



How to Sequence a whole genome

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Why do we want to know the sequence of an entire genome??



To know all the genes – then proteins, then pathways... We can understand:

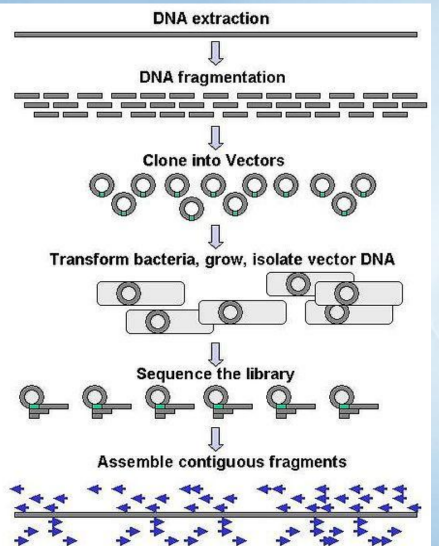
- the biochemistry of the organism
- diseases

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Regulation

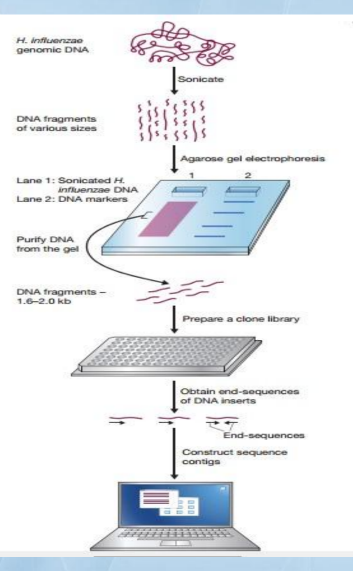


Genome Sequencing (Shotgun)



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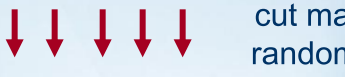


Method to sequence longer regions

~500 bp

genomic segments

~500 bp



cut many times at random (*Shotgun*)

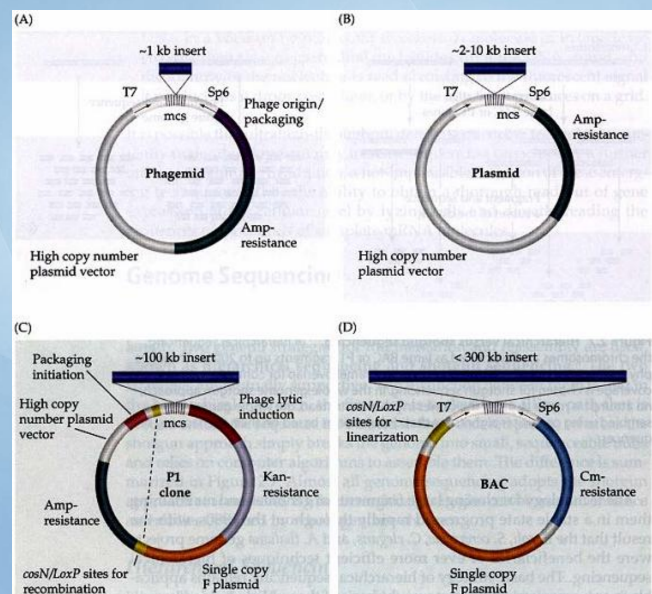
Get one or two reads from each segment

Cloning vectors for genome sequencing

Jul Univers

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Sizes of inserts in sequencing vectors



