



Swiss Institute of
Bioinformatics

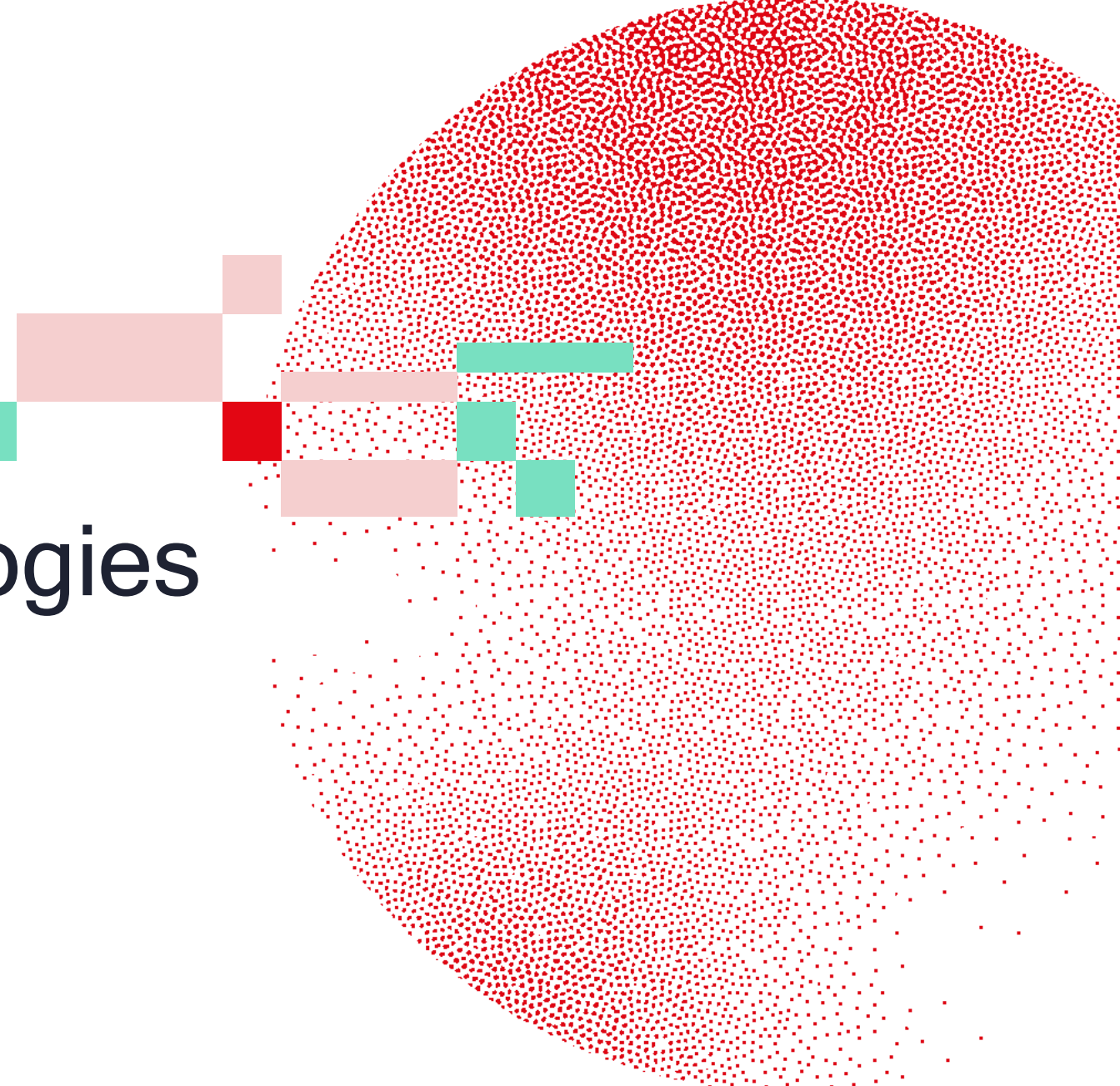
INTRODUCTION TO SEQUENCING DATA ANALYSIS

Sequencing Technologies

Deepak Tanwar
Geert van Geest

November 19-21, 2025

Adapted from previous year courses



Learning objectives

Understand the principles behind major DNA/RNA sequencing technologies

Identify applications of each sequencing technology in research

Evaluate technology limitations, especially those affecting read length and sequencing accuracy

Select appropriate sequencing methods for different genomic and transcriptomic analyses based on the experimental need

What is sequencing?

Quiz: 4

What is the primary function of sequencing technologies?

Sequencing technology refers to the various methods used to determine the order of nucleotides (the building blocks of DNA and RNA) in a strand of genetic material.

