

Cycle

« Analyse de données de séquençage à haut-débit »

Module 1/5: Analyses ADN

8 et 9 Mars 2023

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Jour 1 :

Matin

- Cours
 - NGS Introduction
 - Reads Quality Control + Cleaning
- TP FastQC + multiqc + cleaning

<https://training.galaxyproject.org/training-material/topics/sequence-analysis/tutorials/quality-control/tutorial.html>

Pause midi

Après-midi

- Cours
 - Reads mapping on reference
- TP Mapping

<https://training.galaxyproject.org/training-material/topics/sequence-analysis/tutorials/mapping/tutorial.html>

Jour 2 :

Matin

- TP Mapping
 - Deep dive into Bowtie2 alignment parameters
- Cours
 - Genome assembly

Pause midi

Après-midi

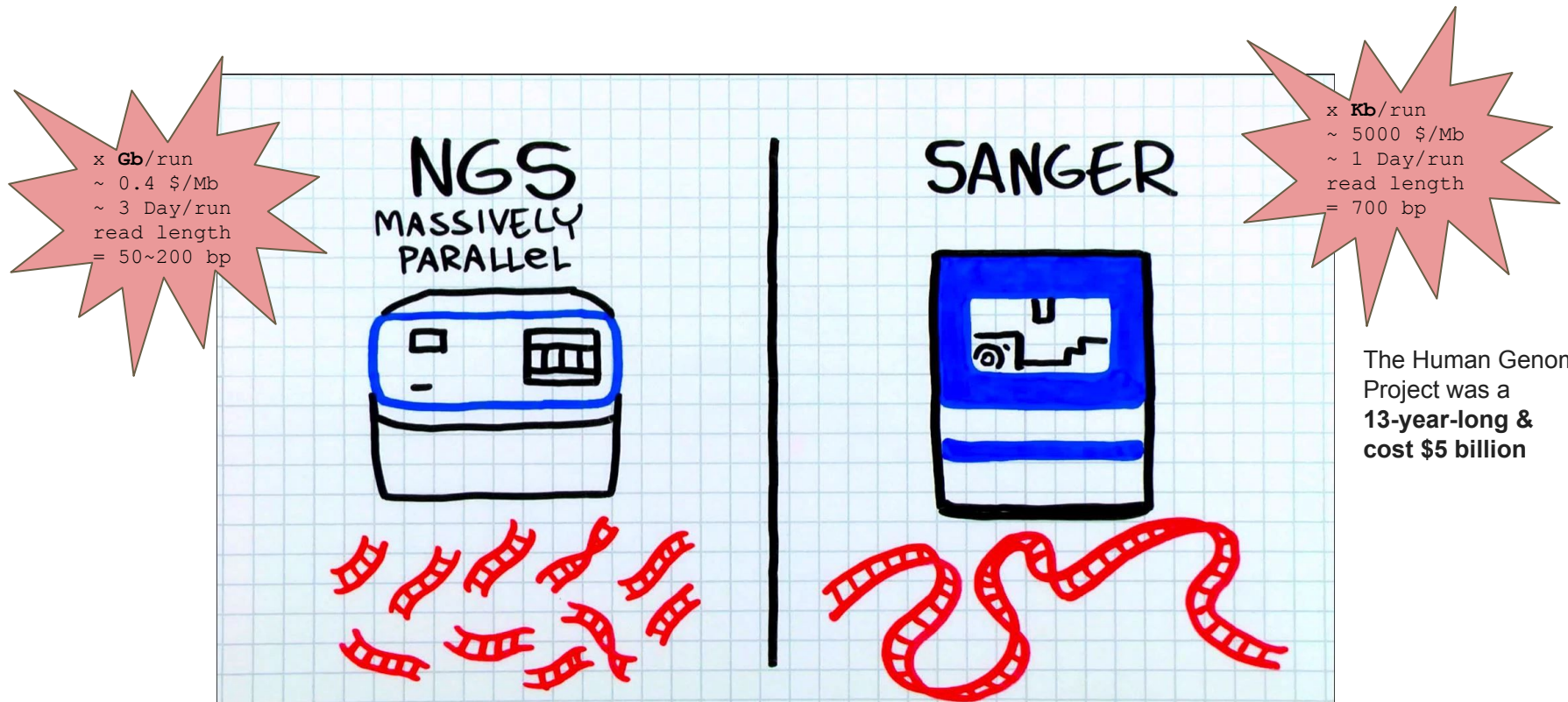
- TP Genome assembly
 - <https://training.galaxyproject.org/training-material/topics/assembly/tutorials/debruijn-graph-assembly/tutorial.html>
 - <https://training.galaxyproject.org/training-material/topics/assembly/tutorials/assembly-quality-control/tutorial.html>

Jour 1 :

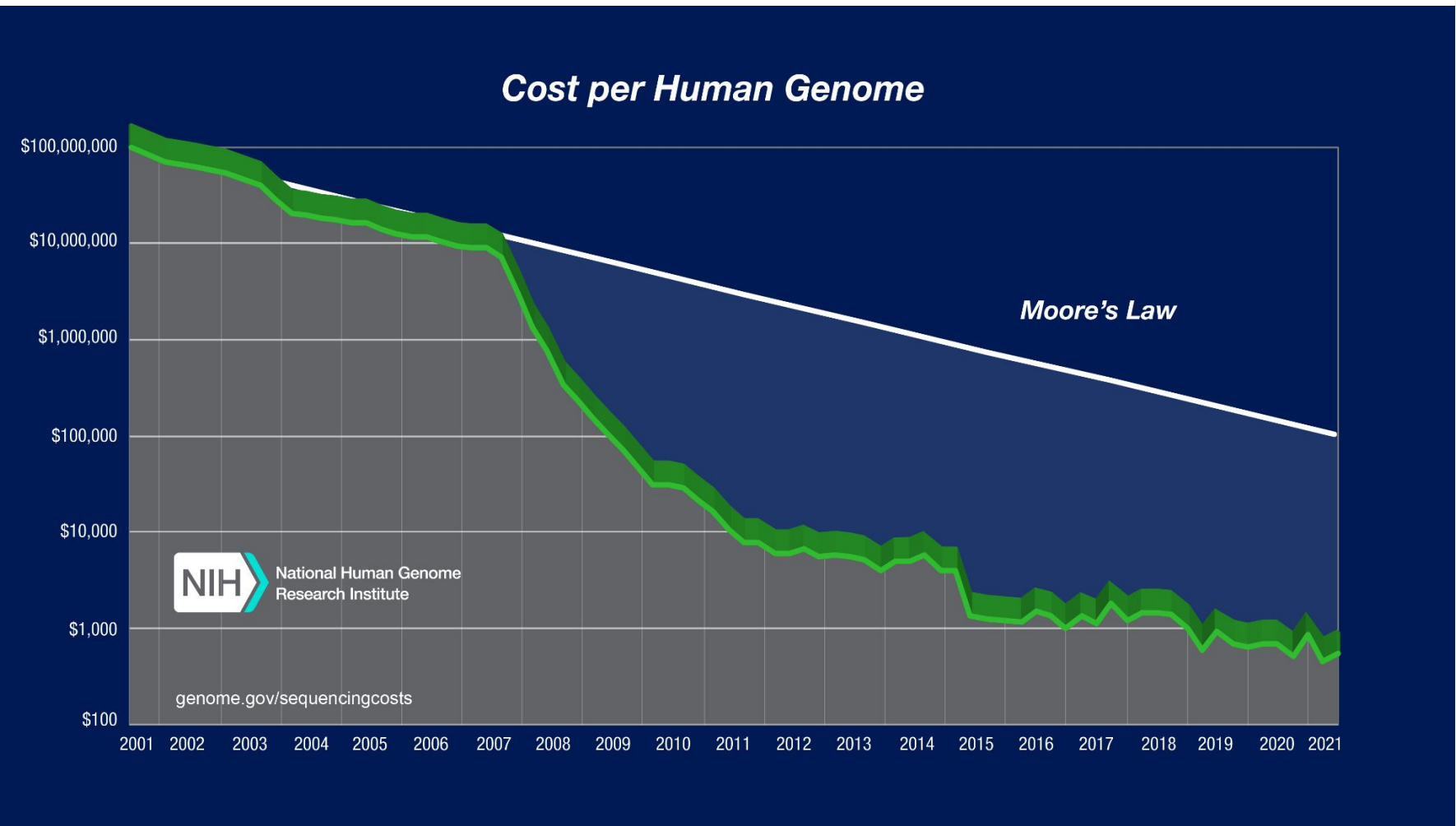
- NGS Introduction
 - What is NGS?
 - Sequencers
 - Applications
 - NGS workflow
 - Output data

What is Next-Generation Sequencing (NGS)?

"Next-generation sequencing (NGS), also known as high-throughput sequencing, is the catch-all term used to describe a number of different modern sequencing technologies. These technologies allow for sequencing of DNA and RNA much more quickly and cheaply than the previously used Sanger sequencing, and as such revolutionised the study of genomics and molecular biology"



What is Next-Generation Sequencing (NGS)?



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Illumina sequencing

Illumina sequencing works by simultaneously identifying DNA bases, as each base emits a unique fluorescent signal, and adding them to a nucleic acid chain

Ion Torrent: Proton / PGM sequencing (thermofisher)

Ion Torrent sequencing measures the direct release of H⁺ (protons) from the incorporation of individual bases by DNA polymerase and therefore differs from the previous two methods as it does not measure light.



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MGI sequencing (BGI group)



Sequencers – Illumina



Benchtop Sequencers		Production-Scale Sequencers				
		iSeq 100	MiniSeq	MiSeq Series	NextSeq 550 Series	NextSeq 1000 & 2000
Popular Applications & Methods		Key Application	Key Application	Key Application	Key Application	Key Application
Large Whole-Genome Sequencing (human, plant, animal)						
Small Whole-Genome Sequencing (microbe, virus)	●	●	●	●	●	●
Exome & Large Panel Sequencing (enrichment-based)					●	●
Targeted Gene Sequencing (amplicon-based, gene panel)	●	●	●	●	●	●
Single-Cell Profiling (scRNA-Seq, scDNA-Seq, oligo tagging assays)					●	●
Transcriptome Sequencing (total RNA-Seq, mRNA-Seq, gene expression profiling)					●	●
Targeted Gene Expression Profiling	●	●	●	●	●	●
miRNA & Small RNA Analysis	●	●	●	●	●	●
DNA-Protein Interaction Analysis (ChIP-Seq)				●	●	●
Methylation Sequencing					●	●
16S Metagenomic Sequencing		●	●	●	●	●
Metagenomic Profiling (shotgun metagenomics, metatranscriptomics)					●	●
Cell-Free Sequencing & Liquid Biopsy Analysis					●	●
Run Time	9.5-19 hrs	4-24 hours	4-55 hours	12-30 hours	11-48 hours	
Maximum Output	1.2 Gb	7.5 Gb	15 Gb	120 Gb	330 Gb*	
Maximum Reads Per Run	4 million	25 million	25 million [†]	400 million	1.1 billion*	
Maximum Read Length	2 × 150 bp	2 × 150 bp	2 × 300 bp	2 × 150 bp	2 × 150 bp	

Sequencers – Illumina



Benchtop Sequencers



NextSeq 1000 & 2000



NovaSeq 6000



NovaSeq X Series

Popular Applications & Methods	Key Application	Key Application	Key Application
Large Whole-Genome Sequencing (human, plant, animal)		●	●
Small Whole-Genome Sequencing (microbe, virus)	●	●	●
Exome & Large Panel Sequencing (enrichment-based)	●	●	●
Targeted Gene Sequencing (amplicon-based, gene panel)	●	●	●
Single-Cell Profiling (scRNA-Seq, scDNA-Seq, oligo tagging assays)	●	●	●
Transcriptome Sequencing (total RNA-Seq, mRNA-Seq, gene expression profiling)	●	●	●
Chromatin Analysis (ATAC-Seq, ChIP-Seq)	●	●	●
Methylation Sequencing	●	●	●
Metagenomic Profiling (shotgun metagenomics, metatranscriptomics)	●	●	●
Cell-Free Sequencing & Liquid Biopsy Analysis	●	●	●
Run Time	11-48 hours	~13-38 hours (dual SP flow cells) ~13-25 hours (dual S1 flow cells) ~16-36 hours (dual S2 flow cells) ~44 hours (dual S4 flow cells)	~13-21 hours (1.5B flow cells [†]) ~18-24 hours (10B flow cells [†]) ~48 hours (25B flow cells [†])
Maximum Output	360 Gb *	6000 Gb	16 Tb
Maximum Reads Per Run	1.2 billion *	20 billion	26 billion (single flow cells) 52 billion (dual flow cells)
Maximum Read Length	2 × 150 bp	2 × 250 bp**	2 × 150 bp

Sequencers – Illumina (pre-2020)

illumina®



	NextSeq Series	HiSeq Series	HiSeq X Series [†]	NovaSeq 6000 System
Popular Applications & Methods	Key Application	Key Application	Key Application	Key Application
Large Whole-Genome Sequencing (human, plant, animal)	●	●	●	●
Small Whole-Genome Sequencing (microbe, virus)	●	●		●
Exome Sequencing	●	●		●
Targeted Gene Sequencing (amplicon, gene panel)	●	●		●
Whole-Transcriptome Sequencing	●	●		●
Gene Expression Profiling with mRNA-Seq	●	●		●
miRNA & Small RNA Analysis	●	●		●
DNA-Protein Interaction Analysis	●	●		●
Methylation Sequencing	●	●		●
Shotgun Metagenomics	●	●		●

Optimized NGS Sample Tracking and Workflows

See how BaseSpace Clarity LIMS (Laboratory Information Management System) enabled this large genomics lab to standardize lab procedures and cope with increasing sample volumes from diverse clients.

[Read Case Study >](#)

Run Time	12–30 hours	< 1–3.5 days (HiSeq 3000/HiSeq 4000) 7 hours–6 days (HiSeq 2500)	< 3 days	16–36 hours (Dual S2 flow cells) 44 hours (Dual S2 flow cells)
Maximum Output	120 Gb	1500 Gb	1800 Gb	6000 Gb [§]
Maximum Reads Per Run	400 million	5 billion	6 billion	20 billion**
Maximum Read Length	2 × 150 bp	2 × 150 bp	2 × 150 bp	2 × 150 bp

Sequencers – Thermo Fisher Scientific

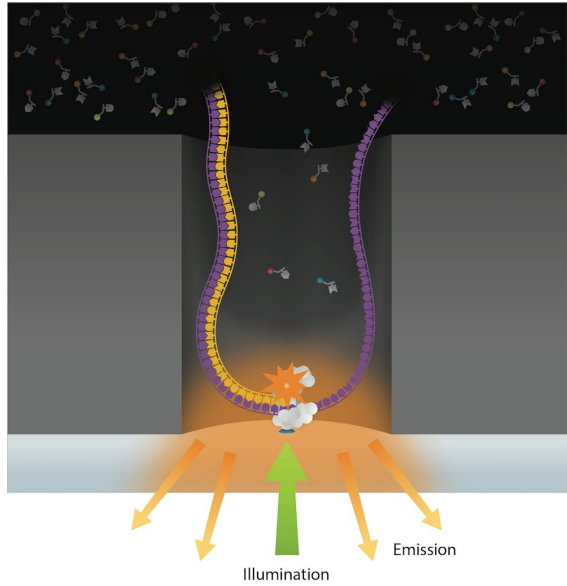
Plateformes de séquençage			
	Système Ion PGM™ pour le séquençage de nouvelle génération	Système Ion S5™ pour le séquençage de nouvelle génération	Système Ion S5™ XL pour le séquençage de nouvelle génération
Avantages	<p>Évolutivité : de 30 Mo à 2 Go</p> <p>Rapidité : séquençage exécuté en 2 à 7 heures, selon la longueur de lecture et la sortie par la puce</p>	<p>Simplicité : solutions de flux de travaux automatisé, de la préparation des échantillons à l'analyse</p> <p>Évolutivité : de 600 Mo à 15 Go</p> <p>Rapidité : séquençage effectué en 2,5 à 4 heures (quelle que soit la sortie par la puce)</p>	<p>Simplicité : solutions de flux de travaux automatisé, de la préparation des échantillons à l'analyse</p> <p>Évolutivité : de 600 Mo à 15 Go</p> <p>Rapidité : de l'ADN aux données en 24 heures</p>
Applications	ARN ciblé de séquençage ADN ciblé	ARN ciblé	ARN ciblé
	Microbien	Microbien	Microbien
		Transcriptome	Transcriptome
		Exome	Exome
		Séquençage de l'ARN	Séquençage de l'ARN

Sequencers - MGI (BGI group)

