Gene Technology & & DNA Sequencing

Lecture-3

For all computational purposes, DNA is represented as a string of 4-letter alphabets - A, T, C, G:

attgctacgttacatcgctgca

How do we get this string representation from a dynamic double-stranded molecule?

DNA Sequencing - determine the precise order of the nucleotides in a sample of DNA

To carry out this task we need to be able to chop the DNA, store it, make copies of it.

To sequence a gene, we need to

- Identify the region of interest
- Isolate it from the organism DNA fragmentation
- move it to another easily manageable organism such as a bacterium for obtaining multiple copies — cloning

Such manipulations are conducted by a toolkit of enzymes:

Restriction endonucleases - used as molecular scissors

DNA ligase - to bond pieces of **DNA** together

- a variety of additional enzymes that modify DNA are used to facilitate the process.

Restriction endonucleases are enzymes that make site-specific cuts in the DNA — chemical scissors

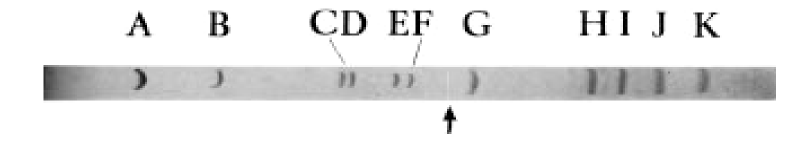
Ability to cut DNA into discrete fragments allows to understand

- how genetic material of an organism is organized
- how expression of genetic information is controlled
- how alteration of genetic information can give rise to genetically inherited disorders, etc.
- in bulk production of pharmaceutically important proteins

First restriction enzyme was isolated from H. influenzae in 1970 by Daniel Nathans and Kathleen Danna

- awarded the Nobel Prize for Medicine in 1978

Radioautogram of ¹⁴C-labeled SV40 DNA cleaved with endonuclease R



SV40 DNA (a tumor virus) - after cutting, or "digesting" it with *H*. *influenzae* restriction enzyme, analyzed the pieces using a polyacrylamide gel electrophoresis.

- 11 distinct DNA bands were visible in the gel, indicating that the enzyme always cut SV40 DNA resulting in the same 11 pieces

Background

How were these restriction endonucleases identified?

Bacteria are under constant attack by viruses, e.g., bacteriophages

To protect themselves, bacteria have developed a method to chop up any foreign DNA, by an enzyme, called endonuclease

- it circulates in the bacterial cytoplasm, waiting for any attacking virus.
- also called restriction enzymes because they restrict the infection of bacteriophages.

Why do the restriction enzymes not chew up the genomic DNA of their host?

Background

A bacterium that makes a particular restriction endonuclease, also synthesizes a companion DNA methyltransferase,

- which methylates the DNA target sequence for that restriction enzyme, thereby protecting it from cleavage.

DNA from an attacking bacteriophage will not have these protective methyl groups and will be destroyed.

Methyl groups (attached to the cytosine in dinucleotide CG) block the binding of restriction enzymes, but do not block the normal reading and replication of the genomic information stored in the host DNA.

DNA Fragmentation

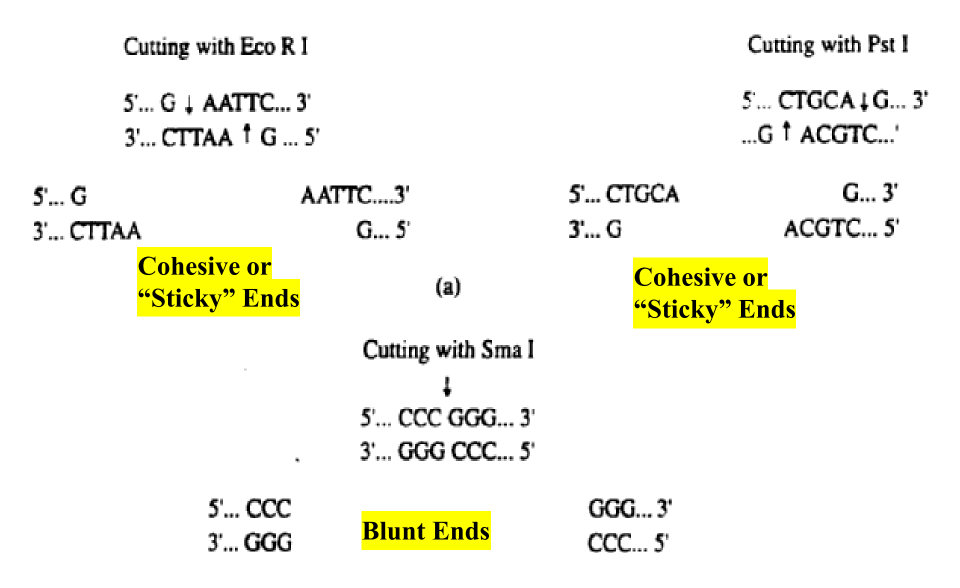
Different endonucleases present in different bacteria recognize different nucleotide sequences

Naming of restriction enzymes - after their host of origin, e.g.,

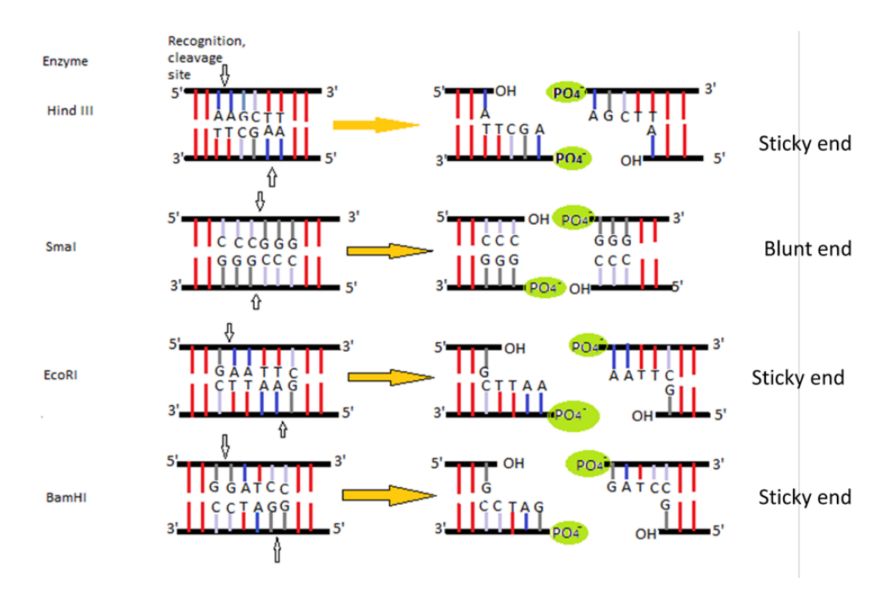
- EcoRI Escherichia coli
- Hind II & Hind III Haemophilus influenzae
- XhoI Xanthomonas holcicola

When cut with a restriction enzyme (RE), the ends of the cut DNA fragment can be cohesive or blunt-ended depending on the enzyme.

