

CALIDAD DE LECTURAS Y LIMPIEZA

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FORMATOS DE DATOS



FASTA

FASTQ

FAST5 (binarios)



FASTA

```
>gi|5524211|gb|AAD44166.1| cytochrome b
LCLYTHIGRNIYYGSYLYSETWNTGIMLLLITMATAFMGYVLPWGQMSFWGATVITNLFSAIPYIGTNLV
EWIWGGFSVDKATLNRFFAFHFILPFTMVALAGVHLTFLHETGSNNPLGLTSDSDKIPFHPYYTIKDFLG
LLILILLLLLLLALLSPDMLGDPDNHMPADPLNTPLHIKPEWYFLFAYAILRSVPNKLGGVLALFLSIVIL
GLMPFLHTSKHRSMMMLRPLSQALFWTLTMDLLTLTWIGSQPVEYPYTIIGQMASILYFSIILAFLPIAGX
IENY
```

FASTQ

```
@SRR001666.1 071112_SLXA-EAS1_s_7:5:1:817:345 length=36
GGGTGATGGCCGCTGCCGATGGCGTCAAATCCCACC
+
IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII9IG9IC
@SRR001666.1 071113_SLXA-EAS1_s_7:5:1:817:345 length=36
GGGTGATGGCCGCTGCCGATACGGACAAATCCCACC
+
IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIH9IG9IB
```

- Línea 1 (comienza con @)
- Línea 2
- Línea 3 (comienza con +)
- Línea 4 —ASCII char

SGR
Sistema General de Regalías

https://en.wikipedia.org/wiki/FASTQ_format

5

ANTES DE EMPEZAR

- ¿Cuántas lecturas tenemos en cada archivo?
- ¿Qué diferencia hay entre los formatos de salida de cada secuenciador?


CALIDAD



FASTQC



<https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>

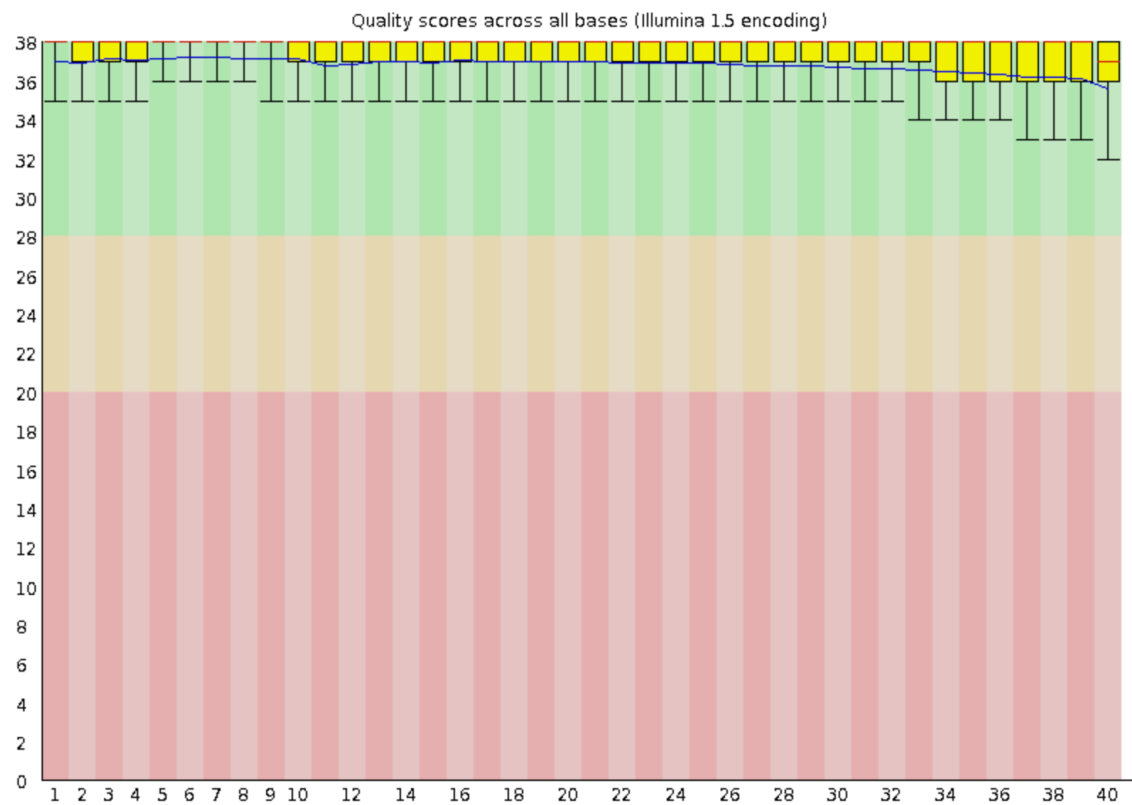




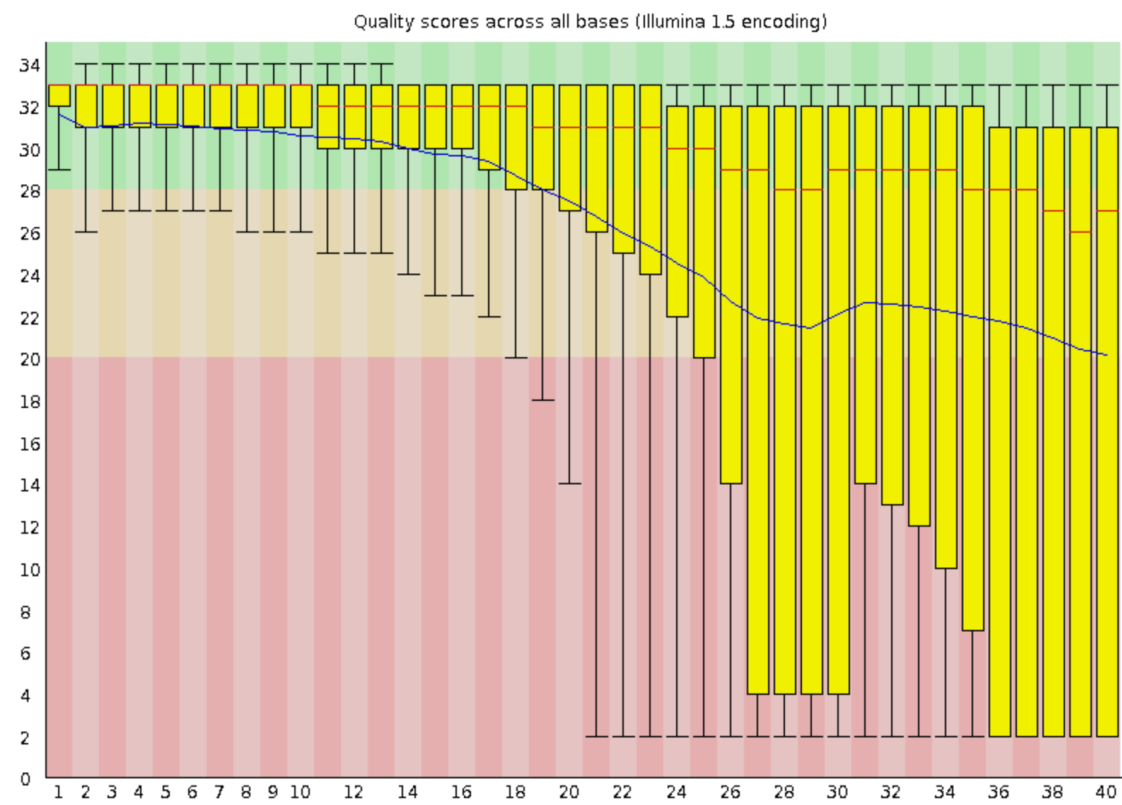
Basic Statistics

Measure	Value
Filename	good_sequence_short.txt
File type	Conventional base calls
Encoding	Illumina 1.5
Total Sequences	250000
Sequences flagged as poor quality	0
Sequence length	40
%GC	45