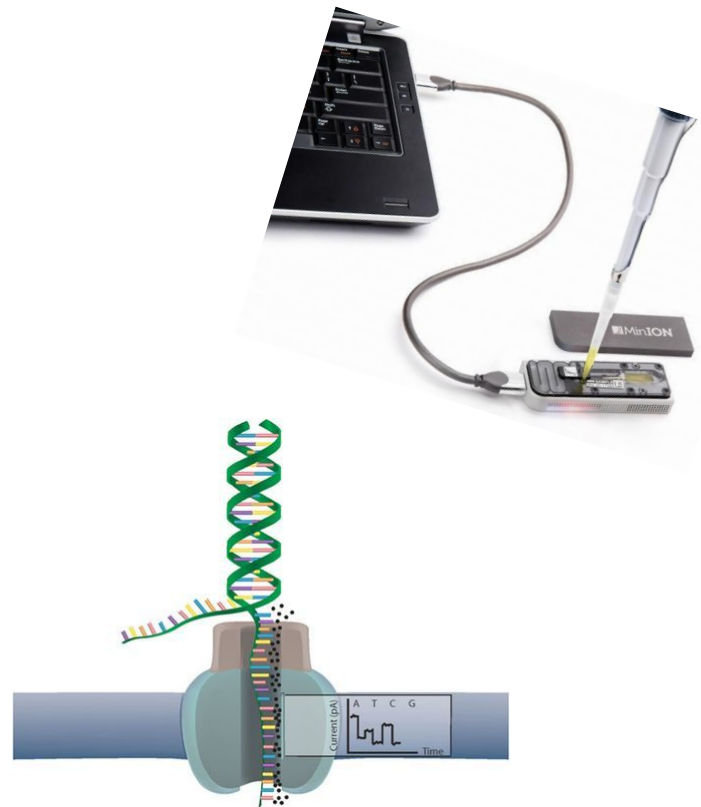


	Sequencers +	Sequencers +	Sequencers +	Sequencers +	Sequencers +	Sequencers +	Sequencers +
Product Model	DNBSEQ-T7	DNBSEQ-T7* For HotMPS Only	DNBSEQ-G400	DNBSEQ-G400* For HotMPS Only	DNBSEQ-G400C*	DNBSEQ-G99	DNBSEQ-G50
Features	Ultra-high Throughput	Ultra-high Throughput	Adaptive	Adaptive	Adaptive	Fast	Effective
Applications	Whole Genome Sequencing, Deep Exome Sequencing, Transcriptome Sequencing, and Targeted Panel Projects.	Whole Genome Sequencing, Deep Exome Sequencing, Transcriptome Sequencing, and Targeted Panel Projects.	WGS, WES, Transcriptome sequencing, etc.	WGS, WES, Transcriptome sequencing, etc.	Small RNA, Pathogen Fast Identification etc.	Targeted oncology panel sequencing, infectious disease sequencing, oncology methylation sequencing, small whole-genome sequencing	Small whole genome sequencing, targeted DNA/RNA panels, low-pass whole genome sequencing
Flow Cell Type	FC	FC	FCL & FCS	FCL	FCL	FC	FCL & FCS
Lane/Flow Cell++	1 lane	1 lane	2 or 4 lanes	4 lanes	4 lanes	1 lane	1 lane
Operation Mode	Ultra-high Throughput	Ultra-high Throughput	High Throughput	High Throughput	High Throughput	Small and Medium Throughput	Medium Throughput
Max. Throughput / RUN	6Tb	4Tb	1440Gb	720Gb	360G	48Gb	150Gb
Effective Reads / Flow Cell	5000M	5000M	300M/550M/1500-1800M	1500-1800M	1500-1800M	80M	500M / 100M
Average run time	24~30 hours for PE150 sequencing	20~22 hrs for PE100 sequencing	FCS: 13~37 hours FCL: 14~109 hours	15.5-50.5 hours	17/30 hours	12 hours (PE150)	9~40 hours
Min. Read Length	PE100	PE100	SE50	SE50	SE50	SE100	SE50
Max. Read Length	PE150	PE100	PE300	PE100	SE100	PE150	PE150

Third-generation sequencing



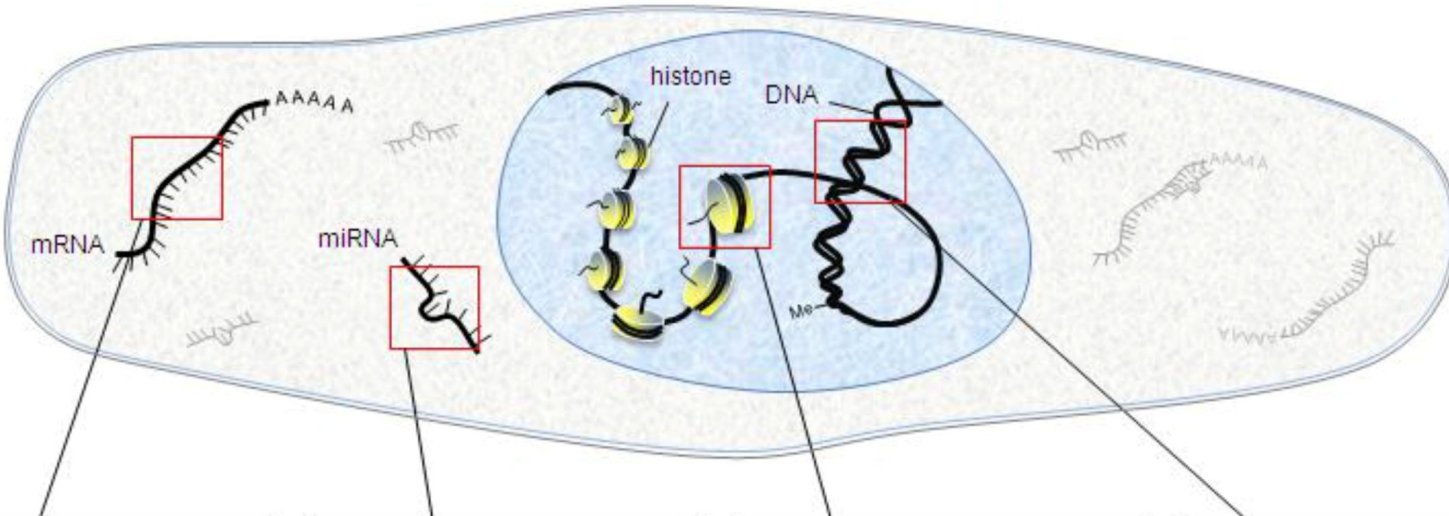
PacBio Sequencing



Nanopore technology (ONT)

- much longer reads (> Kb)
- error rate (~ 10 → 40 %)

Applications



information content

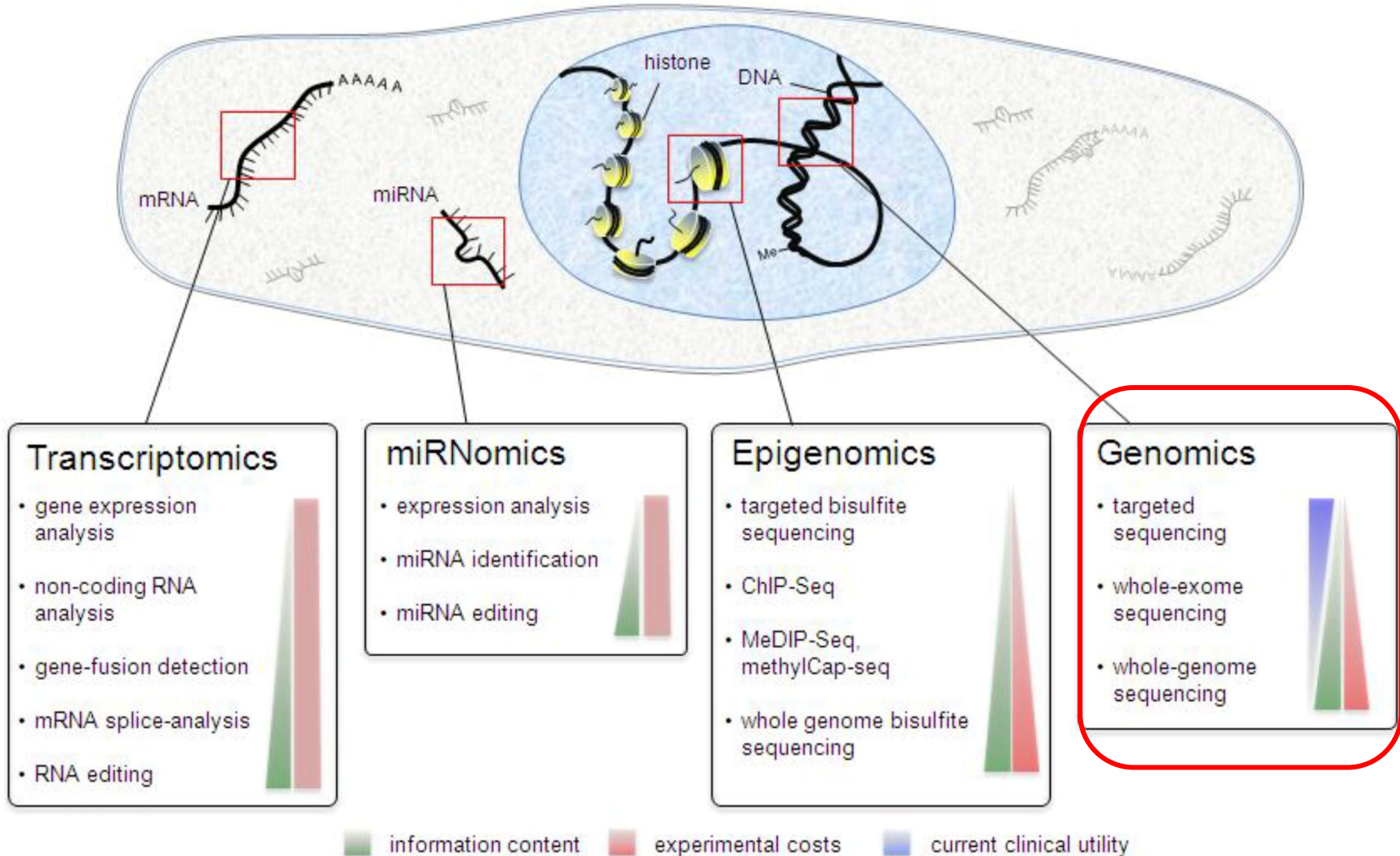


experimental costs



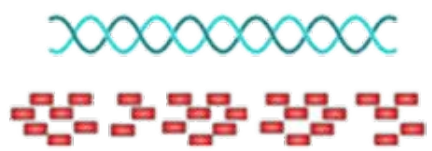
current clinical utility

Applications

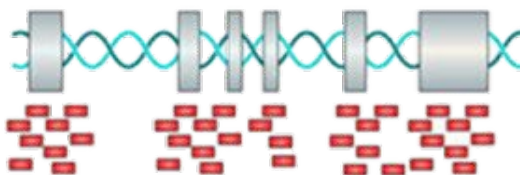


Applications : genomics (DNA-seq)

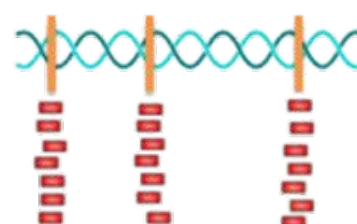
Whole genome sequencing



Whole exome sequencing



Targeted sequencing



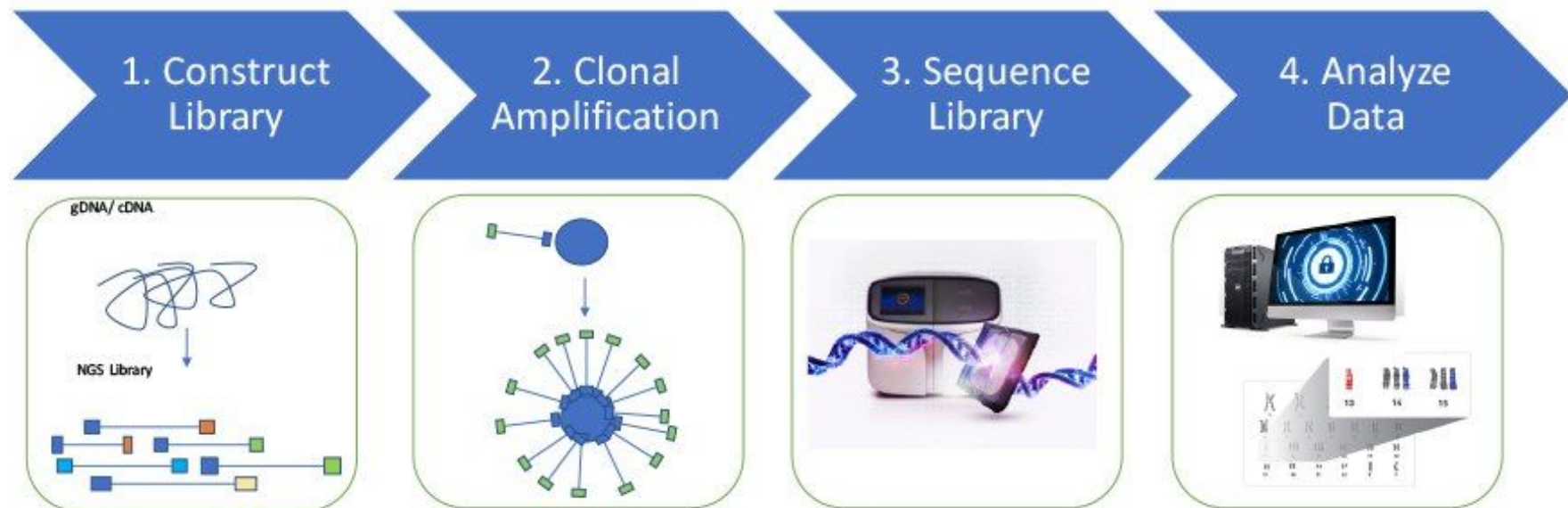
- Sequencing region: whole genome
- Sequencing Depth: >30X
- Covers everything – can identify all kinds of variants including SNPs, INDELs and SV.

- Sequencing region: whole exome
- Sequencing Depth: >50X ~ 100X
- Identify all kinds of variants including SNPs, INDELs and SV in coding region.
- Cost effective

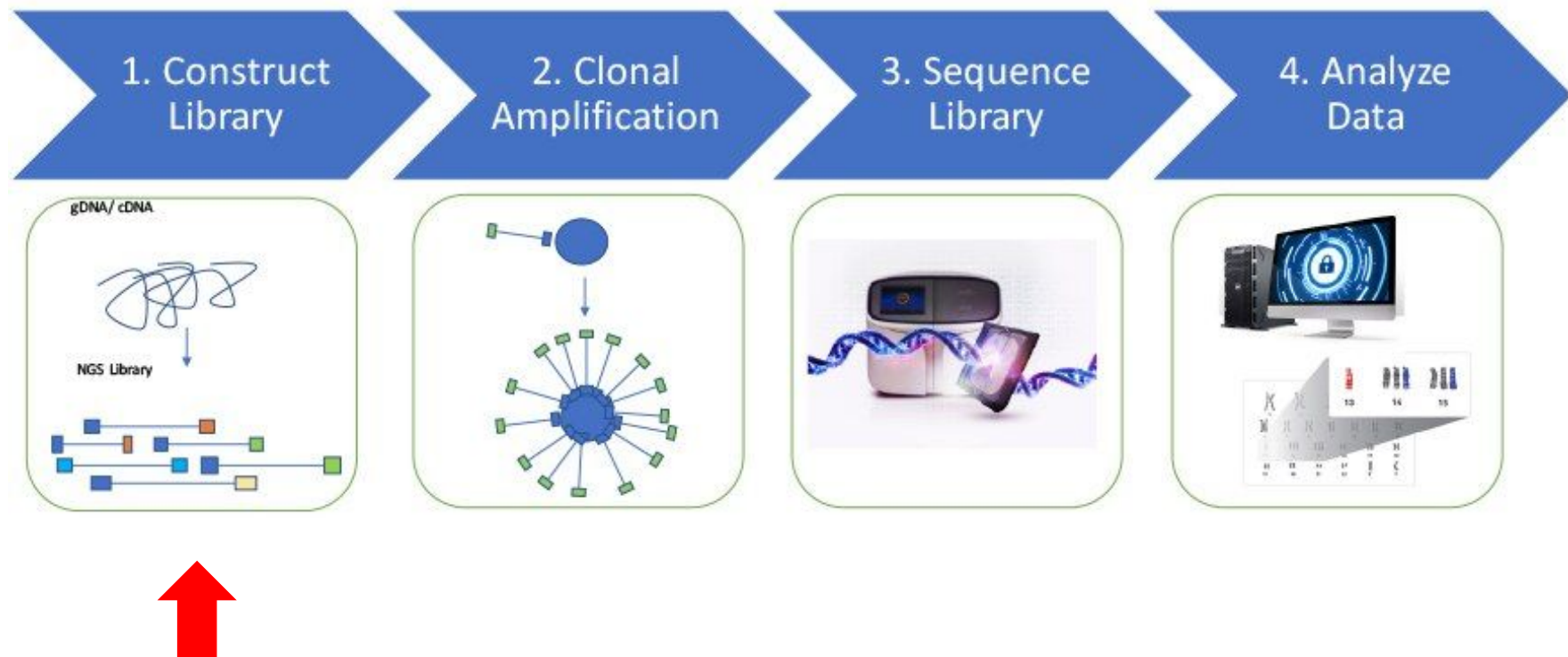
- Sequencing region: specific regions (could be customized)
- Sequencing Depth: >500X
- Identify all kinds of variants including SNPs, INDELs in specific regions
- Most Cost effective

