## NGS - variant analysis

Sequencing and alignment

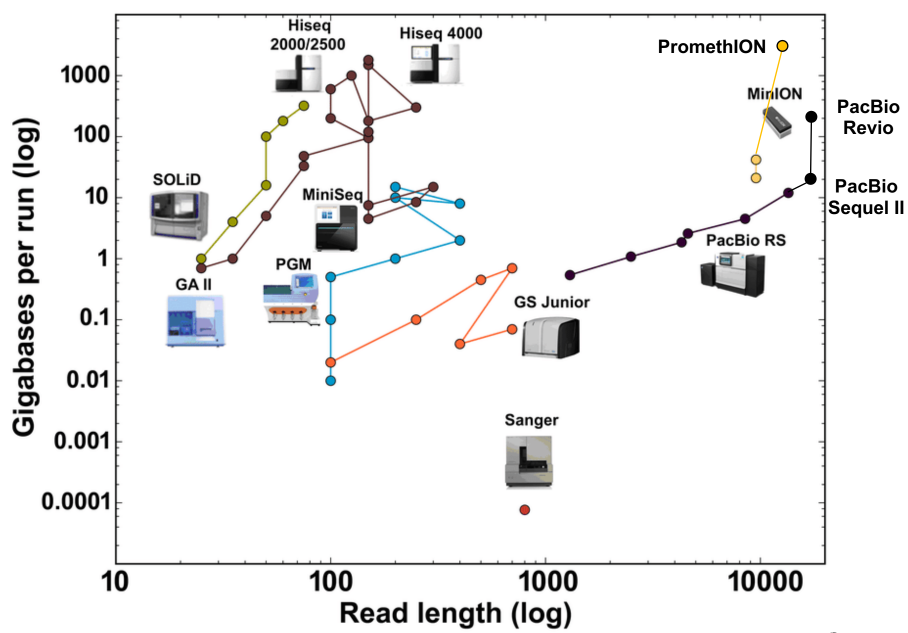


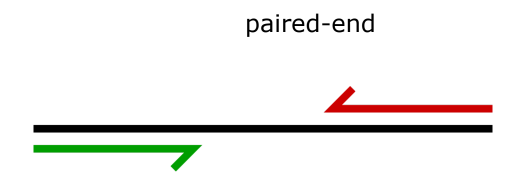
Image from: G. Silva (2016)

#### Illumina sequencing

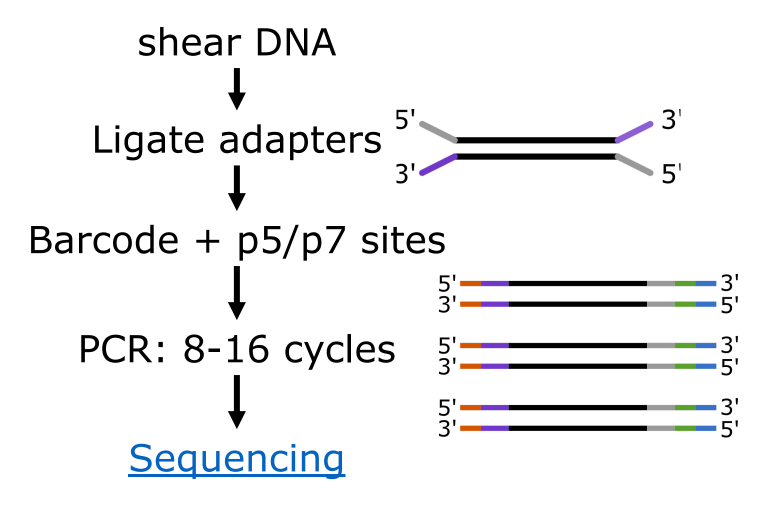
- Sequencing-by-synthesis: 2nd generation sequencing
- Massive throughput: up to 500x10<sup>9</sup> bases/run
- Most used platform today