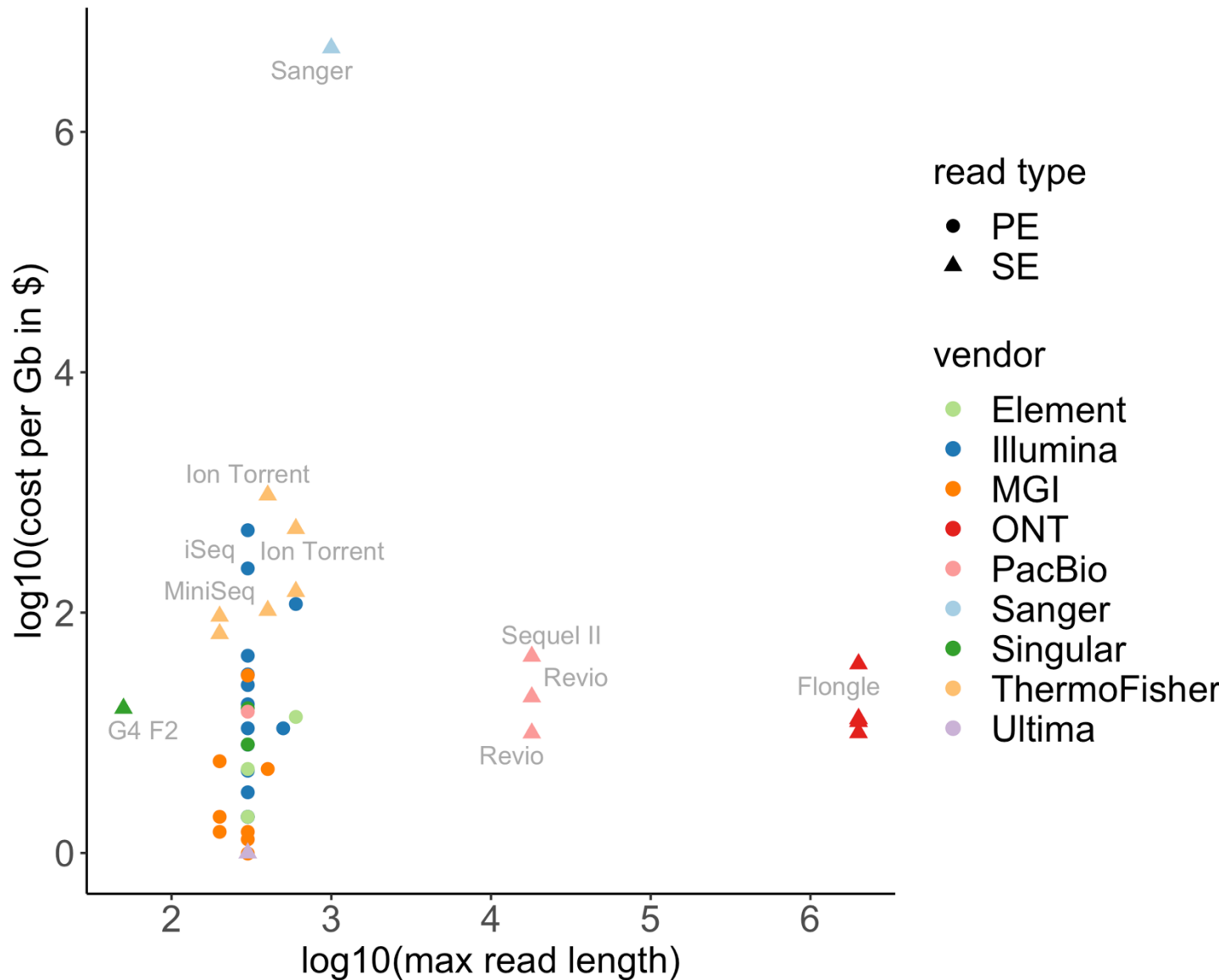
Sequencing technologies

Sanger sequencing

Second generation sequencing

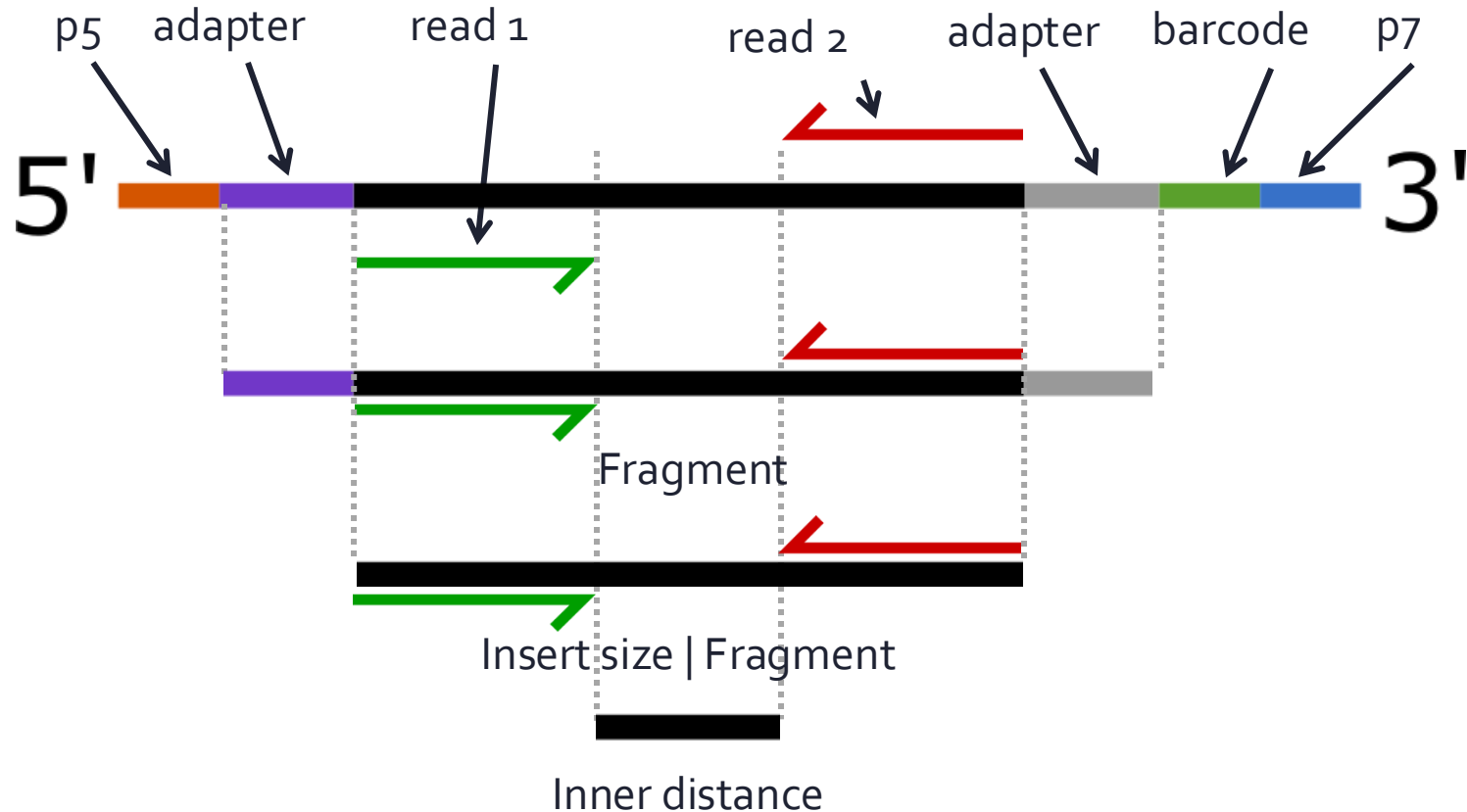
Third generation sequencing

Comparison of Sequencing Technologies by Cost and Read Length



drawn from:
<https://docs.google.com/spreadsheets/d/1GMMfhylK0-q8Xklo3YxIWaZA5vVMuhU1kg41g4xLkXc/> Albert
Vilella

Some definitions



Some more definitions..

Library: fragments from one (c)DNA sample that share a barcode

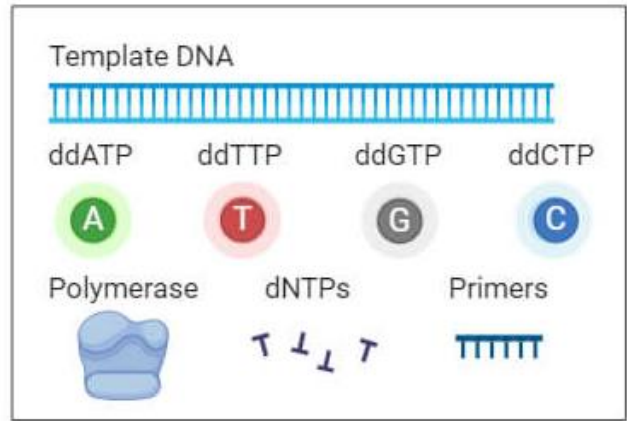
Sequencing run: complete cycle of generating reads on a machine

Flow cell: physical platform where sequencing reactions take place. Used once in a sequencing run.

Lane: compartment within the flow cell. An Illumina flow cell often has multiple lanes (2 or 4)