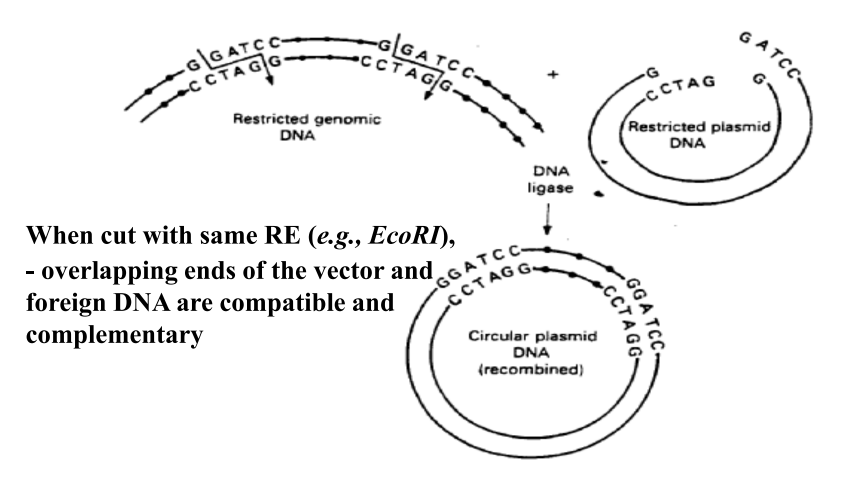
Generation of Cohesive & Blunt-ended Fragments



Cleavage patterns of *HindIII*, *SmaI*, *EcoRI* and *BamHI*



Restriction enzyme digestion of genomic DNA and plasmid vector DNA



A plasmid is a small, circular, double-stranded DNA molecule that is distinct from a cell's chromosomal DNA.

Features of Restriction Enzymes

- Length of recognition sequence dictates how frequently the enzyme will cut a DNA sequence
 Frequency of recognition sites of length, 4, 6, or 8?
- Different REs can have the same recognition site and are called isoschizomers, e.g., SacI & SstI: GAGCTC
- Restriction recognitions sites can be unambiguous, e.g., *Hinf* I recognition site: GANTC it's frequency of occurrence?
- Most recognition sequences are palindromes they read the same on forward and backward strands:

GAGCTC CTAGAG

Can we use the property of palindrome sequence to identify restriction recognition sites?

Applications of Restriction Enzymes

- To prepare a physical map of the genome
- In genetic engineering to assemble customized genomes; create designer bacteria that make insulin, or growth hormones, or add genes for disease resistance to agricultural plants, etc.
- in DNA sequencing

Restriction enzymes and related technology laid the groundwork for the Human Genome Project.

Restriction Map

Restriction map is a description of restriction endonuclease cleavage sites within a piece of DNA

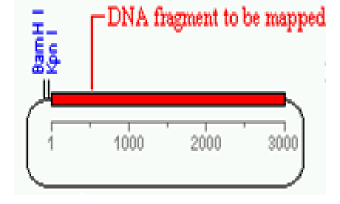
- generating such a map is the first step in characterizing an unknown DNA

Multiple Complete Digest Mapping – creates a map by digesting DNA with multiple REs

- each recognizing a different specific short DNA sequence and producing a separate fingerprint for each clone

Ex: Consider a plasmid that contains a 3000 bp fragment of unknown DNA & unique recognition sites for enzymes Kpn I

& BamH I.



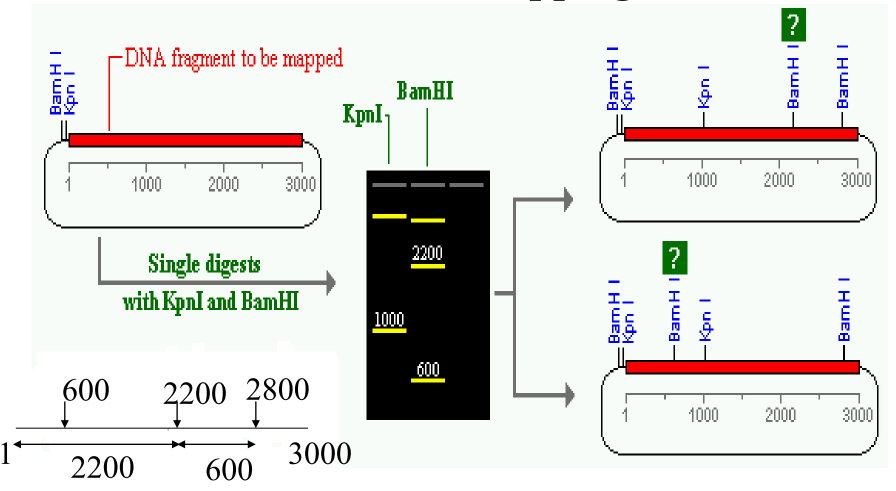
Consider first separate digestions with Kpn I & BamH I:

Kpn I yields 2 fragments: 1000bp & "big"

BamH I yields 3 fragments: 600, 2200 & "big"

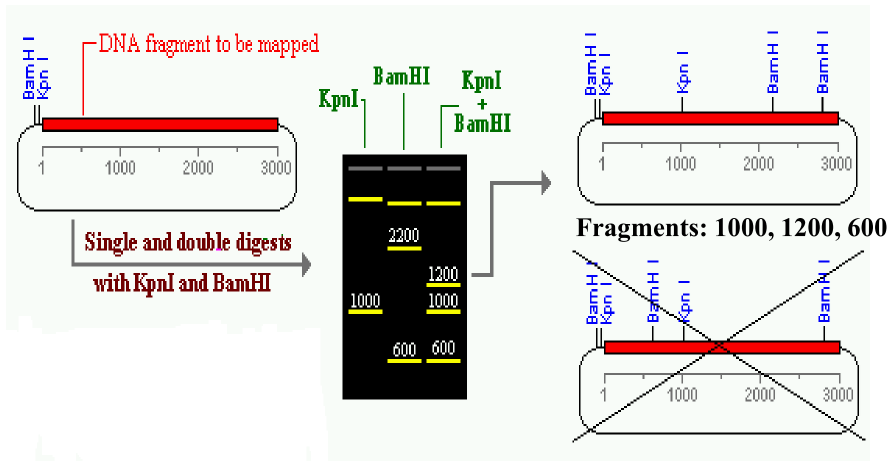
big – part of unknown DNA sequence + vector

⇒ one Kpn I site & two BamH I sites are present in the unknown DNA sequence



One BamH I site is at 2800 bp. Trick to determine the location of 2nd BamH I site is to digest the plasmid with Kpn I & BamH I together

Double digest yields fragments of 600, 1000 & 1200 bp (plus the "big" fragment).



Fragments: 600, 400, 1800

If the above process is conducted with a larger set of enzymes, a much more complete map would result

single digests - are used to determine which fragments are in the unknown DNA, and

multiple digests - to order and orient the fragments correctly.

Can we computationally generate a restriction map of DNA sequence?

Using a Computer to Generate Restriction Maps

For a known DNA sequence, following programs will build a map for various RE recognition sites:

- Mapper available as part of Molecular Toolkit http://arbl.cvmbs.colostate.edu/molkit/mapper/
- Webcutter
 http://www.firstmarket.com/cutter/cut2.html
- RebSite as part of the REBASE Tools http://tools.neb.com/REBsites/index.php3