Vector **P1** YAC BAC Cosmid Plasmid M13 or Phagmid

<u>Size (approx.)</u>

100 Kb 300 -1500 Kb 70 - 300 Kb ~ 40 Kb 2 -10 Kb

~ 1 Kb



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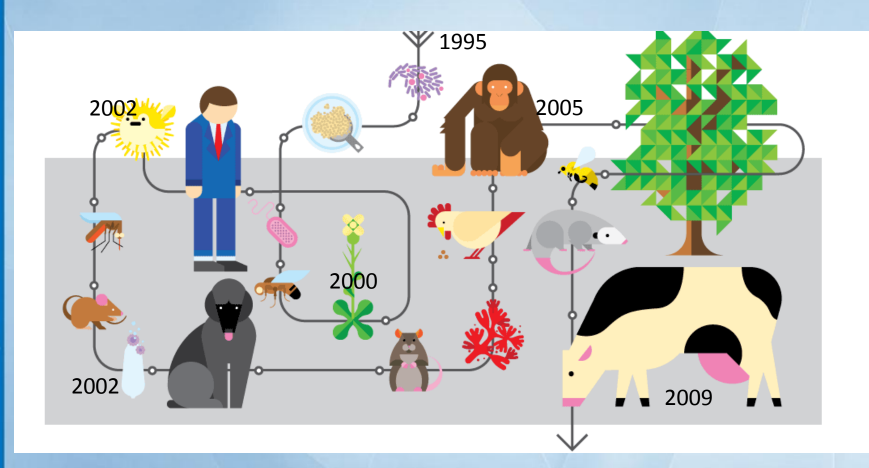


Overlap reads and extend to reconstruct the original genomic region



Will we sequence every species?





Genomes to Date



- 69 higher animals + other model animals
- 55 insects and lower metazoans
- 39 plants
- 563 fungi
- Over 200 protist species and subspecies
- Over 20 000 bacteria species and subspecies
 - Microbial communities in oceans, desserts, hot springs, inside bodies

Sequencing of extinct species

The complete genome sequence of a Neanderthal from the Altai Mountains

Kay Prüfer, Fernando Racimo, Nick Patterson, Flora Jay, Sriram Sankararaman, Susanna Sawyer, Anja Heinze, Gabriel Renaud, Peter H. Sudmant, Cesare de Filippo, Heng Li, Swapan Mallick, Michael Dannemann, Qiaomei Fu, Martin Kircher, Martin Kuhlwilm, Michael Lachmann, Matthias Meyer, Matthias Ongyerth, Michael Siebauer, Christoph Theunert, Arti Tandon, Priya Moorjani, Joseph Pickrell, James C. Mullikin \Rightarrow et al.

Affiliations | Contributions | Corresponding authors

Nature 505, 43–49 (02 January 2014) | doi:10.1038/natu Received 05 September 2013 | Accepted 15 November 2 2013

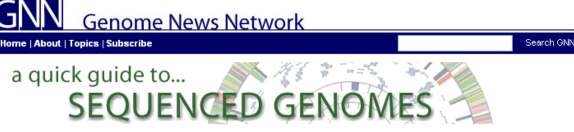


Neanderthal toe bone



Where can I find genome sequences?





The genomes of more than 180 organisms have been sequenced since 1995. The Quick Guide includes descriptions of these organisms and has links to sequencing centers and scientific abstracts.

» See the Complete List of Organisms

Written by Kate Ruder and Edward R. Winstead Created for GNN by Mary S. Gibbs

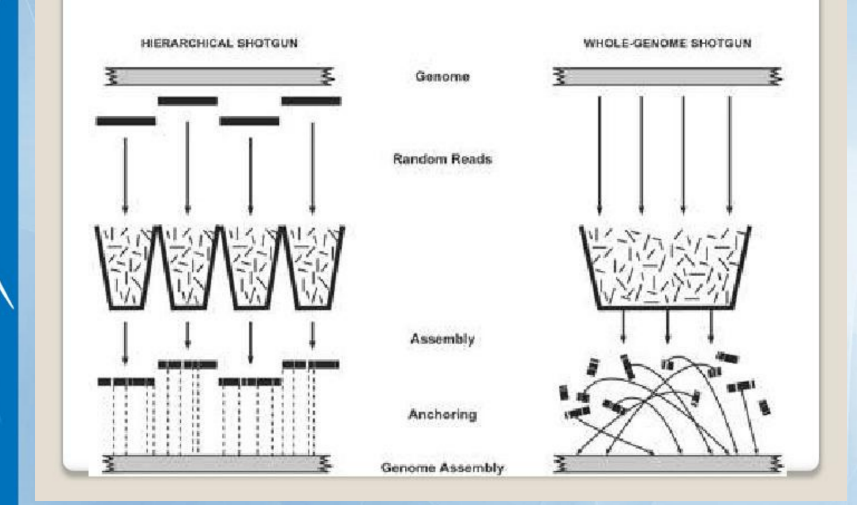
http://www.genomenewsnetwork.org/

Websites "genome browsers" (include annotations of genes) Ensembl genome browser UCSC genome browser NCBI genome browser