✔ Per base sequence quality

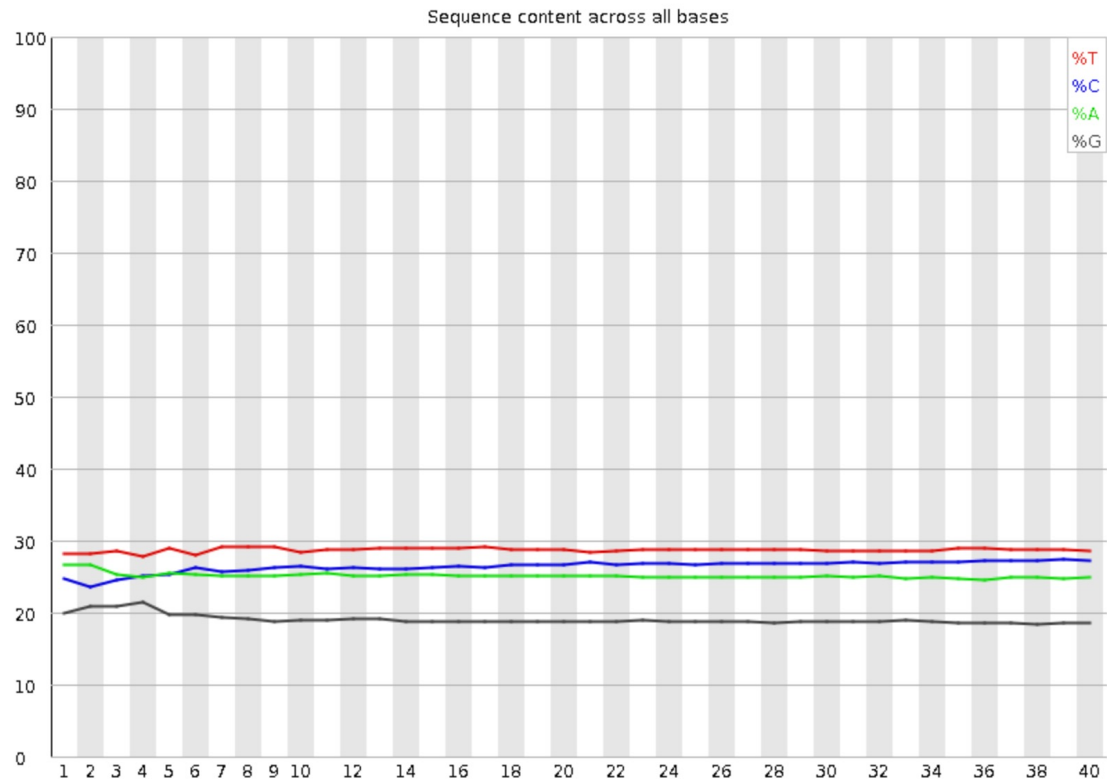


✖ Per base sequence quality

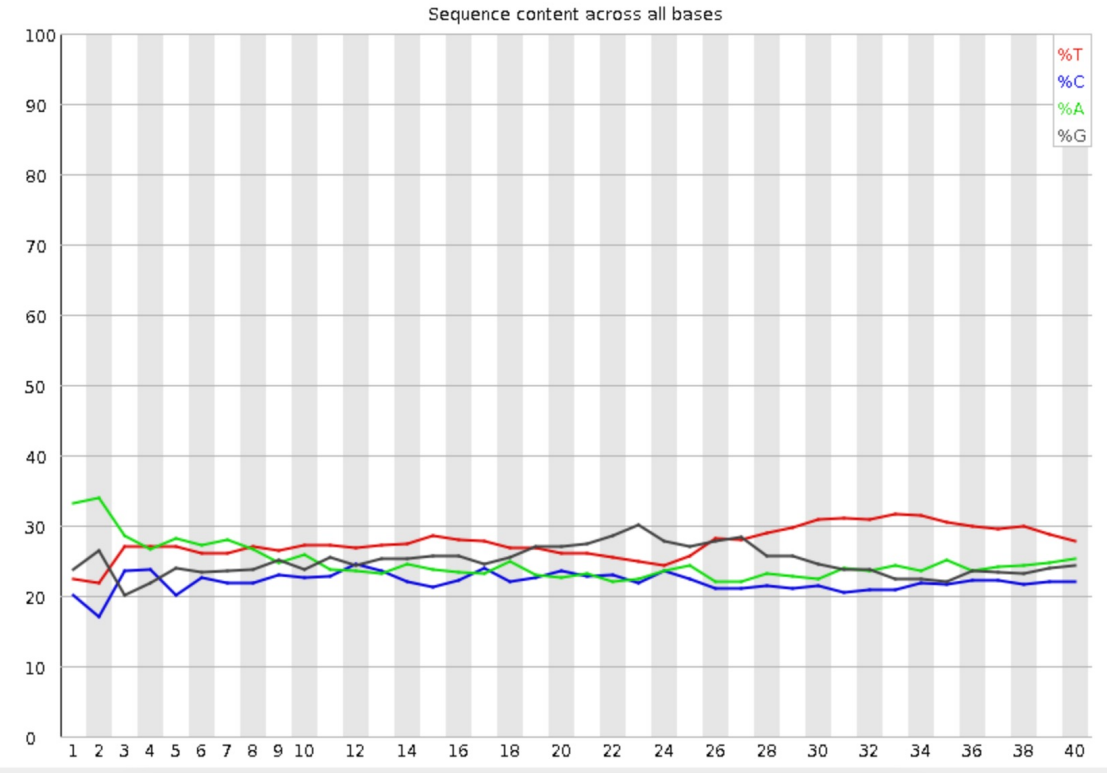




Per base sequence content



Per base sequence content



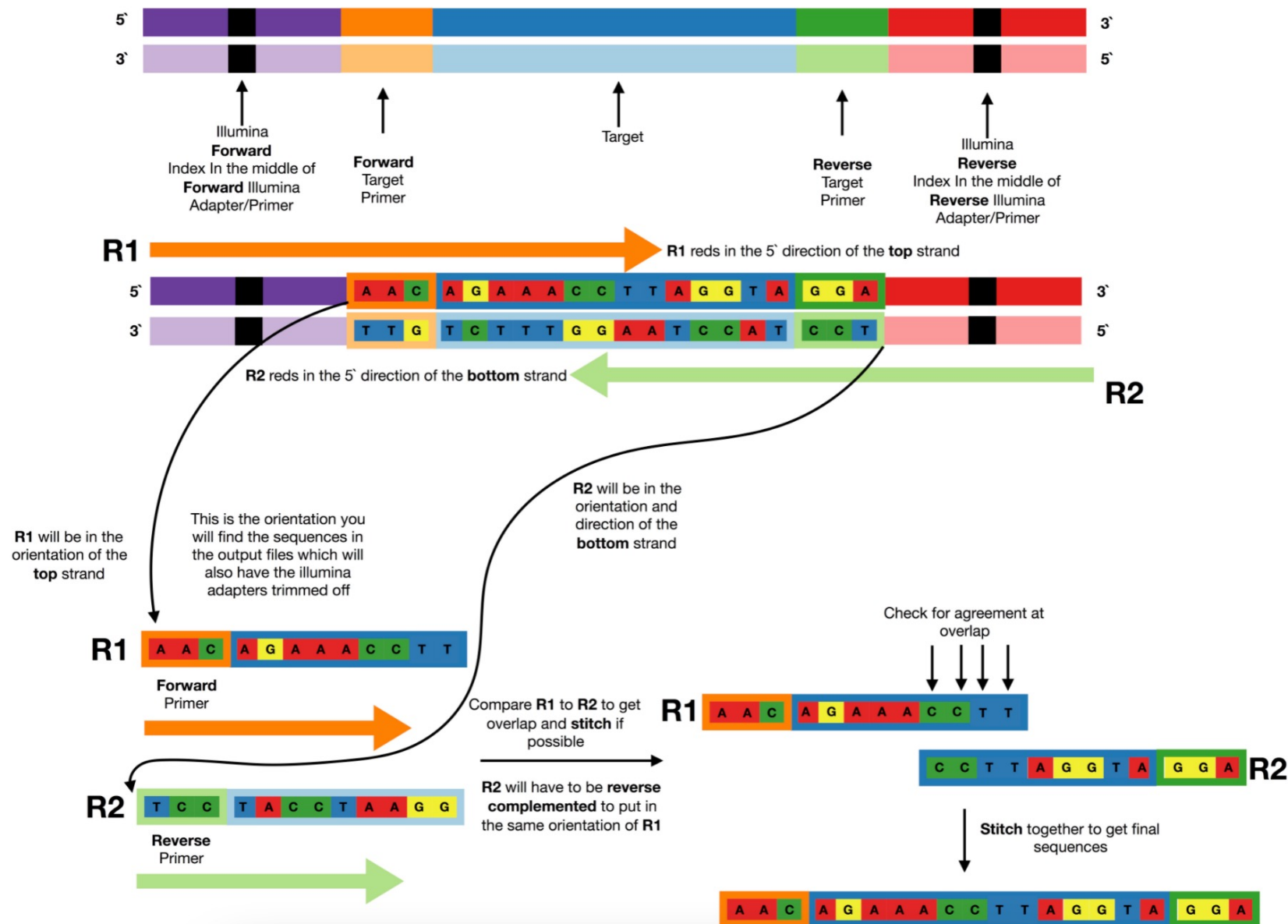
LIMPIEZA



ILLUMINA – ION TORRENT
NANOPORE