REBASE - The Restriction Enzyme dataBASE



REBsites

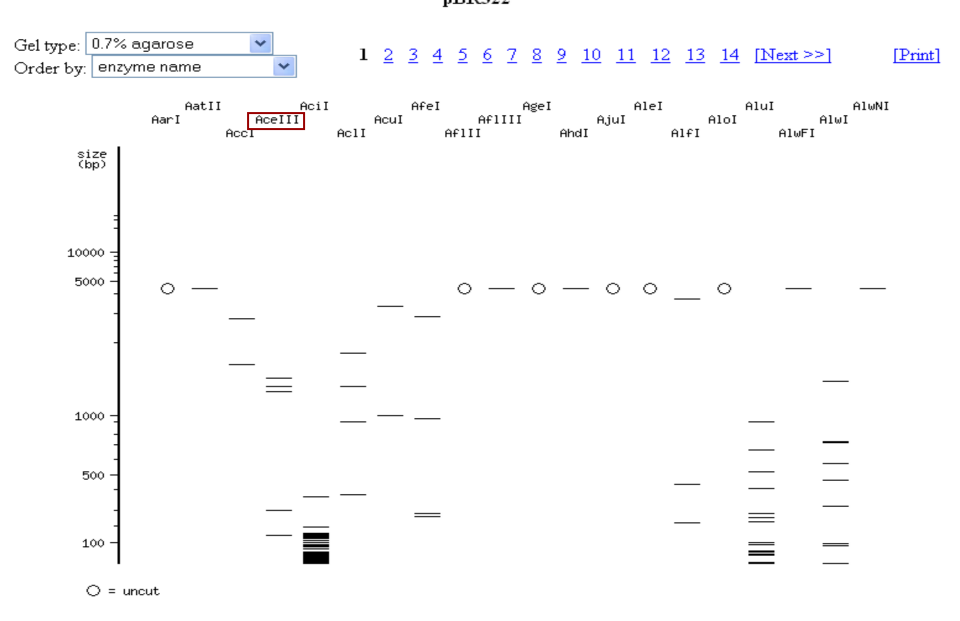


This tool will take a DNA sequence and digest it with one example of each of the known Type 2 restriction enzyme specificities.

The maximum size of the input file is 2 MByte, and the maximum sequence length is 200 KBases.

					1
Local sequence file:		Browse		Standard	
GenBank number:		(Browse GenBank)		sequences:	
Name of sequence:		(optional)		pBR322	
or Paste in your DNA sequenc	e: (plain or FAS	STA format)		Ad2	
				Submit	
	3 T :				
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	Infallt cited:	All specificities Defined oligonucleotide sequence	a·		
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	CI	ear the table below			rototypes
	Name	Oligonucleotide sequence		DASE P	lototypes

REBsites pBR322



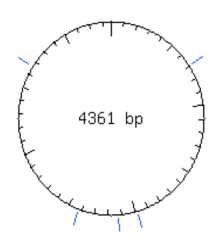
zyme name for a list of fragments. gel was generated by interpolating experimental data. See <u>details</u>. Print

Fragment list

Close

 $p\mathbf{BR322}$ digested with AceIII

[Sites with flanks]



#	Coordinates	Length (bp)
1	3666-697	1393
2	698-1984	1287
3	2426-3665	1240
4	2126-2425	300
5	1985-2125	141



REBASE Enzymes 12/18/2006



Type II Restriction Enzymes

Enzymes	Recognition Sequence	Isoschizomers	<u>Suppliers</u>
<u>AaaI</u>	claccc	<u>yes</u>	-
<u>AacI</u>	GGATCC	<u>yes</u>	-
<u>AaeI</u>	GGATCC	yes	-
<u>AagI</u>	AT ∮ CGAT	yes	-
<u>AamI</u>	-	-	-
<u>AaqI</u>	GTGCAC	<u>yes</u>	-
<u>AarI</u>	CACCTGC (4/8)	-	У
<u>AasI</u>	GACNNNN I NNGTC	<u>yes</u>	У
<u>AatI</u>	AGG¶CCT	yes	У
<u>Aat∏</u>	GACGT ∜ C	<u>yes</u>	У
<u>AauI</u>	T 4 GTACA	<u>yes</u>	-
<u>AbaI</u>	T 4 GATCA	<u>yes</u>	-
<u>AbeI</u>	CCTCAGC (-5/-2)	<u>yes</u>	-
AboORF2079P	AGGCCT	<u>yes</u>	-
<u>AbrI</u>	C#TCGAG	<u>yes</u>	-
<u>AcaI</u>	TTCGAA	<u>yes</u>	-
<u>AcaII</u>	GGATCC	yes	-
<u>AcaIII</u>	TGCGCA	yes	-
<u>AcaTV</u>	GGCC	yes	-
AccI	GT 4 MKAC	yes	У
<u>AccII</u>	cg l cg	yes	У

Assignment

- Write a program to generate a restriction map for a specific RE and compare your results with Mapper.
- Write a program to identify all possible restriction recognition sites in a given DNA sequence.
- Write to program to obtain the reverse strand (forward strand of the DNA sequence is given).

Cloning

What is cloning?

The process of cloning involves the production of multiple copies of a DNA fragment of interest by amplification *in vivo*

- depends upon the ability of vectors to continue their life cycles in bacterial or yeast cells in spite of having foreign DNA inserted into them.

Cloning vector - a DNA molecule that carries foreign DNA into a host cell, replicates inside a bacterial (or yeast) cell and produces many copies of itself and the foreign DNA.

The other option is to make copies just before sequencing in vitro, using PCR.

Types of Vectors

Vector	Insert size (kb)	
Plasmids	<10 kb	
Bacteriophage	9 - 20 kb	
Cosmids	33 - 47 kb	
Bacterial artificial	75 - 125 kb	
chromosomes (BACs)		
Yeast artificial	100-1000 kb	
chromosomes (YACS)		

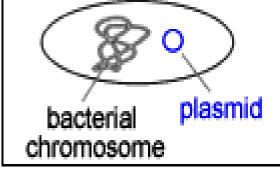
Types of Vectors

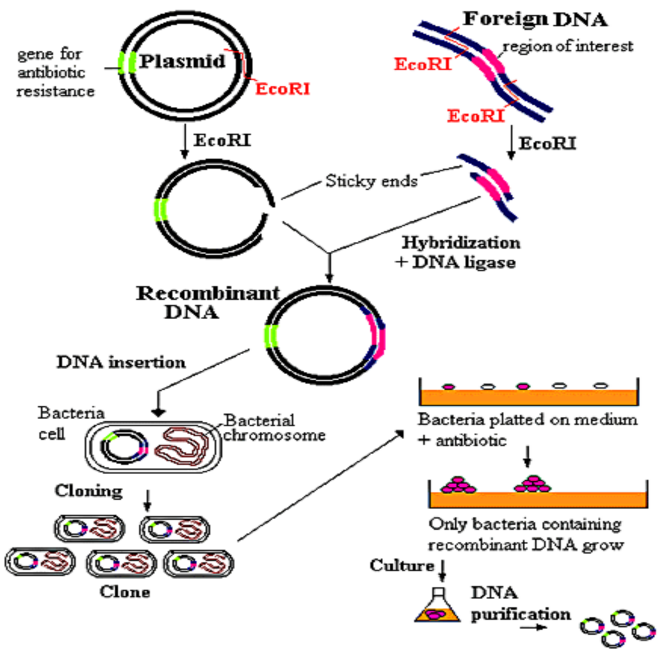
Plasmids - an extra-chromosomal double-stranded circular DNA molecules that replicates autonomously inside the bacterial cell

Plasmids are important as one can:

- (i) isolate them in large quantities,
- (ii) cut & splice them, add DNA of choice,
- (iii) put them back into bacteria, where they replicate along with the bacteria's own DNA,
- (iv) isolate them again to get billions of copies of inserted DNA

<u>Limitation</u>: size of DNA that can be introduced into the cell by transformation (~2 - 10kb)



