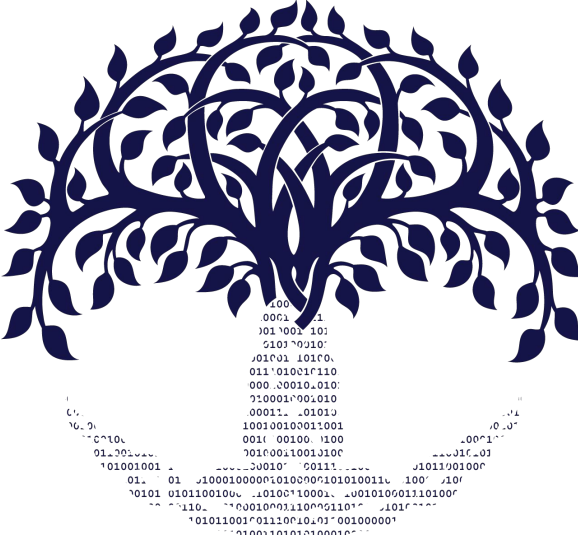


DTU





**DTU Health Technology
Bioinformatics**

Introduction to NGS

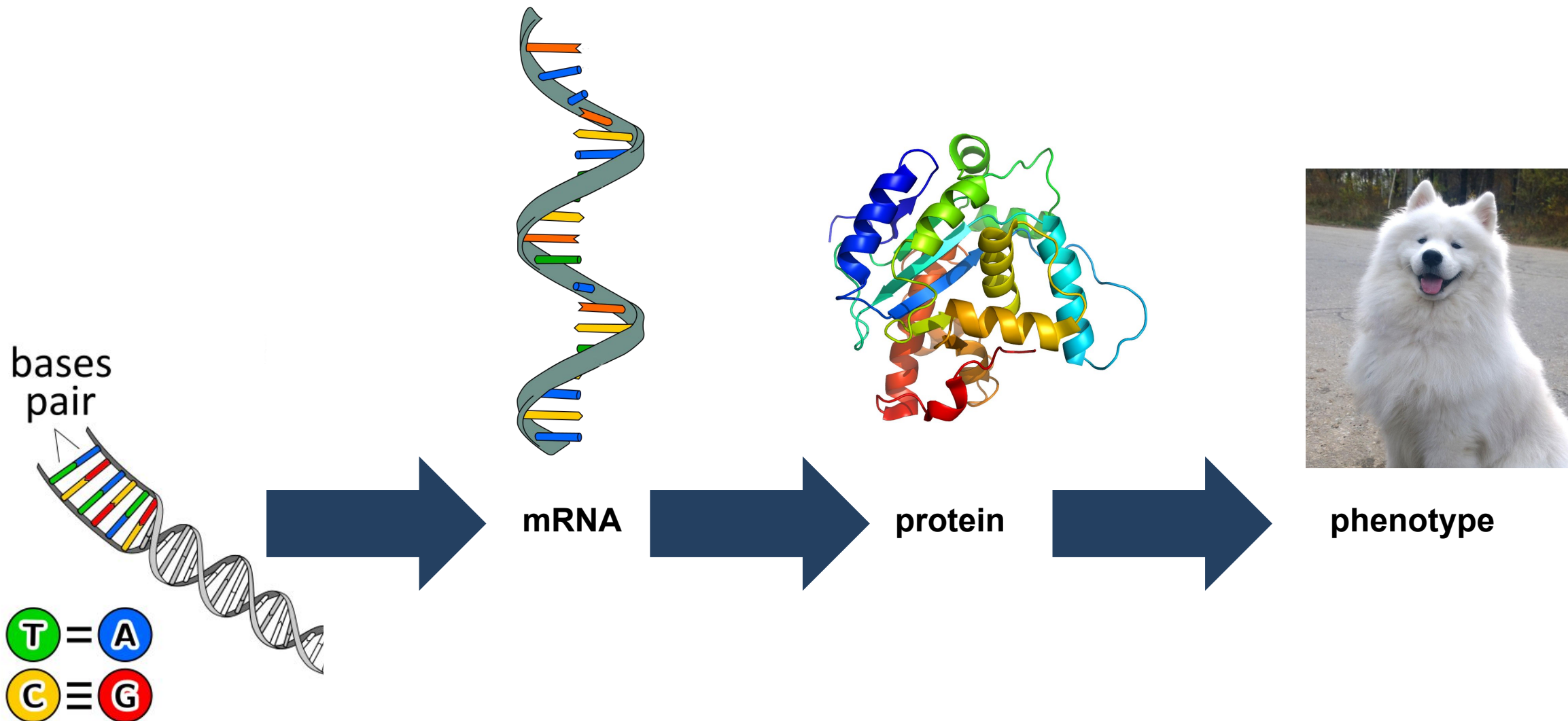
*Gabriel Renaud
Associate Professor
Section of Bioinformatics
Technical University of Denmark
gabriel.reno@gmail.com*

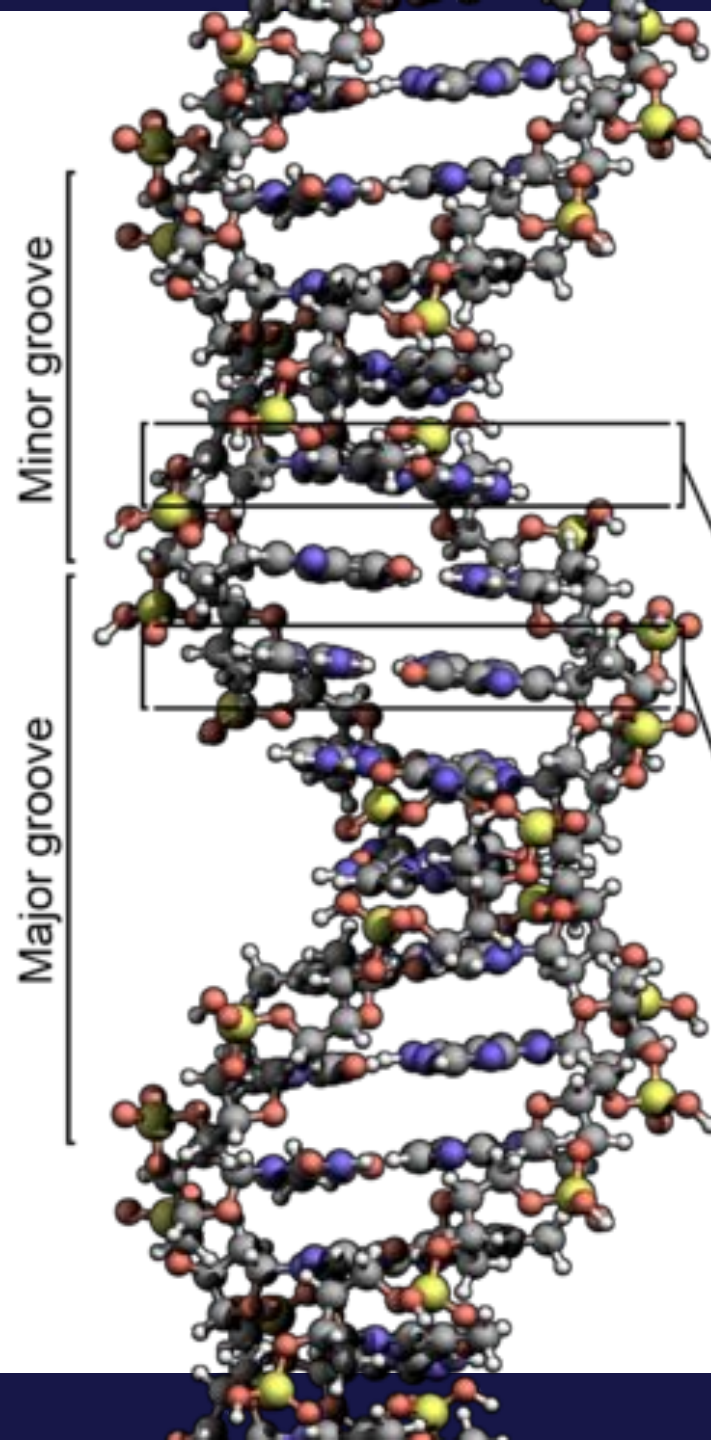
Menu

- What is sequencing? why?
- Basic nomenclature

What is sequencing?

