



Quality control of Illumina data

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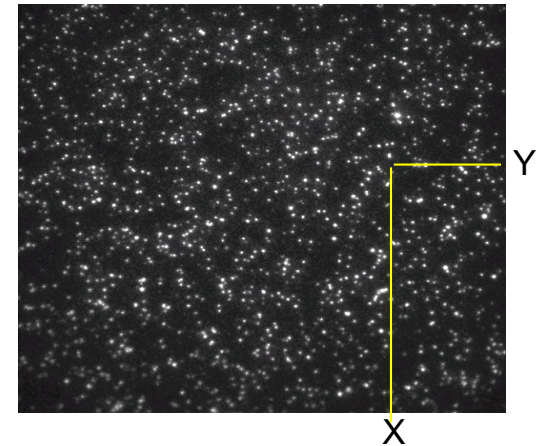
Quality control of Illumina data

- Primary analysis
- Quality control
- Data pre-processing

Quality control of Illumina data

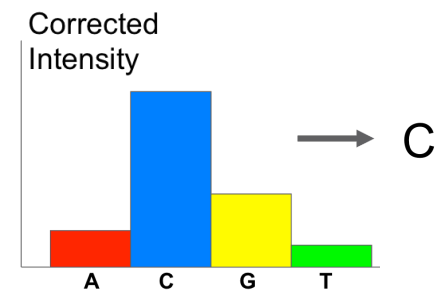
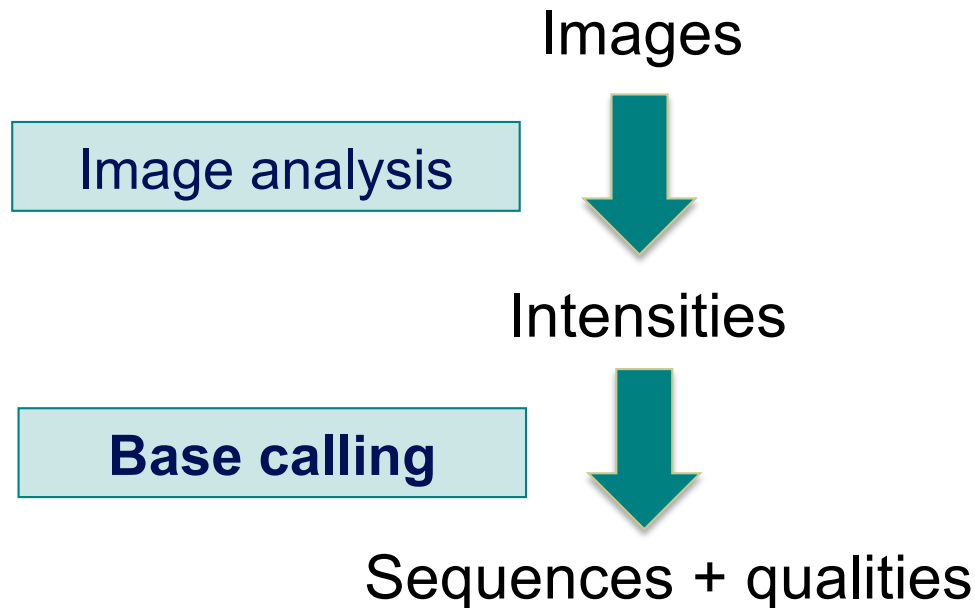
- Primary analysis
- Quality control
- Data pre-processing

Primary analysis



- Determination of cluster position (only for non-patterned flow cells)
- Extraction of intensities for each cluster

Primary analysis



- Intensity correction
 - Take into account \neq intensities per molecule for the 4 bases
- Call the base with the maximum intensity
- Determine “Passing filter” clusters
 - Remove clusters that have “too much” intensity corresponding to bases other than the called base