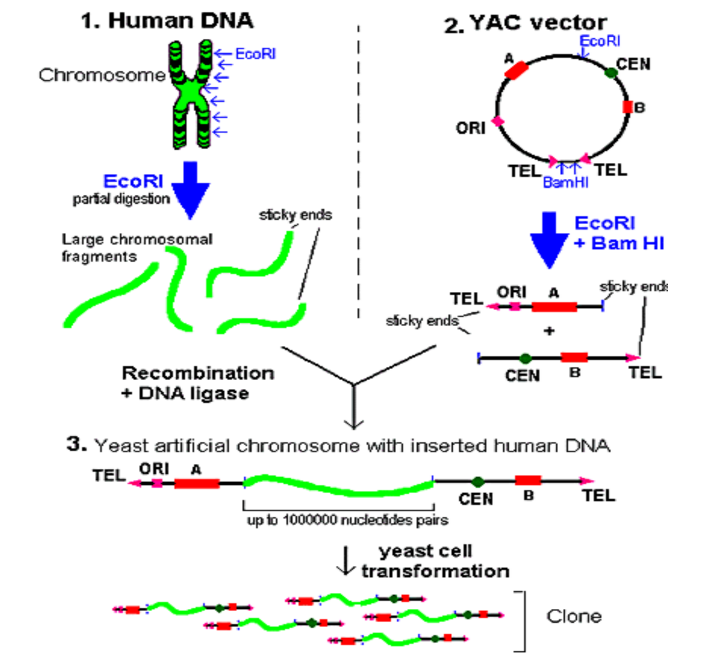


Cloning into a plasmid

- YAC a functional self-replicating yeast artificial chromosome. It includes three specific DNA sequences that enables it to propagate from one cell to its offspring:
- TEL: The telomere which is located at each chromosome end, protects the linear DNA from degradation
- CEN: The centromere which is the attachment site for mitotic spindle fibers, "pulls" one copy of each duplicated chromosome into each new daughter cell.
- ORI: Replication origin sequences, specific DNA sequences that allow the DNA replication machinery to assemble on the DNA to carry out the replication process

It also contains few other specific sequences like:

- A and B: selectable markers that allow easy isolation of yeast cells that have taken up the artificial chromosome.
- Recognition site for two REs: EcoRI & BamHI



Cloning into a Yeast Artificial Chromosome (YAC

Why is it important to be able to clone large sequences?

To map the entire human genome (3x10⁹ bps) would require more than 1000,000 plasmid clones (~10Kb limit).

In principle, the human genome could be represented in about 10,000 YAC clones (~1Mb limit)

DNA Sequencing

DNA Sequencing - determine the precise sequence of nucleotides in a sample of DNA – the order of A, T, G, C

Various types of sequencing:

- Sequencing a region of interest, e.g., gene.
- Whole Genome/Exome Sequencing
- cDNA Sequencing sequencing cDNA libraries of the expressed genes
- High-throughput sequencing next-generation, 3rd & 4th generation sequencing whole Genome/Exome/targeted
- Metagenome sequencing sequencing of environmental samples
- depending on the nature of analysis, type of DNA sample and type of sequencer is used

Sequencing a Region of Interest

First requirement in sequencing a region of DNA is

- to have enough starting template for sequencing.

This is achieved by PCR - Polymerase Chain Reaction

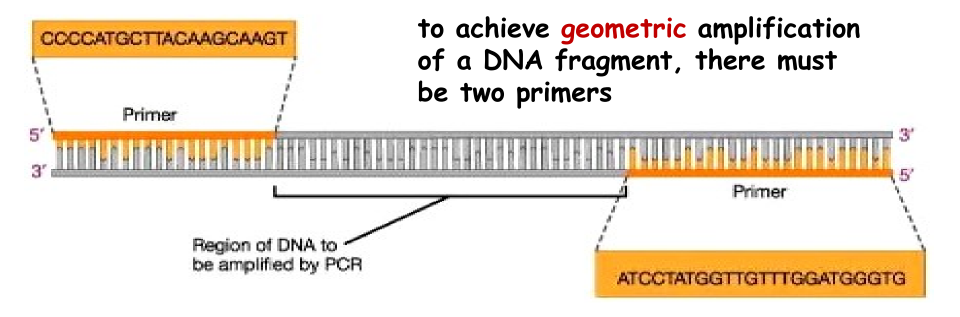
- carried out in an automated cycler for 30 - 40 cycles.

Essential requirements for a PCR:

- a mixture of 4 deoxy-nucleotides in ample quantities dATP, dGTP, dCTP, dTTP
- Taq DNA polymerase
- Primers ?
- Genomic DNA of interest

What is the advantage of using PCR over traditional gene cloning?

Region of DNA to be amplified by PCR



Primers - short single-stranded oligonucleotides which anneal to the DNA template and serve as a starting point for DNA synthesis

Why are primers required?

The Cycling Reactions

Step-1: Denaturation at 94°C

- opens up double stranded DNA, all enzymatic reactions stop.

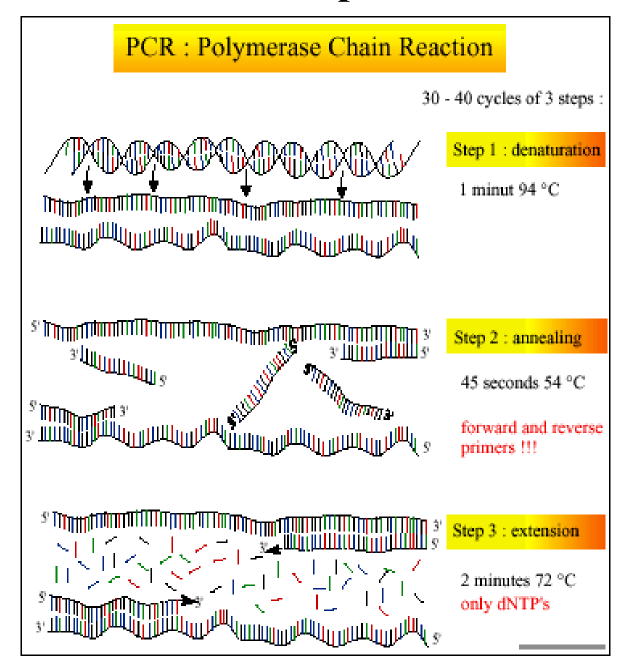
Step-2: Annealing at 54°C

- Primers jiggling around because of Brownian motion, binds to single stranded template once an exact match is found; the polymerase then attaches and start copying the template.

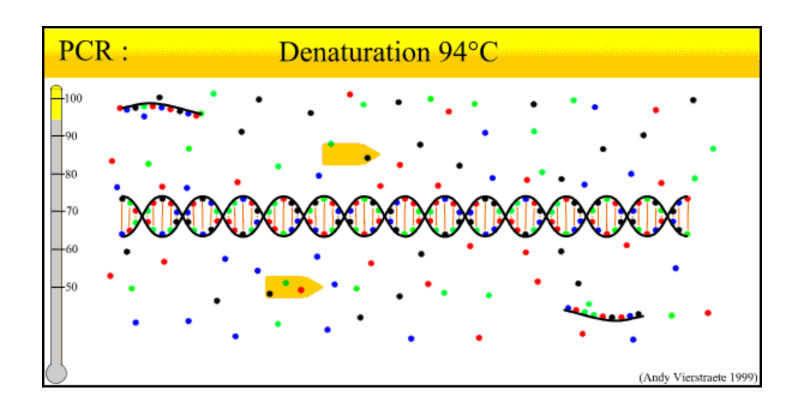
Step-3: Extension at 72°C

- ideal working temperature for the polymerase. Bases complementary to the template are coupled to the primer on 3' side (reading the template from 3' to 5' side)

Different Steps in PCR



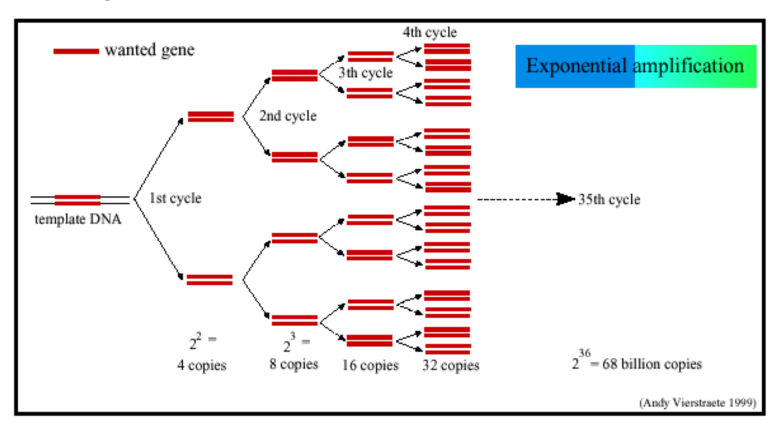
Different Steps in PCR



Exponential amplification of region of interest

Both strands are copied during PCR

- leading to an exponential increase of the number of copies of the region of interest.