- Targeted sequencing : rapid and cost-effective way to detect known and novel variants in selected sets of genes or genomic regions
- Whole exome sequencing : sequencing all of the protein-coding regions of genes in a genome (applications : discover rare-variants, adjacent splice-sites,...)
- Whole genome sequencing : alterations in regulatory sequences and non-coding regions, chromosomal rearrangements,

NGS workflow

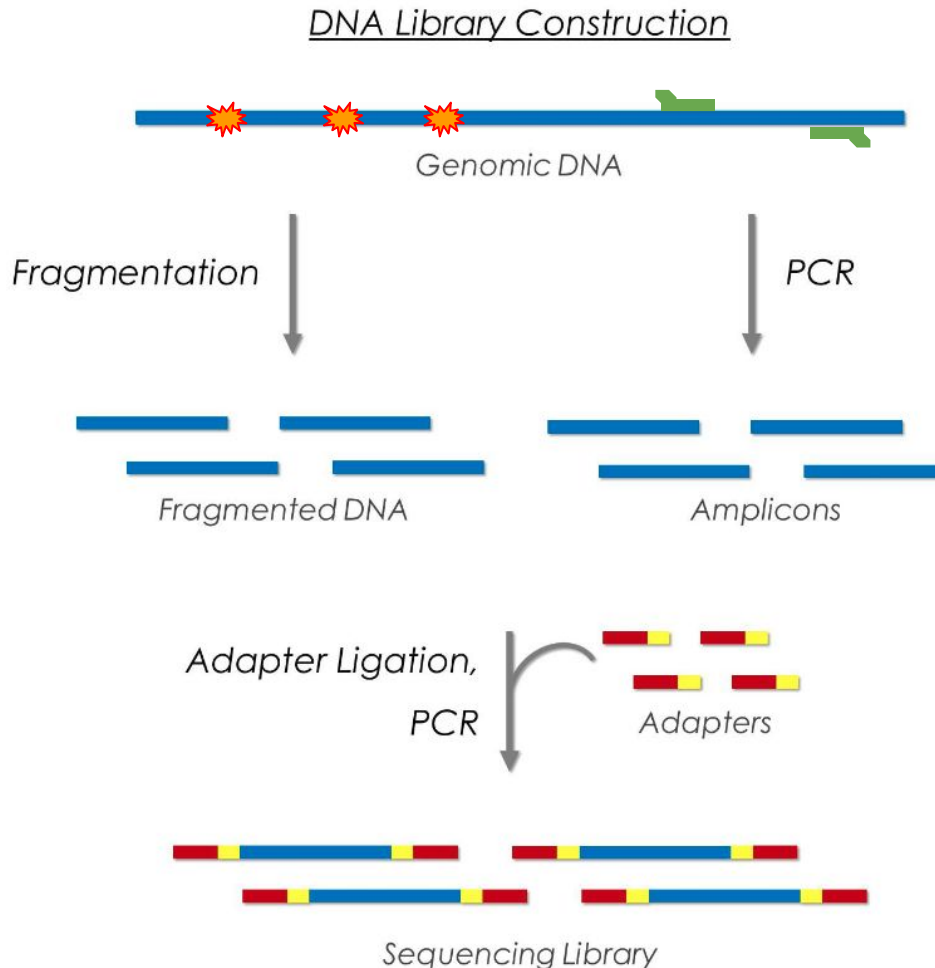


NGS workflow



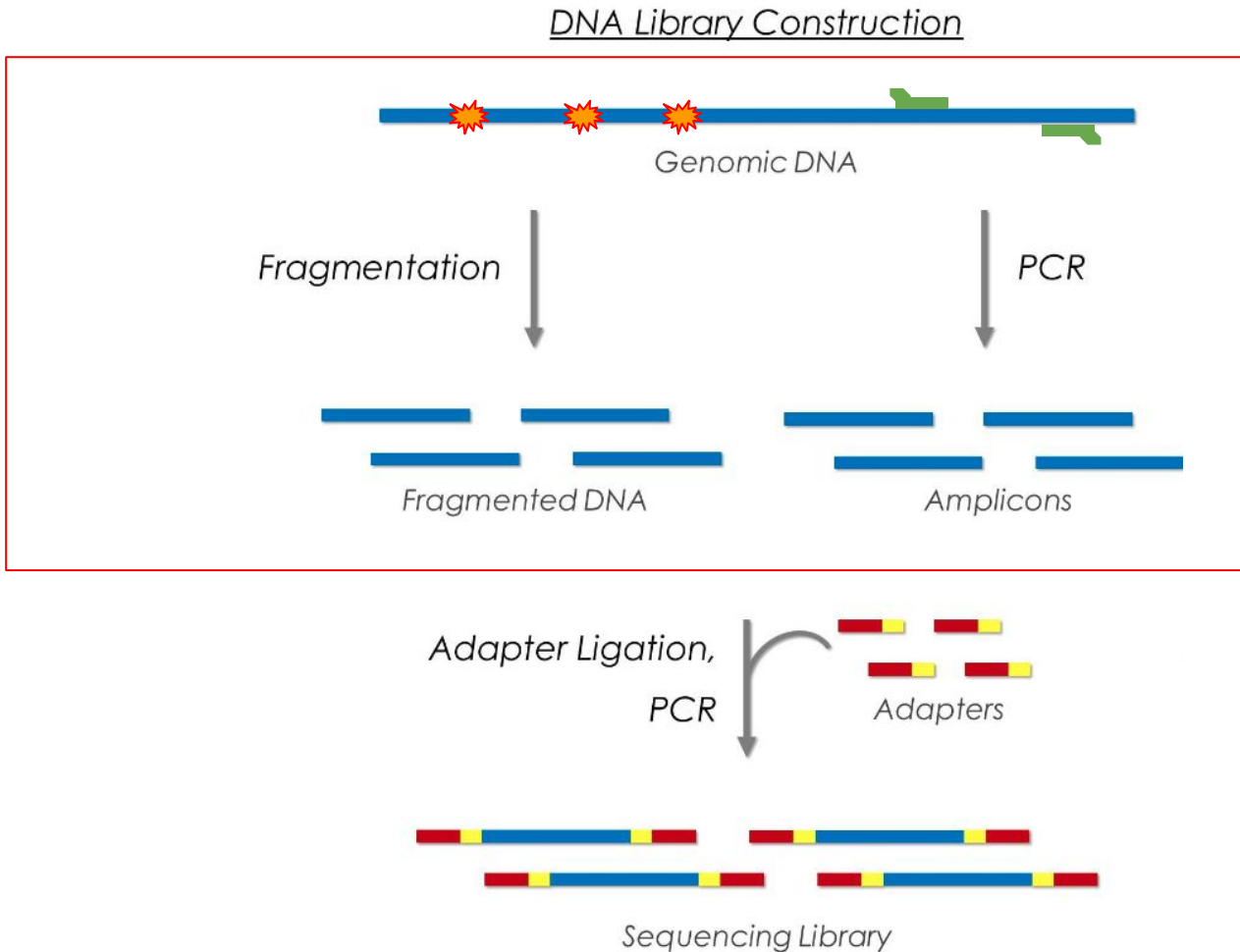
Library construction

A sequencing “library” must be created from the sample. The DNA (or cDNA) sample is processed into relatively short double-stranded fragments (100–800 bp)



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Library construction

Targeted sequencing : enrichment methods

Effective in enrichment and specificity

Simple and fast protocol

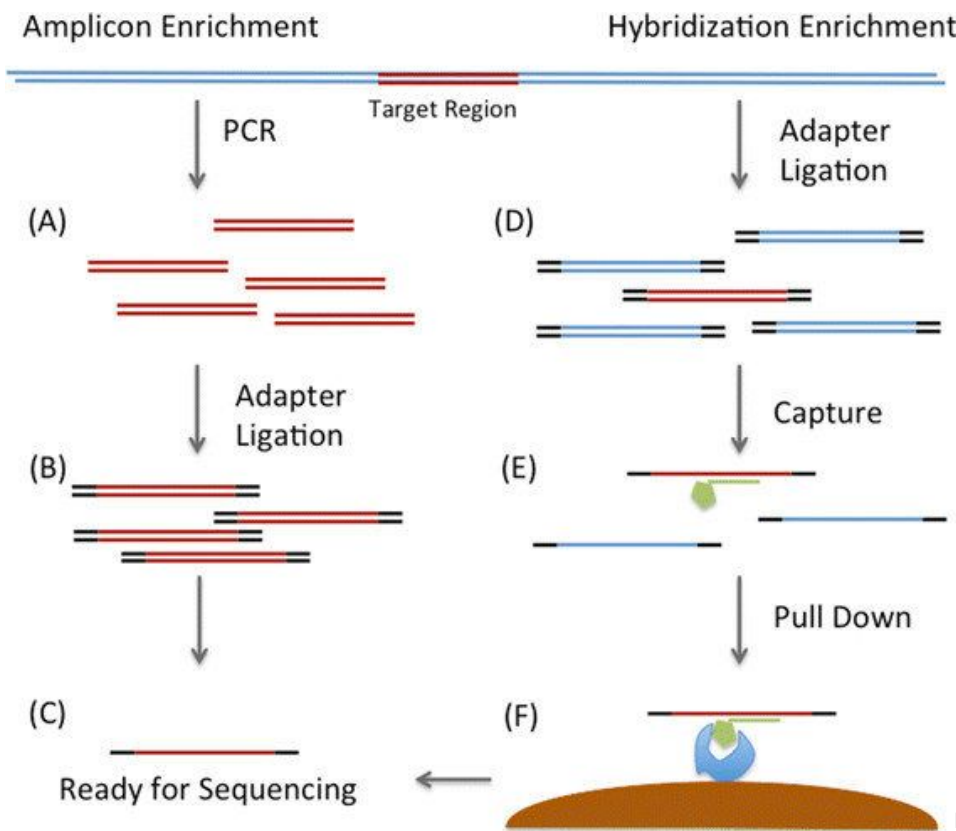
Target from Kb to Mb

Low DNA input (100 ng)

HaloPlex

AmpliSeq

...



Effective in enrichment and specificity

Complex procedure

Larger gene panels

Higher DNA input (>1 µg)

Agilent's SureSelect

Roche/Nimbelgen's SeqCap

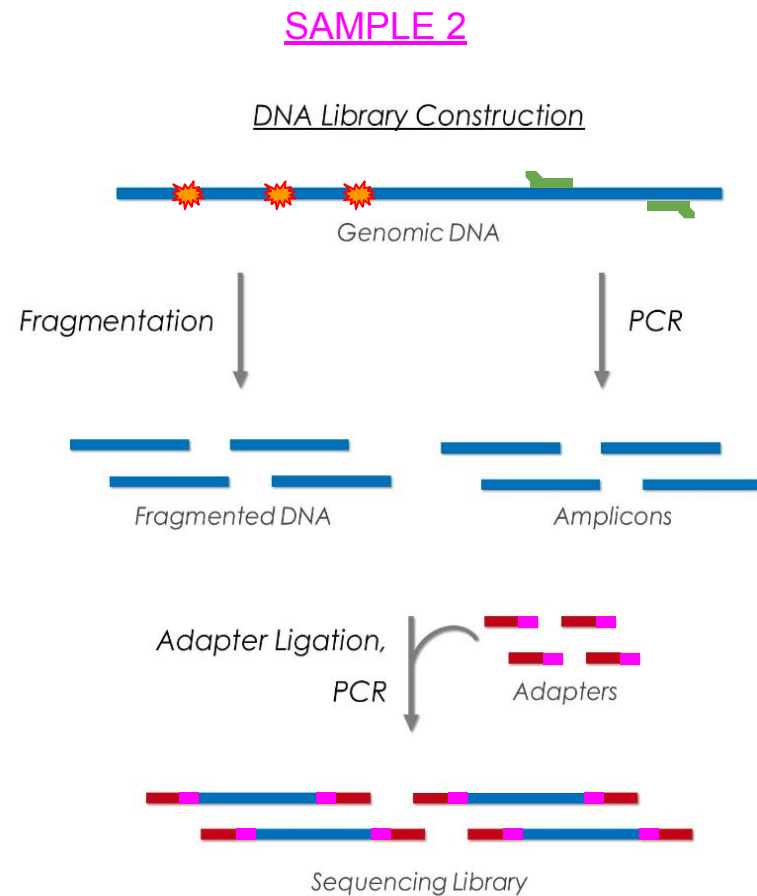
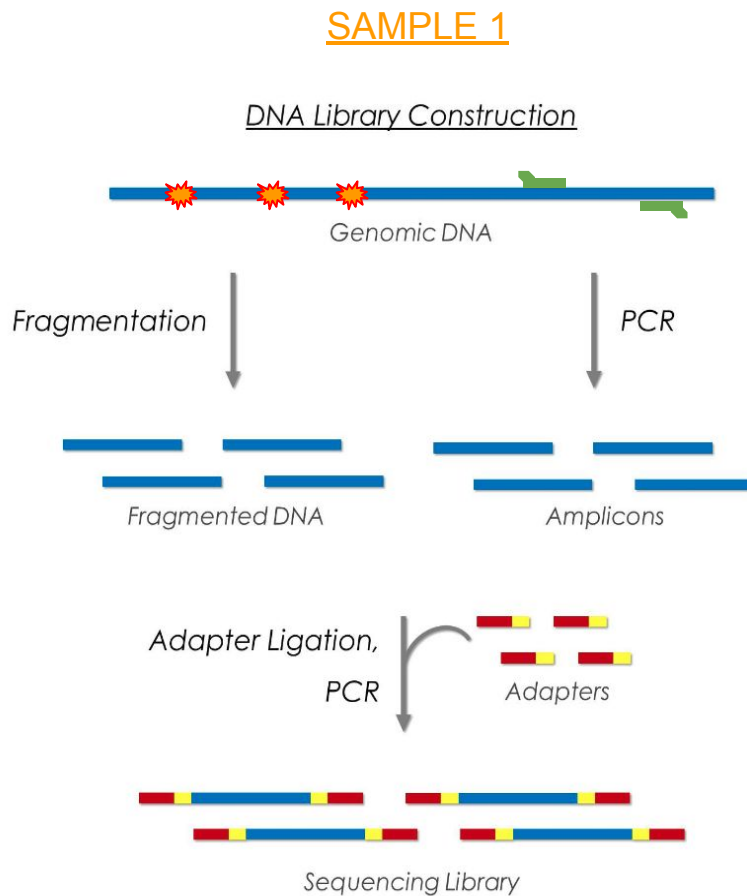
Illumina's TruSeq and Nextera

...

DOI: [10.1186/s13075-014-0490-4](https://doi.org/10.1186/s13075-014-0490-4)

Library construction

Multiplex sequencing using DNA barcoding



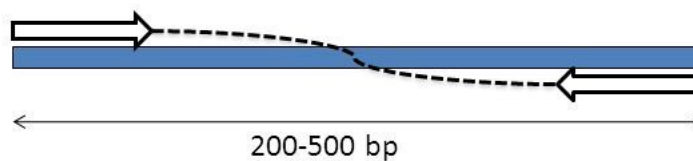
Single-end vs paired-end

- **Single-End Read:** When sequencing process only occurs in 1 direction
- **Paired-End Read:** When sequencing process occurs in both directions
- **Mate-pair Read:** Short fragments consisting of two segments that originally had a separation of several kilobases in the genome.