Phred quality scores

- Prediction of the probability of error in base calling

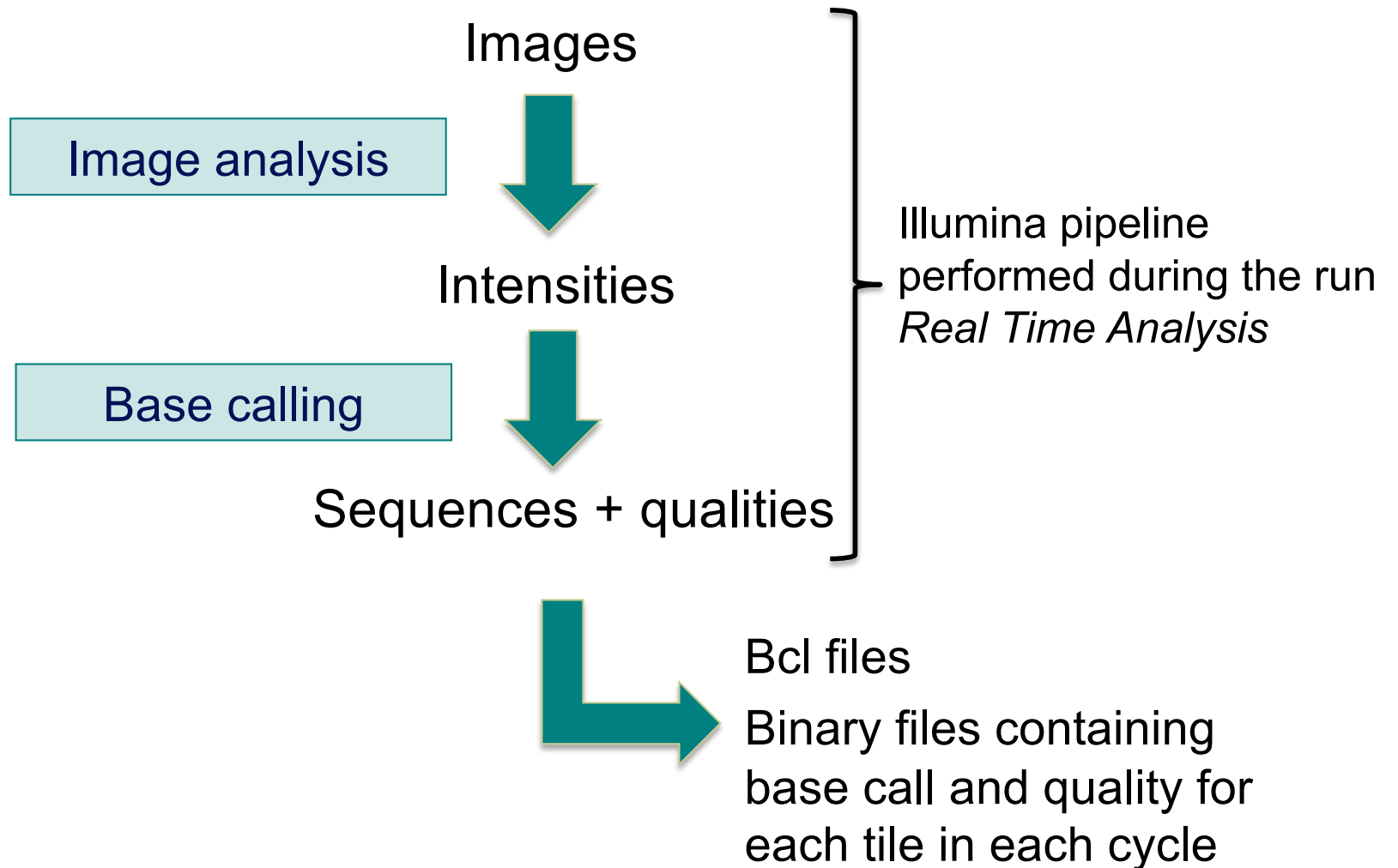
$$Q = -10 \log_{10} P$$

Q : quality score