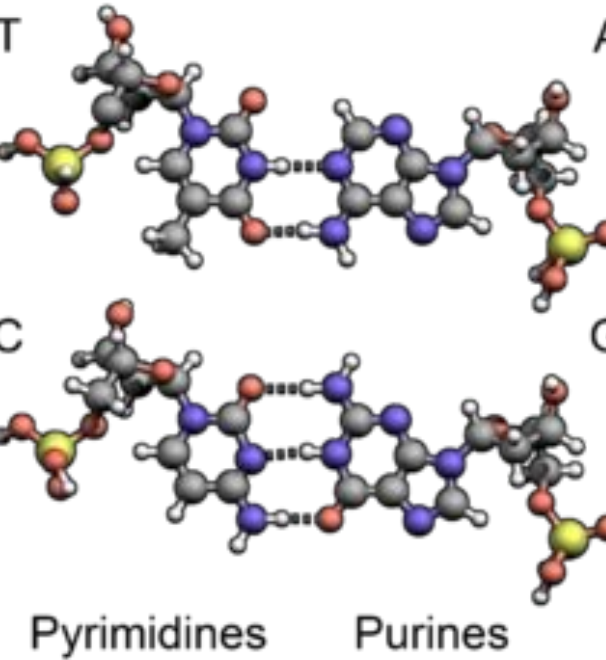
Remember high school?



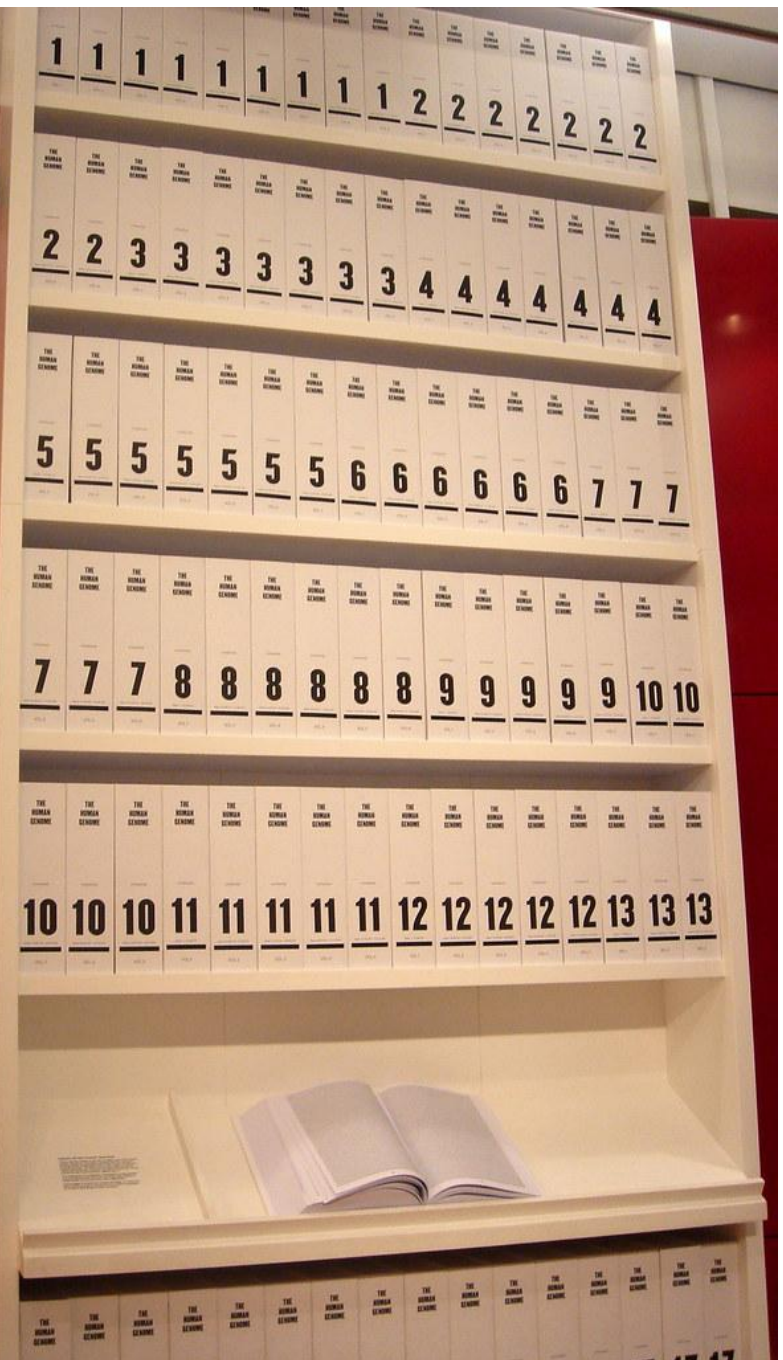


READING

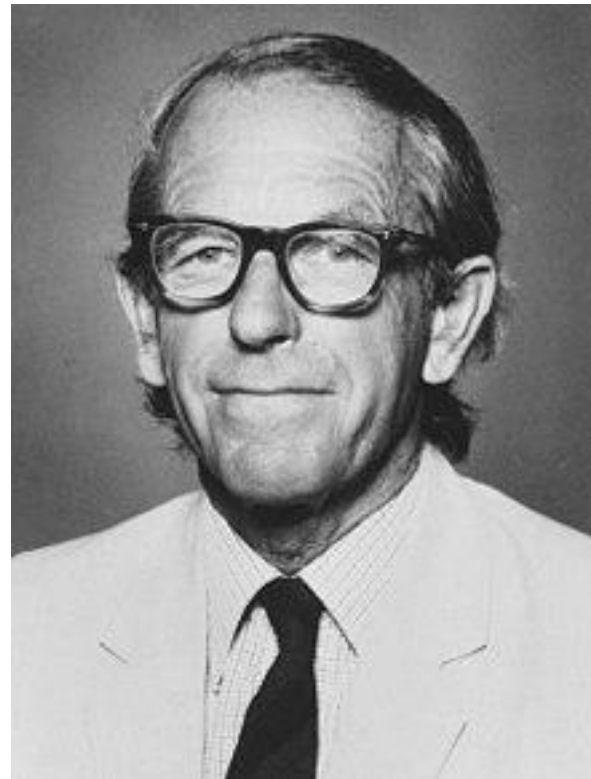
AGCAATCTCAATTACA



Human genome 3 billion letters

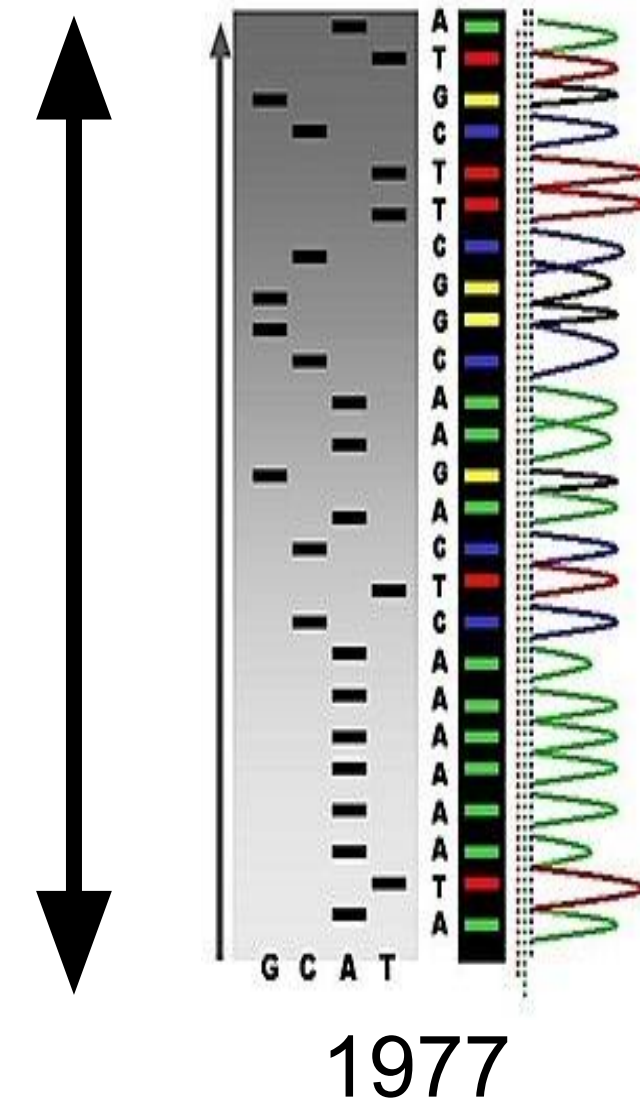


If we study Next-Generation Sequencing, why “next”? What was before?