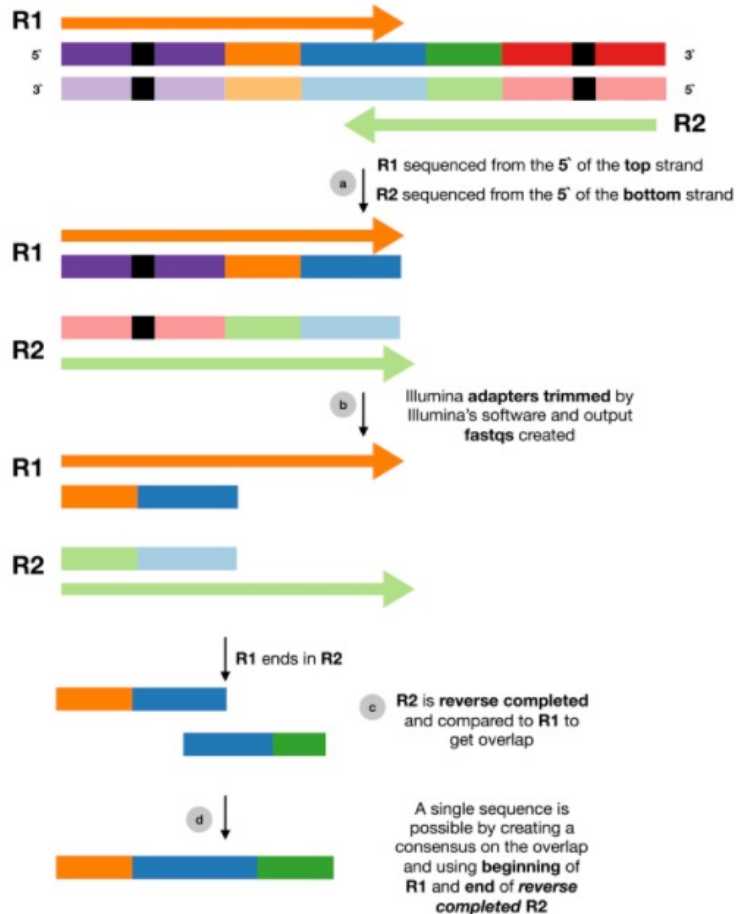
LECTURAS ILLUMINA O ION TORRENT



1

R1EndsInR2

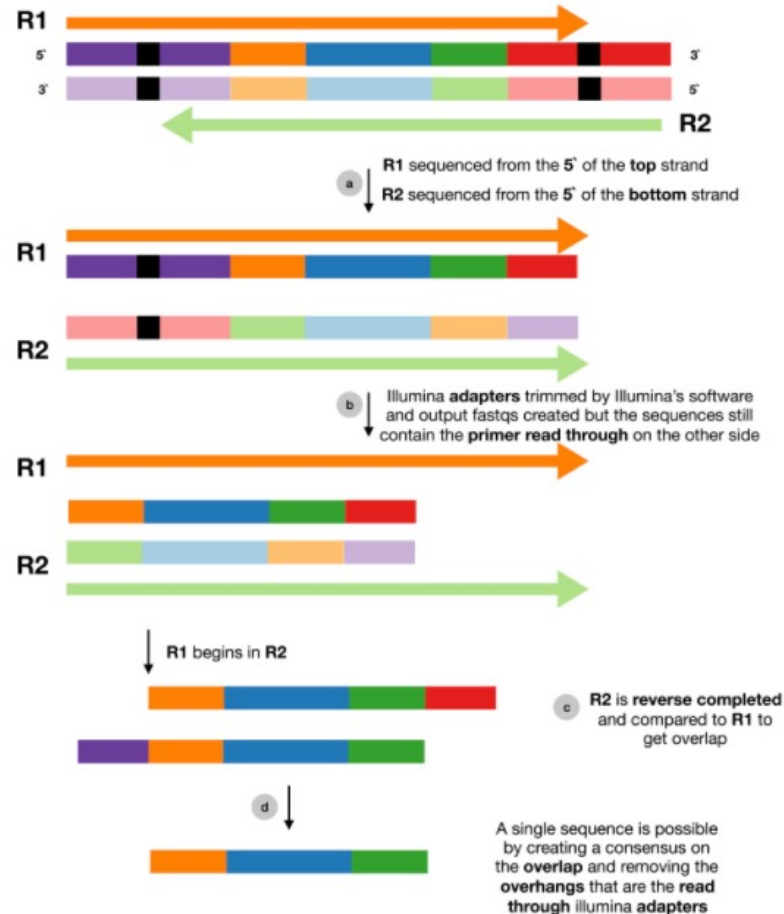
This situation happens when **target sequence plus primers** are **more than read length** but **shorter than 2** times the read (e.g. **target size of 200** on **2x150** sequencing run)