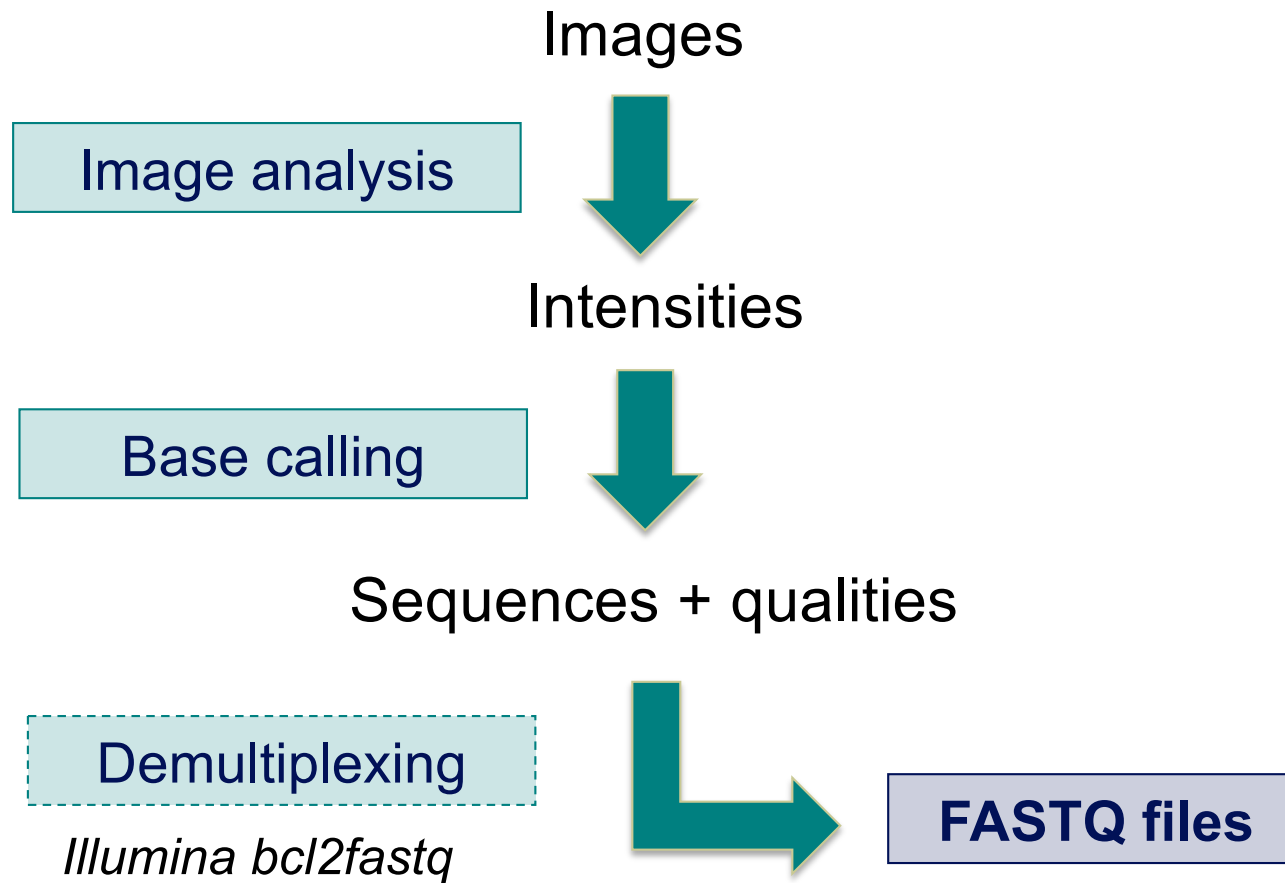
P : error probability

Quality Score	Error Probability
Q40	0.0001 (1 in 10,000)
Q30	0.001 (1 in 1,000)
Q20	0.01 (1 in 100)
Q10	0.1 (1 in 10)