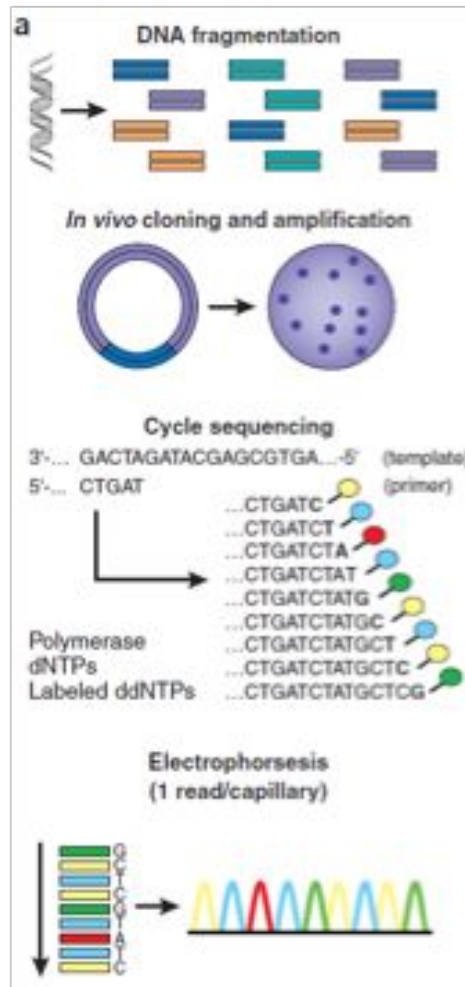
1000 bases
x
96

Frederick Sanger 1918 - 2013



1977

First generation: Sanger



- Fragment DNA
- Clone into plasmid and amplify
- DNA polymerase and only 1 primer
- Sequence using labeled dinucleotides which cap seqs.
- Run capillary electrophoresis/gel and “read” DNA code
- Low output, long reads (~800-1200 nt), high quality
- Produces 96 reads / run

Why sequence?



AGGATTATTGGTACT



AGGATTATTGGTACT



AGGATTAT**C**GGTACT



AGGATTATTGGTACT



AGGATTATTGGTACT



AGGATTATCGGTACT



AGGATTATCGGTACT



AGGATTATTGGTACT



AGGTTTATTGGTACT



AGGATTATCGGTACT



AGGATTATCGGTACT



AGGTTTATTGGTACT



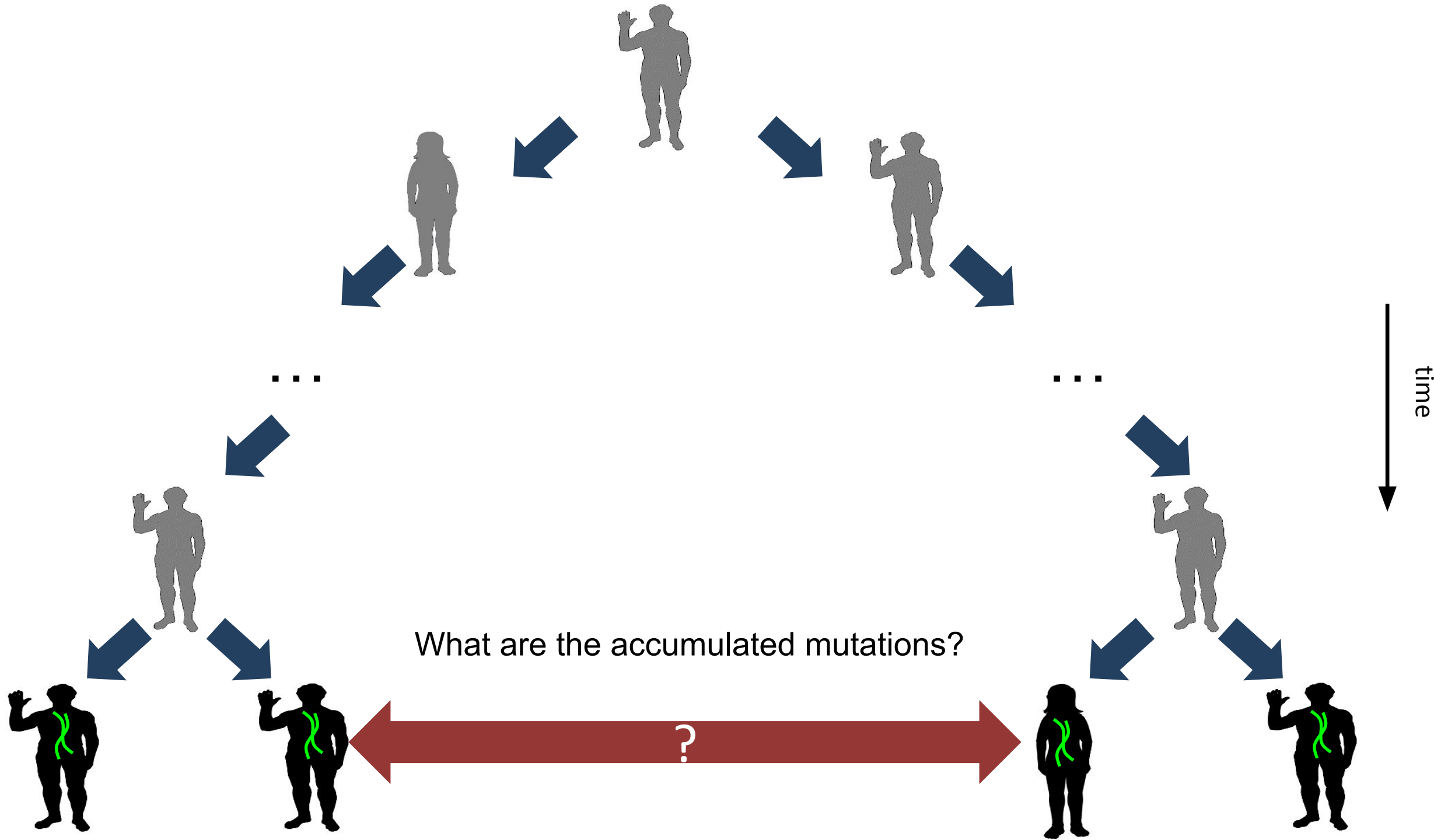
AGGTTTATTGGTAGT



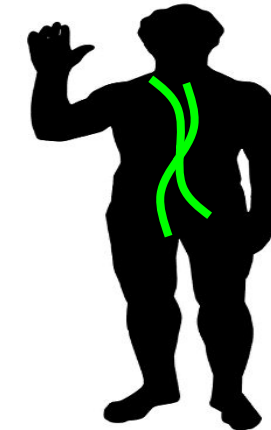
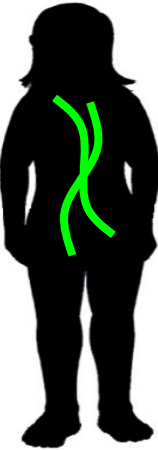
AGGATTATCGGTACT



AAGATTATCGGTACT

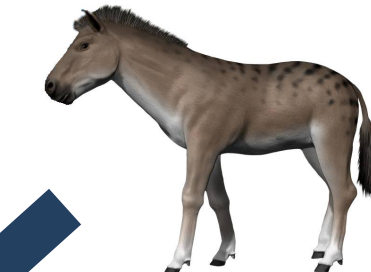


Investigate differences within a species

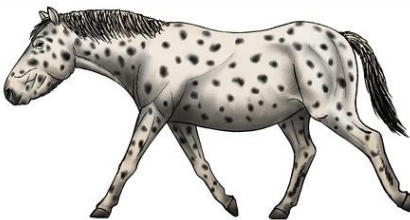


AGGTTTATTGGTAGT
AAGATTATCGGTACT

Investigate differences between species



?



time

"Nothing in Biology Makes Sense Except in the Light of Evolution"

Theodosius Dobzhansky, 1973

"Nothing in Biology Makes Sense Except in the Light of Evolution"

Theodosius Dobzhansky, 1973

NGS

"Nothing in Biology Makes Sense Except in the Light of Evolution"

me, I made that up just now

What can we use it for?