2

R1BeginsInR2

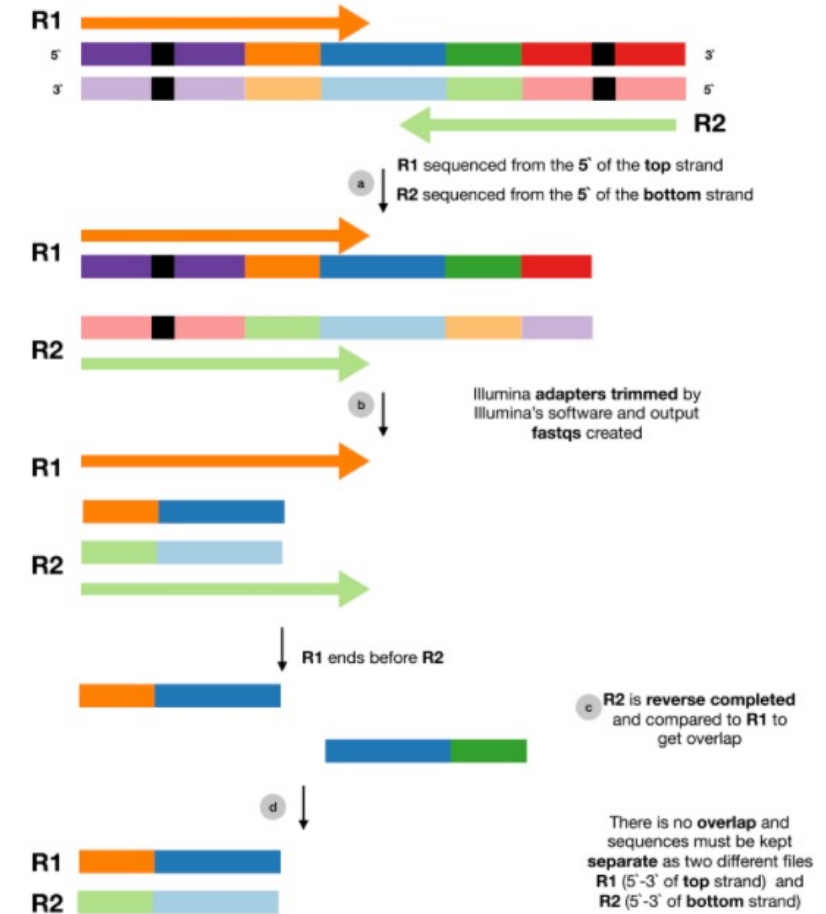
This situation happens when **target sequence plus primers** are **less than read length** (e.g. **target size** of 100 on **2x150** sequencing run)



3

NoOverlap

This situation happens when **target sequence plus primers** are **more than 2** times read length (e.g. **target size** of 400 on **2x150** sequencing run)

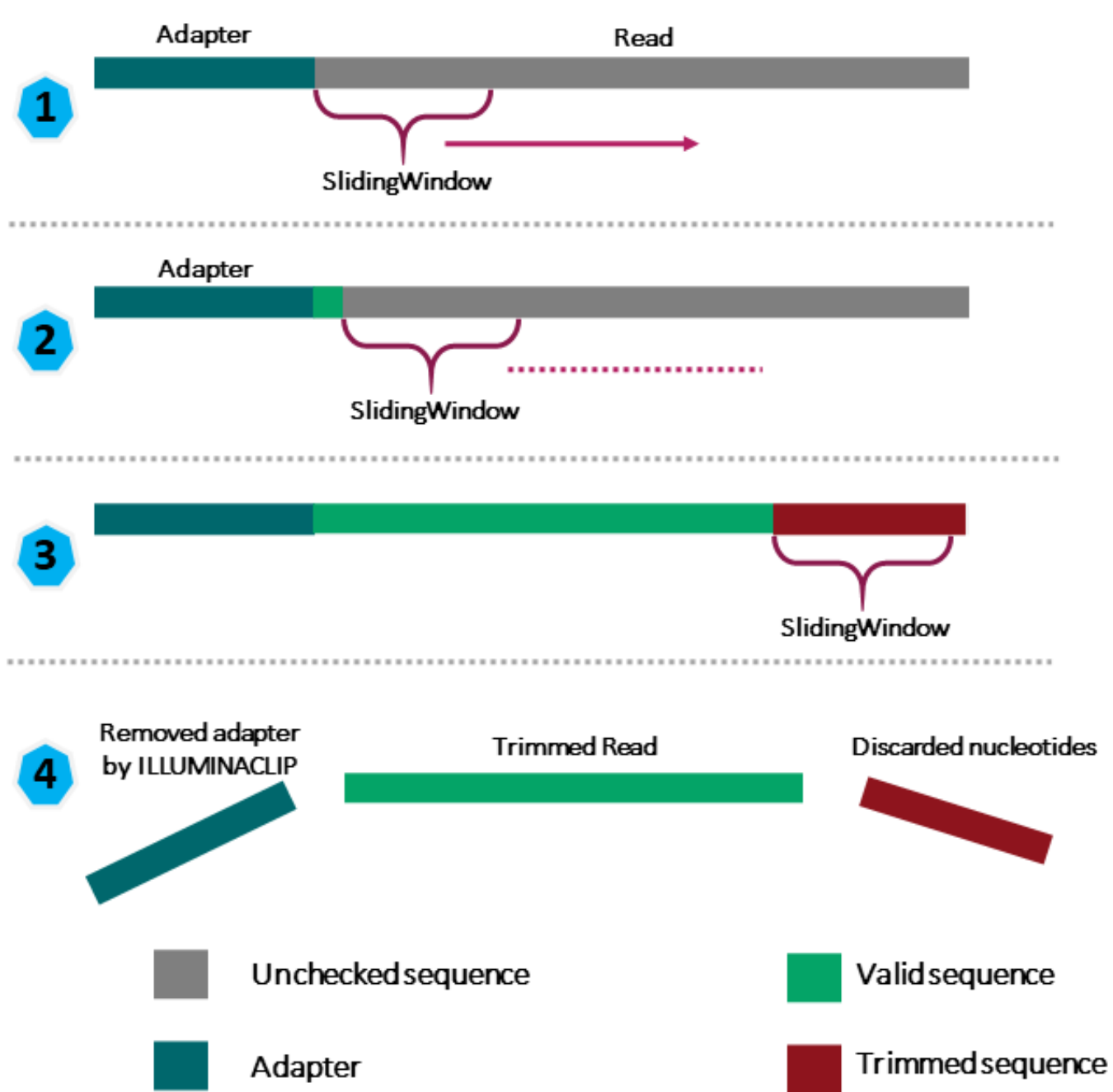


¿QUÉ LIMPIAMOS?

