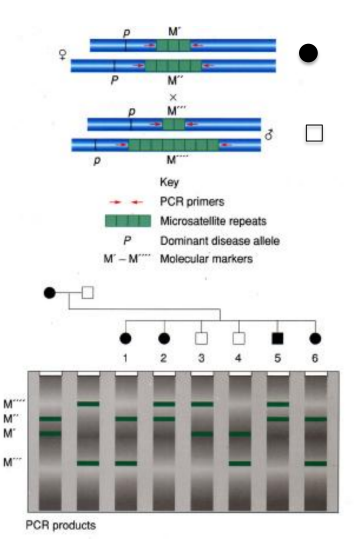
VNTR (Variable Number of Tandem Repeats) Minisatellites or Microsatellites (STRs or SSRs)

Principle of polymorphism detection (ex. VNTR)

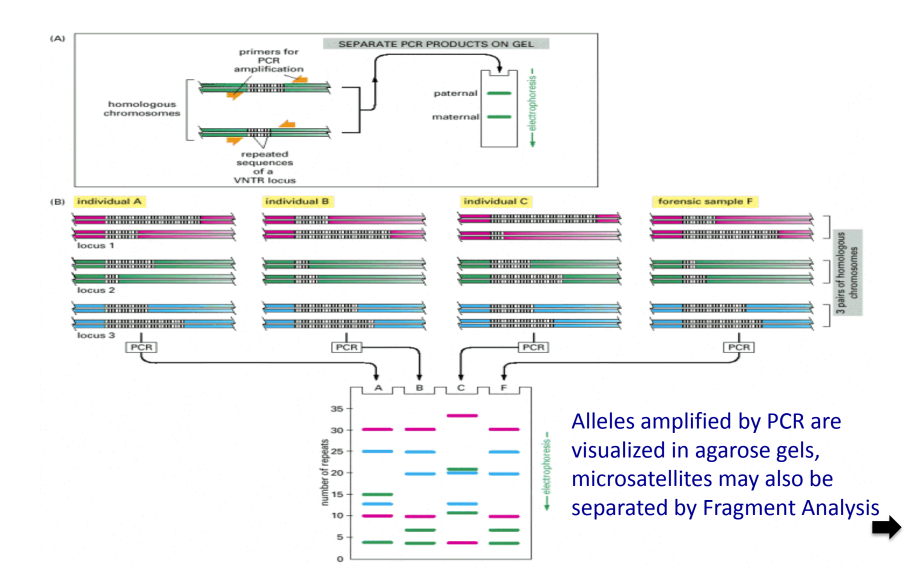


Alleles amplified by PCR are separated in agarose gels

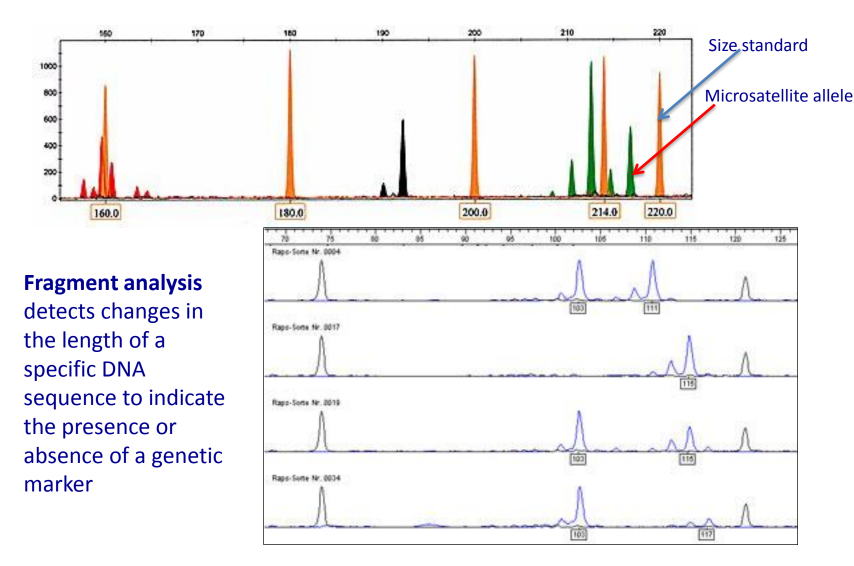
Polymorphism detection Case study



VNTR detection by PCR using multiple primers Multiplex reaction

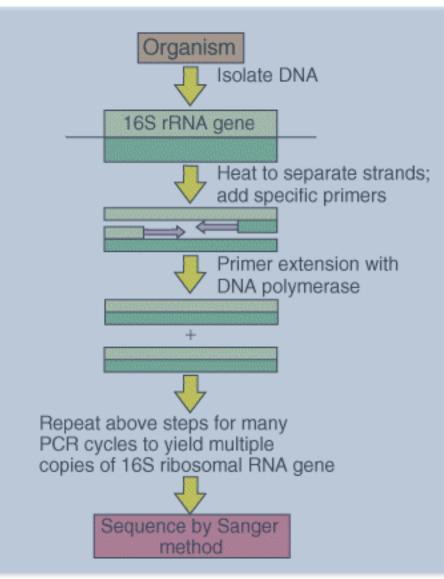


Microsatellites detected by capilary electrophoresis - Fragment Analysis



Phylogenetic studies

Ex. Amplification of 16S rRNA



Other DNA regions frequently used in phylogenetic studies: rRNA 16S **18S** 25S/26S ITS, internal transcribed spacer, between 16S and 23S in bacteria or between 18S and 25S/26S in plants trnL matK *rbcL* Cox1 •••

Brock, Molecular Biology of Microorganisms