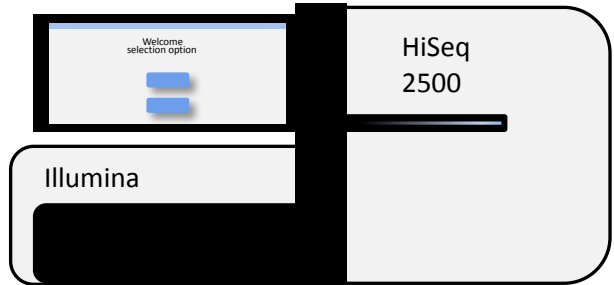
- Whole genome re-sequencing
- Population genomics
- Diagnostics
- Cancer genomics
- Ancient genomes
- Metagenomics
- RNA sequencing
- Single cell sequencing
- Genomic Epidemiology
- anything with DNA



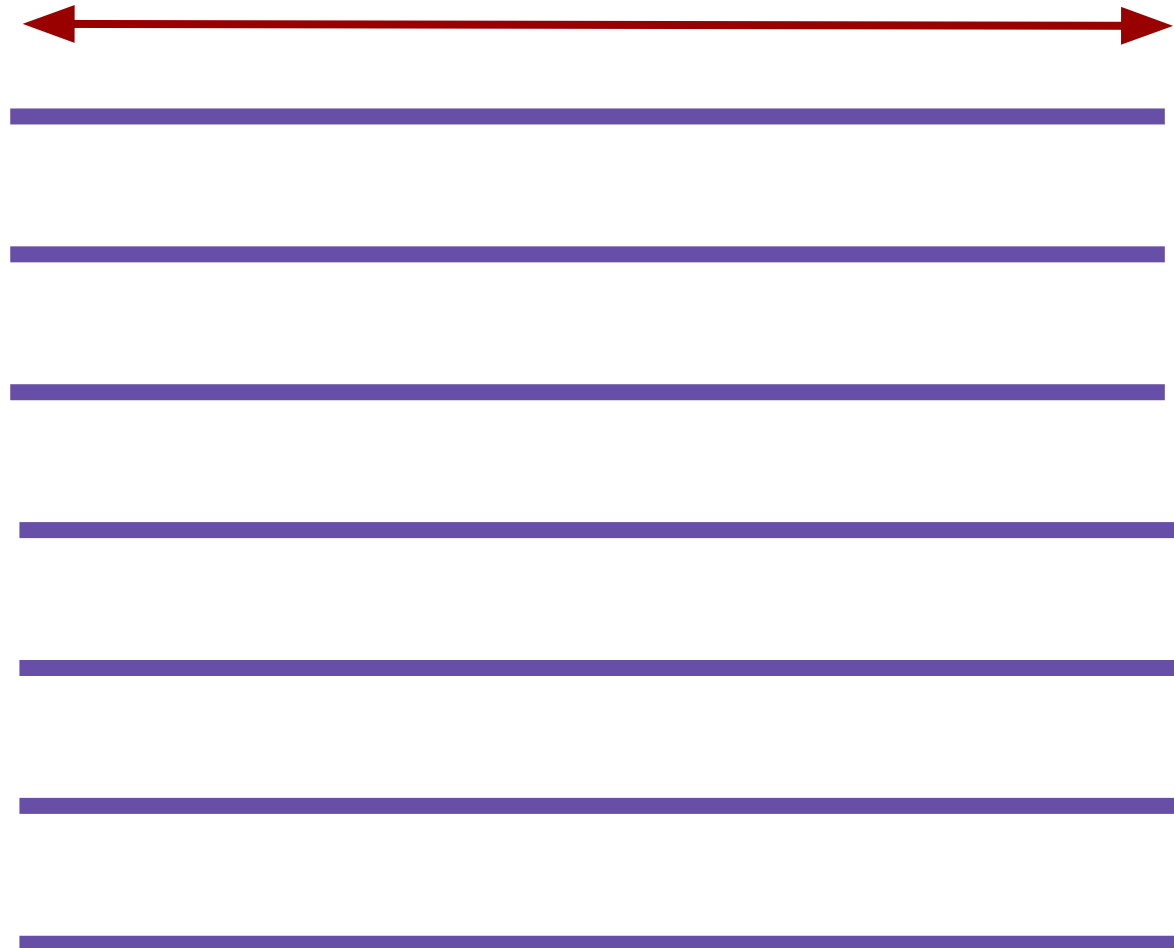
Basic concepts

3 key concepts

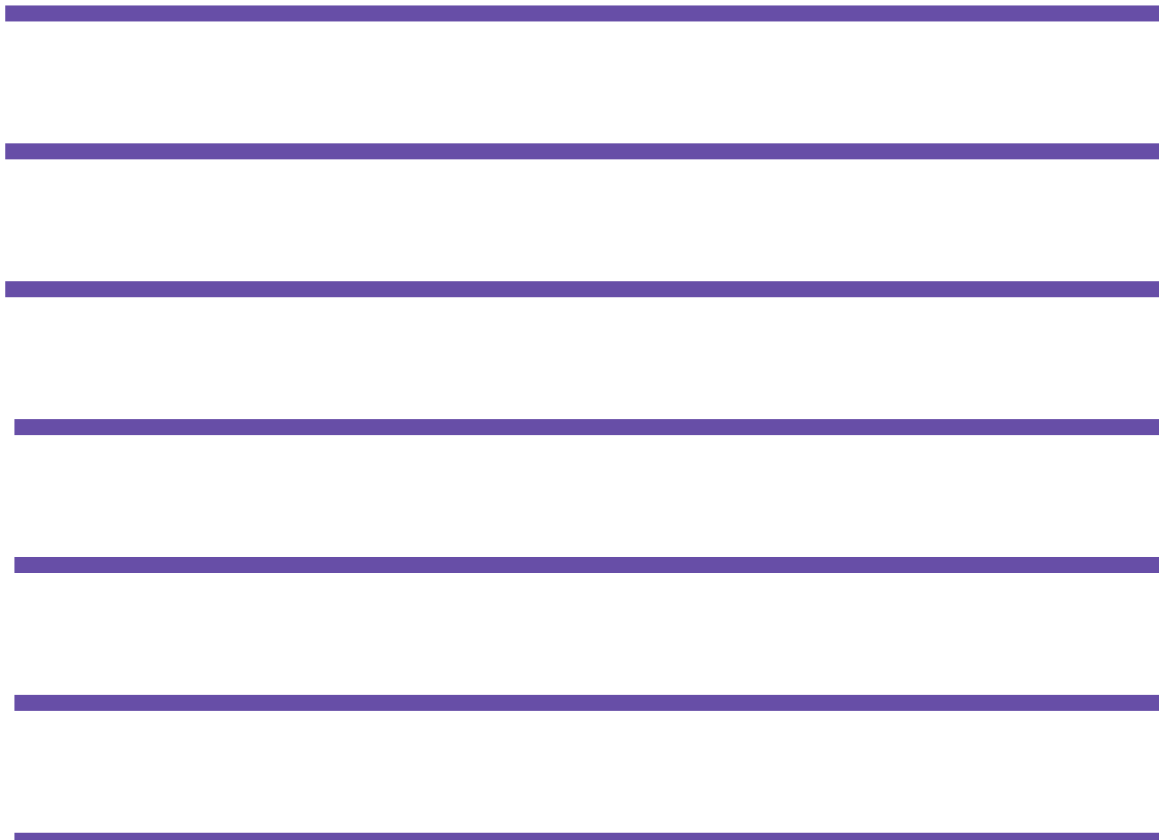
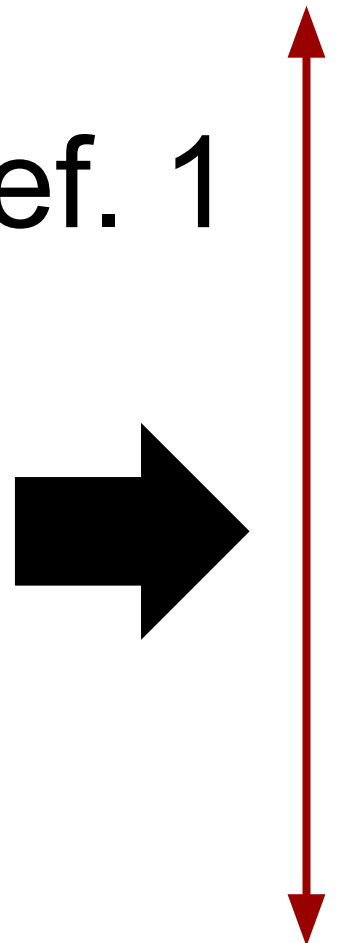
- Read length
- Throughput
- Types of errors