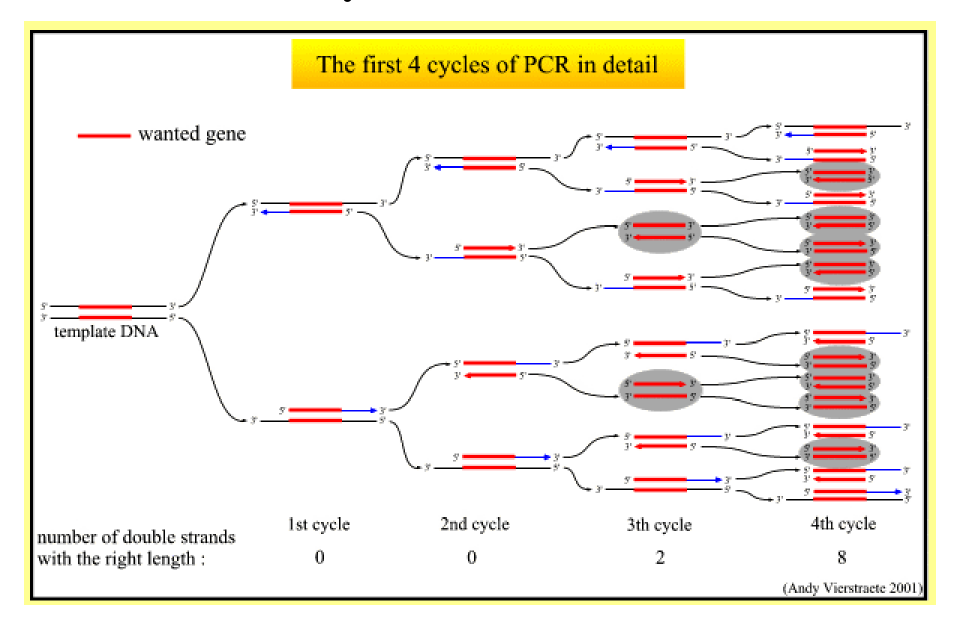
Verification of PCR Product

Is the template copied during PCR and is it the right size?

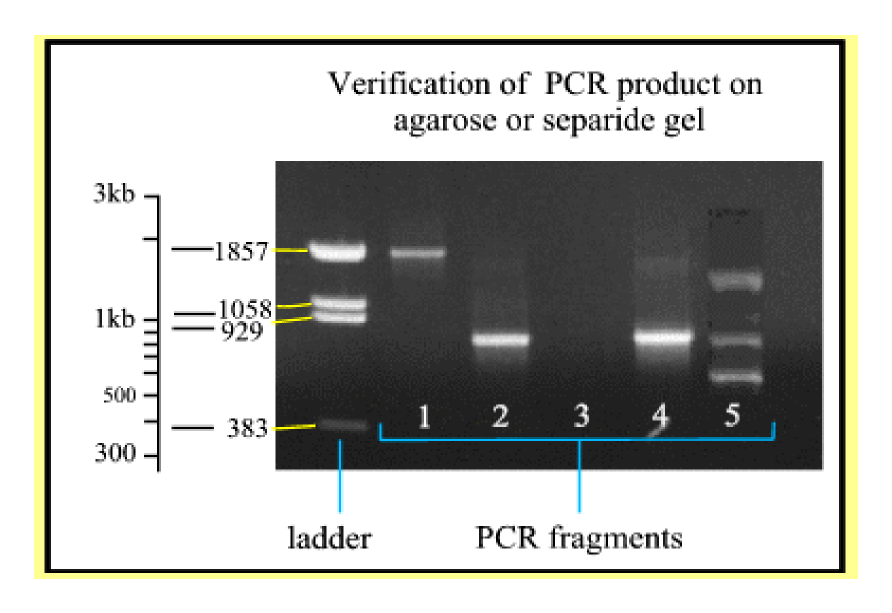
Before the PCR product is used in further applications, it has to be checked if:

- 1. A product is formed
- 2. The product is of the right size
- 3. Only one band is formed

First 4 cycles of a PCR reaction



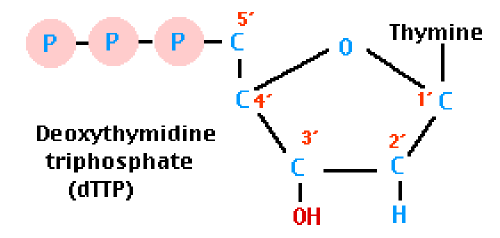
Verification of the PCR product



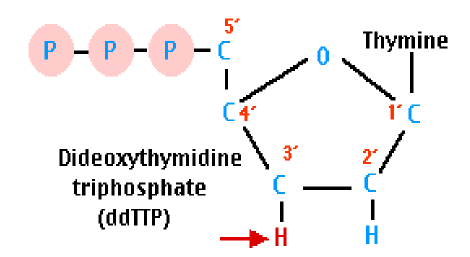
For sequencing, we don't start from gDNA (like in PCR) but mostly from PCR fragments or cloned genes.

Amplified PCR product is supplied with

- a mixture of all four <u>normal</u> (deoxy) nucleotides in ample quantities
 - dATP
 - dGTP
 - dCTP
 - dTTP
- Taq DNA polymerase



- a mixture of all four <u>dideoxynucleotides</u>, each present in limiting quantities and each labeled with a "tag" that fluoresces a different color:
 - ddATP
 - ddGTP
 - ddCTP
 - ddTTP

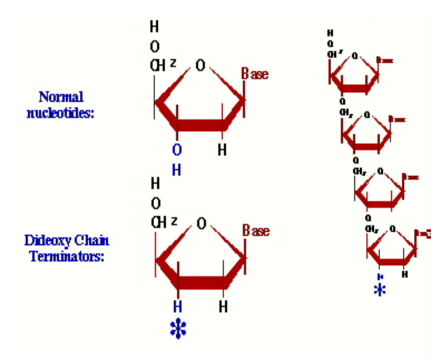


This method of DNA sequencing is called dideoxy method, or chain termination method, or Sanger's method.

Dideoxy method: DNA is synthesized from 4 deoxynucleotide triphosphates.

Each new nucleotide is added to 3'-OH group of the last nucleotide added.

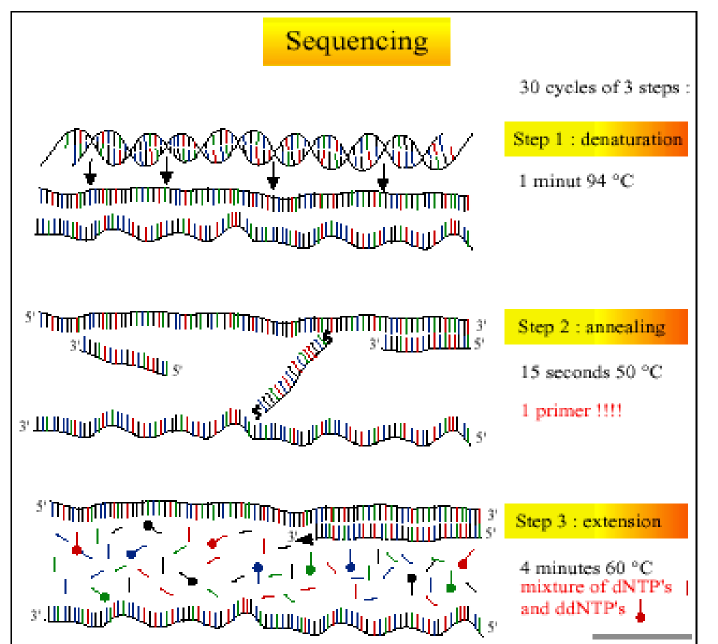
When a dideoxynucleotide, ddNTP, is added to the growing DNA strand, chain elongation stops because there is no 3'-OH for the next nucleotide to be attached to.