## Illumina sequencing

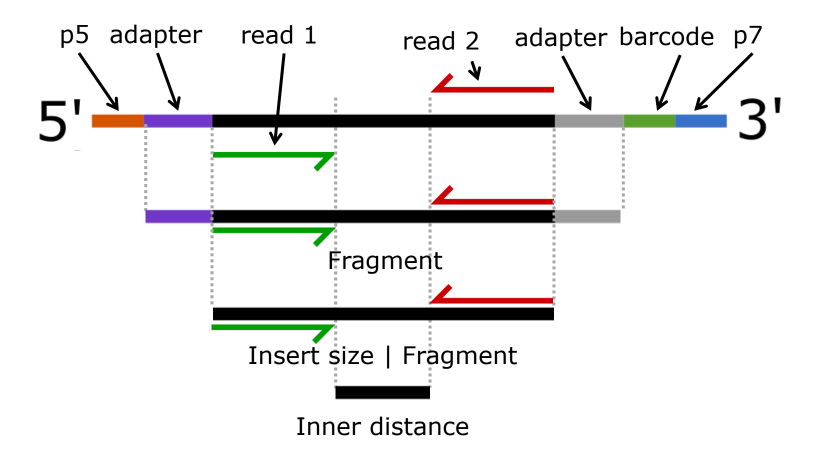
- 50 300 bp
- Paired-end (or single-end)



## Illumina libray prep

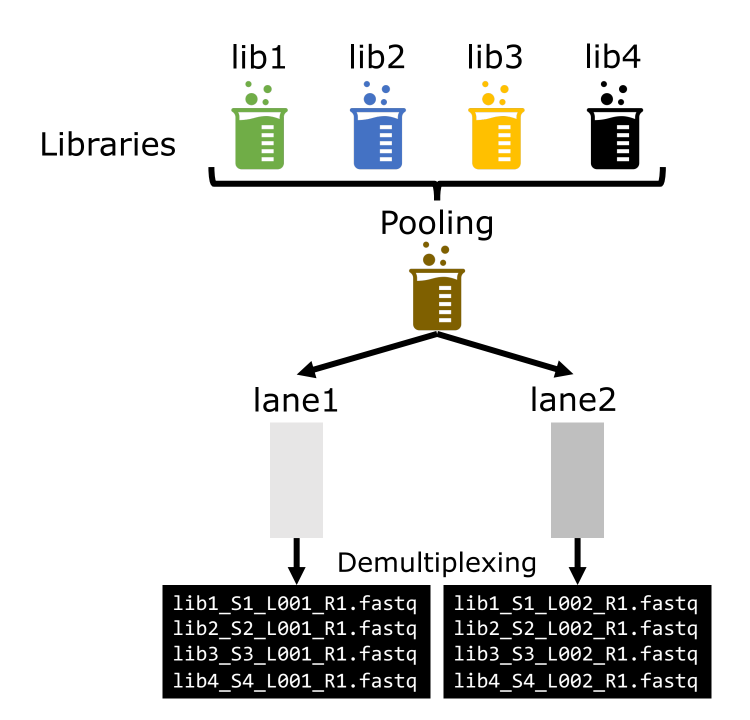


#### Some definitions



#### 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)

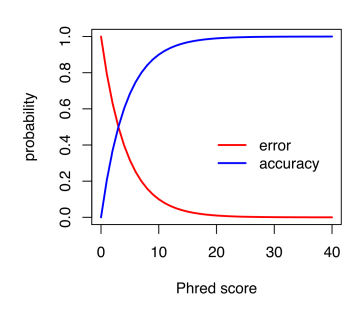


## fastq

fasta + basequality (fasta + q = fastq)

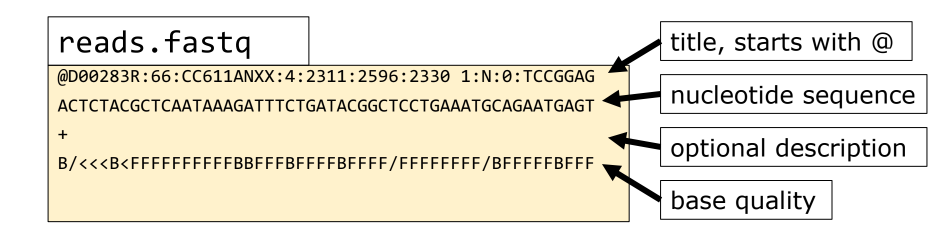
 $BASEQ = -10log_{10} \Pr\{base \ is \ wrong\}$ 