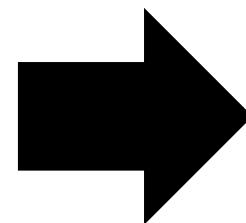
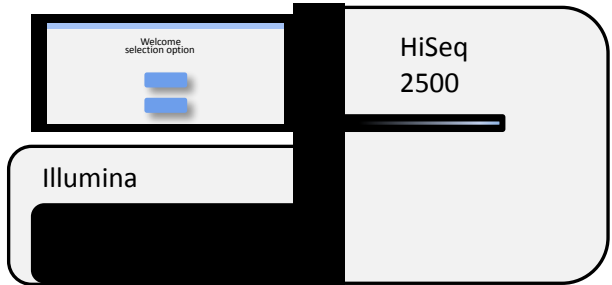
read length



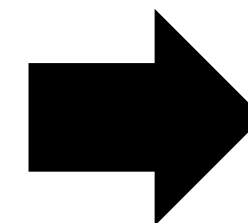
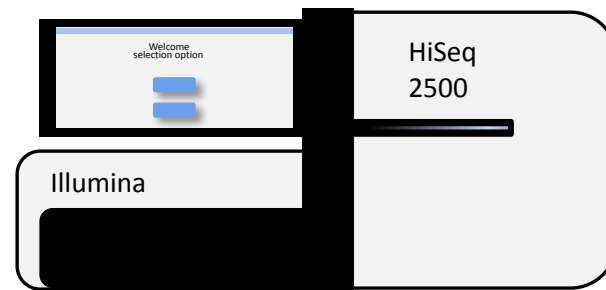
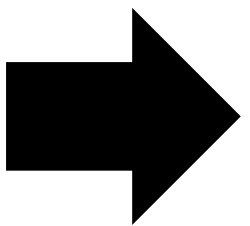
throughput def. 1



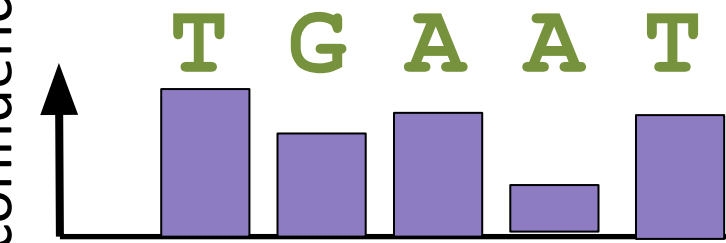
throughput def. 2



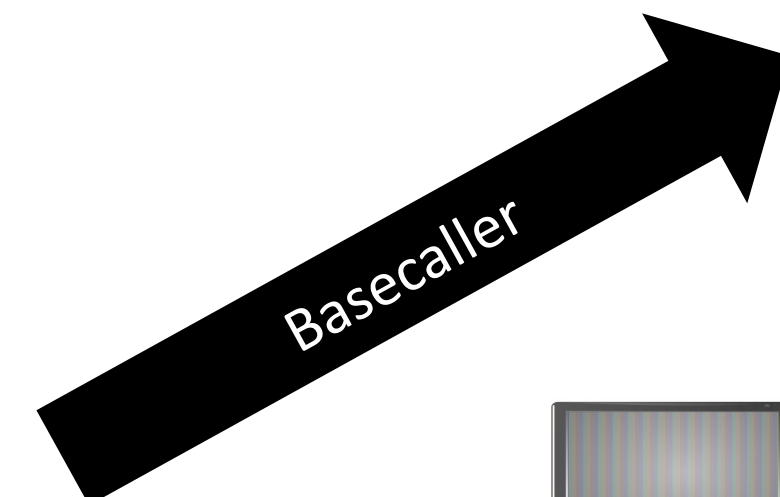
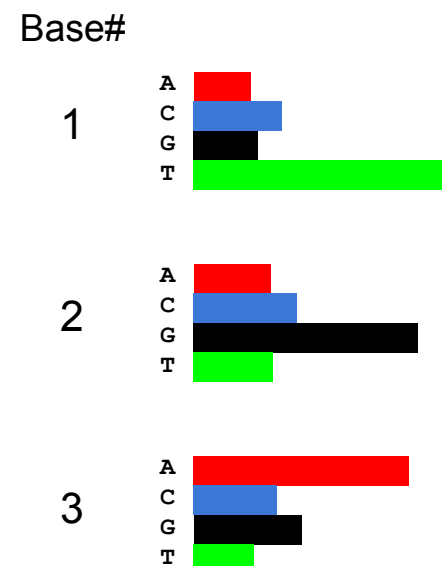
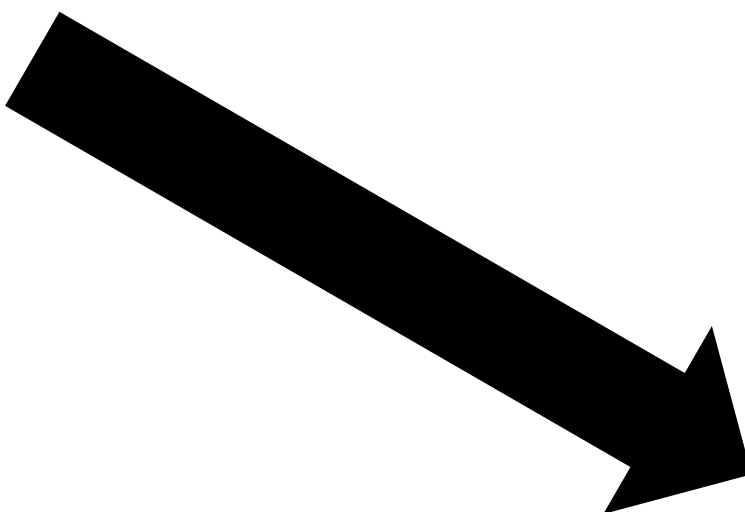
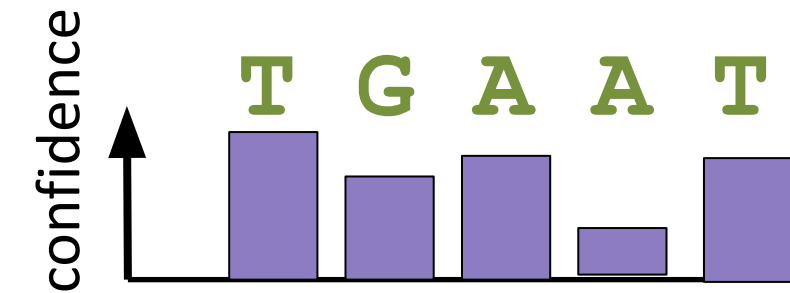
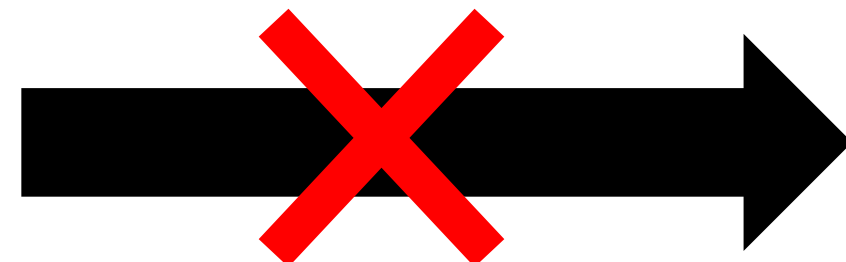
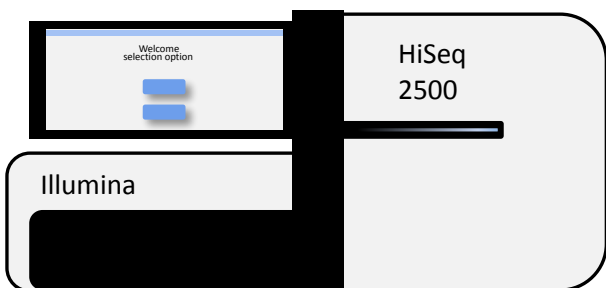
template