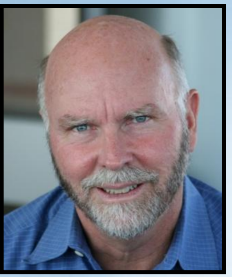


Hierarchical vs. Whole Genome

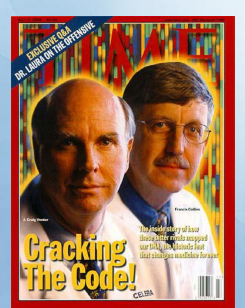








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The Human Genome Project



- The Mission of the HGP: To understand the human genome and the role it plays in both health and disease.
- Initiated 1990 -Completion originally planned for 2005 (expected to take 15 years)
- \$3-billion

<u>movie</u>

Finished sequence anticipated Spring, 2003, after 50 years of Watson and Crick publication (Nature 171: 737-738, April 25, 1953)



VOL CXLIX No. 51,432 Copyright () 2000 The New York Times TUESDAY, JUNE 27, 2000

Genetic Code of Human Life Is Cracked by Scientists

6 guaninia r. thymina of the intertwining

double helix of DRA

JUSTICES REAFFIRM MIRANDA RULE, 7-2; **A PART OF 'CULTURE'**

By LINDA GREENHOUSE

WASHINGTON, June 26 - The Supreme Court reaffirmed the Miranda decision today by a 7-to-2 vote that erased a shadow over one of the most famous rulings of modern times and acknowledged that the Miranda warnings "have become part of our national culture."

The court said in an opinion by Chief Justice William H. Rehnquist that because the 1966 Miranda deciaton "announced a constitutional rule," a statute by which Congress had sought to overrule the decision was itself unconstitutional.

Miranda had appeared to be in jeopardy, both because of that longignored but recently rediscovered law, by which Congress had tried to overrule Miranda 32 years ago, and because of the court's perceived hostility to the original decision.

The chief justice said, though, that the 1968 law, which replaced the Miranda warnings with a case-by-case test of whether a confession was voluntary, could be spheld only if the Supreme Court decided to overturn Miranda, But with Miranda having "become embedded in routine police practice" without causing any measurable difficulty for prosecutors. there was no justification for doing so, he said. [Excerpts, Page Al8.]

Justices Antonin Scalia and Clarence Thomas cast the dissenting votes.

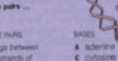
The decision overturned a ruling last year by the federal appeals court in Richmond, Va., which beld that Congress was entitled to the last word because Miranda's presumption that a confession was not voluntary unless preceded by the warnings was not required by the Constitution.

The decision today - only 14 pages long, in Chief Justice Rehnquist's typically spare style - brought an abrupt end to one of the odder episodes in the court's recent history, an intense and strangely delayed refighting of a previous generation's battle over the rights of criminal suspects. Miranda v. Arizona was a hallmark of the Warren Court, and Chief Justice Rehnquist, despite his record as an early and tenacious critic of the decision, evidently did not want its repudiation to be an imprint of his own tenure.

There was considerable drama in the courtroom today as the chief justice announced that he would deliver the decision in the case. Dickerson v. United States, No. 99-5525. The announcement meant that he was the majority opinion's author. Given his statements over more than 25 years about Miranda's lack of constitutional foundation, there was the The 3 billion hese pairs

The Book of Life

Rungs between the strands of the double helix



Science Times A special testar

- · Putting the genome to work.
- Some information has already paid research dividends.
- # Two research methods, two results

 More articles, charts and photos of the genome effort. From Mendel to hei

to genome. Section D

head of the Human Genome Project, right, with J. Craig Venter, head of Celera Genomics, after the announcethey had finished the first survey of

Francis S. Collins,

ment vesterday that the human genome.

.... that make up the set of chromosomes in our cells, have been sequenced.

By ordering the base units, scientists hope to locate the genes and determine their functions.

The New York Timer



Marks New Medical Era, Risks and All By NICHOLAS WADE. WASHINGTON, June 26 - In an

A SHARED SUCCESS

2 Rivals' Announcement

achievement that represents a pinnacle of human self-knowledge, two rival groups of scientists said today that they had deciphered the hereditary script, the set of instructions that defines the human organism.

"Today we are learning the language in which God created life," President Clinton said at a White House ceremony altended by members of the two tealns and, via satellite, Prime Minister Tony Blair of England. (Excerpts, Page D8.)