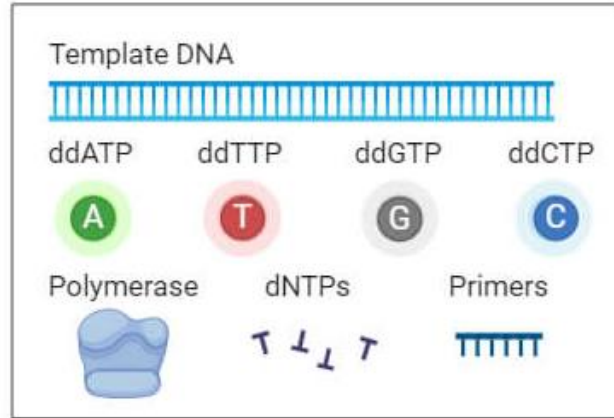
Sanger sequencing

Reagents

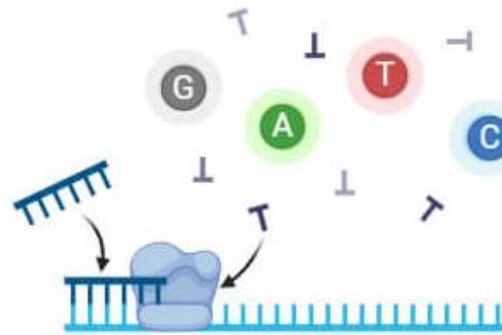


Sanger sequencing

Reagents

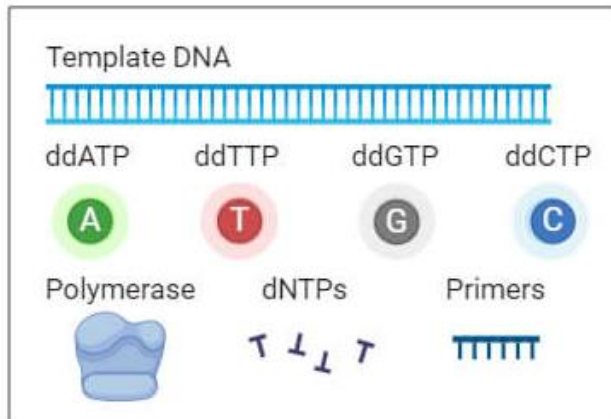


① Primer annealing and chain extension

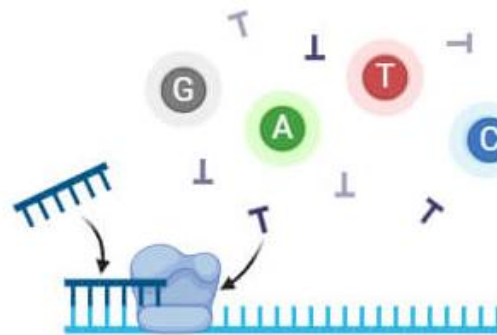


Sanger sequencing

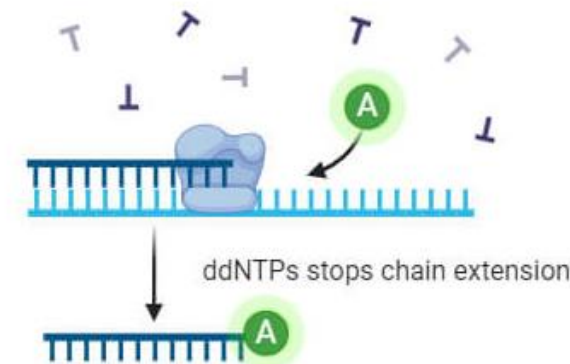
Reagents



① Primer annealing and chain extension

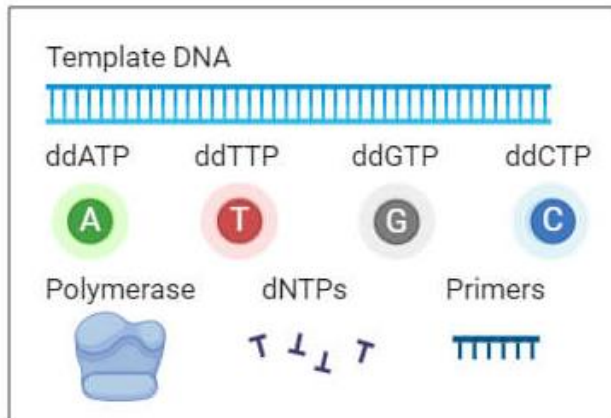


② ddNTP binding and chain termination

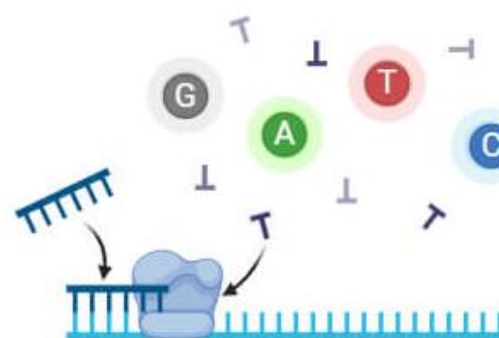


Sanger sequencing

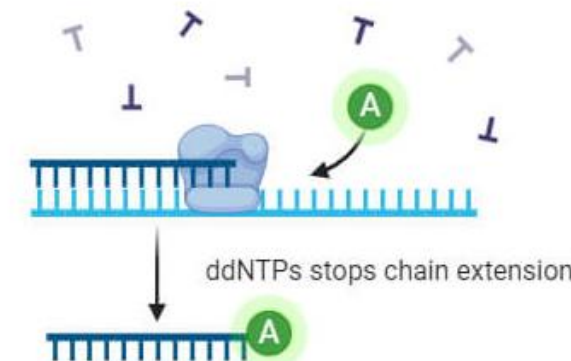
Reagents



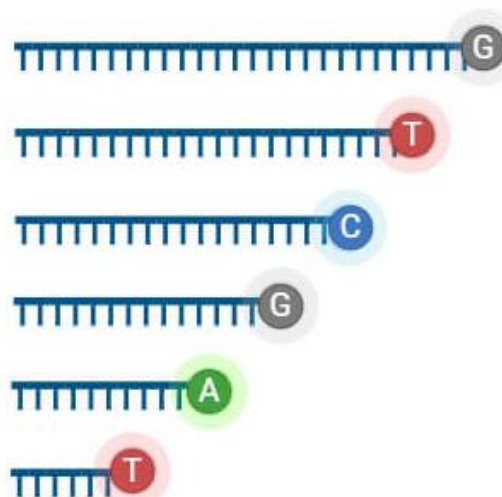
① Primer annealing and chain extension



② ddNTP binding and chain termination

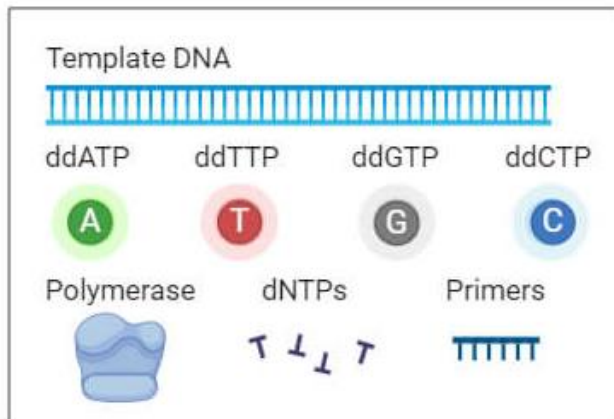


③ Fluorescently labelled DNA sample

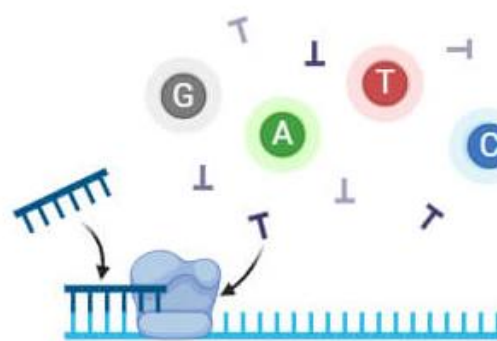


Sanger sequencing

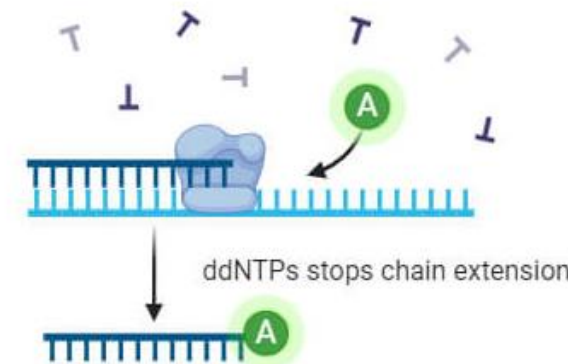
Reagents



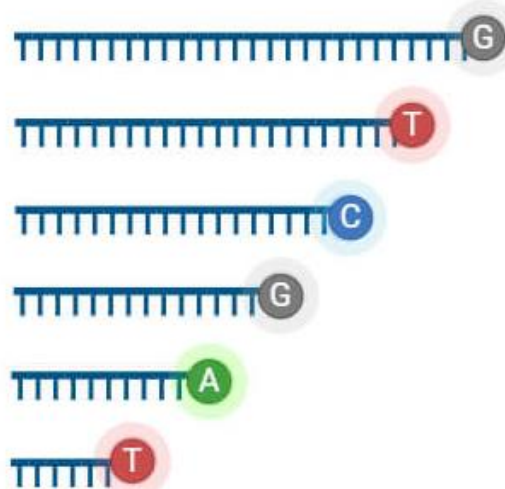
① Primer annealing and chain extension



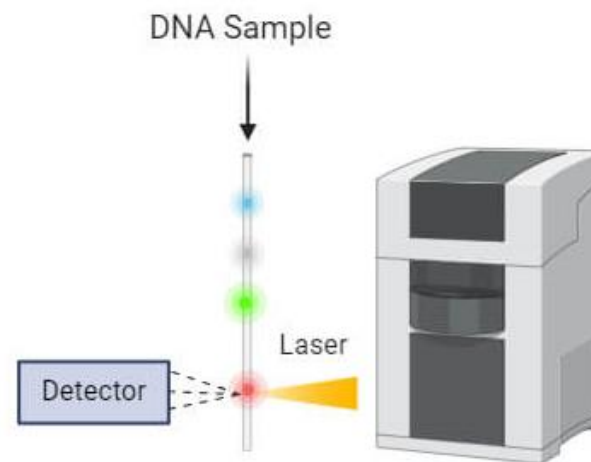
② ddNTP binding and chain termination



③ Fluorescently labelled DNA sample

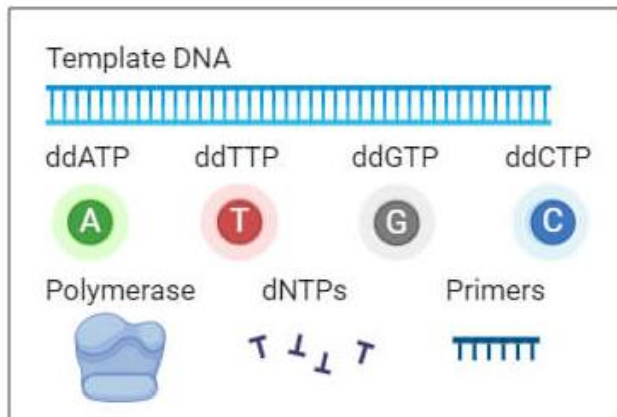


④ Capillary gel electrophoresis and fluorescence detection

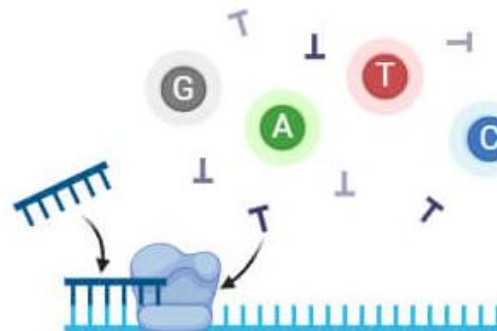


Sanger sequencing

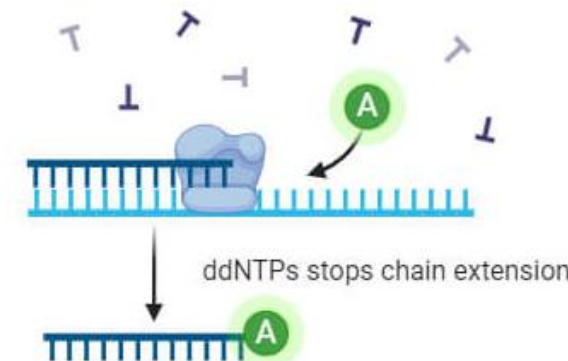
Reagents



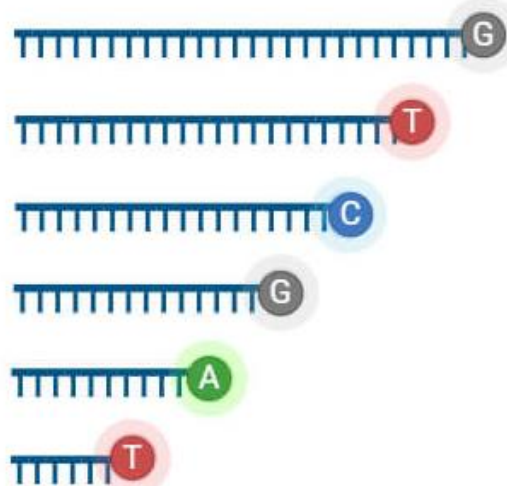
① Primer annealing and chain extension