confidence



Key concept: basecalling

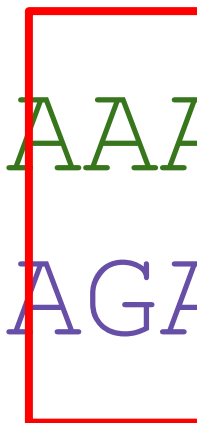


mismatch

template

AGCAATCTCAATTACAAAATATAACACCAACAAA

AGCAATCTCAATTACAGATATAACACCAACAAA



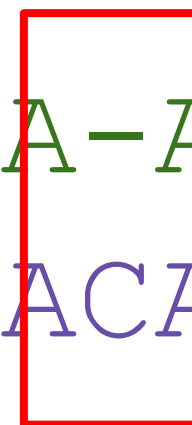
read

insert

template

AGCAATCTCAATTACA-AATATACACCAACAA

AGCAATCTCAATTACACAAATATACACCAACAA



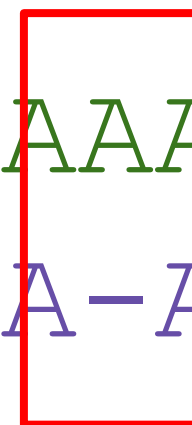
read

deletion

template

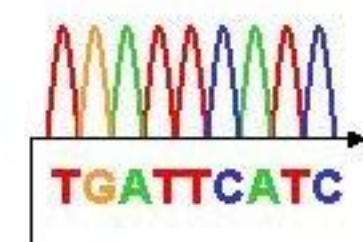
AGCAATCTCAATTACAAAATATAACACCAACAA

AGCAATCTCAATTACA-ATATAACACCAACAA



read

1977 1980 1983 1986 1989 1992 1995 1998 2001 2004 2007 2010 2013 2016 2019



Sanger