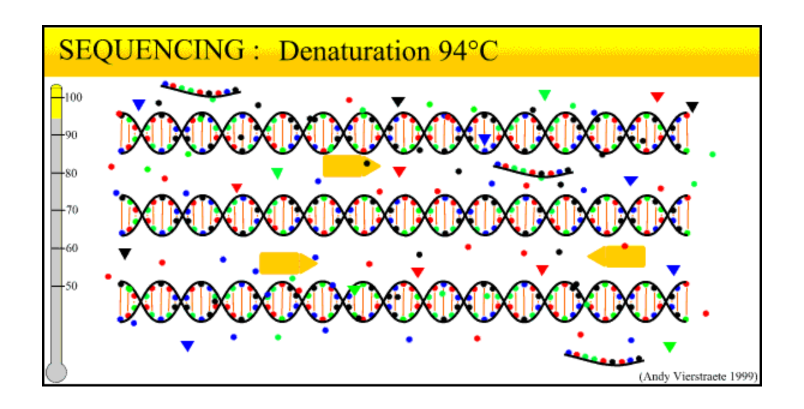
Steps in PCR Sequencing

- I The sequencing reaction
 - Denaturation at 94°C
 - Annealing at 50°C
 - Extension at 60°C ← instead of 72°C
- II Separation of the fragments
- III Detection on an automated sequencer
- IV Assembling the sequenced parts

Different steps in Sequencing

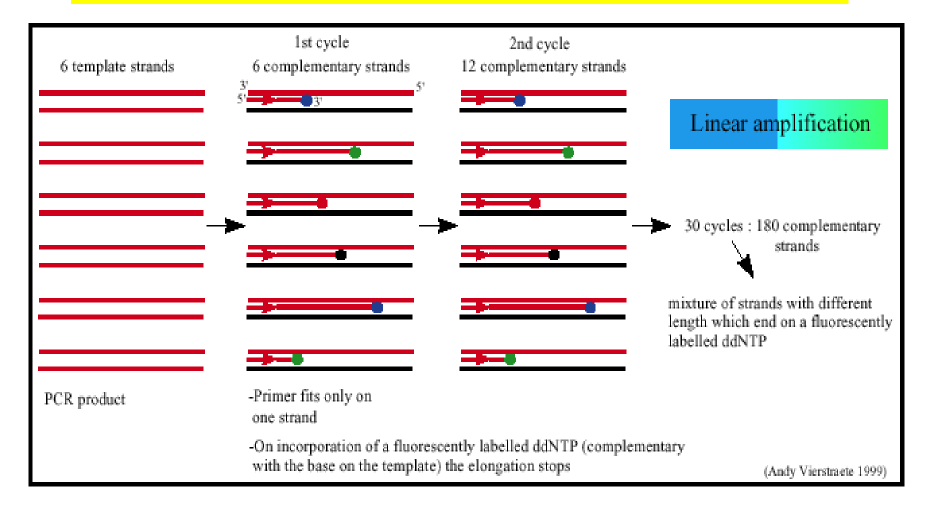


Different steps in Sequencing



Since only one primer is used, only <u>one strand</u> is copied during sequencing – results in <u>linear increase</u> of the No. of copies.

 \Rightarrow large amount of DNA in the starting mixture is required.

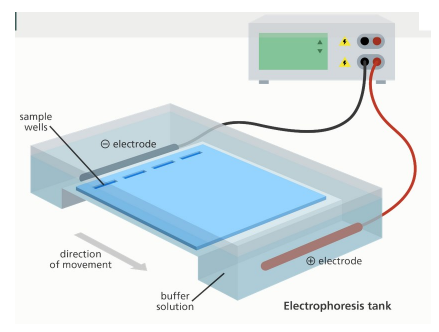


II Separation of the molecules:

The mixture of strands of different lengths with fluorescently labeled ddNTP at one end are loaded on an acrylamide gel for separation

- gel electrophoresis.

smallest strand travels the fastest

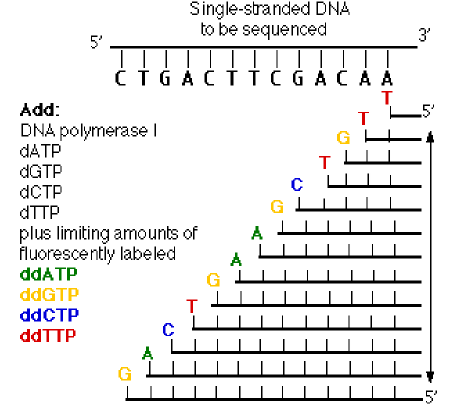


 strands of different length migrate at different rates and thus are separated based on their size

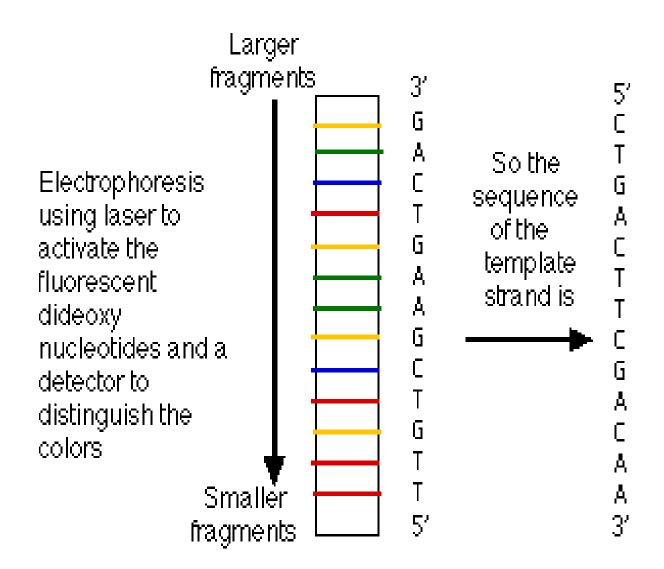
Separation of molecules with electrophoresis

Very good resolution

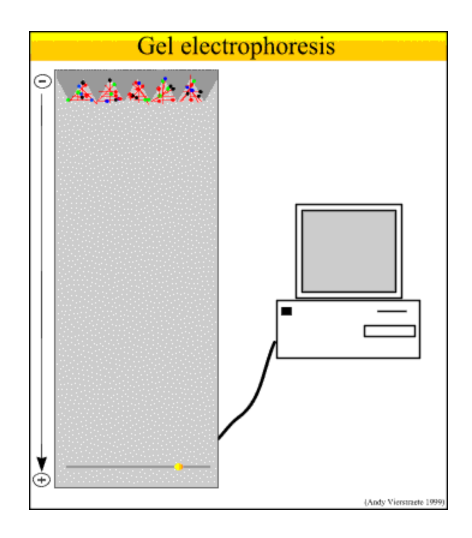
Each ddNTP fluoresces a different color when illuminated by a laser beam and an automatic scanner provides a printout of the sequence.



