Long-read sequence analysis

QC & alignment

Raw file formats

• ONT:

- POD5 (new, apache arrow)
- FAST5 (HDF5)
- Base calling: MinKNOW (guppy)/dorado/third party

PacBio:

BAM (binary sequence alignment format SAM)

fastq



fastq

fasta + basequality (fasta + q = fastq) $BASEQ = -10log_{10} \Pr\{base\ is\ wrong\}$ $\Pr\{base\ is\ wrong\} = 10^{\frac{-BASEQ}{10}}$ $Acurracy = 1 - \Pr\{base\ is\ wrong\}$

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

 $-10log_{10} (0.05) = 13$
 $-10log_{10} (0.5) = 3$

Question 9

Read quality control

- Number of reads
- Read length (mean and spread)
- Base quality
- Overrepresented sequences
- GC content
- Demultiplexing statistics
- Run duration/location dependency
- Others?

Question 10

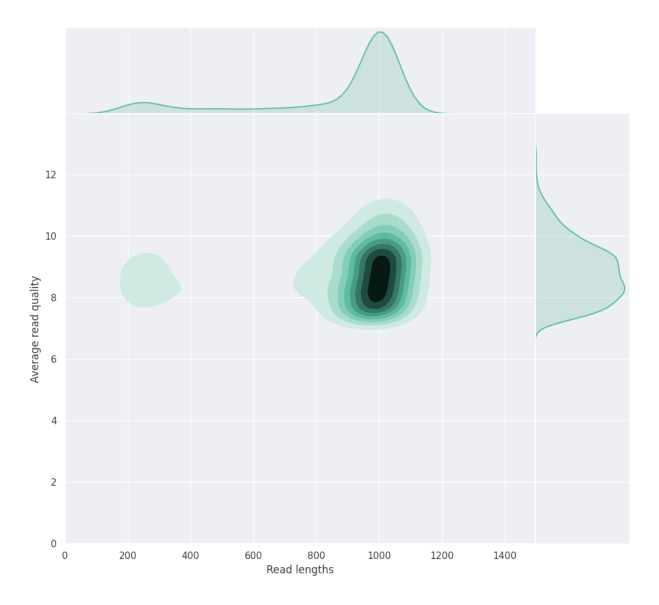
Read quality software

- Software of manufacturer: SMRT Link; MinKNOW
- NanoPlot (https://github.com/wdecoster/NanoPlot)
 - Takes many input formats
 - Basic statistics
- PycoQC (https://github.com/a-slide/pycoQC)
 - Specific for ONT
 - Requires so-called sequencing_summary file
- FastQC

(https://www.bioinformatics.babraham.ac.uk/projects/fastqc/)

- Works also for long reads
- Familiar output to most people

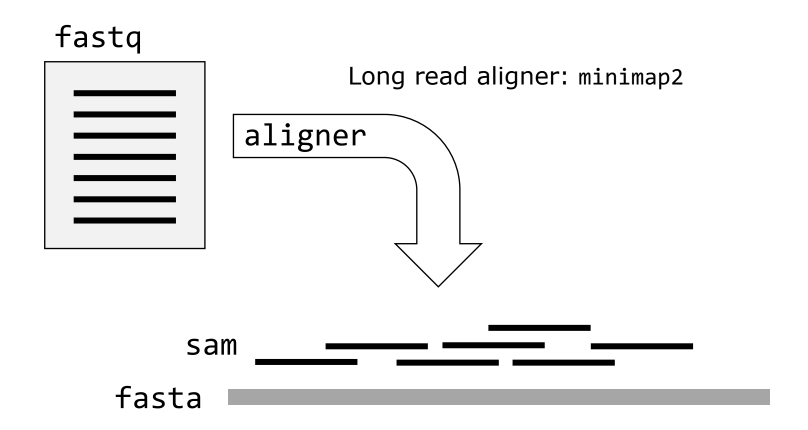
Read lengths vs Average read quality plot

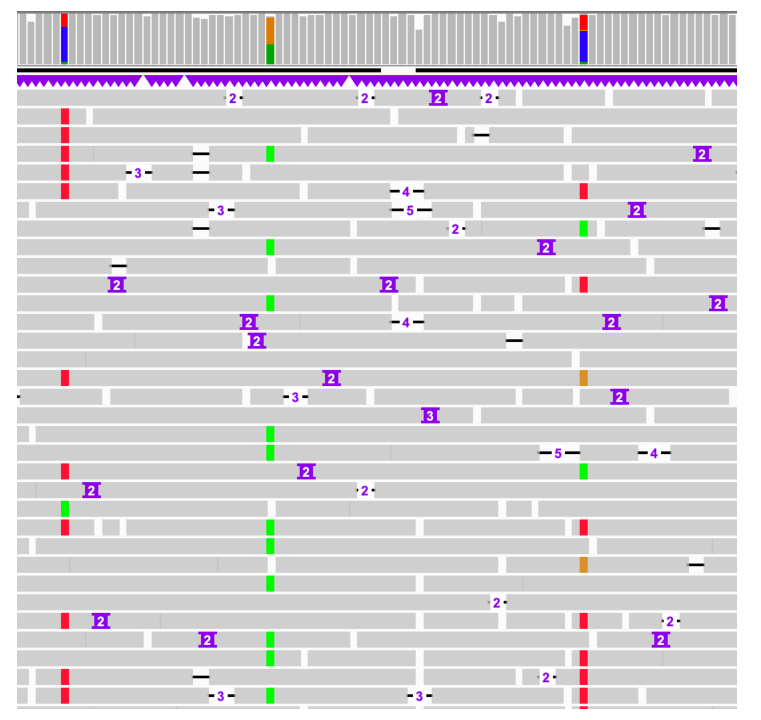


Quality trimming

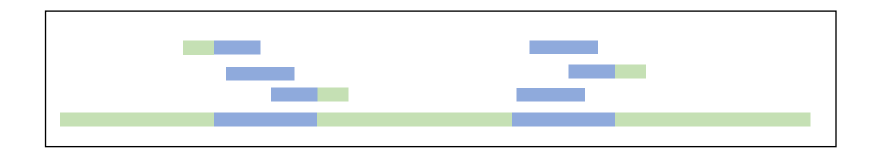
- Removal of:
 - Low quality sequences
 - Adapters/barcodes
- Oxford nanopore: On-instrument (guppy)
- PacBio:
 - On-instrument
 - During CCS generation (pbccs)

Read alignment





Mapping quality



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

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

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

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

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

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Question 11

samtools

- Convert .sam files into (a.o.)
 - .bam (compressed .sam)
 - .fastq
- Subset alignments based on:
 - flag
 - region
- Ordering
- Mark alignment duplicates
- And many other things

Long-reads & fastq

- fastq format is limited to:
 - base
 - base-quality
- Long-read technologies -> need to store more information:
 - PacBio: (unaligned) bam
 - ONT: fast5

Group work preference

Fill out the google form:

https://forms.gle/xWYQQcSoKsbHKJYu6