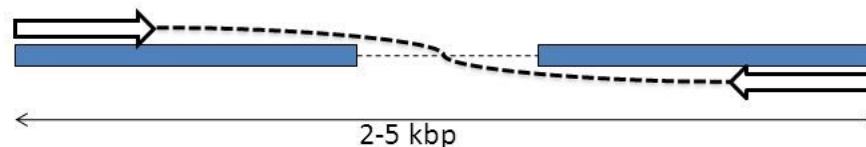
Single-End Reads - 5' or 3' (random)



Paired-End Reads - 5' and 3'



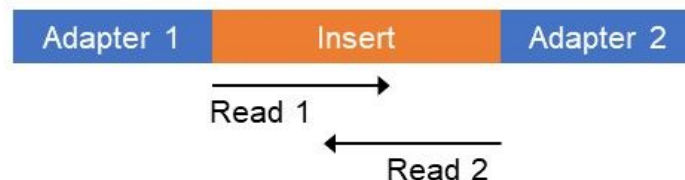
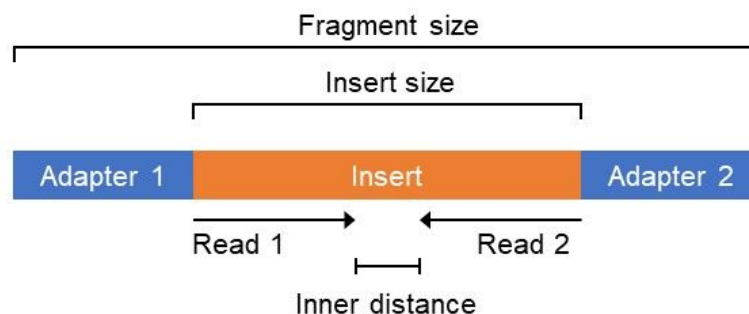
Mate-Pair Reads - 5' and 3'



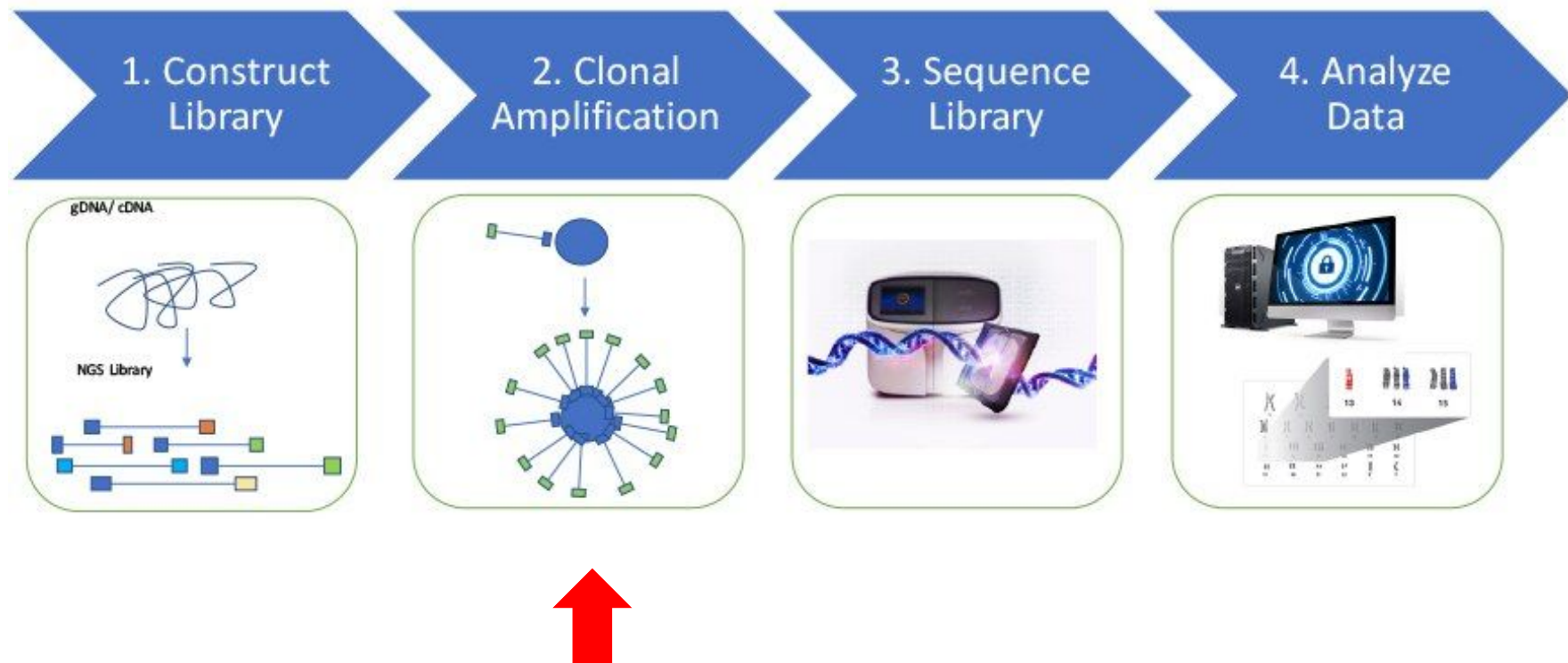
source: <http://slideplayer.com/slide/7847747/25/images/7/Types+of+Sequencing+Libraries.jpg>

Paired-end

- The **insert size** is the size of the piece of DNA of interest, without the adapters.

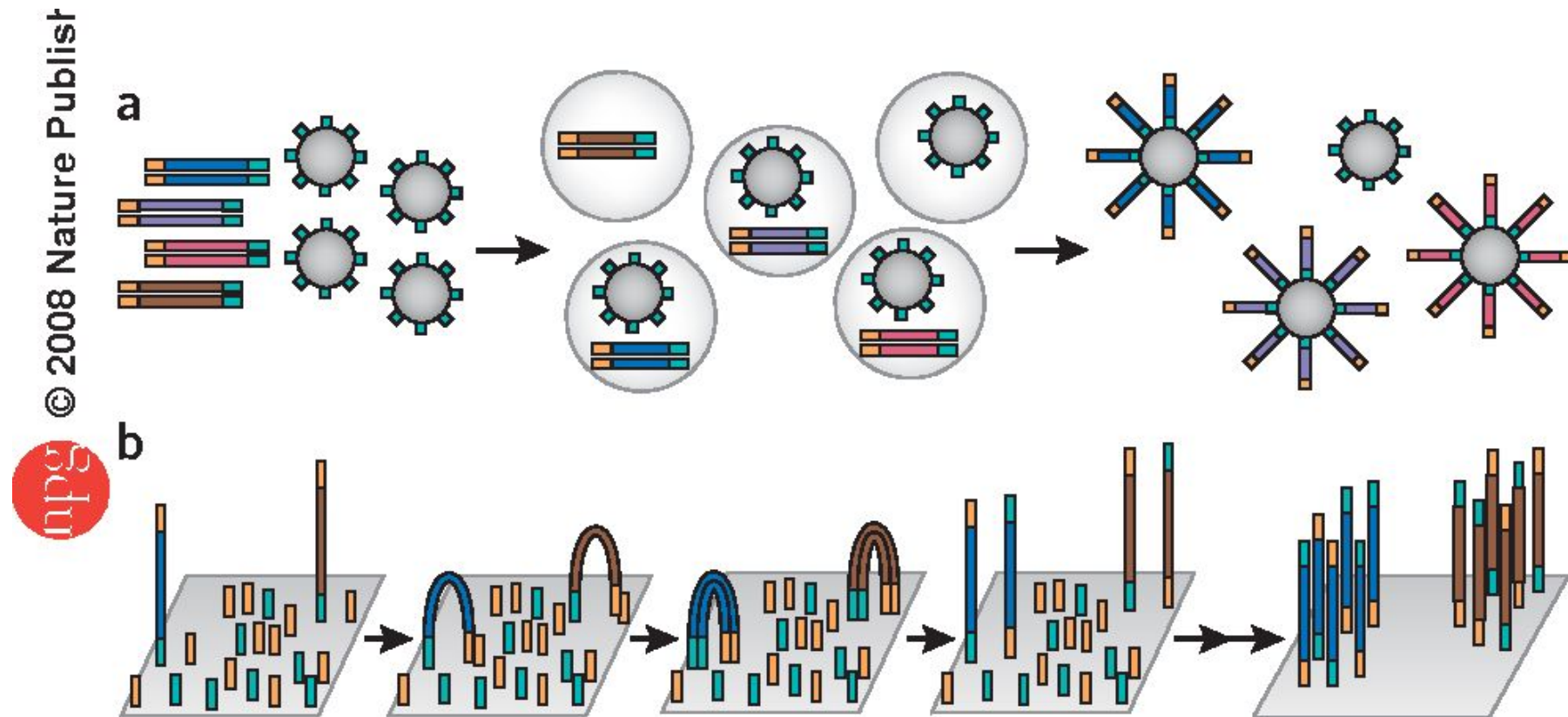


NGS workflow



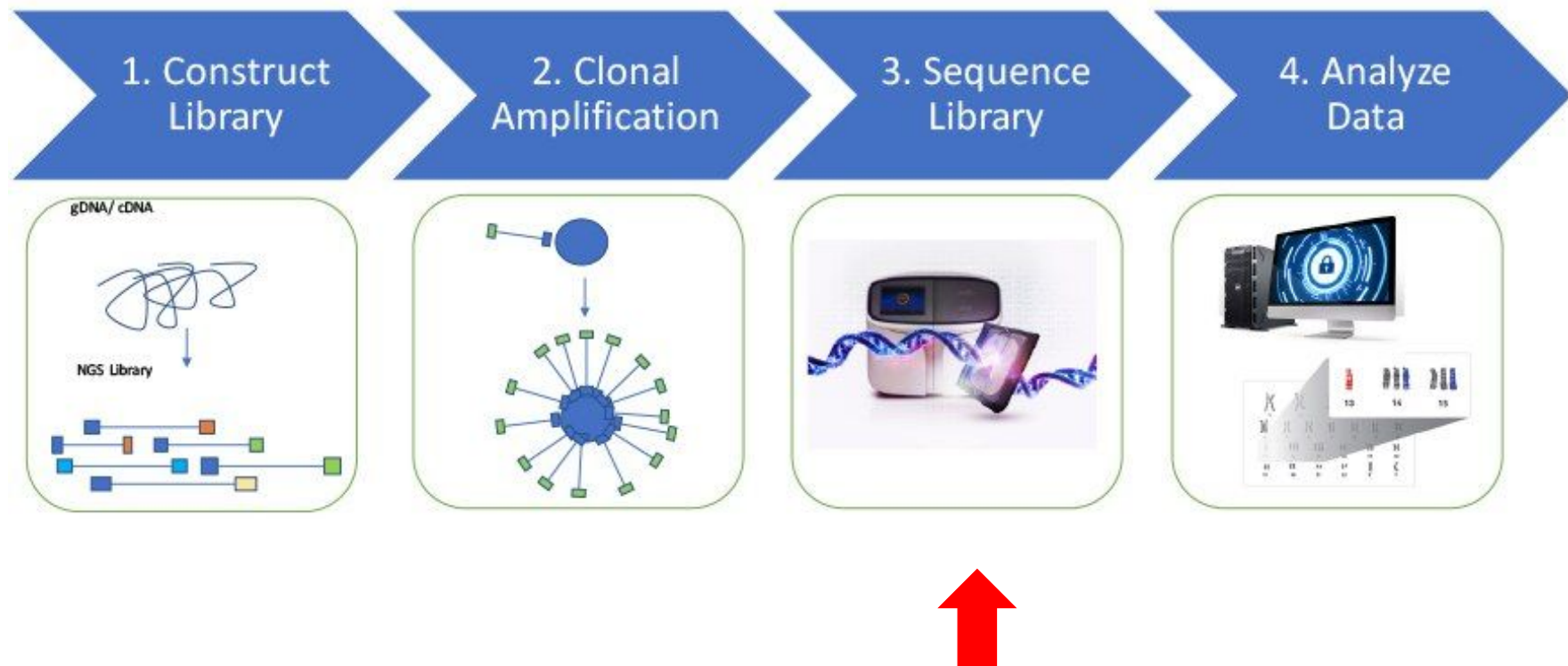
Clonal amplification

Prior to sequencing, the DNA library must be attached to a solid surface and clonally amplified to increase the signal that can be detected from each target during sequencing.



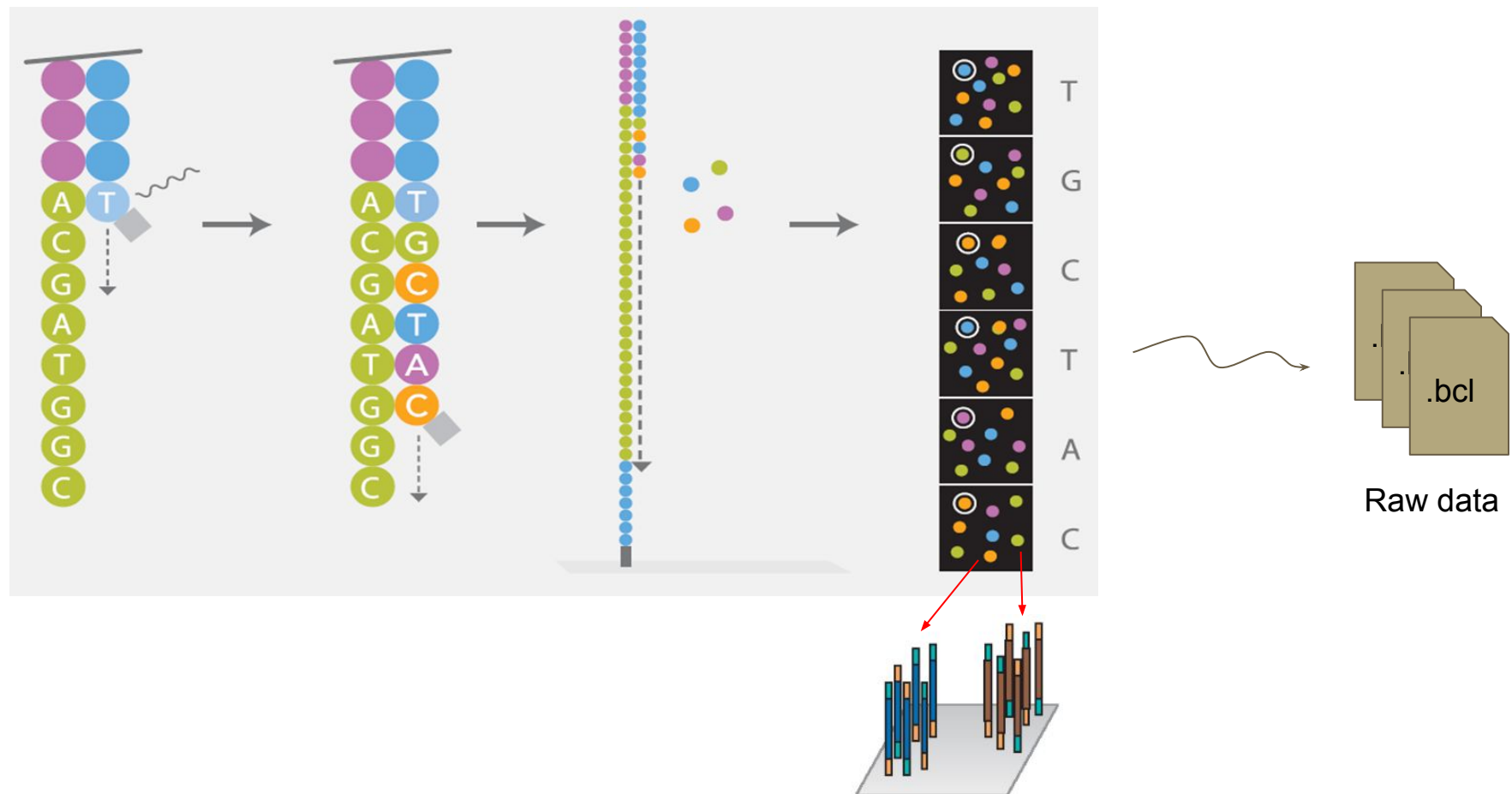
(a) thermofisher platforms rely on emulsion to amplify clonal sequencing features. (b) The Illumina technology relies on bridge PCR^{21,22} (aka 'cluster PCR') to amplify clonal sequencing features.