- Secuencias de adaptadores de PCR o secuenciación
- Regiones de baja calidad
- Lecturas de baja calidad (total o ventanas)
- Lecturas muy cortas

TRIMMOMATIC

- ILLUMINACLIP
- SLIDINGWINDOW
- LEADING
- TRAILING
- CROP
- HEADCROP
- MINLEN

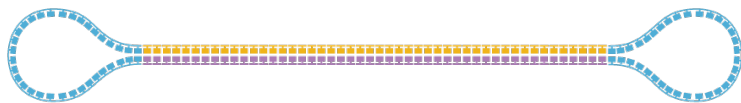


HiFi READS

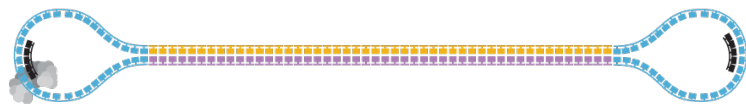
Start with high-quality double stranded DNA



Ligate SMRTbell adapters and size select



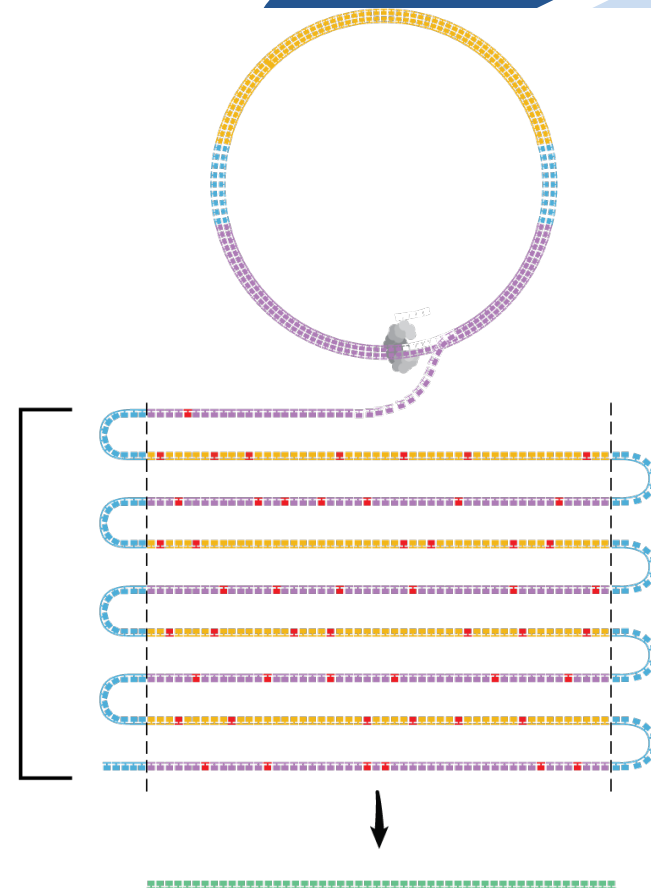
Anneal primers and bind DNA polymerase



Circularized DNA is sequenced in repeated passes

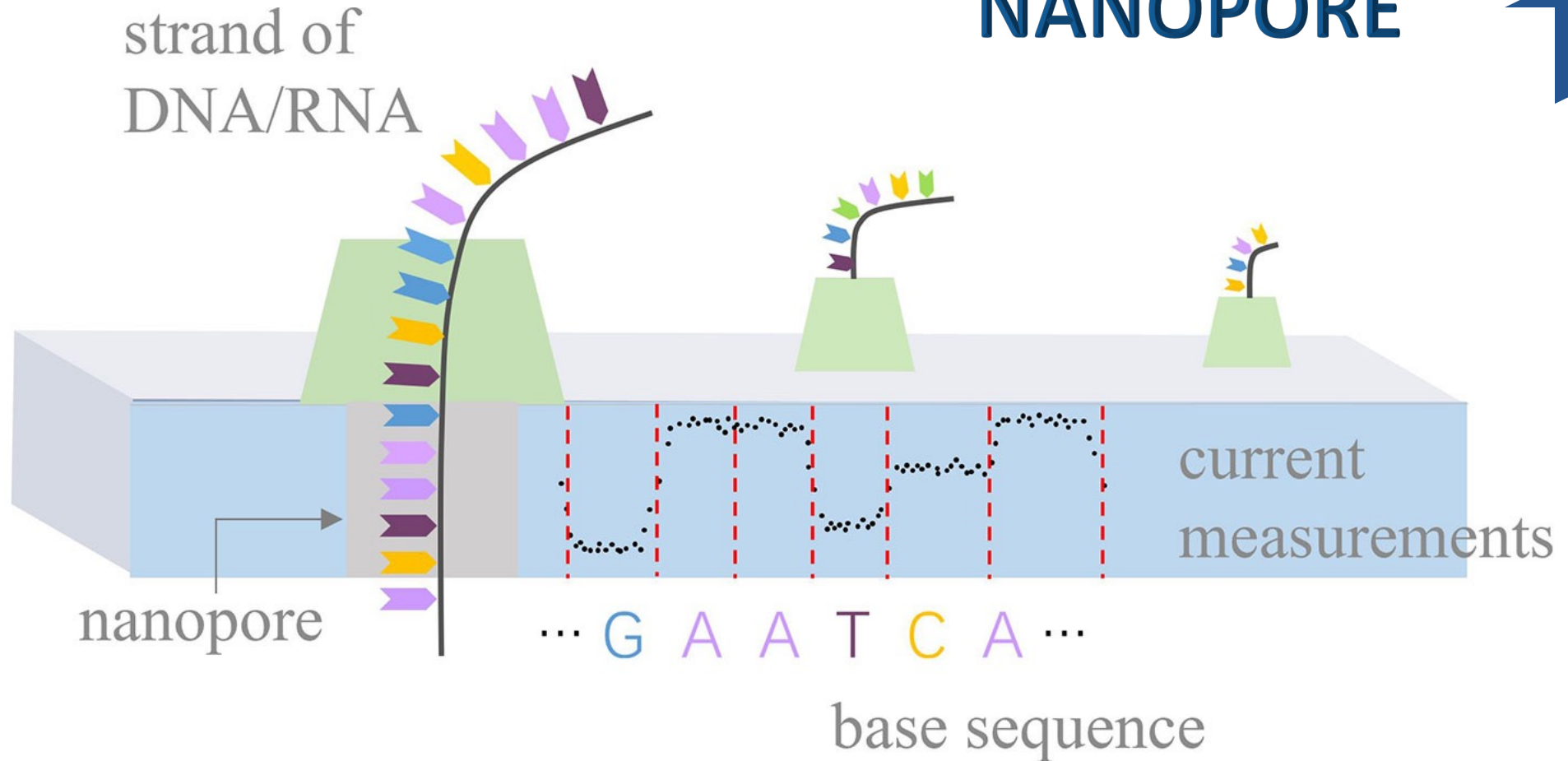
The polymerase reads are trimmed of adapters to yield subreads