Separation of Molecules with Electrophoresis

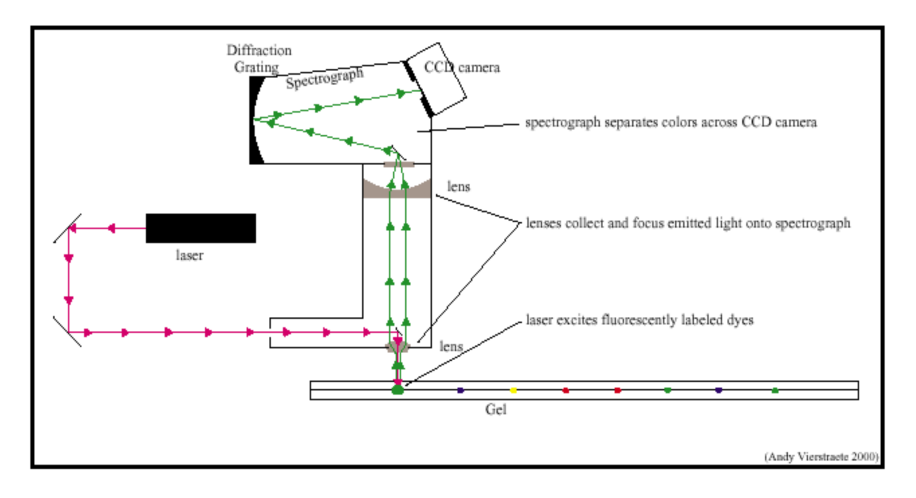


Separation of the Molecules with Electrophoresis

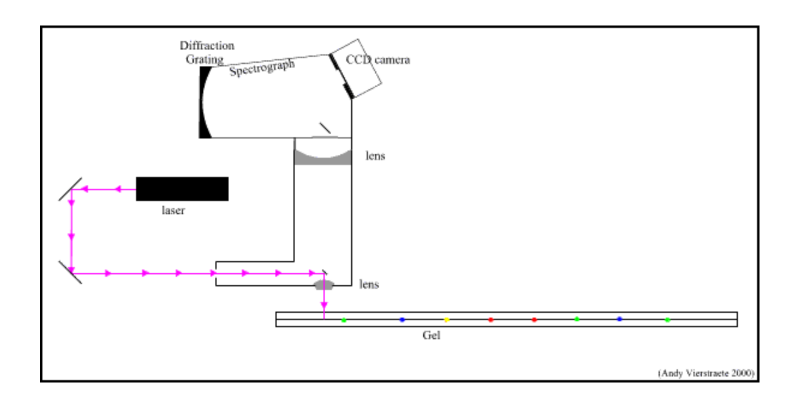


III Detection on an automated sequencer:

Fluorescently labeled fragments that migrate through the gel pass a laser beam at the bottom of the gel.



Scanning & Detection System on a Sequencer



Plot of the colors detected in a 'lane' of the gel (one sample), scanned from smallest fragments to largest - chromatogram.

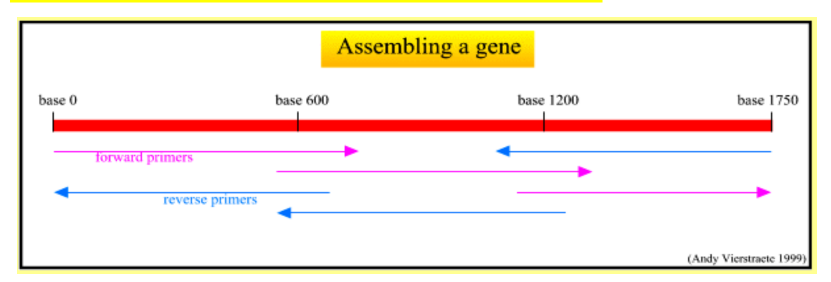
The computer interprets the colors by printing the nucleotide sequence across the top of the plot and also provides a quality score.

IV Assembling the sequenced parts of a gene:

For publication, a gene sequence has to be confirmed in both directions using forward & reverse primers

Since it is only possible to sequence ~ 700-800 bases in one run, a gene of, say, 1800 bases, is sequenced with internal primers.

- the sequenced fragments are assembled using a computer program to obtain complete gene sequence.



By Sanger's method, we can sequence a fragment of DNA ~ 1000bp long.

But what about longer pieces?

Human genome is 3 billion bases long, arranged on 23 pairs of chromosomes.

Sequencing machine reads just a drop in the ocean!

Solution: Break the entire genome into <u>manageable</u> pieces and sequence them.

Two approaches used for sequencing the Human genome:

- Publicly funded Human Genome Project (HGP) clone-byclone or hierarchical shotgun sequencing method
- Privately Funded Sequencing Project Celera Genomics –
 whole genome shotgun sequencing method

Hierarchical shotgun sequencing approach:

- genomic DNA is cut into pieces of about 150 Mb
- inserted into **BAC/YAC** cloning vectors,
- transformed into *E. coli*/yeast where they are replicated and stored.

BAC/YAC inserts are isolated & mapped to determine the order of each cloned 150 Mb fragment

- referred to as the Golden Tiling Path

Begun formally in 1990, Human Genome Project was a 13-yr effort coordinated by the U.S. DAE and NIH.

- completed in 2003

Each BAC/YAC fragment in the Golden Path is

- fragmented randomly into smaller pieces,
- each piece is cloned into a plasmid and sequenced on both strands.

These sequences are aligned so that identical regions overlap.

Contiguous pieces are then assembled into finished sequence once each strand has been sequenced about 5 times to produce 10× coverage of high-quality data.

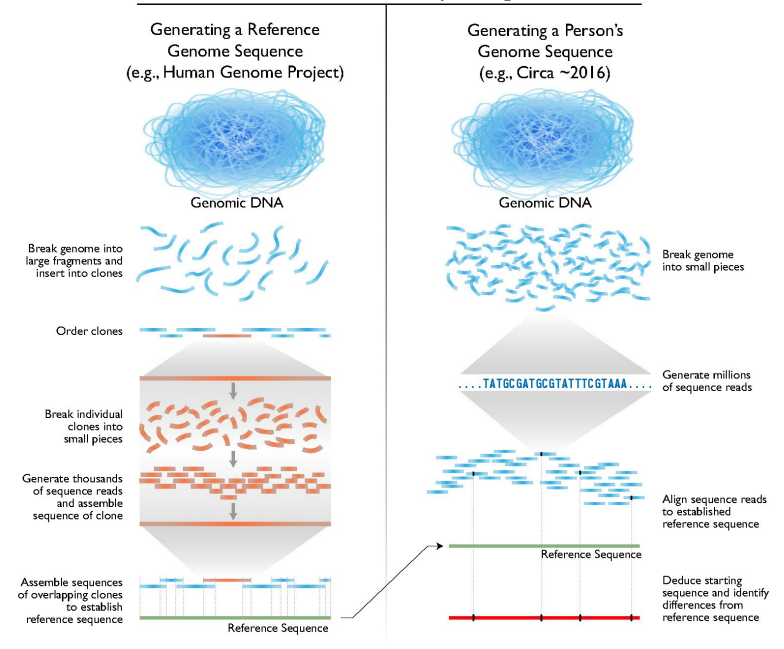
Whole genome shotgun sequencing (WGS)

- method developed and preferred by Celera Genomics
- skips the entire step of making libraries of BAC clones

Blast apart the entire human genome into fragments of 2 - 10 kb and sequence them.

Challenge is then to assemble these fragments into the whole genome sequence.