NGS workflow

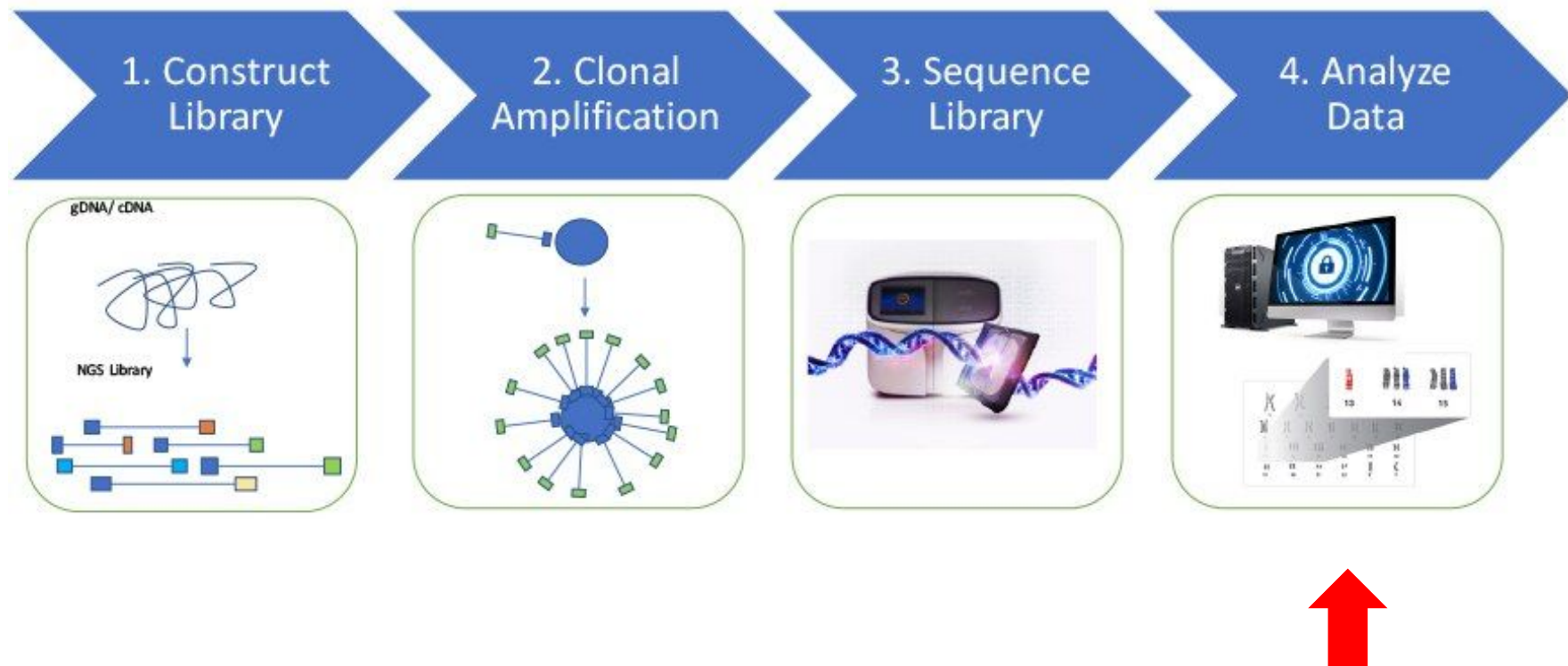


Sequencing

Illumina technology

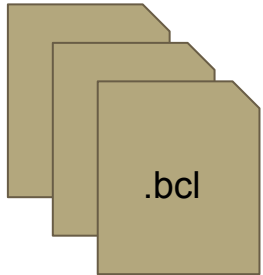


NGS workflow



Data analyses

Extracting reads, Demultiplexing



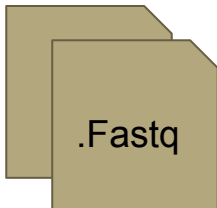
+

Sample Sheet

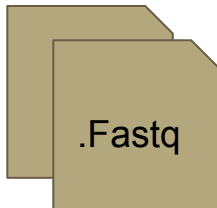
[Header]	4							
IEMFileVersion								
Experiment Name	Project1							
Date	4/16/2016							
Workflow	GenerateFASTQ							
Application	NextSeq FASTQ Only							
Assay	TruSeq LT							
Description								
Chemistry	Default							
[Reads]								
	151							
	151							
[Settings]								
Adapter	AGATCGGAAGAGCACACGTCTGAACCTCCAGTCA							
AdapterRead2	AGATCGGAAGAGCGTCGTAGGGAAAGAGTGT							
[Data]								
Sample_ID	Sample_Name	Sample_Plate	Sample_Well	I7_Index_ID	index	Sample_Project	Description	
Sample_1				A002	CGATGT			
Sample_2				A004	TGACCA			
Sample_3				A005	ACAGTG			
Sample_4				A006	GCCAAT			

bcl2fastq

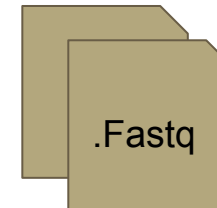
sample 1



sample 2

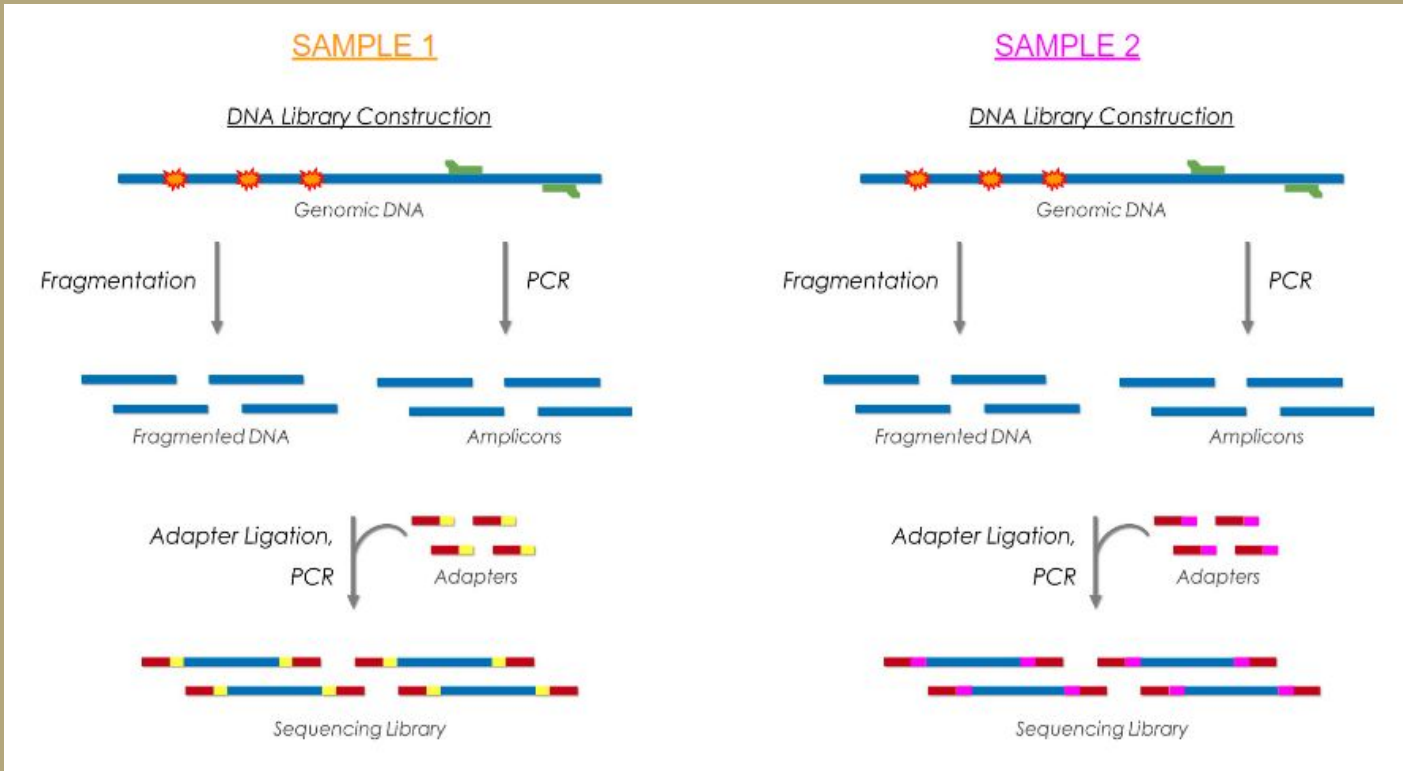


sample 3

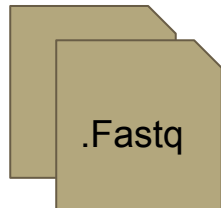


Data analyses

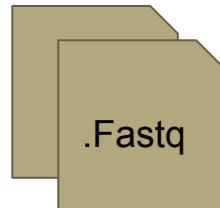
Extracting reads, Demultiplexing



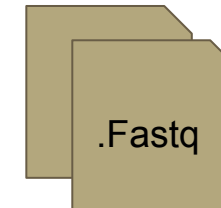
sample 1



sample 2

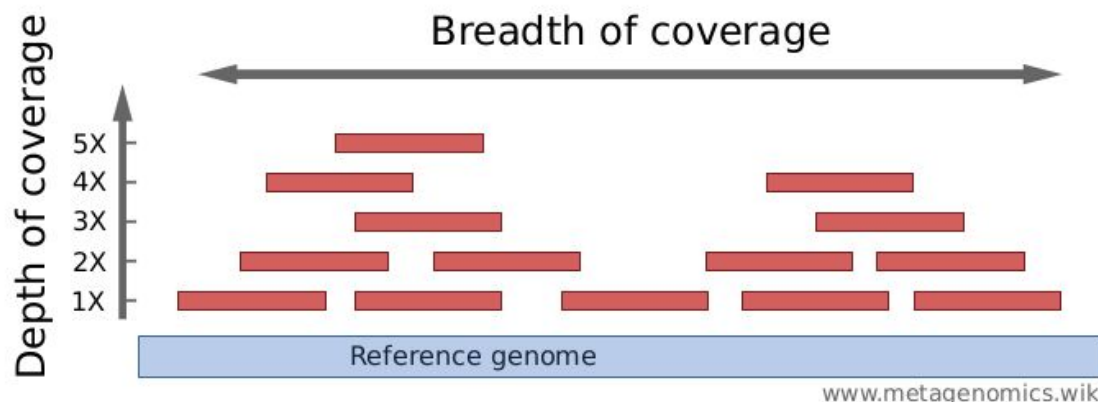


sample 3



Coverage and depth of coverage

- **Depth of coverage** = average number of reads covering a base (X)
 - Example: 30X for normal sample, 100X for tumor sample
- **(Breadth of) Coverage** = percentage of the targeted regions covered by at least X read
 - For example: 90% of a genome is covered at 1X depth; and still 40% is covered at 4X depth.



Source :

- Élodie Girard , 5ème Ecole de bioinformatique AVIESAN-IFB 2016 ,http://www.france-bioinformatique.fr/sites/default/files/V01_ITMO_2016_EG_from_fastq_to_mapping_1.pdf

Fastq file format

READ

1. Identifier 2. Sequence 4. Quality scores (as ASCII chars)

```
@SRR062641.6751359
CGCCCGGCCAATCATTGGTTTAAGTCACTAAGTTGAGGCCTTTACAGCAAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCTCATCT
+
CBLNPGJQQQJPPQPQRPQRGPPPRRQQRSPGRQQQLRRRMEPQQPMJHQQEHKMMFIIRH?SIIHKNJIKRLJJIKHEABHIFGCGGEFCGDGDCE

@SRR062634.16249693
CTAAGTTGAGGCCTTTACAGCAAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCTCATCTGTGCACCCAGCATTGCCAGAACAGGGC
+
ALKMOOOOPPQJQOPPPPPQPPPPRJQRQQQQQRPQPRQQPFQSQQPRLIMHKSNRJQORMFELRPQNQRQJQRRPQQLIRKDMKQJPN8CFDGCCB

@SRR062634.20060465
CTCCCAGCTTCCAACAGACCCTGTCCCAGCTCCCTCCAAGCTGAGTGTTGGCCTGATAACCTACCAAGTGGAGCGAGGGGAACCCGAGGACTGCCAAGGGCA
+
D?KMPQECPQQNPQIQIGR@DPERQHEKBED=HCHG8EHFDCD6<329@<:69A<6, ;<967>;=C:>AA8BBED#####="#####="#####="#####="#
```

Fastq files (Paired-end)

2 files : R1, R2

Reads1.fq

```
@ERR229776.100000840
CTAGGAAGCGTAGTCCTGGGTCATCTCCCTATTAAATACTGTTGGGAATGTTAGTA
+
BAEEAGEED96EHFE@BF><>EAAC;EBH<K<6:HJGFFHBC>DDIKG4AIHFFD@0/=

@ERR229776.100020365
CATTATTCATAGTAGCCAAAAAGTGGAAACAGTCAAAATATCCGTCACTGAATTGACC
+
1.*//,. /&((&3=;B@F860C>@51(3:).6GG-68C*:CG) #B4/=HDJ6;79)<@C/
@ERR229776.100104918
TATTTCTGAAATTCCATTAAATATTCAGACTGCAGTTGACTGCGGTAAGTAAA
+
CEEEEFFEDAEGGGFDHGFFHGIIHHIIIGKHBKJJIGHFHKILJKLEJLJJIFJMJK
```

Reads2.fq

```
@ERR229776.100000840
TTCTGGTCAGTAAGACCTAAAAGTTAAATACTAGCGATTTACACACCTAAATGATT
+
CFIEEG@FFFGFJHJ>HHKLLJIJLJILJHKAKJJKJJJJLMKJMKJJJKJ
@ERR229776.100020365
CCTAAAATGGGTGTTTCGTATATTCAACATGCTGGAACCACCACTATCTGAT
+
4B@EDFF=(/CHBHEHCE6@ED8E@I6HJB6E:6%@C46FFIBGCIGKD,DN=CBBE@
@ERR229776.100104918
TCTTTCTTTGTTTTCTGAGATGTCTTGTTGAGGTCTGTTATG
+
CFIGGGKHHHFHHFIJIIJKLIIHJIIKJJKIJKLLKFJJMHJJLFJMJKJJ
```

1 interleaved paired file

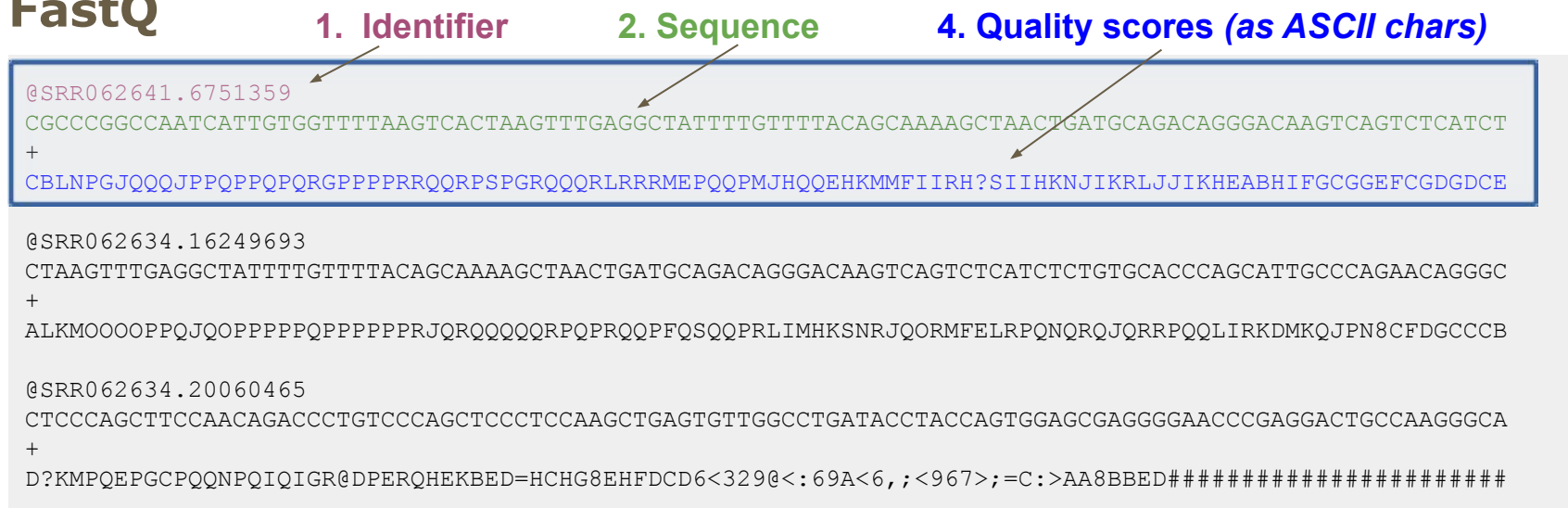
Reads.fq

```
@SRR531199.1 ILLUMINA_0130:3:1101:1249:1993 length=101
TTTCAGAGTAGTTGGTACCCAATTGGAAGATGTGACCCACTTCGATACCGCGCTTGAG
+
dfffffffdfeffdadfffffeeffdeffefffffffddeefeydfefefe[e
@SRR531199.1 ILLUMINA_0130:3:1101:1249:1993 length=99
ANNNNNNCTCGGTATNAACTGGGNNNNGATGTTGAACCTGGTAAAGTCGAAGATCTG
+
BBBBBBSZTUVWO]YB_[cbabbWB BBBBVUUUggadcdedbedcddfffdegegegef
@SRR531199.2 ILLUMINA_0130:3:1101:1463:1964 length=101
NTGAGTAGCTCAATGCGCTGACGCCAATAGCTATACCAACGACTGCCAGATTATGTT
+
BXSSRU[X[Wcc_cccccccccccc_cccccccccccc_ccccccccccccccccccccc
@SRR531199.2 ILLUMINA_0130:3:1101:1463:1964 length=99
AAGTGACCCATCGCGATAAGTGCCTGCGCAGTAAANAGCANCTGTTNGATGCTGGCTTA
+
ggggggggggggggggggggf g f gggggggggggggggg^BbbbaBbbaz]BZ[ccccfggggg
@SRR531199.3 ILLUMINA_0130:3:1101:1366:1970 length=101
NAAGTCGCGGCACCCCTATCGTGGCTTCCGGCGTACGCCATTCAATGCGGCCGCC
+
B[[X[YVcc_cccc_cc_____] [[V^^^^^V[[SXWUX[\]]Z^^^B
@SRR531199.3 ILLUMINA_0130:3:1101:1366:1970 length=99
TGGTCAATACAAGCCGAATACCTGCATCATGCGNGGAANAATTGCGCGCCGTTTC
+
ggfegggggggdeggggf g cgggagggggg gega^Bb`^]B[Y[[Zffffh_afefe
```

Sequencing reads file formats

FastQ

READ



FastA

```
>SRR062641.6751359
CGCCCGGCCAATCATTGTGGTTTAAGTCACTAAGTTGAGGCTATTTGTTTACAGCAAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCTCATCT

>SRR062634.16249693
CTAAGTTGAGGCTATTTGTTTACAGCAAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCTCATCTGTGCACCCAGCATTGCCAGAACAGGGC

>SRR062634.20060465
CTCCCCAGCTTCAAACAGACCCGTCCCAGCTCCCTCAAGCTGAGTGTGGCCTGATACTACCAGTGGAGCGAGGGGAACCCGAGGACTGCCAAGGGCA
```

Mais aussi: FAST5, BAM, ...