The teams' leaders, Dr. J. Craig Venter, president of Celera Geno mics, and Dr. Francis S. Collins, director of the National Human Genome Research institute, praised each other's contributions and signaled a spirit of cooperation from now on, even though the two efforts will remain firmly independent.

The human geoome, the ancient script that has now been deciphered. consists of two sets of 23 giant DNA molecules, or chromosomes, with each set - one inherited from each parent - containing more than three billion chemical units.

The successful deciphering of this vast genetic archive attents to the extraordinary pace of biology's advance since 1953, when the structure of DNA was first discovered ant presages an era of even brisker

By NATALIE ANGIER

Collins, director of the National Hu-

"We only have to do this once, read- by calling the genome a "portrait of

Though scientists underscore the man Genome Research Institute. importance of their accomplishment

A Pearl and a Hodgepodge: Human DNA

15 February 2001

www.nature.com

human genome

Nuclear fission Five-dimensional energy landscapes Seafloor spreading The view from under the Arctic ice Career prospects Sequence creates new opportunities

naturejobs genomics special



Vol. 291 No. 5507 Pages 1145-1434 \$9

THE HUMAN GENOME

AMERICAN ASSOCIATION FOR THE ADVANCEMENT OF SCIENCE



Human genome content



•Total length 3 billion bp ~ 30,000 genes (coding seq) functions of more than half of them are unknown

•30,000 genes but > 500,000 known proteins (possibly exceed 2 million)

Gene sequences < 5%
Exons ~ 1.5% (coding)
Introns ~ 3.5% (noncoding)