$$-10log_{10} (0.01) = 20$$
  
-10log<sub>10</sub> (0.1) = 10  
-10log<sub>10</sub> (0.5) = 3

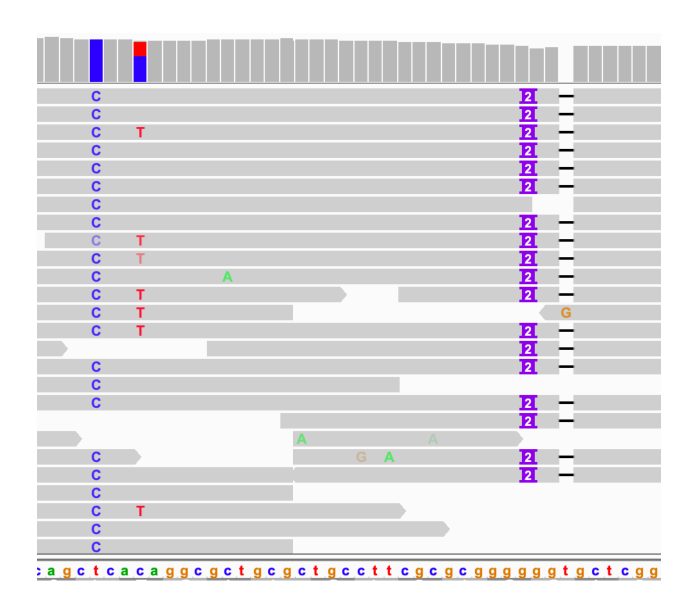


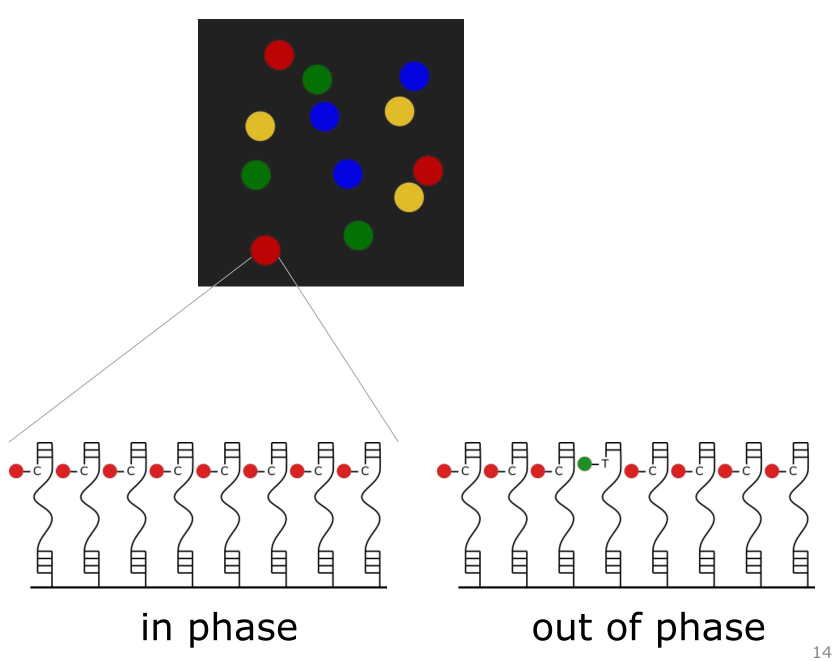
# Question

## fastq files



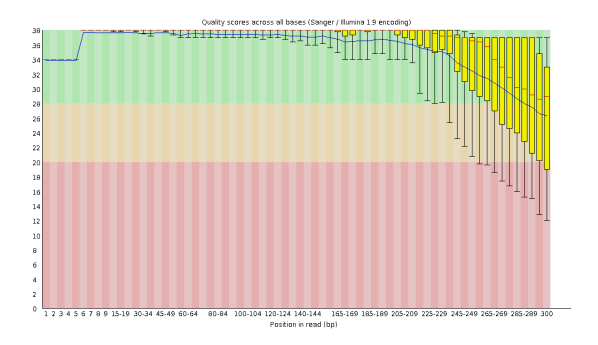
# Question





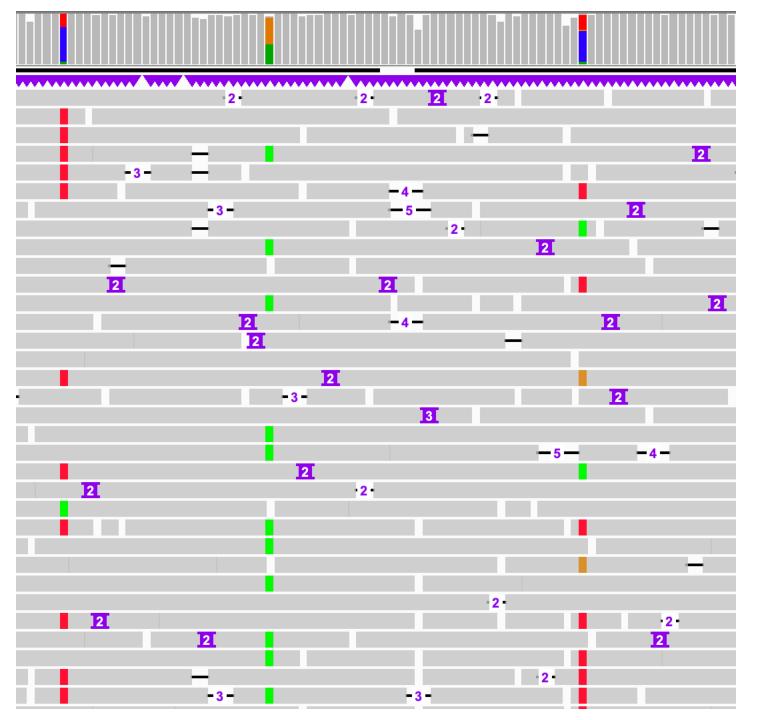
#### Illumina - limitations

- Bridge amplification
- Lengths are limited by out-of-phase of signal



# Long reads (3rd generation)

- Crux: maximizing signal from a singlemolecule base read-out
- Single molecule, so no out-of-phase signal