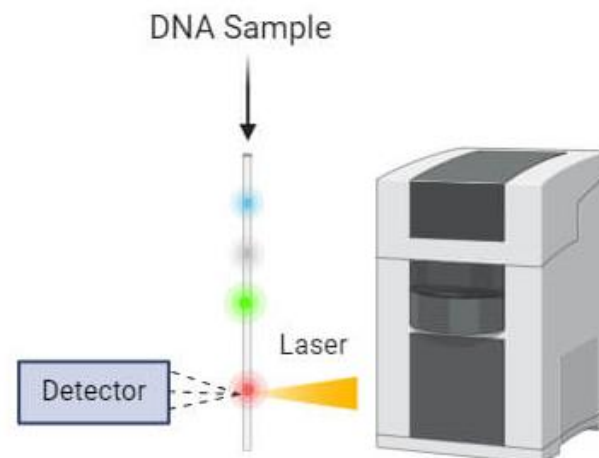
② ddNTP binding and chain termination



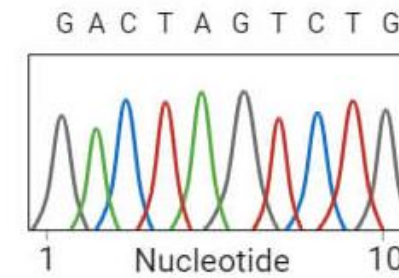
③ Fluorescently labelled DNA sample



④ Capillary gel electrophoresis and fluorescence detection



⑤ Sequence analysis and reconstruction



Sanger sequencing applications

Clinical Applications:

Genetic disease diagnosis (identifying mutations in specific genes)

Research Applications:

Validation of next-generation sequencing results

Forensic Applications:

DNA profiling for identification

Sanger sequencing applications

Clinical Applications:

Genetic disease diagnosis (identifying mutations in specific genes)

Research Applications:

Validation of next-generation sequencing results

Forensic Applications:

DNA profiling for identification

We are not covering Sanger sequencing
in this course

Second generation sequencing

454 Pyrosequencing

- Discontinued due to technological advancements

Ion Torrent (semiconductor sequencing)

- This technology is faster and can be more cost-effective, but it generally has shorter read lengths and slightly lower accuracy compared to Illumina
- Up to ± 400 bp read length
- Homopolymers, such as TTTTT are impossible to sequence

Illumina (sequencing by synthesis)

Illumina sequencing

Massive throughput: up to 16×10^{12} bases/run (NovaSeq X) = $\sim 9,000$ whole exomes

50 - 300 bp

Paired-end (or single-end)