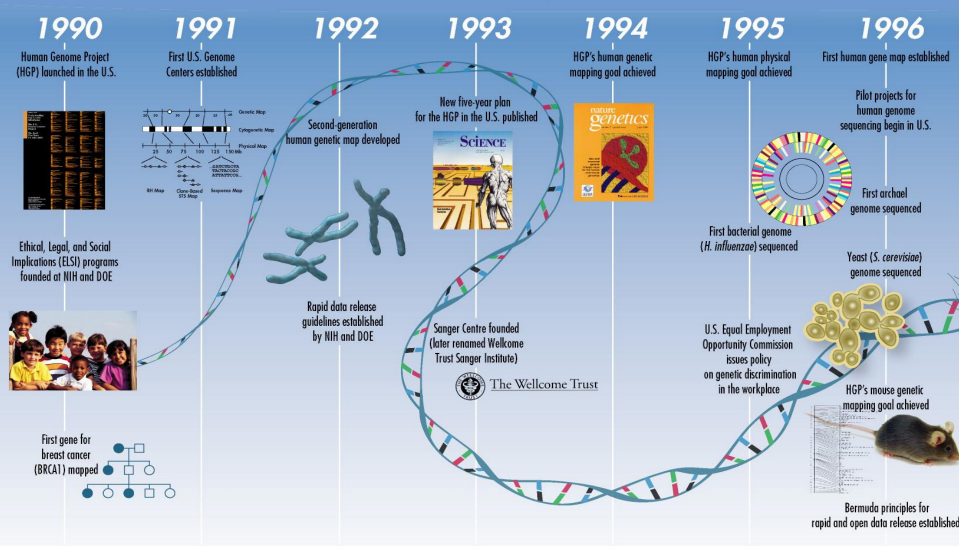
Intergenic regions (junk) > 95%

•The human genome is nearly the same (99.9%) in all people

•Almost half of all human proteins share similarities with other organisms

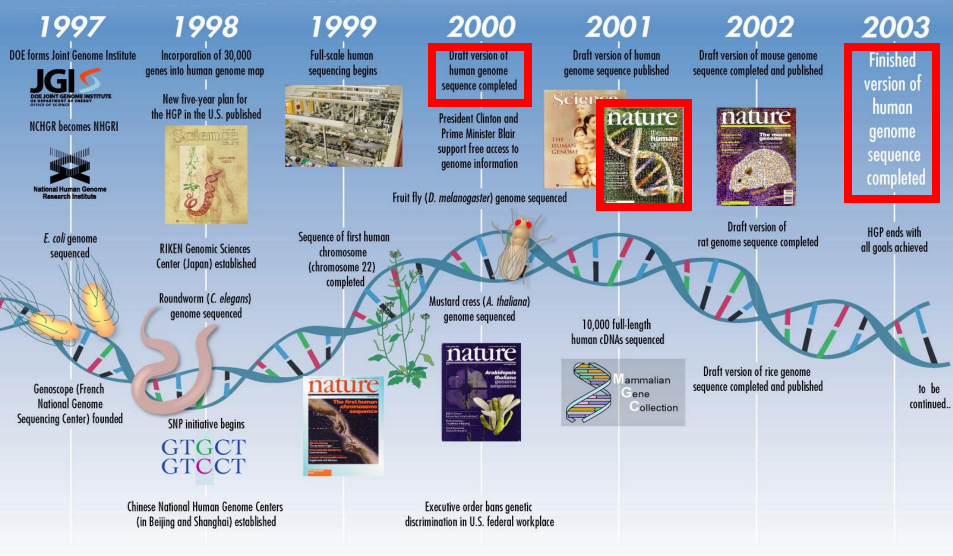












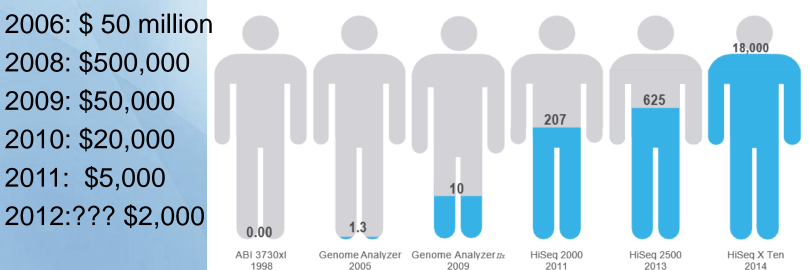




In 2005, a single sequencing run could produce roughly one gigabase of data.

By 2014, the rate climbed to a 1.8 terabases of data in a single sequencing run—an astounding 1000× increase.

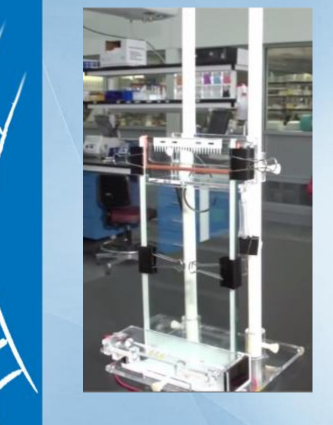
The HiSeq X[™] Ten, released in 2014, can sequence over 45 human genomes in a single day for approximately \$1000 each

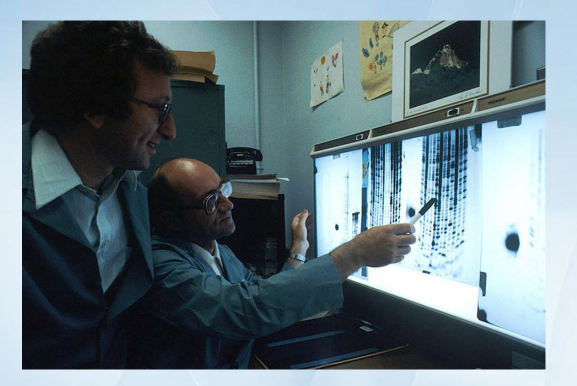


Human Genomes Sequenced Annually











Next Generation Sequencing (NGS)



- Roche/454
 - (GS FLX+/GS Junior)
- Illumina Genome Analyzer
 - (HiSeq/MiSeq/NextSeq)
- Life Technologies
 - (3500 Genetic Analyzer, Ion Torrent Proton/PGM)
- Pacific Biosciences
 - (PACBIO RSII)
 - Applied Biosystems
 - (SOLiD, 3730x/ DNA Analyzer)











Sequencing Principles



Sequencing by Synthesis

- Sanger/Dideoxy chain termination (Life Technologies, Applied Biosystems)
- Pyrosequencing (Roche/454)
- Reversible terminator (Illumina)
- Ion torrent (Life Technologies)
- Zero Mode Waveguide (Pacific Biosciences)

Sequencing by Oligo Ligation Detection

SOLiD (Applied Biosystems)