Jour 1 :

- NGS Introduction
- Reads Quality Control

Reads quality

- Errors when reading bases
 - Depends on sequencing technologie
 - Error rate tends to increase with read size
- ⇒ For each position in the read
- One base (A/T/C/G)
 - One error probability

Phred Quality Score (for a base)

Phred quality scores Q : logarithmically related to the base-calling error probabilities P

$$Q = -10 \log_{10} P$$

Phred Quality Score	Probability of incorrect base call	Base call accuracy
10	1 in 10	90%
20	1 in 100	99%
30	1 in 1000	99.9%
40	1 in 10,000	99.99%
50	1 in 100,000	99.999%
60	1 in 1,000,000	99.9999%

Quality score encoding: ASCII table

READ

	1. Identifier	2. Sequence	4. Quality scores (as ASCII chars)
	@SRR062641.6751359	CGCCCGGCCAATCATTGTGGTTAACGTTGAGGCTATTTGTTTACAGCAAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCTCATCT	
	+	CBLNPGJQQQJPPQPQPRGPPPQRQRLRRMEEQQPMJHQEHKMMFIIRH?SIIHKNJIKRLJJIKHEABHIFGCGGEFCGDGDCE	
	@SRR062634.16249693	CTAAGTTGAGGCTATTTGTTTACAGCAAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCTCATCTGTGCACCCAGCATTGCCAGAACAGGGC	
	+	ALKMOOOOPPQJQOPPPPPQPPPPRJQRQQQQQRPQPRQQFQSQQPRLIMHKSNRJQORMFELRPQNQRQJQRRPQLIRKDMKQJPN8CFDGCCC	
	@SRR062634.20060465	CTCCCAGCTTCCAACAGACCCTGTCCCAGCTCCCTCCAAGCTGAGTGTGGCCTGATACTACCAGTGGAGCGAGGGGAACCCGAGGACTGCCAAGGGCA	
	+	D?KMPQEPEGCPQQNPQIIGR@DPERQHEKBED=HCHG8EHFDCD6<329@<:69A<6, ;<967>;=C:>AA8BBED#####	

Decimal	Hex	Char	Decimal	Hex	Char	Decimal	Hex	Char	Decimal	Hex	Char
0	0	[NULL]	32	20	[SPACE]	64	40	@	96	60	`
1	1	[START OF HEADING]	33	21	!	65	41	A	97	61	a
2	2	[START OF TEXT]	34	22	"	66	42	B	98	62	b
3	3	[END OF TEXT]	35	23	#	67	43	C	99	63	c
4	4	[END OF TRANSMISSION]	36	24	\$	68	44	D	100	64	d
5	5	[ENQUIRY]	37	25	%	69	45	E	101	65	e
6	6	[ACKNOWLEDGE]	38	26	&	70	46	F	102	66	f
7	7	[BELL]	39	27	'	71	47	G	103	67	g
8	8	[BACKSPACE]	40	28	(72	48	H	104	68	h
9	9	[HORIZONTAL TAB]	41	29)	73	49	I	105	69	i
10	A	[LINE FEED]	42	2A	*	74	4A	J	106	6A	j
11	B	[VERTICAL TAB]	43	2B	+	75	4B	K	107	6B	k
12	C	[FORM FEED]	44	2C	,	76	4C	L	108	6C	l
13	D	[CARRIAGE RETURN]	45	2D	-	77	4D	M	109	6D	m
14	E	[SHIFT OUT]	46	2E	.	78	4E	N	110	6E	n
15	F	[SHIFT IN]	47	2F	/	79	4F	O	111	6F	o
16	10	[DATA LINK ESCAPE]	48	30	0	80	50	P	112	70	p
17	11	[DEVICE CONTROL 1]	49	31	1	81	51	Q	113	71	q
18	12	[DEVICE CONTROL 2]	50	32	2	82	52	R	114	72	r

Quality score encoding

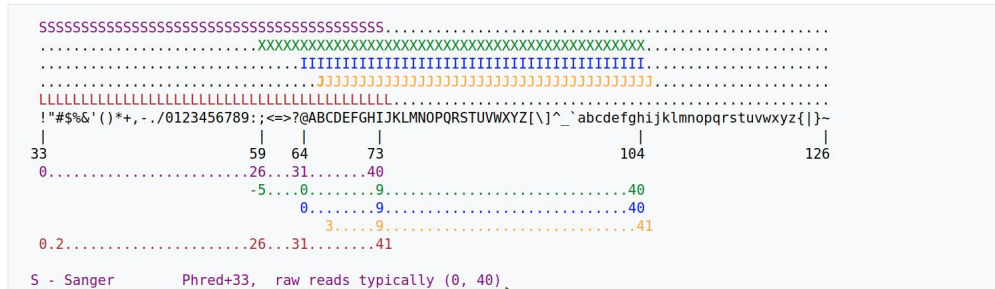
For history reasons, more than one coding convention

S - Sanger Phred+33, raw reads typically (0, 40)
 X - Solexa Solexa+64, raw reads typically (-5, 40)
 I - Illumina 1.3+ Phred+64, raw reads typically (0, 40)
 J - Illumina 1.5+ Phred+64, raw reads typically (3, 41)
 with 0=unused, 1=unused, 2=Read Segment Quality Control Indicator (**bold**)
 (Note: See discussion above).
 L - Illumina 1.8+ Phred+33, raw reads typically (0, 41)
 P - PacBio Phred+33, HiFi reads typically (0, 93)

Source : <https://fr.wikipedia.org/wiki/FASTQ>

Galaxy : Always uses Sanger coding
=> conversion tool (*groomer*)

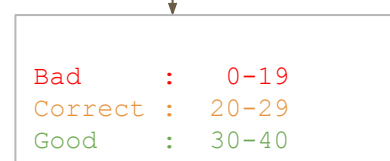
Example for score interpretation using sanger encoding



Phred Quality Score	Probability of incorrect base call	Base call accuracy
10	1 in 10	90%
20	1 in 100	99%
30	1 in 1000	99.9%
40	1 in 10,000	99.99%
50	1 in 100,000	99.999%
60	1 in 1,000,000	99.9999%

S - Sanger Phred+33

```
@SEQ:ID
ACTGTACGATCGATCGCATGATCAGTACGTCGTACCAAGAT
+
!"#$%&'() *+, -./0123456789:;<=>?@ABCDEFGHI
|           |           |           |
0.....1.....2.....3.....4
01234567890123456789012345678901234567890
```



Quality Control (QC)

Quality Control (QC) is important to:

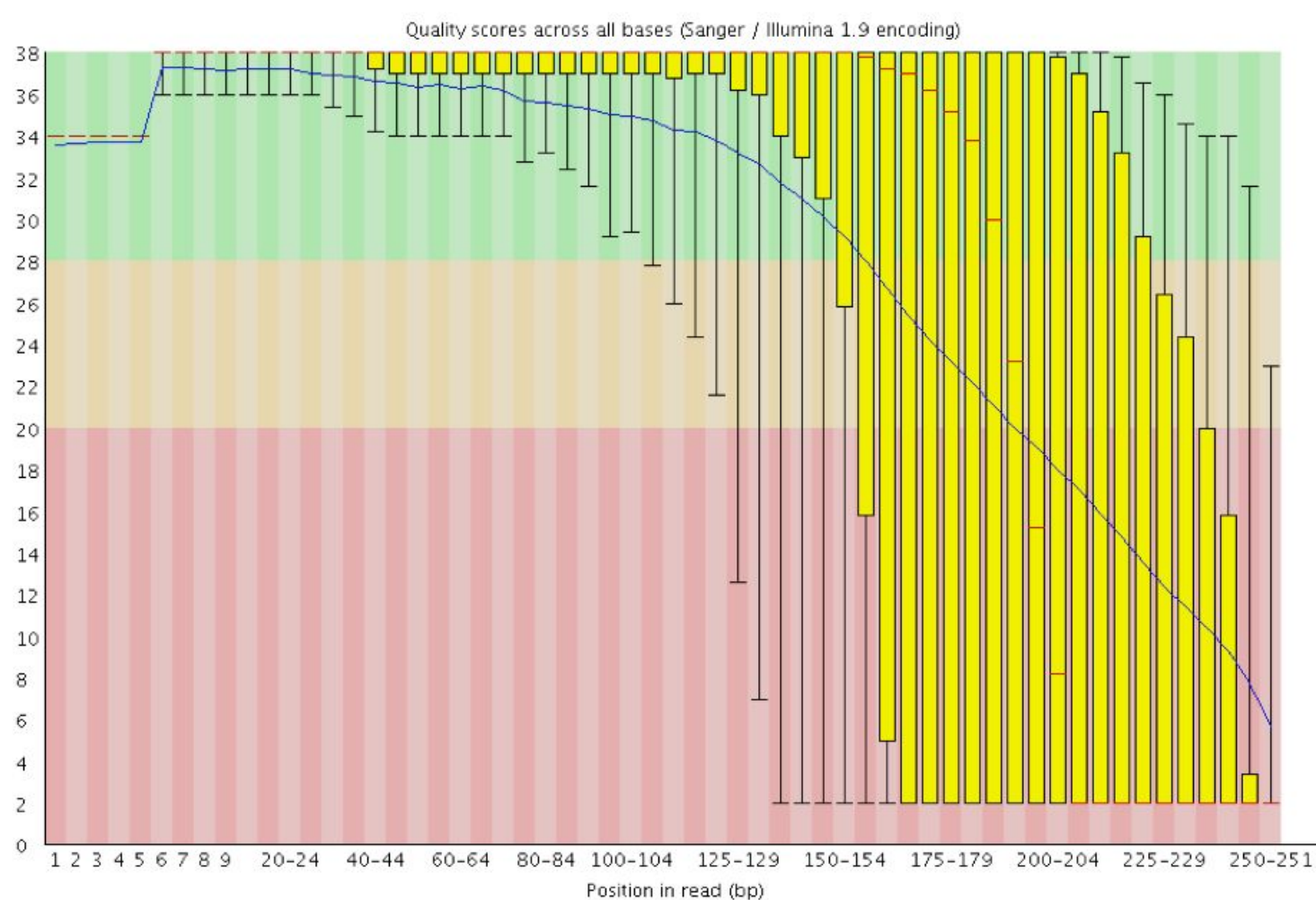
- Check if your sample sequencing went well
- Know when you need to sequence again (sequencing platform QC fail)
- Identify potential problems that can be fixed, or not
- Follow the impact of preprocessing steps

⇒ FastQC (<https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>)

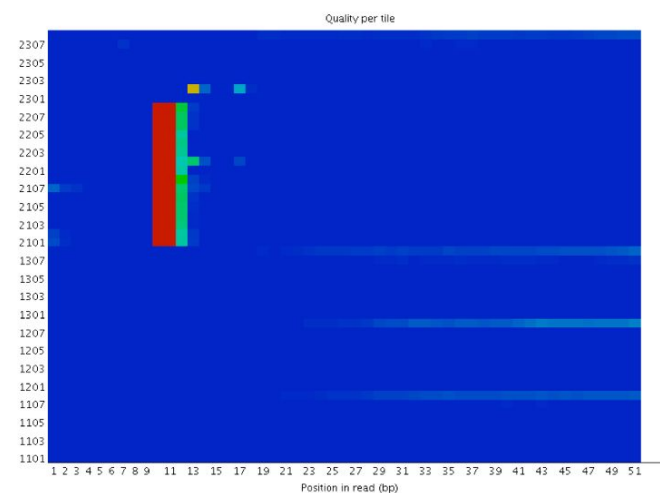
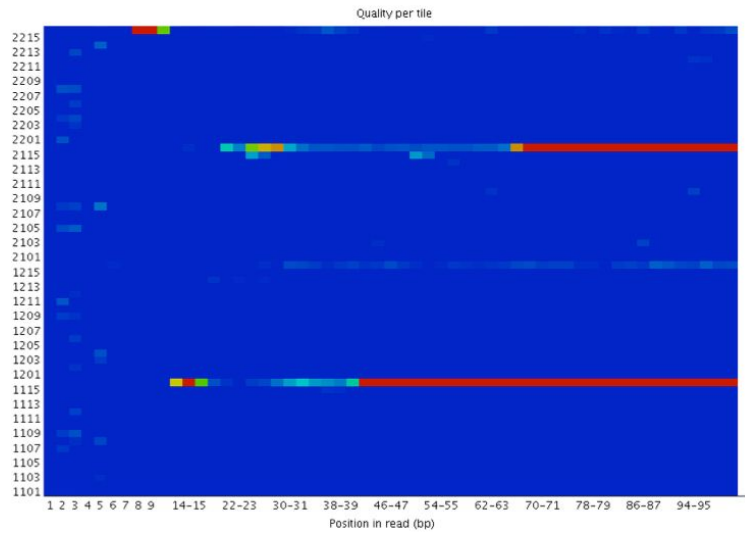
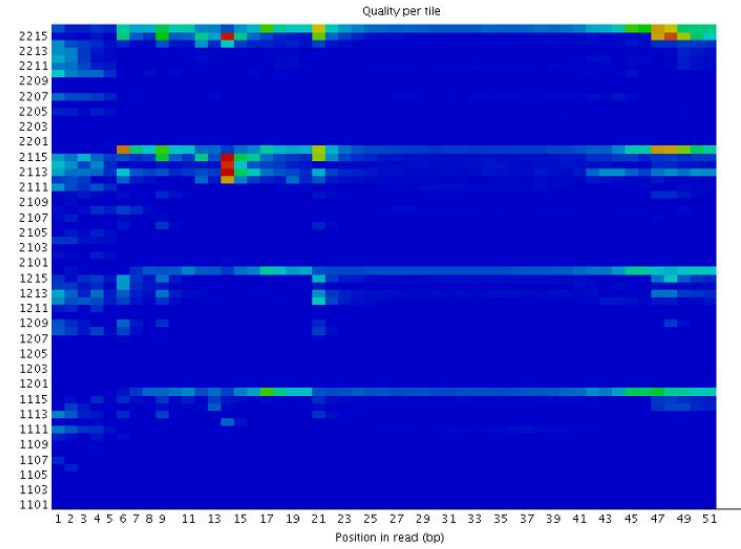
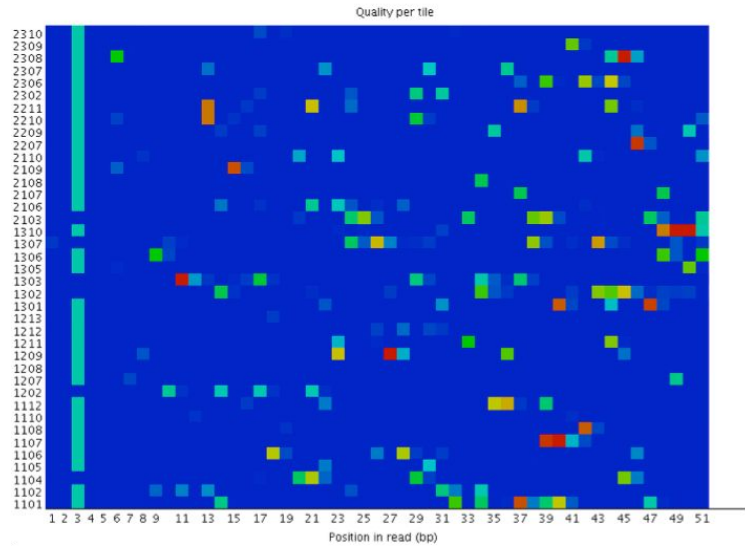
+ MultiQC (<https://multiqc.info/>) when comparing multiple datasets