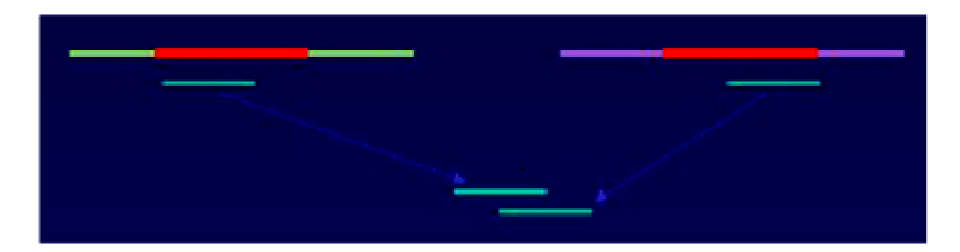
Human Genome Sequencing



Whole Genome Shotgun Method

What makes the task of assembling the genome fragments especially challenging

- repeats in the genome (~50% in human genome).



Whole Genome Shotgun Method

Adding to the challenge is the sheer computational complexity of the task.

Size of H. genome = 3×10^9 bp. Given length of read ~500 bps, for desired coverage of 10x, No. of reads required is:

RequiredReads = GenomeLength * DesiredCoverage /
ReadLength
= 6 * 10⁷

With 60M reads to assemble, we need algorithms that run in near linear time (O(nlogn))

Whole Genome Shotgun Method

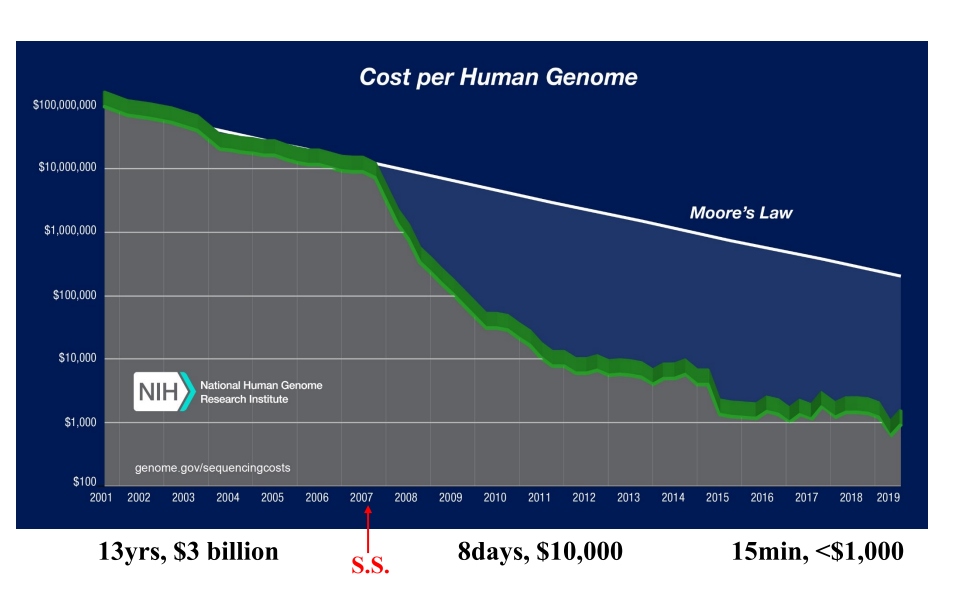
Which method is better?

Depends on the size and complexity of the genome

Note: Celera had access to the HGP data but the HGP did not have access to Celera data.

Which method is preferable for sequencing the genome of a novel coronavirus — SAR-CoV-2? Why?

High-throughput / Next-Generation Sequencing



DNA sequencing beating Moore's law

HTS/NGS Sequencing

High-throughput sequencing (HTS) technologies have revolutionized the way biologists acquire and analyze genomic data - massively parallel sequencing

	Roche GS FLX+	Illumina HiSeq 2000	SOLiD™ 4	Ion Torrent PGM
Bases per run	700Mb	600 Gb	100 GB	1 Gb
Time per run	23h	~11 days	~14 days	4.5 h
Reads per run	1 Million	6 Billion (paired-end) 3 Billion (single)	1.4 Billion	Millions
Read length	~700 bp	2 x 100 bases	2 x 50 bases	35–400 bases

- can generate tens of gigabases per week, at a cost 200-fold less than previous methods.

Sequencing Machines: Overview



Data Analysis

The scale and nature of data produced by all NGS platforms place substantial demands on IT at all stages of sequencing, including data tracking, storage, and quality control:

- **base calling** by proprietary software
- Quality check and filtering or reads
- aligning sequencing data to Reference genome, if available, or a *de novo* assembly is conducted.

Once the sequence is aligned to a reference genome, the data needs to be analyzed in an experiment-specific fashion.

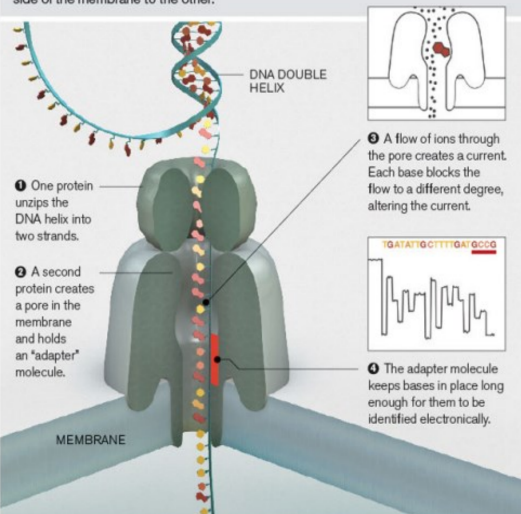
Sequence alignment & assembly is an active area of computational research

Third Generation Sequencing (TGS)

- 'Long read sequencing' read length: ~ 10 60Kb
- Single molecule sequencing nanostructure-based sequencing
- No PCR step involved
- Faster and portable
- Under active development
- e.g., PacBio Single molecule real time sequencing (SMRT) and Oxford Nanopore

Oxford Nanopore - MinION

DNA can be sequenced by threading it through a microscopic pore in a membrane. Bases are identified by the way they affect ions flowing through the pore from one side of the membrane to the other.







HTS Applications

One of the most prominent applications of NGS is re-sequencing:

- whole genome resequencing, e.g., genetic variations
- target-region resequencing, e.g., heritable disorders
- exome resequencing, e.g. cancer
- genome-wide analysis of single nucleotide variations and other structural variations, multiple individuals, or strains, cancer sequencing, population-based sampling of a species, migration patterns of a virus, e.g., SARS-CoV-2, etc.

Any human individual's genome available in NCBI?

How would you go about sequencing SARS-CoV-2 genome, 29903 bases long?

What technique is used for diagnostic testing of COVID-19?

While sequencing a novel genome for the first time, how are primers identified?

Real time RT-PCR used for diagnostic testing of COVID-19

It is a laboratory technique combining reverse transcription of RNA into DNA (called complementary DNA or cDNA) and amplification of specific DNA targets using polymerase chain reaction (PCR).

It is primarily used to measure the amount of a specific RNA.

This is achieved by monitoring the amplification reaction using fluorescence, a technique called real-time PCR or quantitative PCR (qPCR).

- routinely used for analysis of gene expression and quantification of viral RNA in research and clinical settings.

Can we now answer these Qs:

- How is the SARS-CoV-2 genome sequenced?
- How does one identify the coordinates of N gene on it? i.e., how to construct a physical map of a genome?
- How does one select which regions in this gene would give specificity for the presence of SARS-CoV-2?
- How is the specific probe regions extracted and amplified for detection?
- Is it possible to store the DNA sample for re-testing? How?

References:

- 1. Concepts in Biotechnology, ed. D. Balasubramanyam
- 2. REBASE: restriction enzymes and methyltransferases, Nucleic Acids Research, Vol. 31 (1), 418–420 (2003)