- Two frequently used platforms:
  - PacBio SMRT sequencing
  - Oxford Nanopore Technology



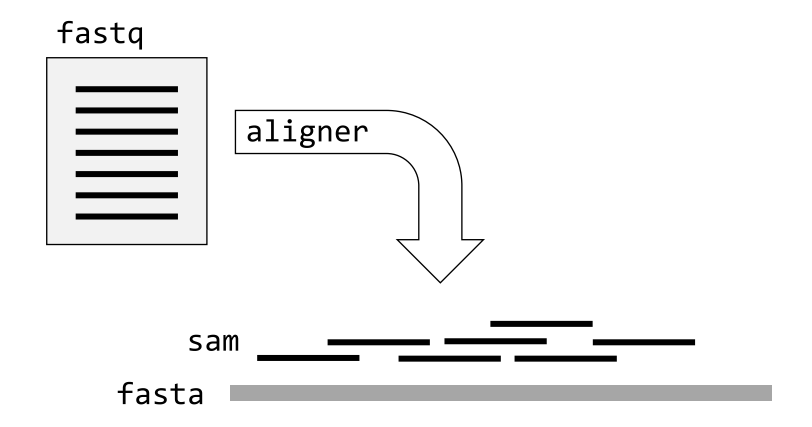
#### Long reads

- More error -> difficulties for variant analysis
- But:
  - PacBio CCS: high baseQ + no bias
  - Long reads can have higher mapping qualities
  - Long reads improve haplotyping

#### What to sequence?

- Whole genome/metagenome
- Reduced representation:
  - Bait capture/whole exome sequencing
  - Restriction Enzyme based (e.g. RAD seq)
  - Amplicon sequencing
  - RNA-seq