454



Illumina



SOLiD



Ion Torrent



PacBio

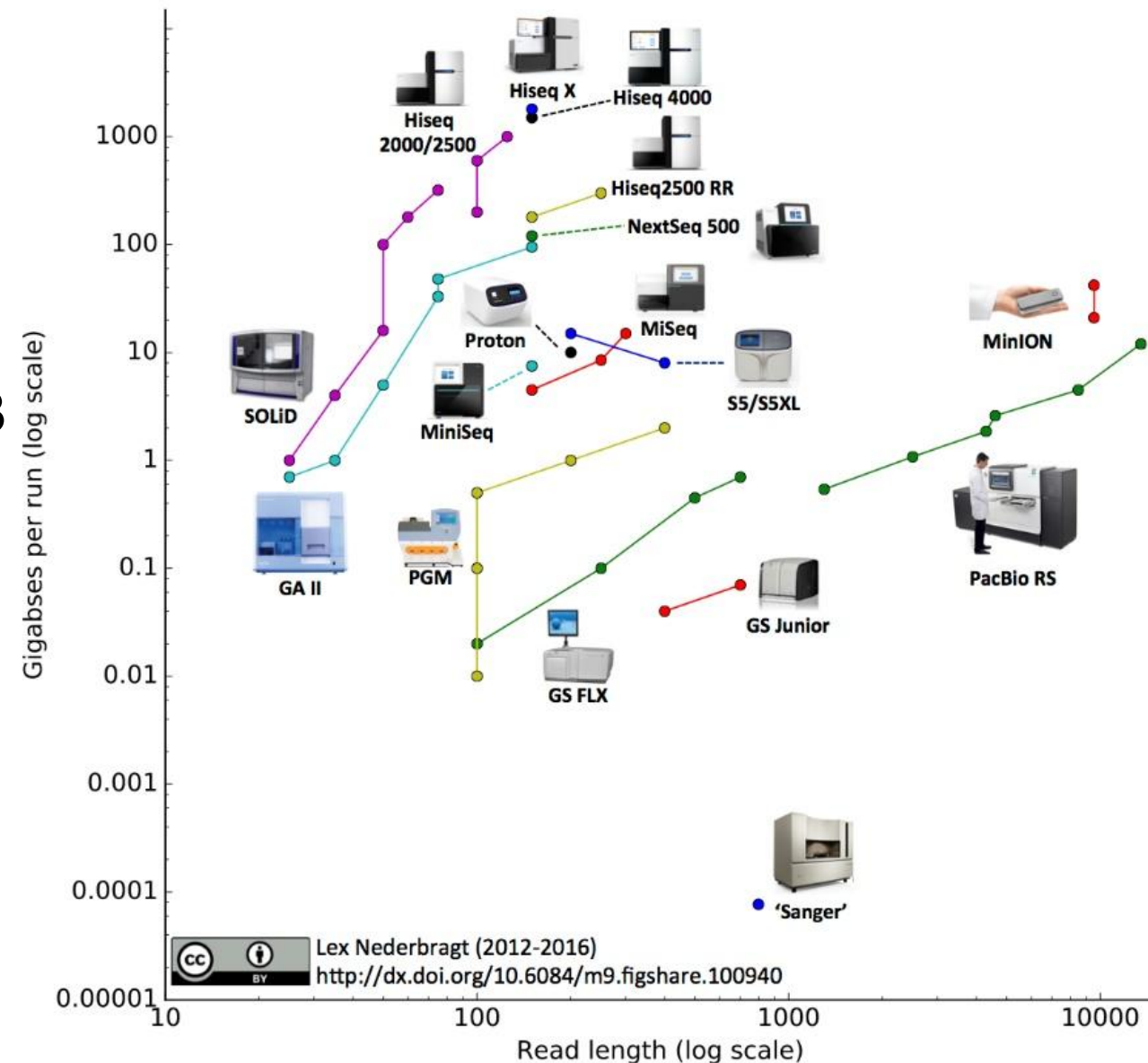


Oxford
Nanopore

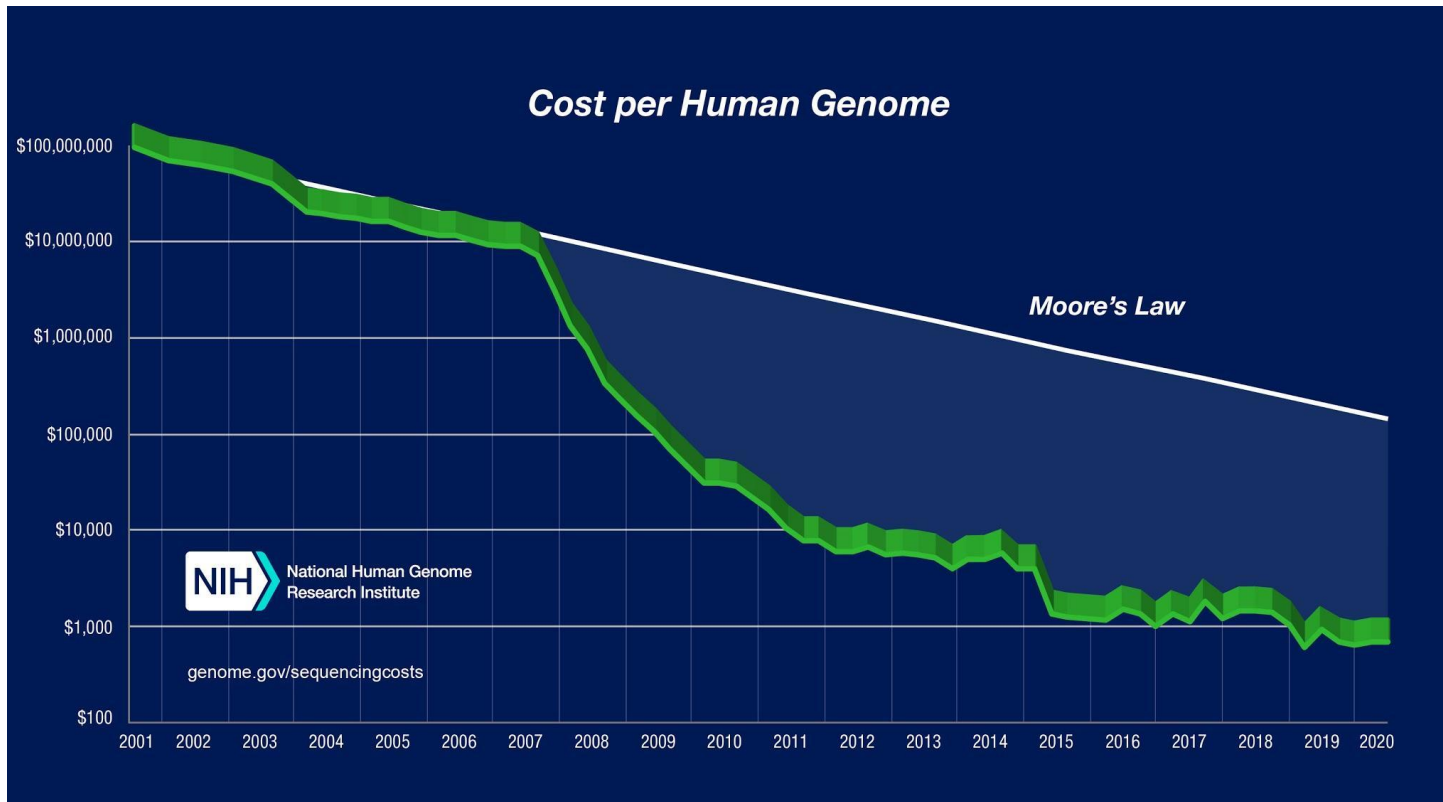
BGI

1st generation of NGS

- 454 Life Sciences.
 - Bought by Roche 2007.
- Illumina/BGI is currently cheapest per GB
- Long-read sequencing is revolutionizing assembly



Sequencing costs



- Computer speed and storage capacity is **doubling every 18 months** and this rate is steady
- DNA sequence data **is doubling faster than computer speeds!**

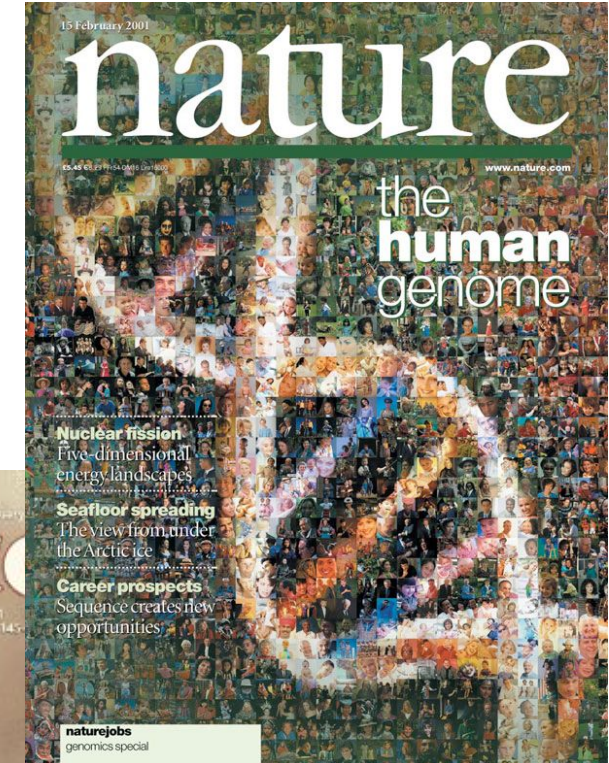
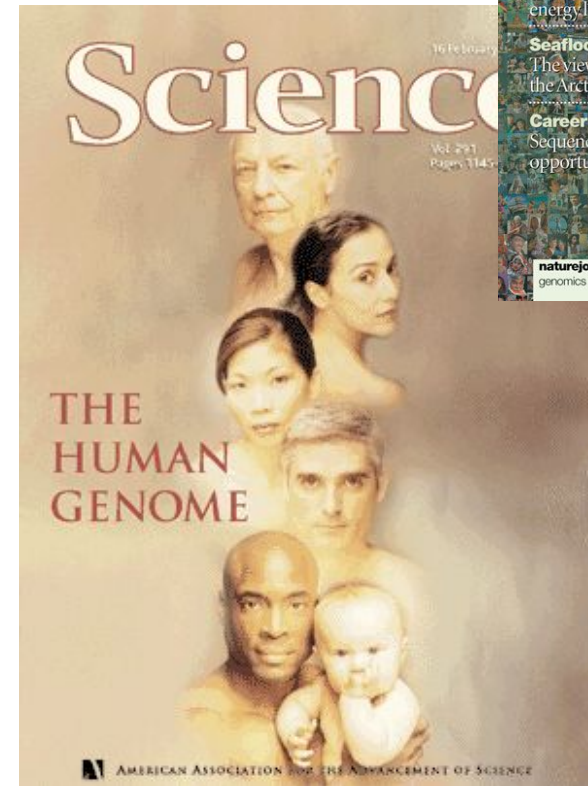