Multiplexing

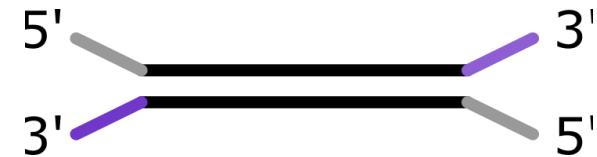


Illumina library prep

shear + size select DNA



Ligate adapters



Barcode + p5/p7 sites

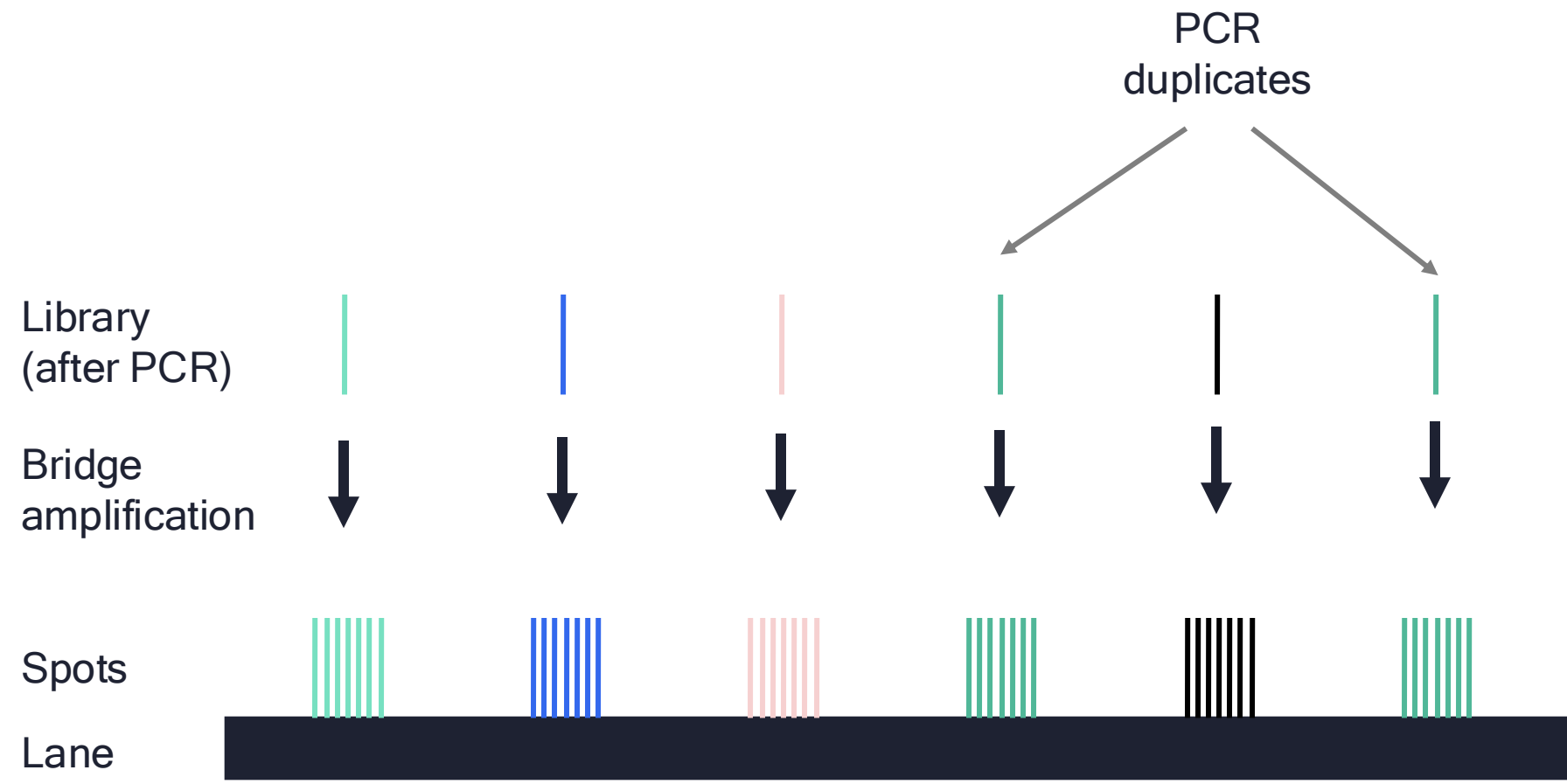


PCR: 8-16 cycles



Sequencing

Illumina Sequencing



Each spot represents one read pair

Illumina Sequencing by synthesis

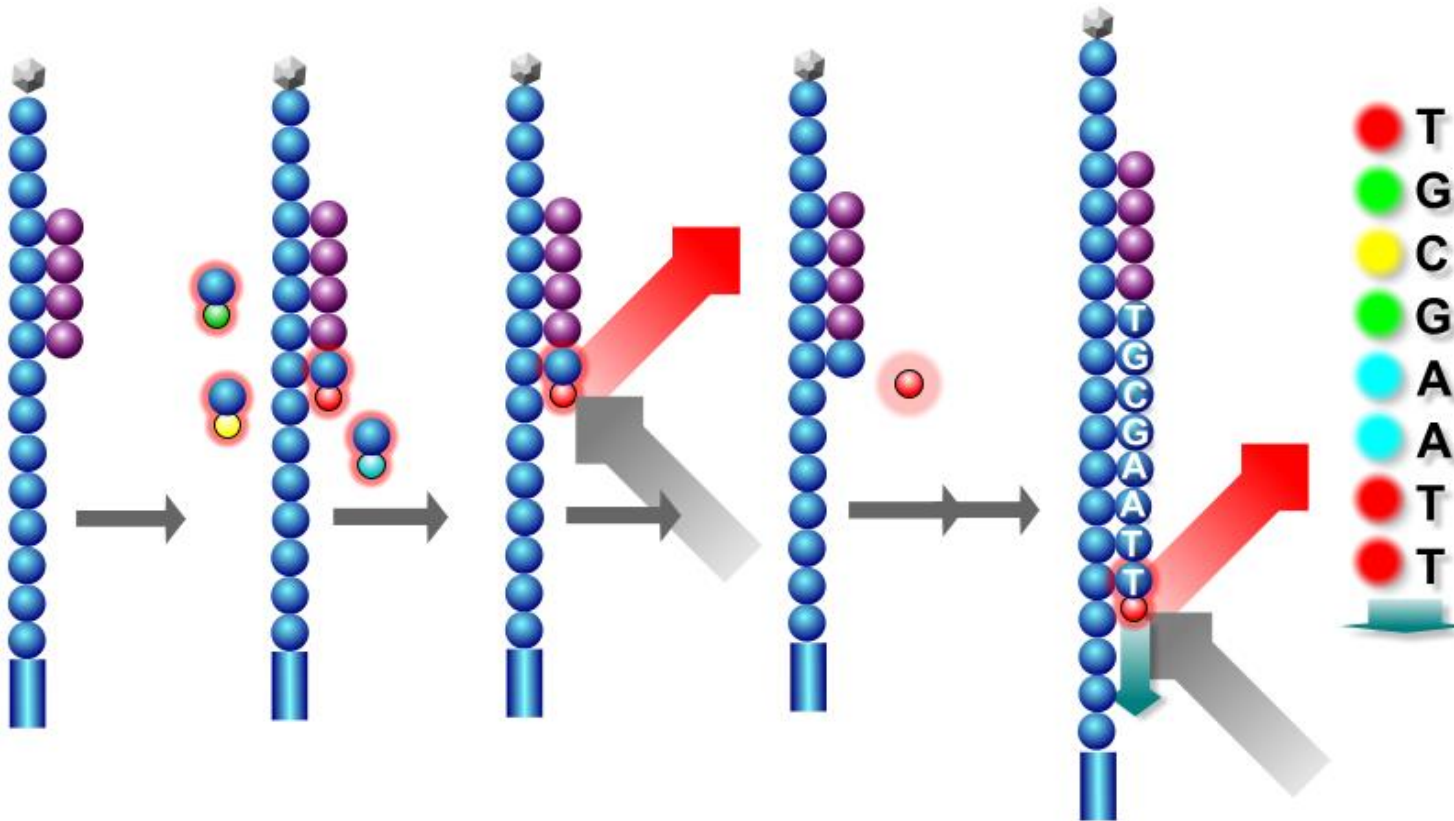


Image: Abizar Lakdawalla CC BY-SA 3.0

What are you using Illumina for?

Second generation sequencing applications

Transcriptome characterization

e.g. RNA-seq - Gene expression, splicing, isoform detection

Second generation sequencing applications

Epigenome characterization

e.g. *ATAC-seq*, *Bisulfite-seq* - Chromatin accessibility, DNA methylation

Second generation sequencing applications

DNA-protein interactions

e.g. *ChIP-seq* - Transcription factor binding, histone modifications

Second generation sequencing applications

Whole genome sequencing & assembly

e.g. short- and long-read WGS - De novo genome assembly, reference genome improvement

Second generation sequencing applications

Variant detection

e.g. *Exome-seq*, *WGS* - SNPs, indels, CNVs for disease association and diagnosis

Second generation sequencing applications

Metagenome characterization

e.g. 16S rRNA sequencing, shotgun metagenomics - Microbiome studies, environmental genomics

Second generation sequencing applications

Targeted sequencing

e.g. Amplicon-seq, hybrid capture panels - Focused gene panels for diagnostics

Second generation sequencing applications

Single-cell sequencing

e.g. *scRNA-seq*, *scATAC-seq* - Cell heterogeneity, developmental lineages, immune profiling

Second generation sequencing applications

Spatial transcriptomics

e.g. 10x Visium, Slide-seq - Gene expression with spatial resolution in tissues

Second generation sequencing applications

Single-cell epigenomics

e.g. scATAC-seq, scChIP-seq, scMethyl-seq - Chromatin accessibility, histone marks, methylation at single-cell level

Second generation sequencing applications

Multi-omics at single-cell level

e.g. *SHARE-seq*, *10x Multiome (RNA + ATAC)*, *CITE-seq (RNA + protein)*

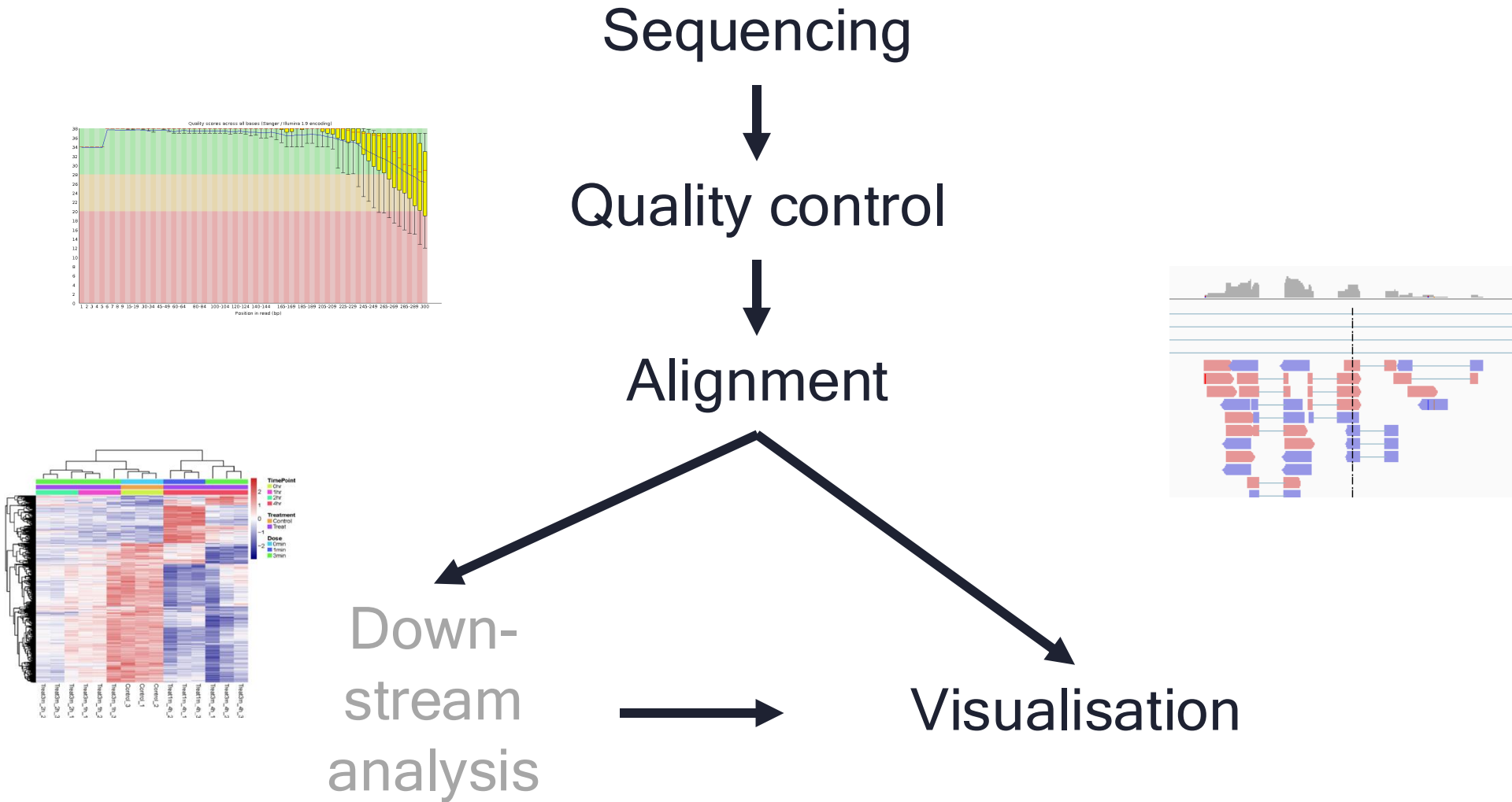
Applications:

Linking transcriptome with epigenome or proteome

Understanding gene regulation networks

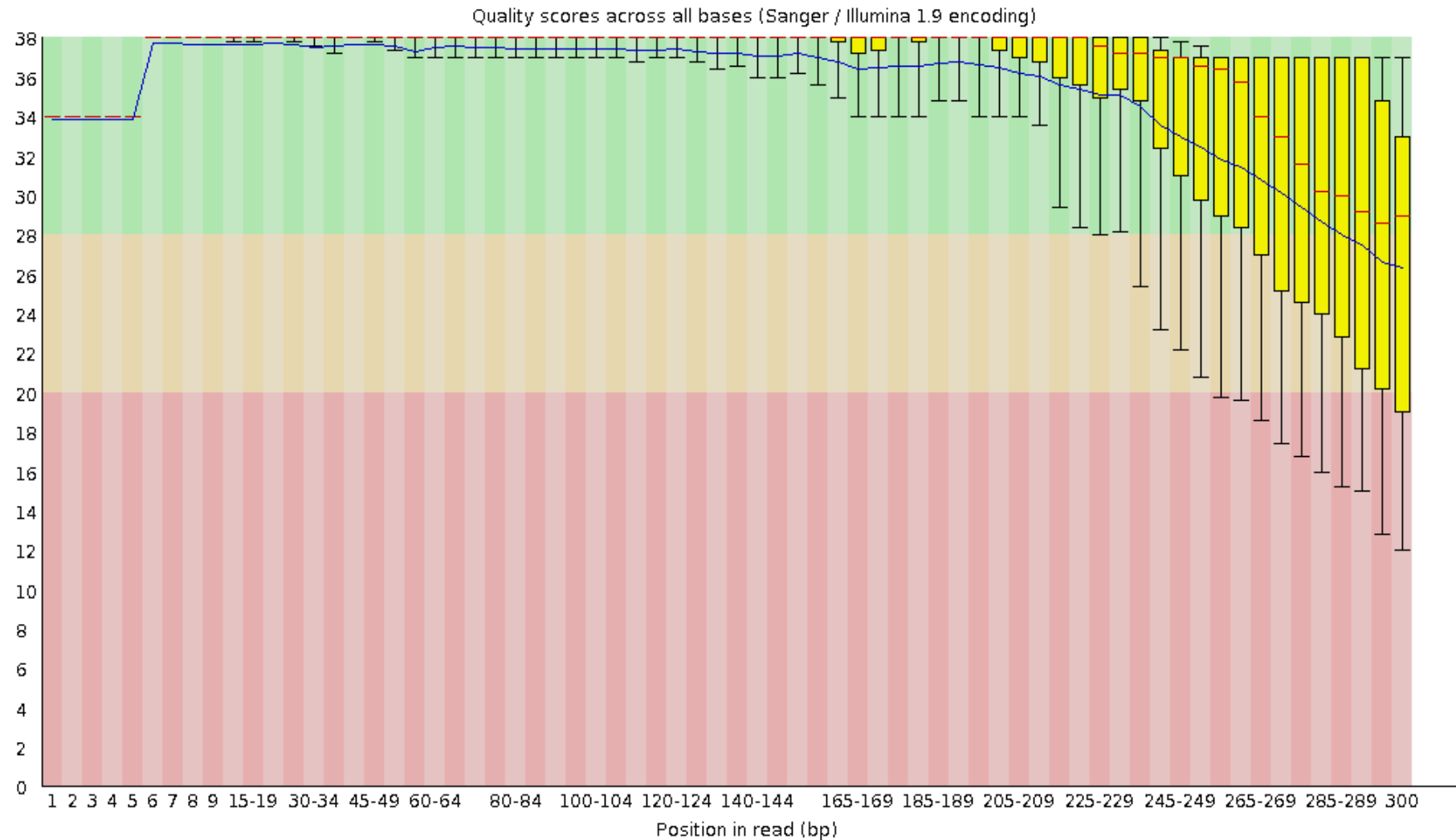
Immune and tumor microenvironment studies

Simple workflow of data analysis

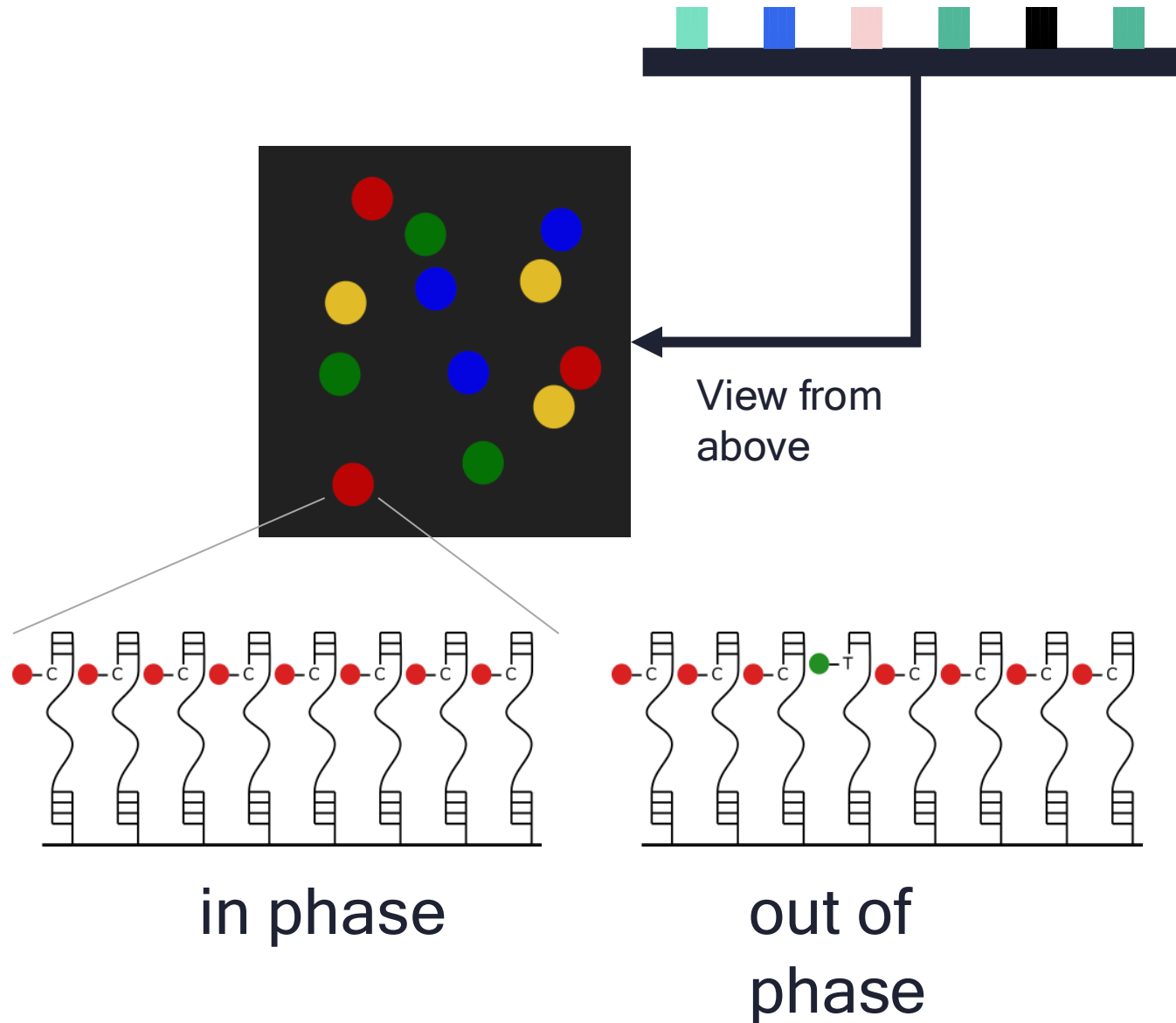


Illumina - limitations

Sequence quality declines towards the end



Phase Sequencing



Illumina - limitations

Maximum read length: 300 bp

Read length is limited by out-of-phase signal

How to reconstruct:

- » Repeats?
- » Isoforms?
- » Structural variation?
- » Genomes?

Quiz: 5

What is a common limitation of Illumina sequencing?

Third generation sequencing: Long reads

Crux: maximizing signal from a single-molecule base read-out

Single molecule, so no out-of-phase signal

Two frequently used platforms:

- » Oxford Nanopore Technology
- » PacBio SMRT sequencing

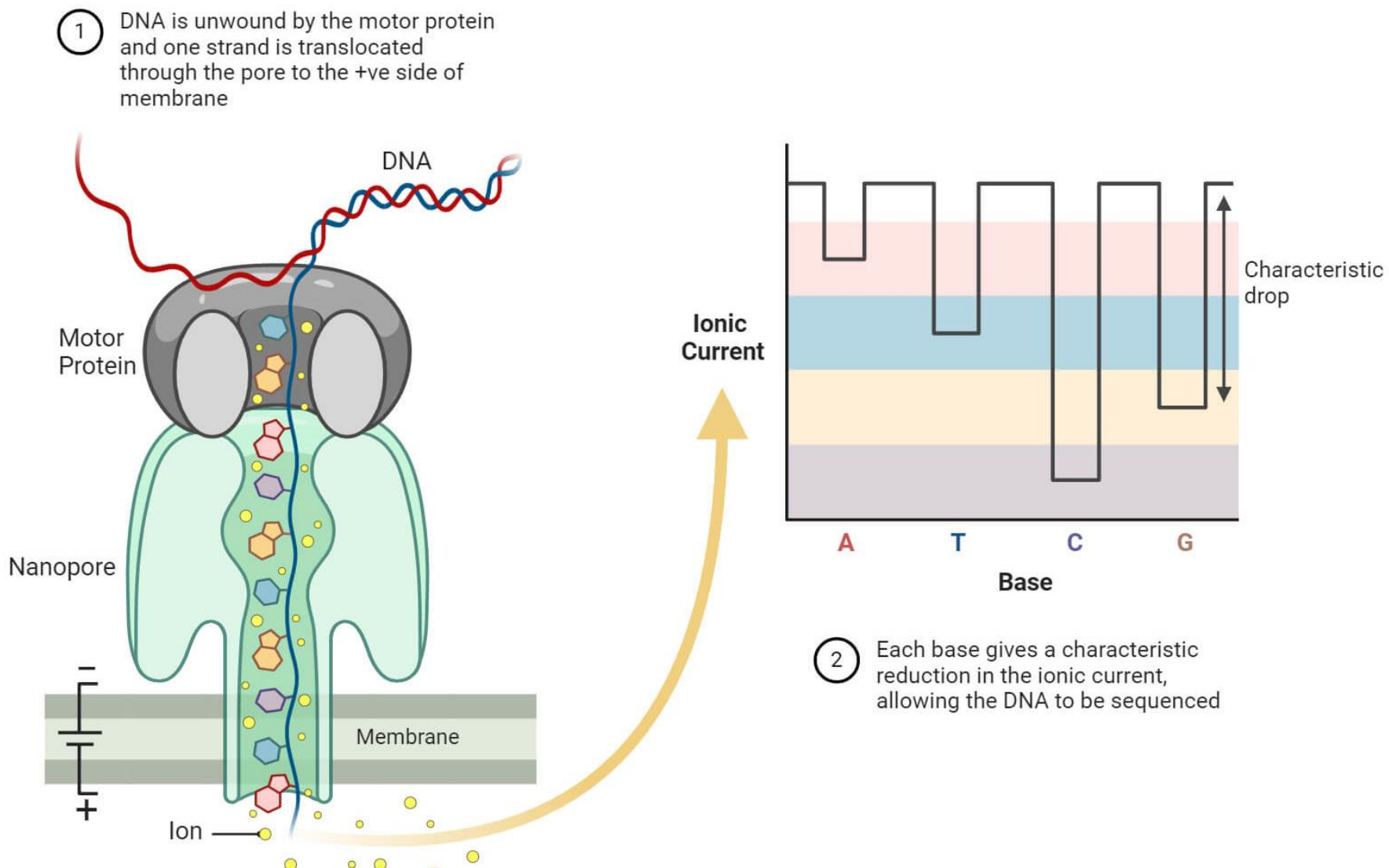
Oxford Nanopore technology

Based on changes in electrical current

Well-known for its scalability and portability

~95-97% accuracy

Oxford Nanopore technology principle



- ② Each base gives a characteristic reduction in the ionic current, allowing the DNA to be sequenced

Oxford Nanopore technology sequencers



MinION



GridION



Flongle



PromethION

PacBio sequencing: Single Molecule, Real-Time (SMRT)

Technology: Single Molecule, Real-Time (SMRT) sequencing.

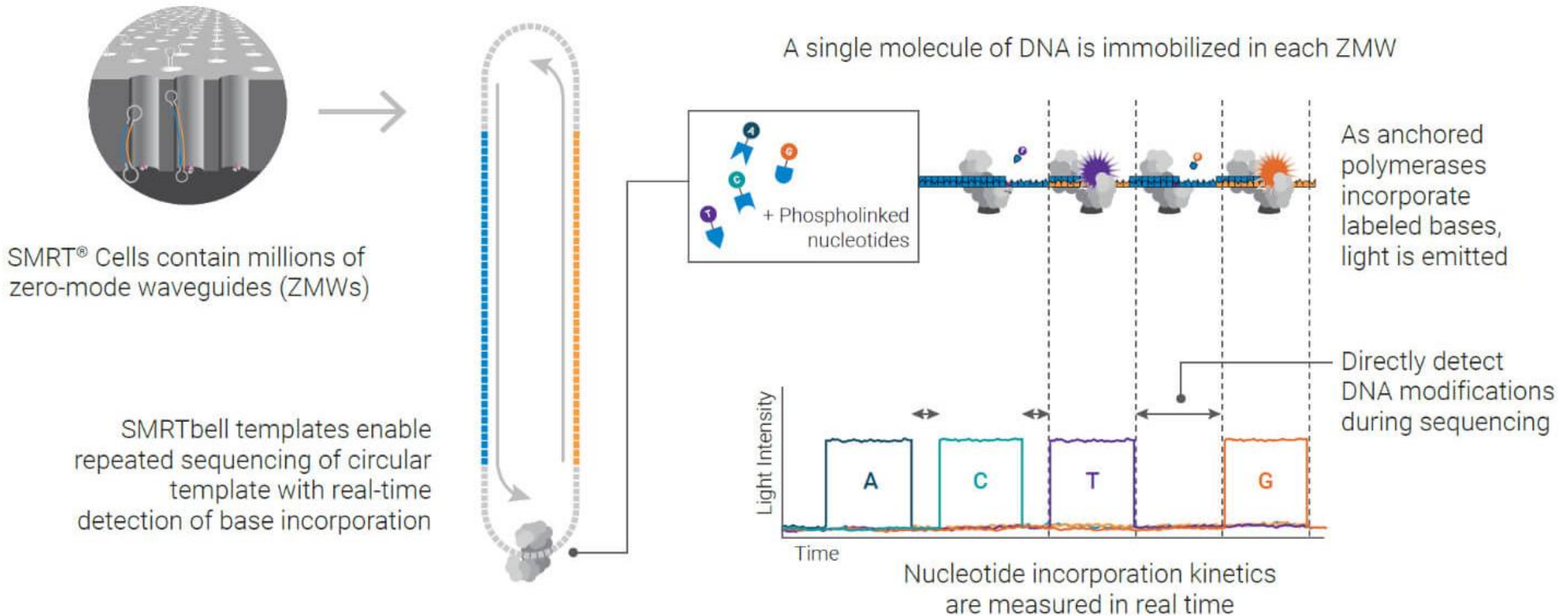
Process: Uses zero-mode waveguides (ZMWs) to observe single DNA molecules in real-time.