## Read alignment (phred)

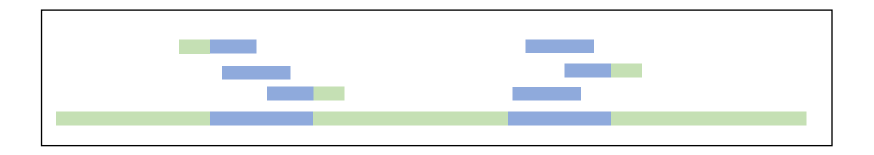




#### Software

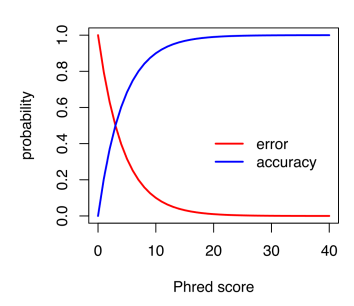
- Basic alignment:
  - bowtie2
  - bwa-mem
- Long reads:
  - minimap2

## Mapping quality



 $MAPQ = -10log_{10} \Pr\{mapping \ position \ is \ wrong\}$ 

$$-10log_{10} (0.01) = 20$$
  
$$-10log_{10} (0.5) = 3$$



# Question

#### sam

sequence alignment format



#### sam header

```
@HD VN:1.0 SO:coordinate

@SQ SN:U00096.3 LN:4641652

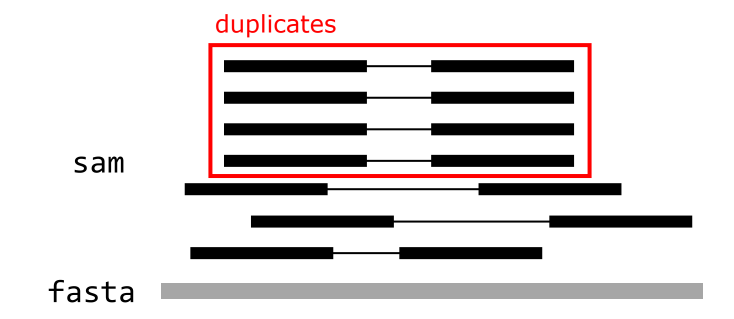
@PG ID:bowtie2 PN:bowtie2 VN:2.4.1 CL: bowtie2-
align-s --wrapper basic-0 -x ref.fasta -1 reads_1.fastq -2
reads_2.fastq"
```

SAM column	example
read name	SRR519926.5
flag	89
reference	chr20
start position	61
mapping quality	42
CIGAR string	150M
reference name mate is mapped	=
start position mate	476
fragment length	515
sequence	CATCACCATTCCCAC
base quality	@>4:4C@89+&9CC@
optional	AS:i:-2
optional	XN:i:0

Z /

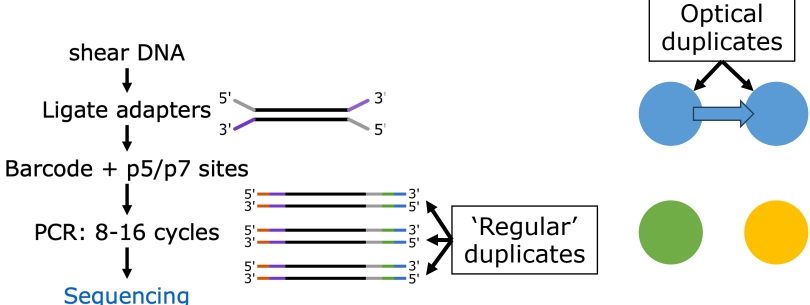
# Question

# Marking duplicates



## Marking duplicates

- 'Regular' duplicate: originates from PCR during library prep
- Optical duplicate: originates from bridge amplification



#### Marking duplicates

- Variant calling: each read is an independent observation of the genome
- Duplicates (can) have the same molecular origin -> not independent
- In a high-quality library, removing duplicates probably doesn't have a big effect on variant analysis

Ebbert MTW et al. (2016) Evaluating the necessity of PCR duplicate removal from next-generation sequencing data and a comparison of approaches. BMC Bioinformatics.

#### Unique Molecular Identifiers

- UMI added before PCR reaction
- Detect PCR duplicates and PCR errors