Loss of base call accuracy with increasing sequencing cycles

Source: <https://sequencing.qcfail.com>



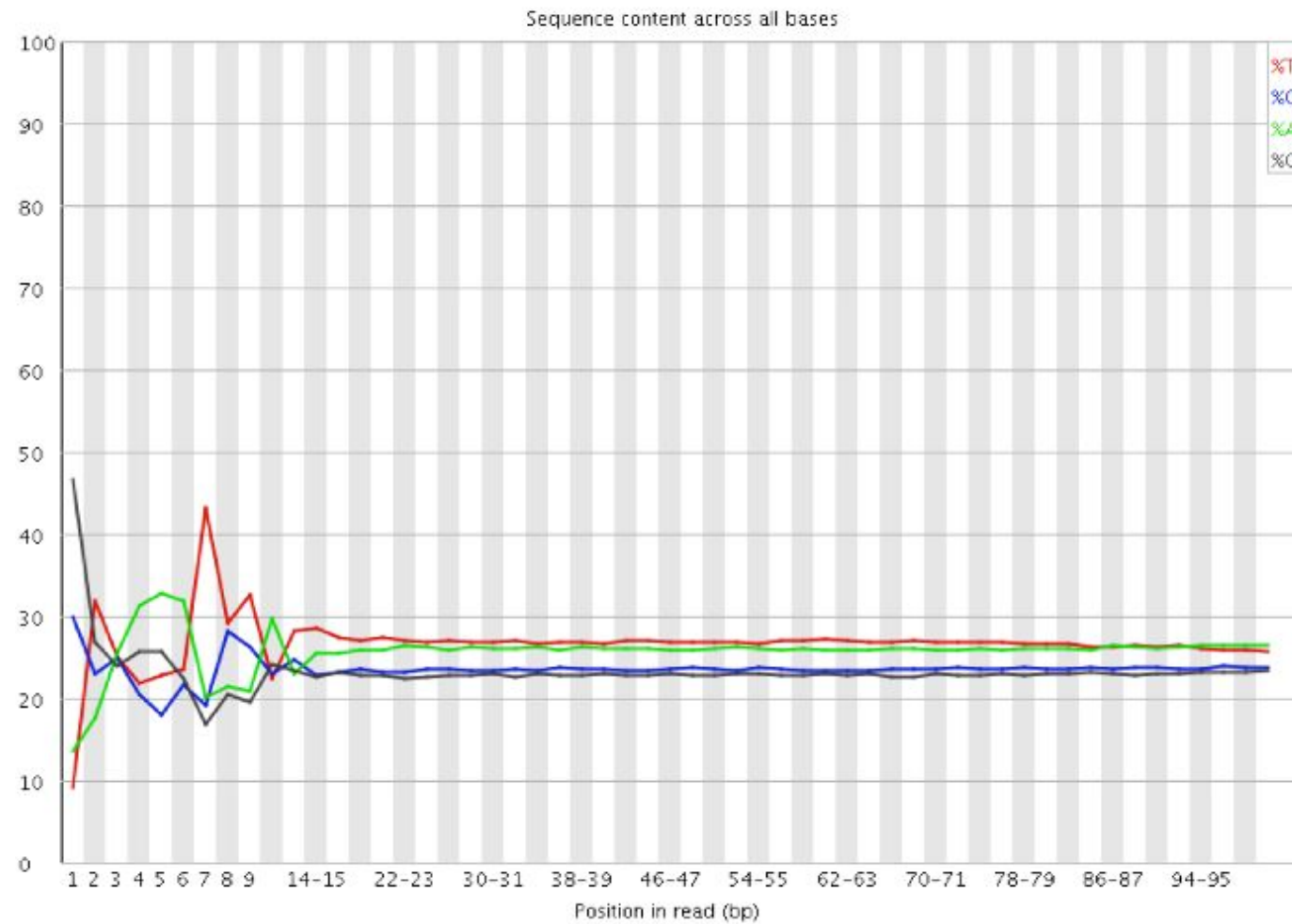
Position specific failures of flowcells



Source: <https://sequencing.qcfail.com> 15

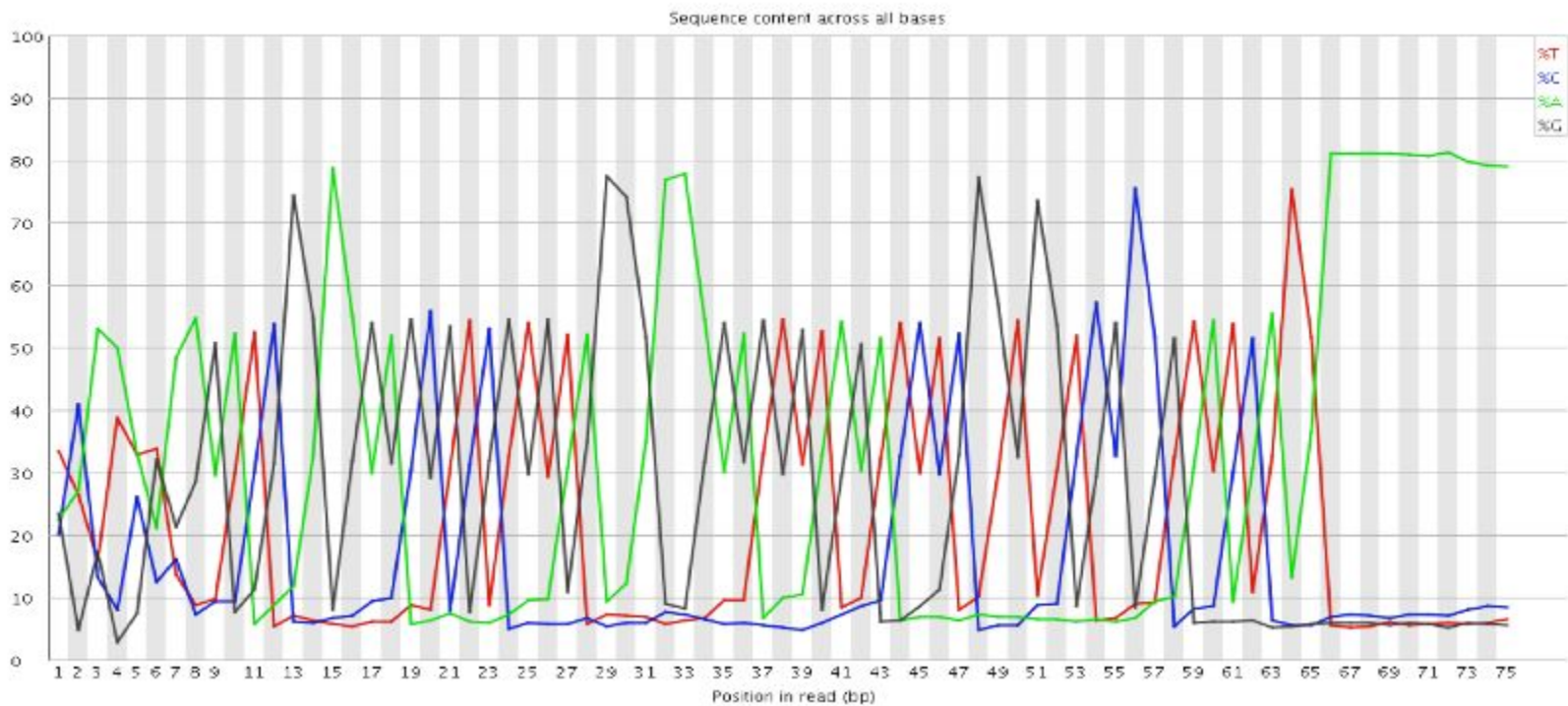
Positional sequence bias in random primed libraries

Source: <https://sequencing.qcfail.com>



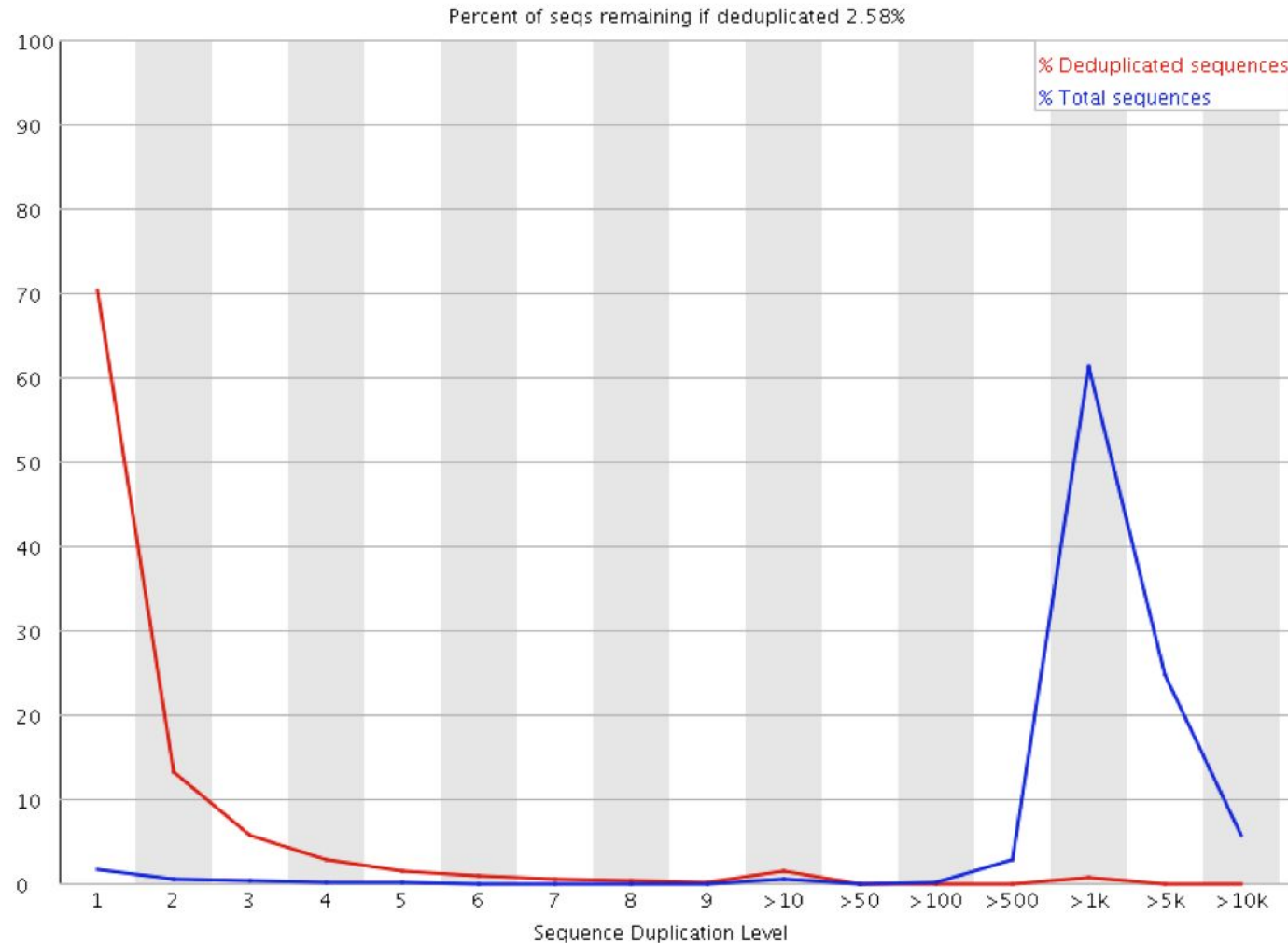
Contamination with adapter dimers

Source: <https://sequencing.qcfail.com>

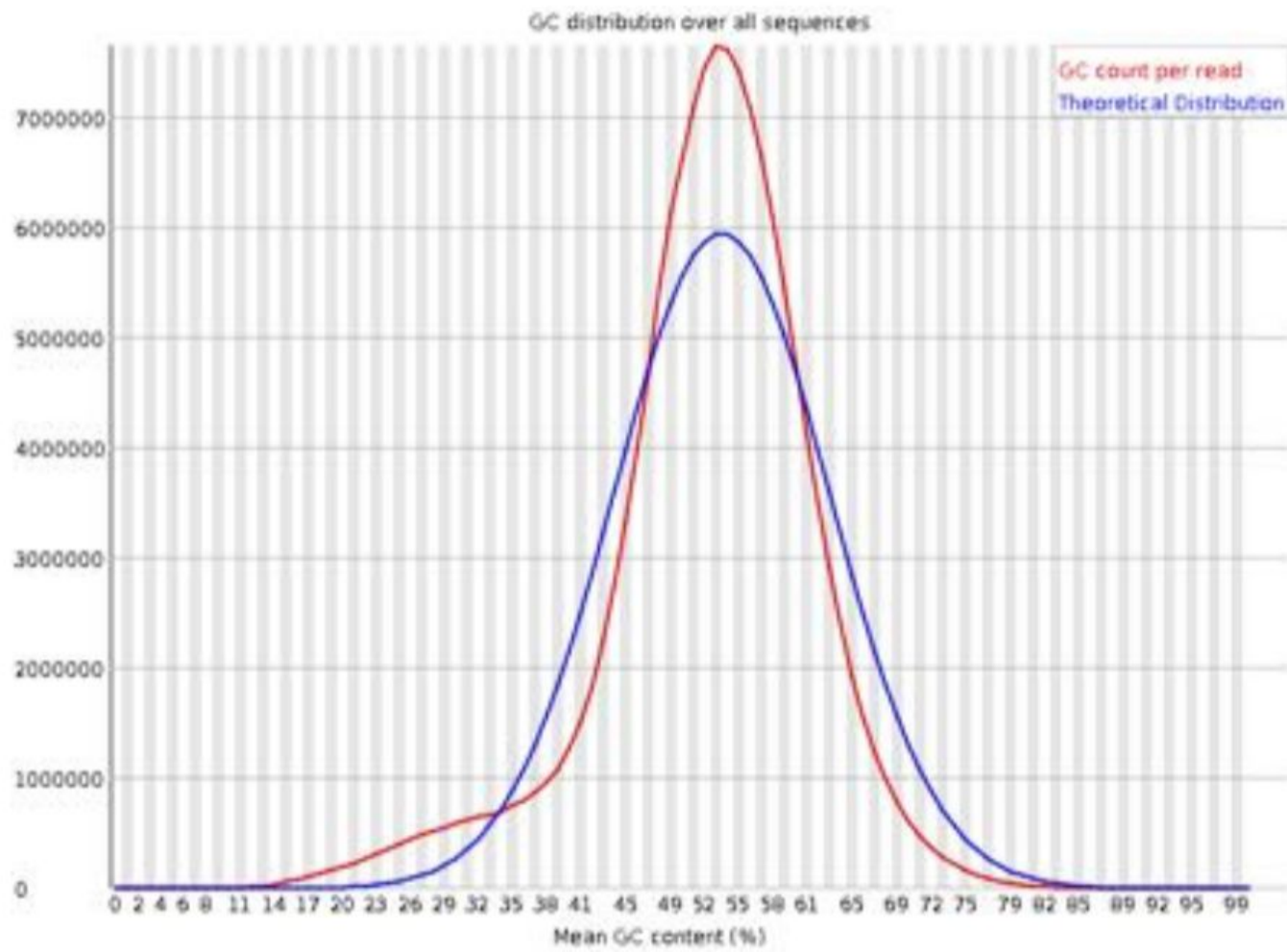


Libraries contain technical duplication

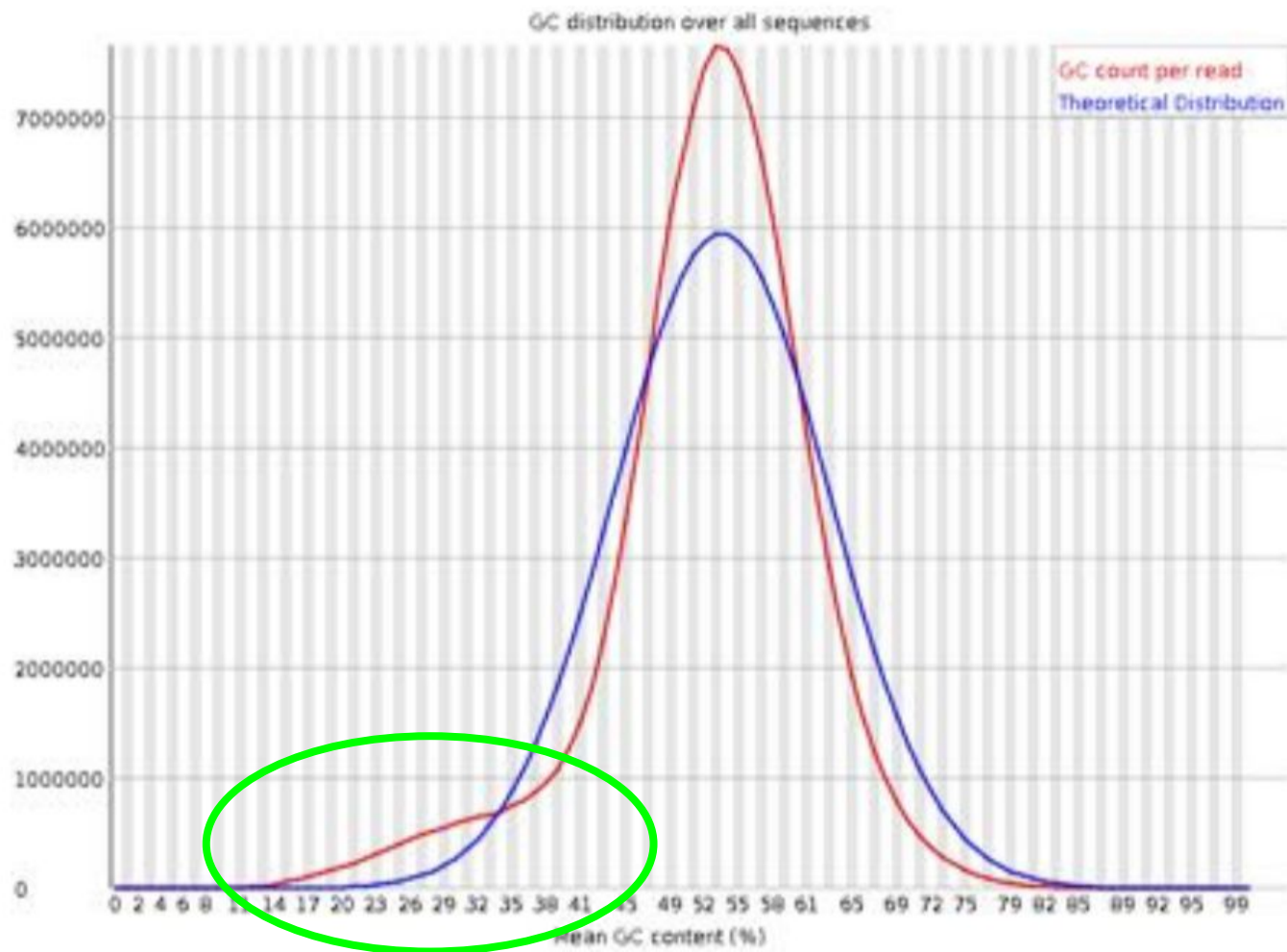
Source: <https://sequencing.qcfail.com>



GC content / Contamination ?