Direct reading of DNA sequence

- Nanopore sequencing
- Electron microscope

3rd generation sequencing 3rd generation sequencing



Platform Company	3730xl ABI	5500xl SOLiD ABI	454 FLX Titanium Roche	HiSeq 2000 Illumina	GAIIx Illumina	MiSeq Illumina	Ion Torrent Life Tech.
Amplification	Biol/PCR	EmPCR	EmPCR	BrPCR	BrPCR	BrPCR	EmPCR
Detection	Fluor.	Fluor.	Fluor.	Fluor.	Fluor.	Fluor.	pH
Run Time (days)	0.08	8	0.5	8	14	1.1	0.08
Max. Aver. Length (bp)	900	60x2	700	101x2	151x2	151x2	100
Max. TP/run (Gbp)	0.00008	310	0.8	600	100	1	0.1
Max.Reads/Run(Million)	0.000096	5,167	1	3,000	320	3	1
TP per 24hr (Gbp)	0.00064	45	1	75	7	1	2.4
Raw Error range (%)	0.01	0.01	1-3	0.1	0.1	0.1	(1)*
Equip.Cost (xUS\$1,000)	150	600	300	690	350	125	60
Cost per Mbp (US\$)	4,000	0.05	8	0.02	0.1	0.7	10

SBS: Sequencing by synthesis, SbL: Sequencing by ligation, PS: Pyrosequencing, EmPCR: Emulsion PCR, Biol: Biological cloning, Fluoresc.: Fluorescence, BrPCR: Bridge PCR, TP: Throughput.

Table 1. Comparison of current sequence technologies.

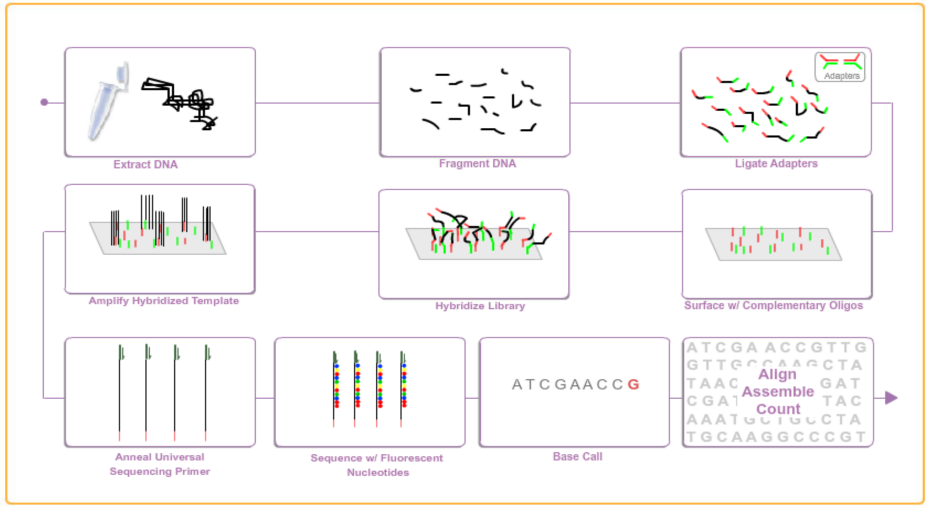
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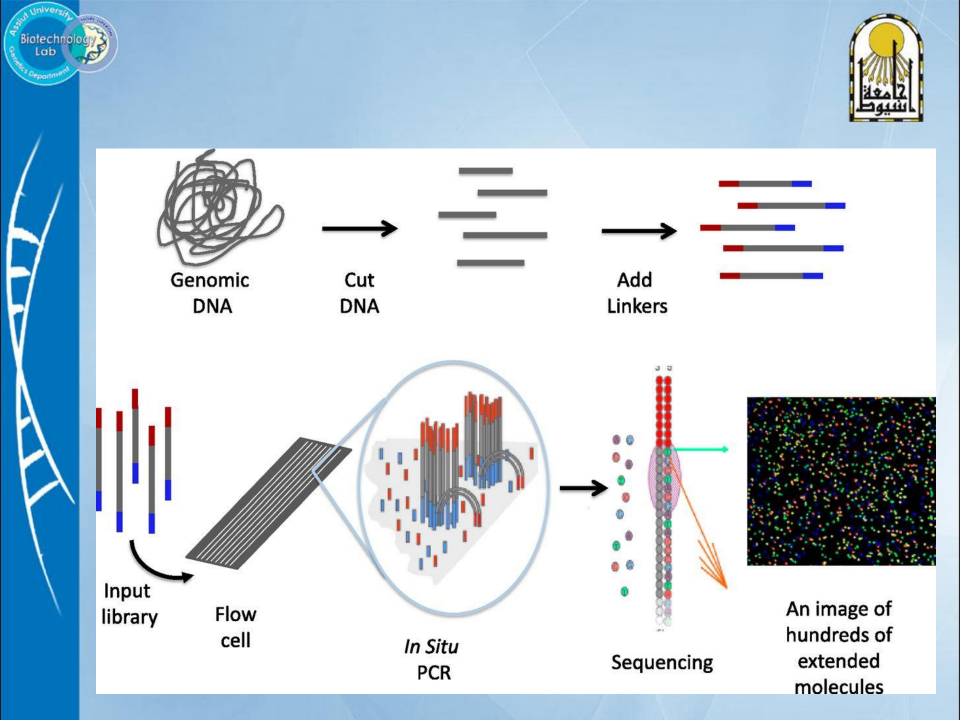