Consensus is called from subreads



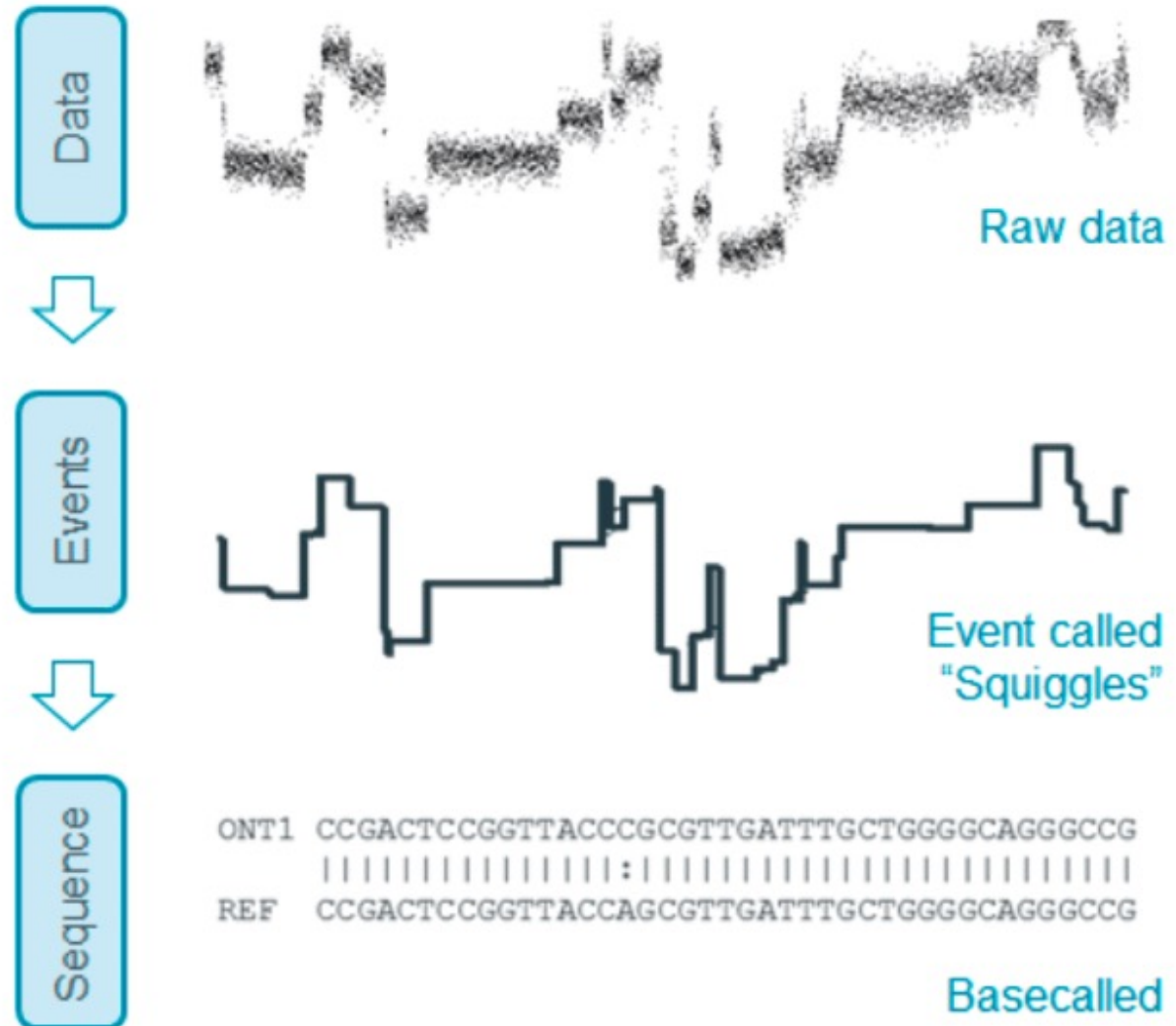
HiFi READ
(>99% accuracy)