Accuracy: High accuracy due to real-time detection of nucleotide incorporation.

De Novo Genome Assembly: Ideal for assembling complex genomes, including those with repetitive regions

Epigenetic Studies: DNAm

PacBio sequencing: SMRT



PacBio sequencing: Circular Consensus Sequencing (CCS)

Technology: Uses SMRTbell libraries for circularized DNA.

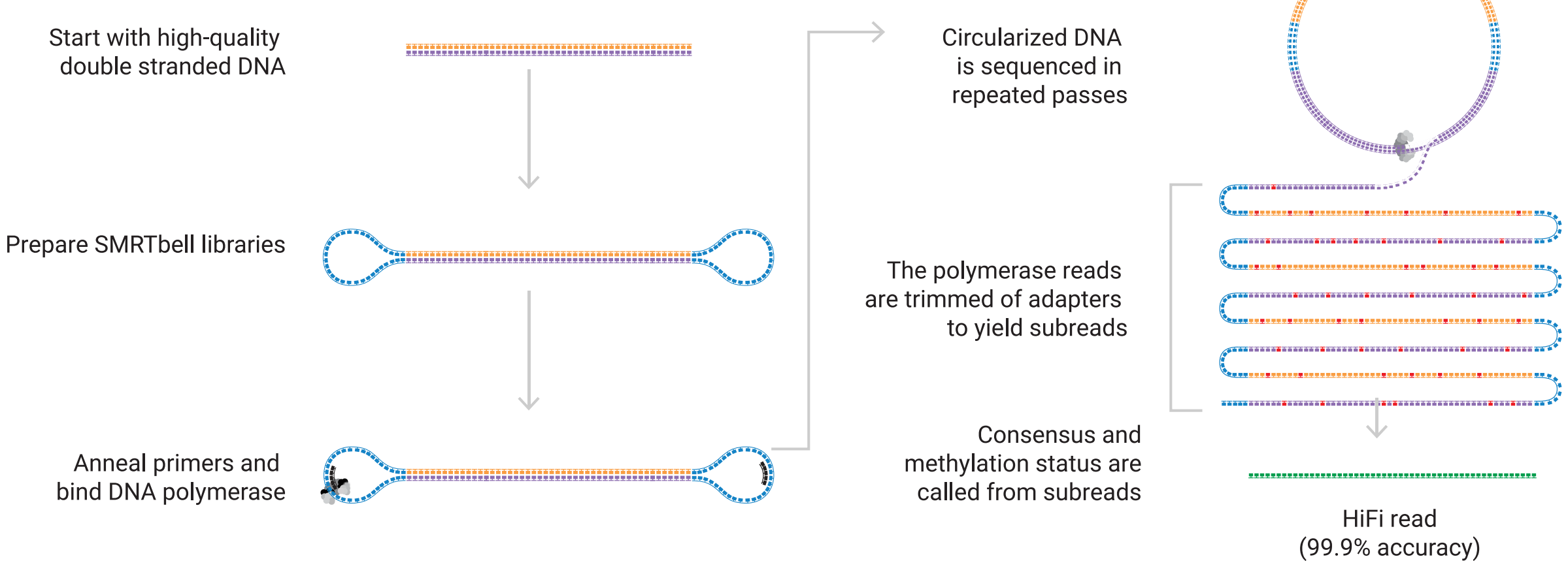
Process: DNA is sequenced in repeated passes, generating long reads.

Output: HiFi reads with 99.9% accuracy, ideal for detailed genomic studies.

High-Accuracy Genome Assembly: Produces highly accurate long reads (HiFi reads) for assembling complex genomes

Rare Disease Research: Helps in identifying genetic variants associated with rare diseases

PacBio sequencing: CCS



Quiz: 6 - 7

Which sequencing method uses changes in electrical current to identify bases?

...

What feature makes PacBio's HiFi reads highly accurate?

How to choose your sequencing method?

Read length
Accuracy
Availability
Costs
Throughput

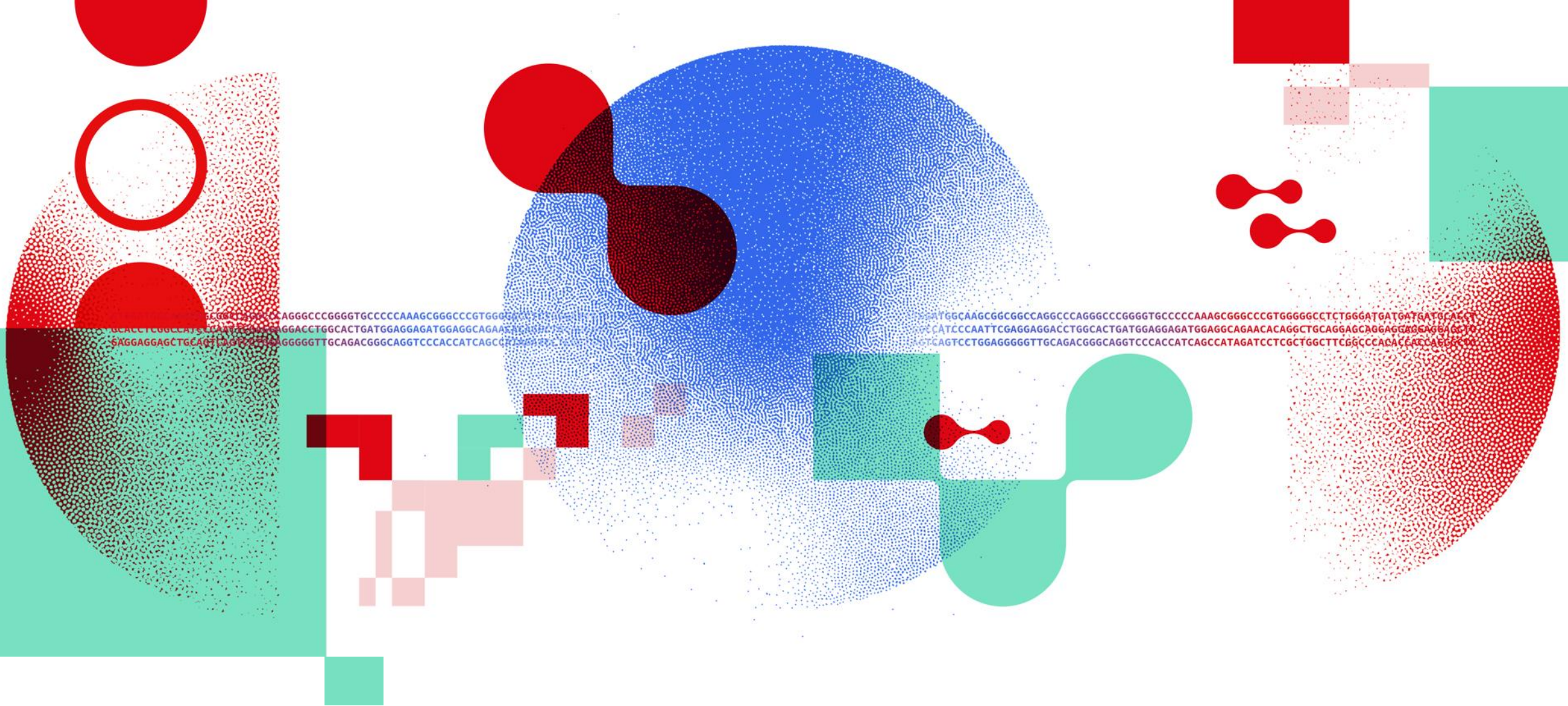


Summary

Sanger sequencing: A legacy method used for mutation identification and validation, especially in clinical and forensic applications.

Second generation (Next-Gen) sequencing: Focused on Illumina (most widely used due to high throughput and cost-efficiency).

Third generation sequencing: Includes Oxford Nanopore (portable, moderate accuracy) and PacBio SMRT (very high accuracy with HiFi reads).



Thank you

DATA SCIENTISTS FOR LIFE

sib.swiss