Sequencing by Synthesis (SBS) Overview

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Illumina sequencing can sequence billions of template strands simultaneously

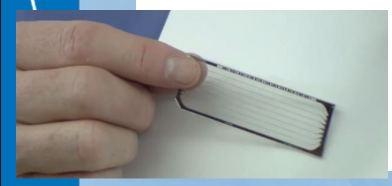




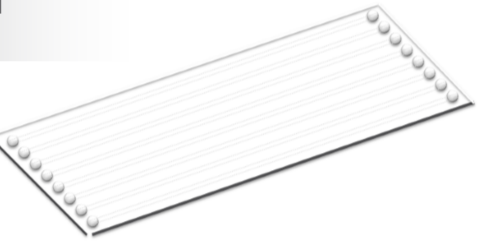
The flow cell - a core component

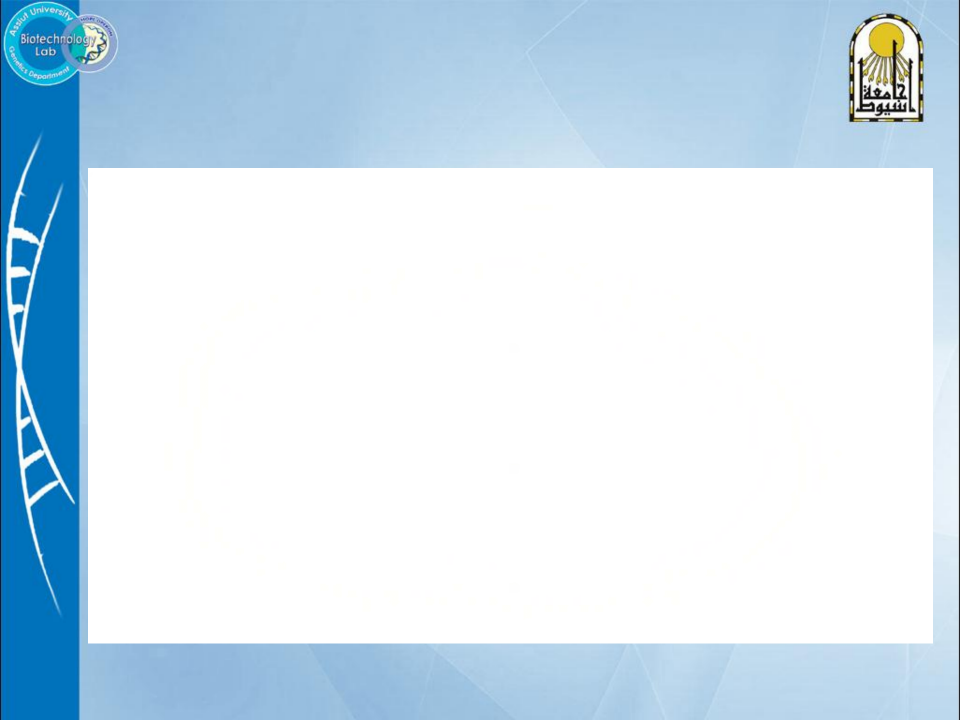
EVERYTHING EXCEPT SAMPLE PREPARATION IS COMPLETED ON THE FLOW CELL

template annealing (1 - 96 samples) template amplification sequencing primer hybridization Sequencing-by-synthesis reaction generation of fluorescent signal