NANOPORE

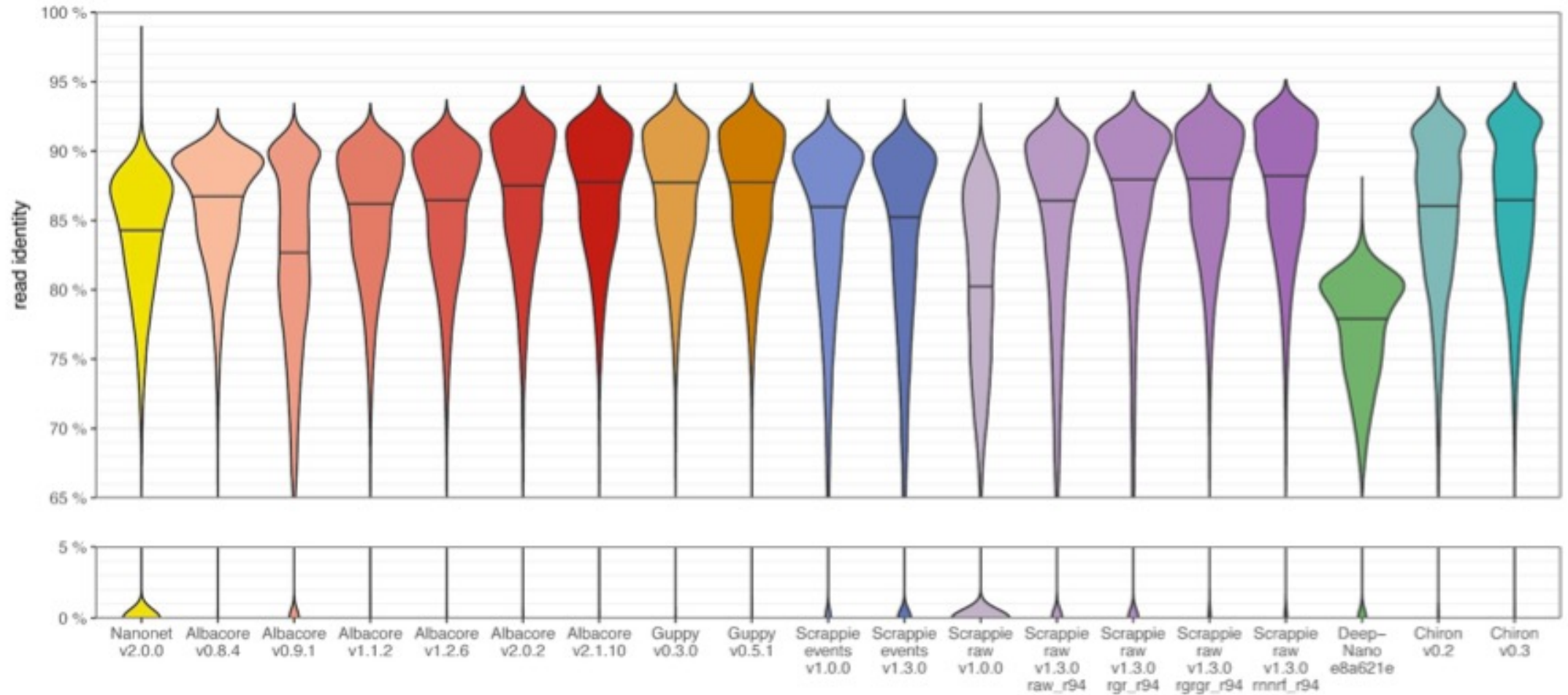


LLAMADO DE BASES

FAST5 (HDF5) a FASTQ



ONT Read calling



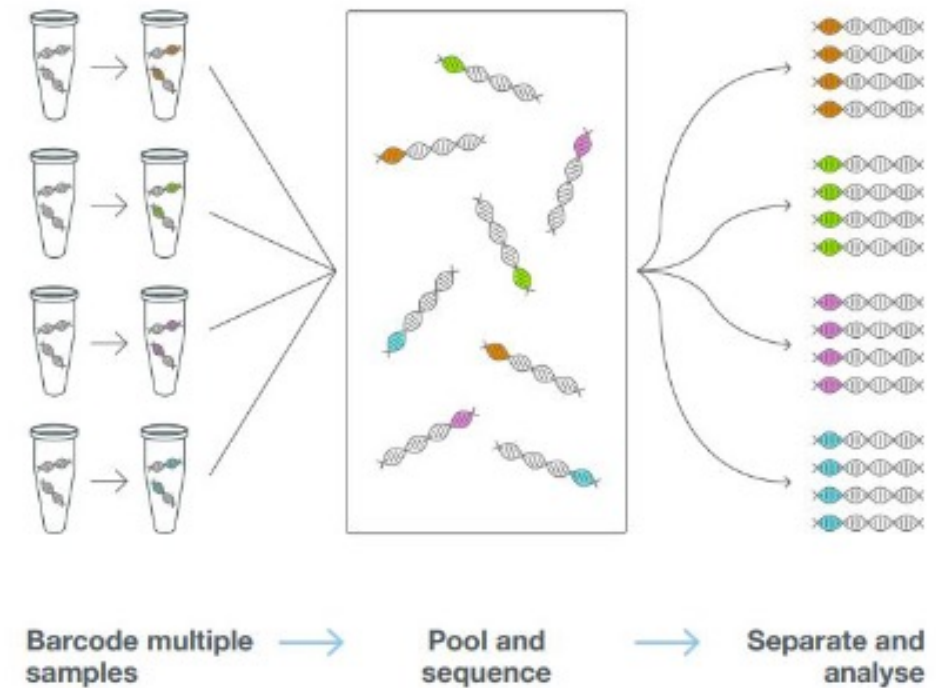
summary_file.txt

filename	FAK47038_aa36ef836fd50817477a5770772dffc63bfed2eb_30
read_id	188e2a0b-780c-440d-9223-61d8979dd002
run_id	aa36ef836fd50817477a5770772dffc63bfed2eb
batch_id	0
channel	70
mux	3
start_time	9688.985500
duration	1.610500
num_events	1288
passes_filtering	TRUE
template_start	9689.318000
num_events_template	1022
template_duration	1.278000
sequence_length_template	545
mean_qscore_template	11.462492
strand_score_template	3.165753
median_template	79.270927
mad_template	9.512511
scaling_median_template	79.270927
scaling_mad_template	9.512511

ONT demultiplexing

Deepbinner: Demultiplexing barcoded ONT reads with deep convolutional neural networks (CNN). The network is trained to classify barcodes based on the raw nanopore signal.

Guppy
In contrast to Deepbinner, guppy barcoding requires basecalling of all reads and detects barcodes in the sequence



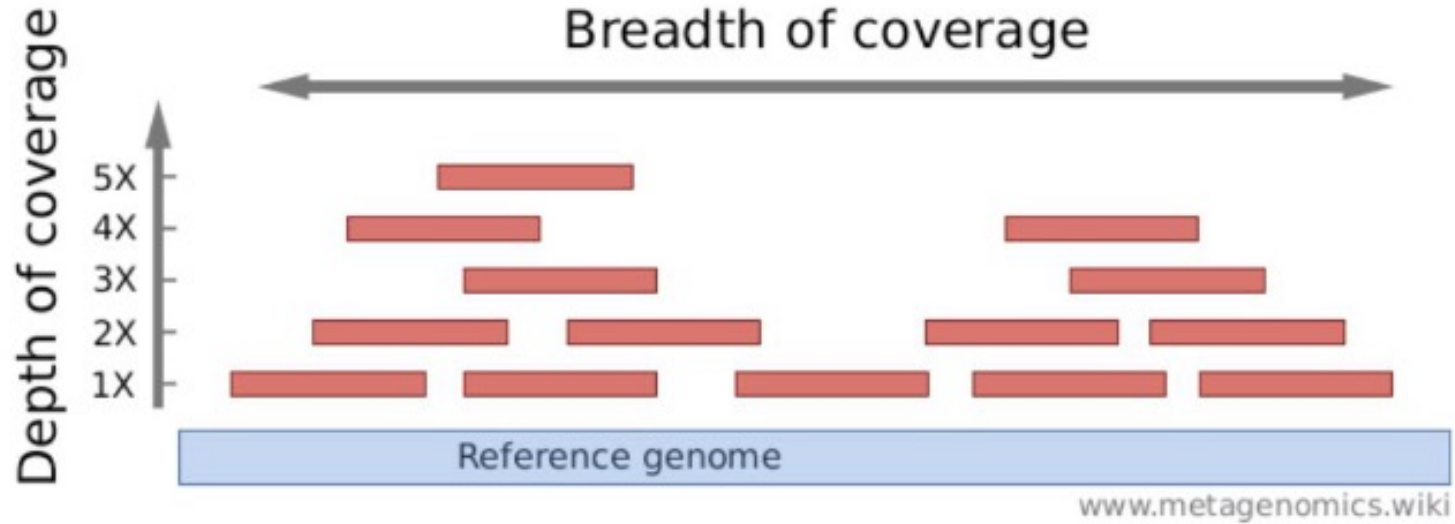
ONT Read calling, cleaning and filtering

Sequencer ONT : raw fast5 files

- Transform fast5 signal in fastq standard format *Guppy, Bonito*
- Optional Demultiplexing and removing adapters *Guppy options*
- Optional Find and remove adapters from reads *Porechop*
- Optional Quality filtering using the *sequencing_summary.txt* information : *Guppy options, filtlong, nanofilt*

Guppy is a neural network based basecaller that in addition to basecalling also performs filtering of low quality reads, clipping of Oxford Nanopore adapters and estimation of methylation probabilities per base

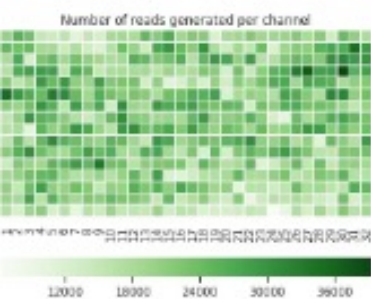
Calculate depth of coverage



depth of coverage estimation :