1990 - 2003
~5.54G USD (adj. for FY 2022)
20 research centers, 6 countries

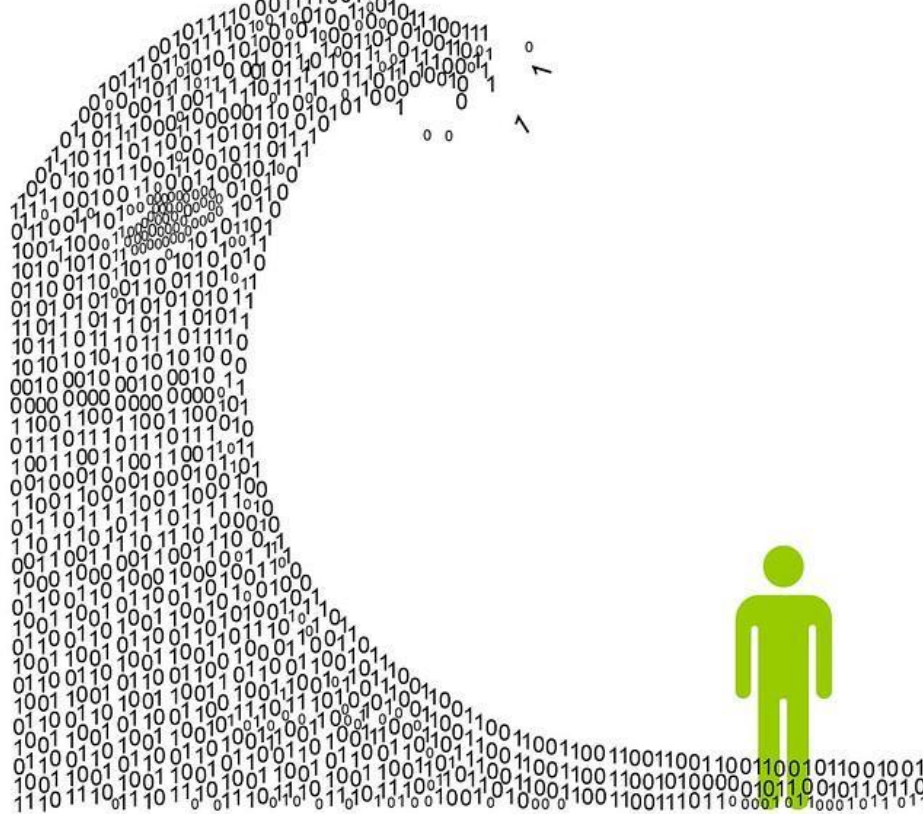
Picture: The Guardian

Human sequencing

- First draft genome of human in 2001, final 2004
- Estimated costs \$5.54 billion USD, time 13 years
- Today:
 - 1000-2000USD for one genome
 - A couple of days!



Storage and analysis

A large, stylized binary sequence is arranged in a shape that resembles a human figure. The sequence consists of binary digits (0s and 1s) and is oriented diagonally, creating a sense of depth and perspective.

- Cost of sequencing is almost less than the cost of **storage and analysis**
- One Illumina NovaSeq system: almost 10,000 human genomes per year!
- A standard human (30-40x) whole-genome sequencing exp. would create 30-150 Gb of data



Distributed data production

- Worldwide >900 centers
- >60 Pb pr year (2014)
- 20,000 Pb pr year (2025)
- Data transfer and storage becomes an issue

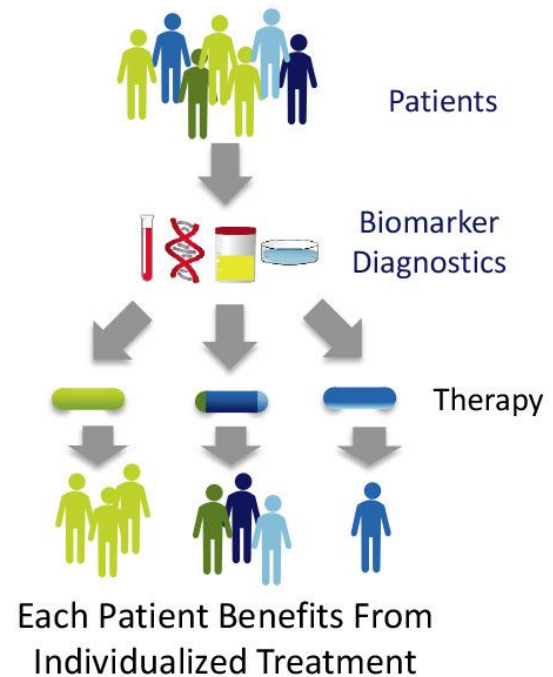


The X Genomes projects

- Human population projects
 - 1000 genomes project (2500 individuals)
 - Genomics England (100k individuals)
 - US Precision Medicine (1 million individuals)
- 100K pathogens project, Earth Microbiome project, Cancer genome project, Plants and animals, Insects,...

NGS in the clinic

- Diagnostics of patients (+ fetus)
- Focused treatment of cancer patients
- Sequencing of bacterial isolates
- Country-wide projects:
 - UK, US, UAE, Qatar, Finland, China, ...
 - DK: Danish regions want to sequence 100k individuals



Personalized medicine

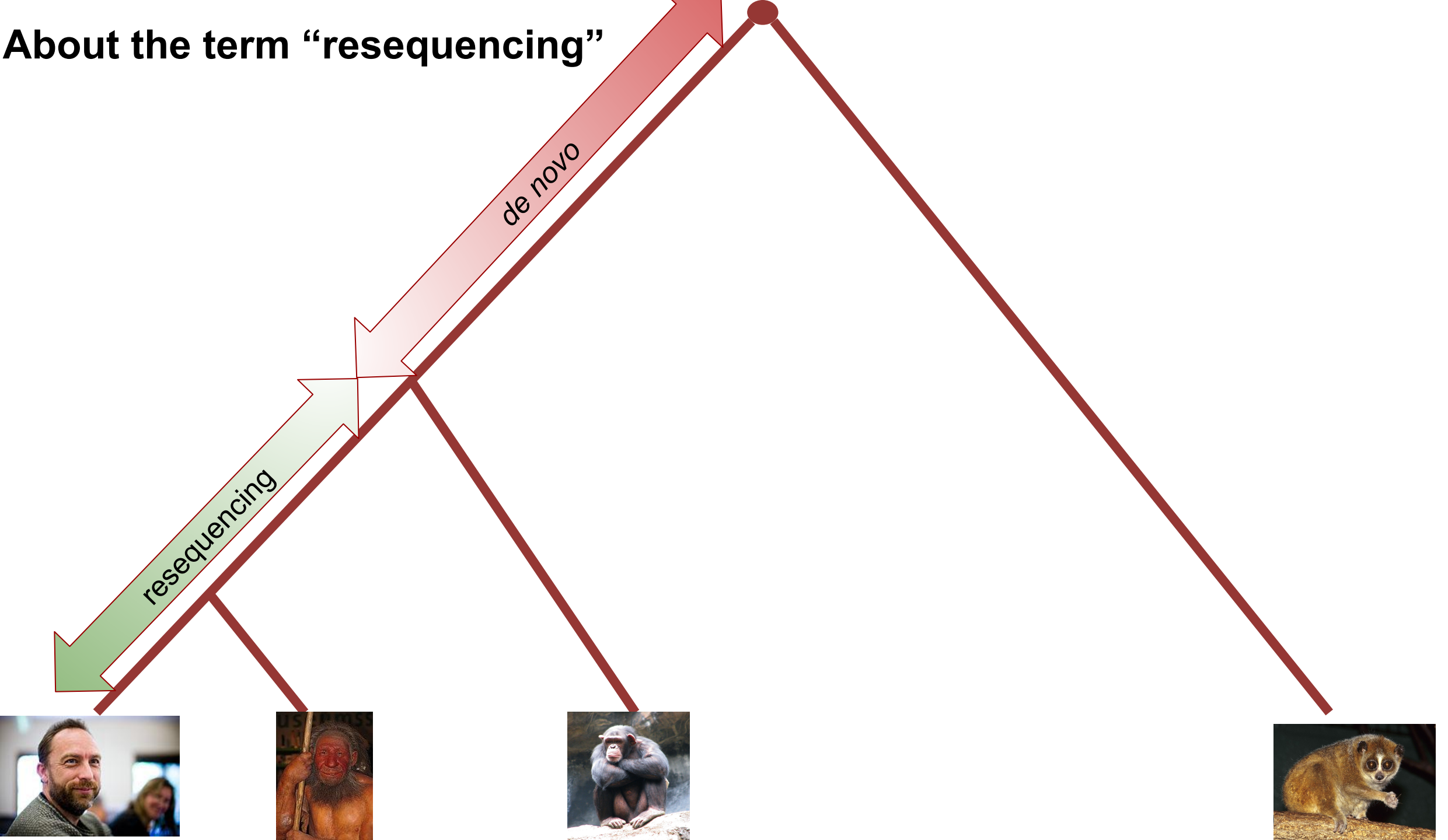


- Giving the same medication to all will not work
- Personalized medicine initiative in DK
- Sequence 100,000 patients on hospitals
- Use extensive registry data
- Current: 100M DKK (estimated 2G DKK)

NGS & bioinformatics

- Extreme data size causes problems
- Just transferring and storing the data
- Cannot do all against all comparisons (N^*N)
- Think in fast and parallel programs
- Cloud computing increasingly used

About the term “resequencing”



Whole genome sequencing