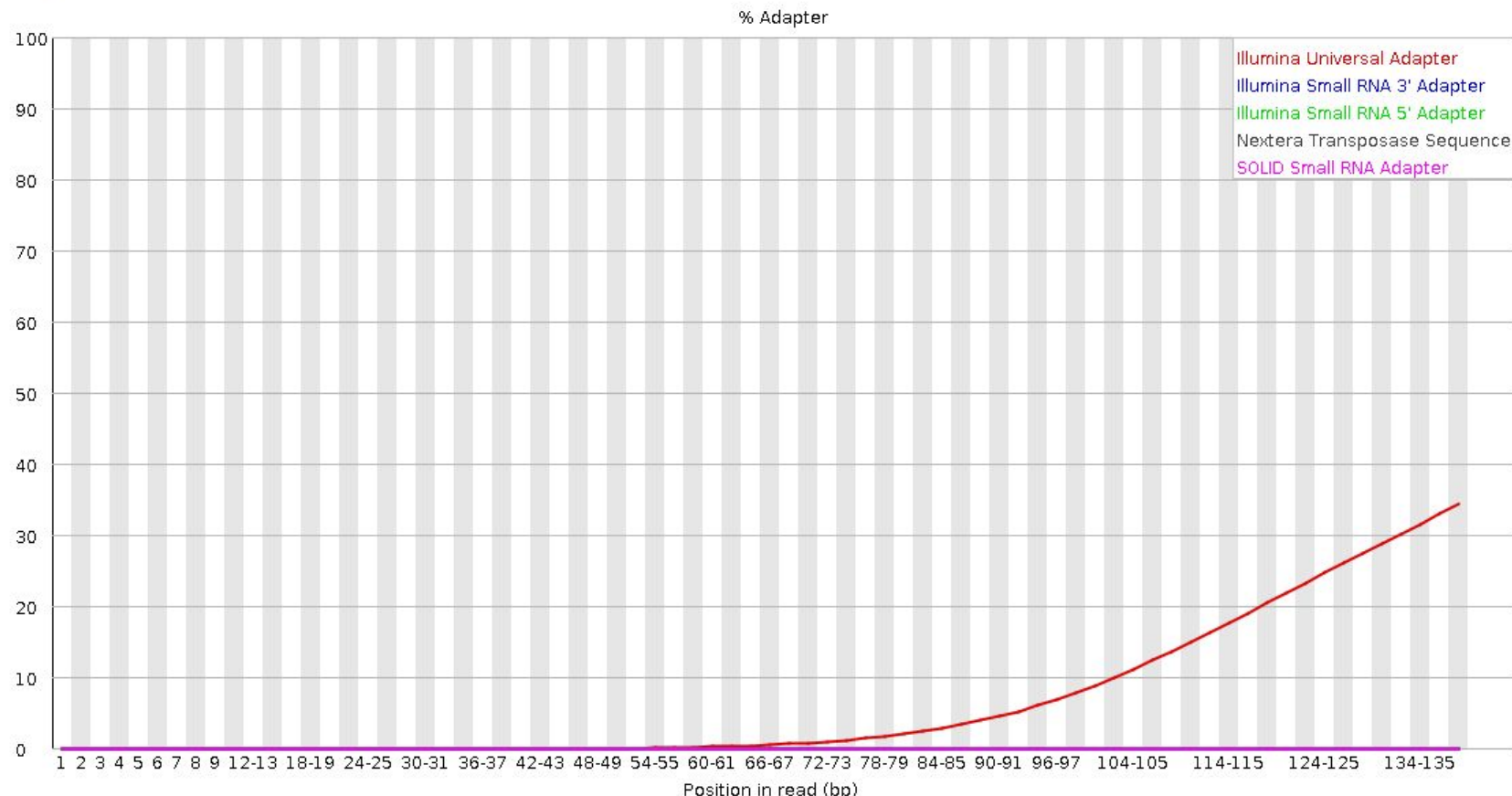


GC content / Contamination ?

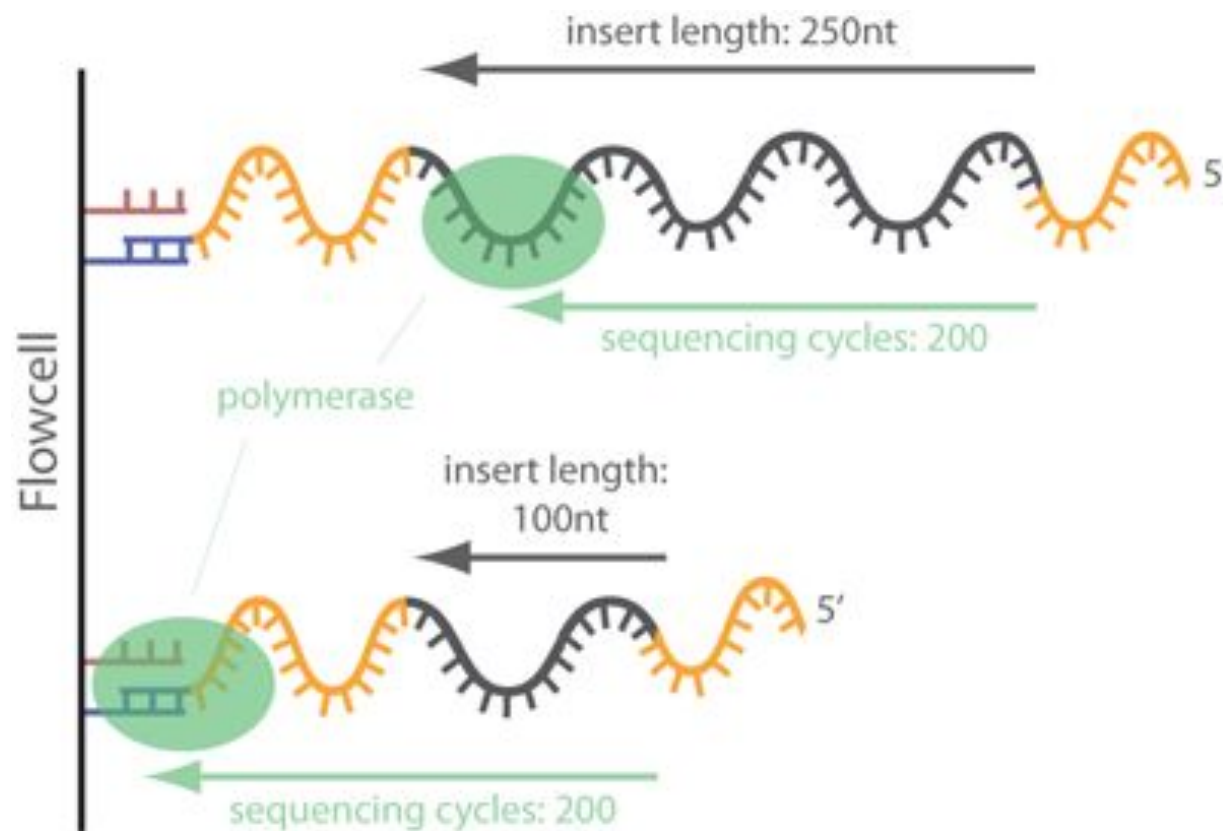


Adapter content

✖ Adapter Content



Adapter content



Jour 1 : Bases de l'analyse NGS pour le RNA-seq

- NGS Introduction
- Reads Quality Control
- Reads Cleaning

Goal: read cleaning

@SRR062641.6751359
CGCCCGGCCAATCATTGTGGTTTAAGTCACTAAGTGTGAGGCTATTTGTTTACAGCAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCATCT
+
CBLNPGJQQQJPPQPQPRGPPPRQRSPGRQQQLRRRMEPQQPMJHQQEHKMMFIIRH?SIHKNJIKRLJJIKHEABHIFGCGBEFCGDGDCE
@SRR062634.16249693
CTAAGTTGAGGCTATTTGTTTACAGCAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCATCTGTGCACCCAGCATTGCCAGAACAGGGC
+
ALKMOOOOPPQJQOPPPPPQPPPPRJQRQQQQQRPQPRQPFQSQQPRLIMHKSNRJQORMFELRPQNQRQJQRRPQQLIRKDMKQJPOFDGCCB
@SRR062634.20060465
CTCCCAGCTTCCAACAGACCCTGTCCCAGCTCCCTCCAAGCTGAGTGTGGCCTGATACTACCACTGGAGCGAGGGGAACCCGAGGACTGCCAAGGGCA
+
D?KMPQECPQQNPQIQIGR@DPERQHEKBEDHCHG8EHFCD6<329@<:69A<6, ;<967>;=C:>AA8BBED#####
@SRR062635.15516129
AAAAAAAAAAAAAAAAAAAAAGGGGGCCCCCTTCCCCCCCAGGGGGGGGACAGGGGGGTGTTGGGCCCGGCCCTGACCACGG
+
EKLMPPPPPOoooooooooooo###



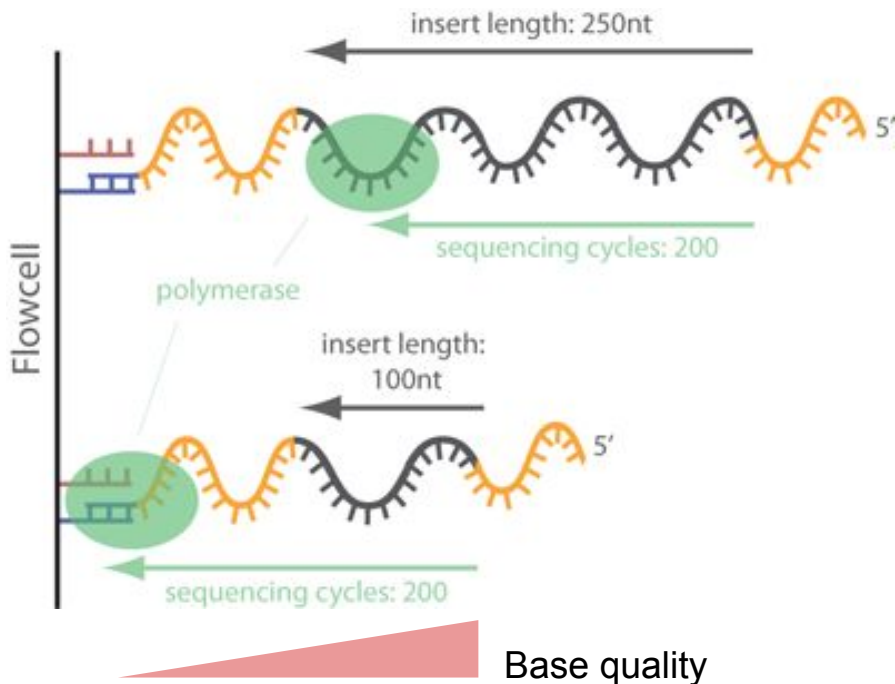
@SRR062641.6751359
CGCCCGGCCAATCATTGTGGTTTAAGTCACTAAGTTGAGGCTATTTGTTTACAGCAAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCATCT
+
CBLNPGJQQQJPPQPPQPRGPPPRRQQRSPGRQQQLRRRMEPQQPMJHQQEHKMMFIIRH?SIHKNJIKRLJJIKHEABHIFGCGGEFCGDGDCE
@SRR062634.16249693
CTAAGTTGAGGCTATTTGTTTACAGCAAAAGCTAACTGATGCAGACAGGGACAAGTCAGTCATCTGTGCACCCAGCATTGCCAGAACAGGGC
+
ALKMOOOOPPQJQOPPPPPQPPPPRJQRQQQQQRPQPRQQPFSQQPLIMHKSNRJQORMFELRPQNQRQJQRRPQQLIRKDMKQJPFDGCCB
@SRR062634.20060465
CTCCCCAGCTTCCAACAGACCCCTGTCCCAGCTCCCTCCAAGCTGAG
+
D?KMPOEPGCPQONPOIOIGR@DPEROHEKBEDHCHG8EHEDCD

RAW

Clean

Reads cleaning

- Cut adaptors at read ends
- Trimming : cut read ends (5' ou 3')
 - Fixed number of bases
 - Individual base quality
 - Mean quality of bases in a sliding window
- Filtering : remove read
 - Size criteria (example < 60bp)
 - Mean base quality for all bases criteria (example < 25)

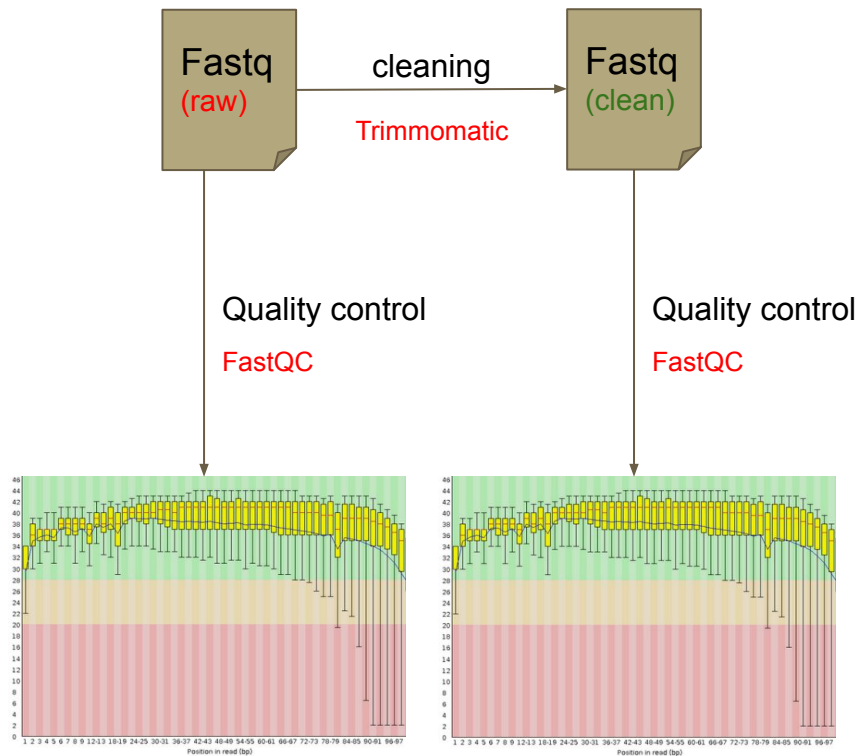


Reads cleaning example

Tool: Trimmomatic

-
- 01 • Clean adaptors
 - 02 • Trimming 5' et 3' on base quality (> 3)
 - 03 • Trimming using sliding window (4 bases, Q < 20)
 - 04 • Filtering on mean read quality (Q < 25)
 - 05 • Filtering on read size (taille < 20)

Workflow



usegalaxy.fr presentation



Practical: Quality Control (QC) & Cleaning

Open Galaxy



Practical:

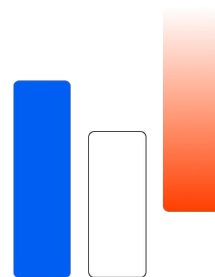
<https://training.galaxyproject.org/training-material/topics/sequence-analysis/tutorials/quality-control/tutorial.html>

TIAAS: <https://usegalaxy.fr/join-training/bililledna/>

Mapping

Practical: Mapping

Open Galaxy



usegalaxy.fr

Practical:

<https://training.galaxyproject.org/training-material/topics/sequence-analysis/tutorials/mapping/tutorial.html>

TIAAS: <https://usegalaxy.fr/join-training/bililledna/>