- Count how much base pairs in all sequenced reads? *total_pb*
- What is the expected genome size? *genome_size*

$$\text{depth_of_coverage} = \text{total_pb} / \text{genome_size}$$

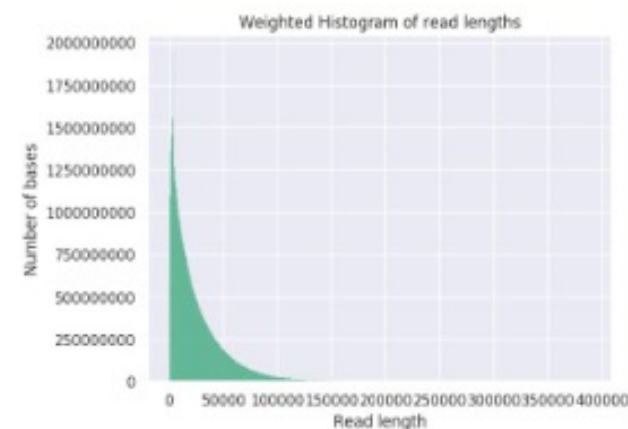
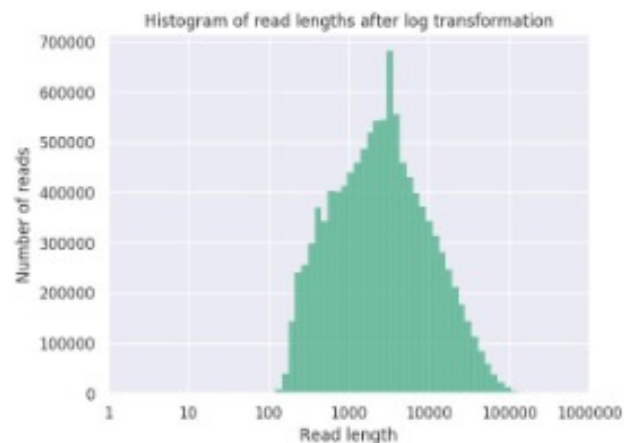
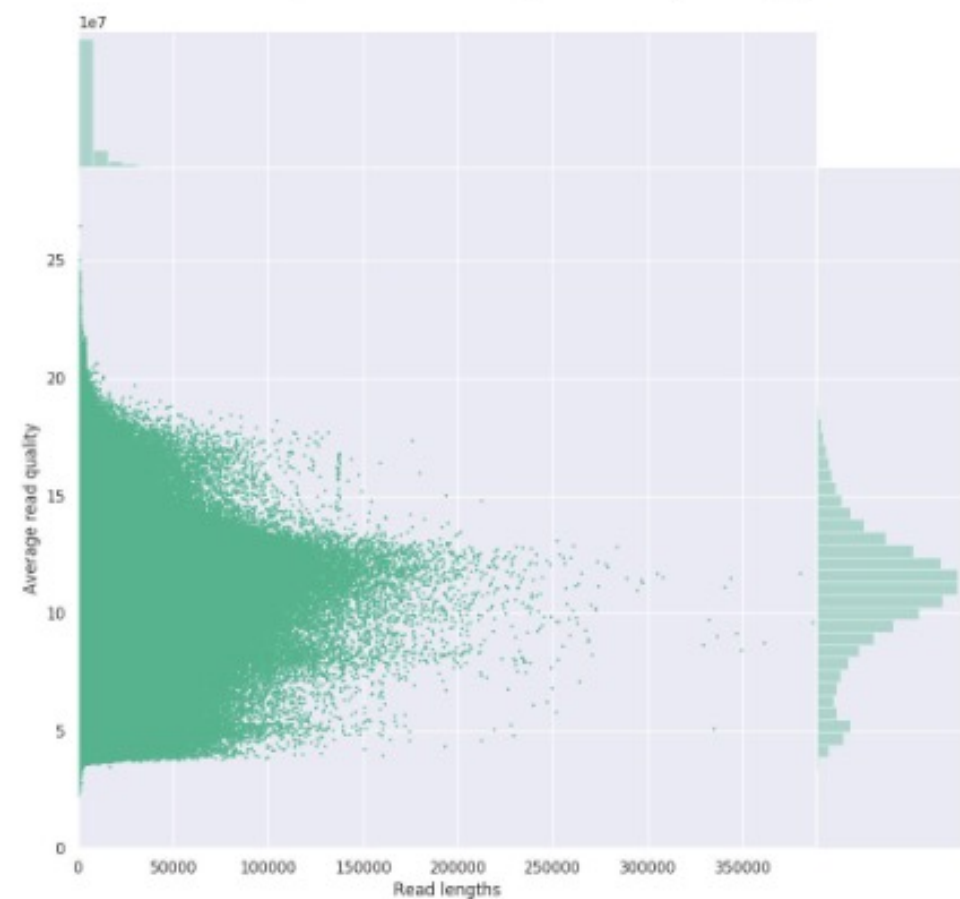


Reads Quality control : *NanoPlot*

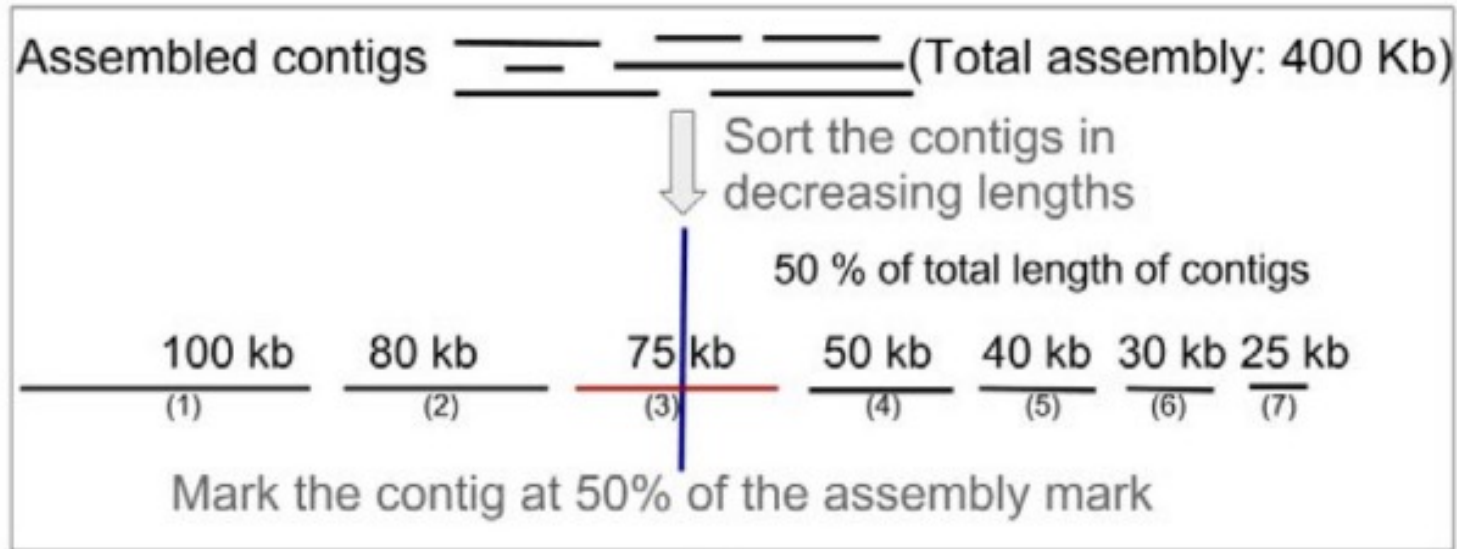
Summary statistics

General summary	
Active channels	512.0
Mean read length	6,315.6
Mean read quality	10.9
Median read length	2,517.0
Median read quality	11.1
Number of reads	10,847,854.0
Read length N50	16,816.0
Total bases	68,510,227,164.0

Read lengths vs Average read quality plot



What is N50 and L50?



- N50, length of the contig at 50% assembly: 75 kb
- L50, number of contigs until 50% assembly: 3

GRACIAS