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Phospholinked Fluorophores

Biotechnology

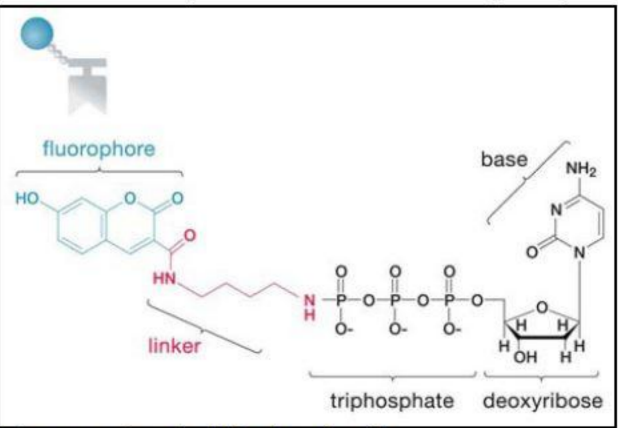


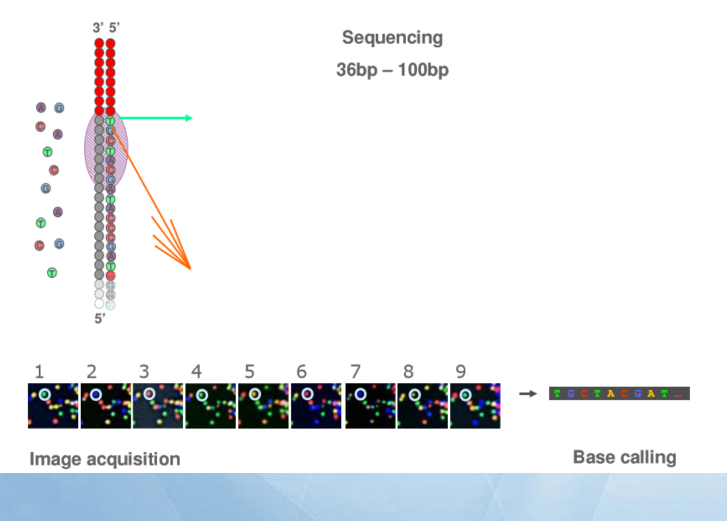
Figure 9. Phospholinked nucleotides

Phospholinked nucleotides have fluorophores attached to the triphosphate chain, which is naturally cleaved when the nucleotide is incorporated.



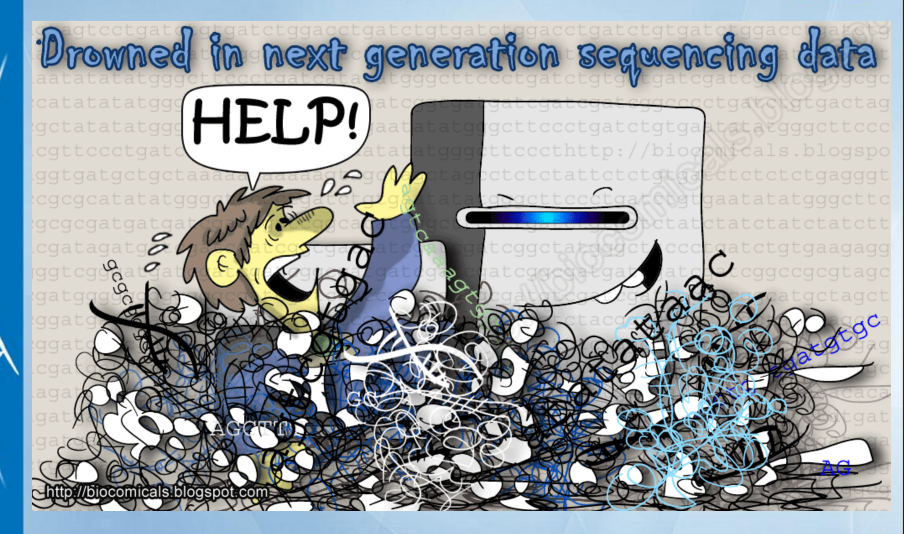


A picture is taken every time a new base is added













Nanopore sequencing (direct reading)



3rd generation sequencing

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.