Primary analysis



Primary analysis



FASTQ file

- Text file containing

- Sequences

- Qualities

Probability that the corresponding base call is incorrect

4 lines per sequence :

```
@HWI-ST1136:97:HS041:7:1101:1681:2104 1:N:0:ACAGTG → 1. @Identifier
CTTTTTATTGAATTCTATGATTCTTGTTAGATTTTCATAATGGCTGCTTA → 2. Sequence
+ → 3.+ optionally followed by same identifier as 1.
@@@DBDDDDFF8:D?EBAEAH,CF:AF9F+2**9?B?1C<<?9*8D?)9*? → 4. Quality
@HWI-ST1136:97:HS041:7:1101:1521:2119 1:N:0:ACAGTG
AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
+
@@@?DDDDFAFHIIHFDAB@B6@B@BBBBBBBBBBBBBB359BBBB8BBB
@HWI-ST1136:97:HS041:7:1101:1669:2145 1:N:0:ACAGTG
CTGCTGTTTTCCAAATGTCCGATGTGTGCTATGACTGACAACTACTTTTC
+
@@<1?DDDFDBFD?FE>+<CCF>FAG++2+<<F**?:?1:C?:8B:9BBBD4
```

(Cock et al. NAR 2009; 38(1): 1767-1771)

Beginning of siLuc3_S12040.fastq file

The screenshot displays the Galaxy France web interface. At the top, the navigation bar includes the Galaxy France logo, a home icon, and menu items for Workflow, Visualize, Shared Data, Help, User, a graduation cap icon, a bell icon, and a grid icon. A green badge in the top right corner indicates 'Using 51%'. The left sidebar contains a 'Tools' section with a search bar and an 'Upload Data' button. Below this are categories for 'Get Data', 'Send Data', 'Collection Operations', 'GENERAL TEXT TOOLS', 'Text Manipulation', 'Filter and Sort', 'Join, Subtract and Group', and 'GENOMIC FILE MANIPULATION'. The main content area features an orange warning box stating 'This dataset is large and only the first megabyte is shown below.' with 'Show all' and 'Save' links. Below the warning is a preview of the FASTQ file content, showing header lines and sequence data. The right sidebar shows a 'History' section with a search bar and a list of datasets under the heading 'RNA-seq data analysis'. One dataset, '1: siLuc3_S12040.fastq.gz', is highlighted in green and has a red circle around its eye icon, indicating it is the current view.

Galaxy France Workflow Visualize Shared Data Help User Using 51%

Tools search tools Upload Data

Get Data
Send Data
Collection Operations
GENERAL TEXT TOOLS
Text Manipulation
Filter and Sort
Join, Subtract and Group
GENOMIC FILE MANIPULATION

History RNA-seq data analysis
1 shown
1.96 GB
1: siLuc3_S12040.fastq.gz

Warning: This dataset is large and only the first megabyte is shown below. [Show all](#) | [Save](#)

```
@HWI-ST1136:52:HS008:4:1101:2560:2035 1:N:0:GCGAAT
GCCGGTGGGGTCGATGCCATGTTTCATCACTGATCAACTCCCAGAACTTGG
+
?@;BBD<?)@<@@:):1:?GFD?:?GF<9*9BG9B99?*0?CCBBBBF9@F
@HWI-ST1136:52:HS008:4:1101:2669:2093 1:N:0:GCCAAT
GCTGTTTGCTTTTGTCTCCCTCTGTCTTAGGAAAAGCCATCTTTAATAT
+
???7DD;=D+?CDD<EEEIEEECFFFCFD<F<AEEE@DIDIIEIEIAD
@HWI-ST1136:52:HS008:4:1101:2690:2156 1:N:0:GCCAAT
TTTGCATTTACGCCTGTAATGTATTATTCTTAATTTATGTAAGGTTTT
+
???DDDDHFDHF<FHIGEHIII9?HBFFF<CHH@FFHCGHIGDIICDGH
@HWI-ST1136:52:HS008:4:1101:2663:2212 1:N:0:GCCAAT
CAAATAGACTACATAATATACGTGGGCAAAAAGGCAATTAAGTGAATCTC
+
?8?DD?A:CCCF??ECFH@,CAFHFGGIHIGCGGE??<FDHGGEGGIE
```

Sequence identifier in FASTQ files

- Begins with @
followed by sequence ID and an optional description
- Illumina sequence identifiers :

Instrument Name Run number Flowcell ID Lane Tile X_pos Y_pos Read Is Filtered Control Number Index Sequence

@HWI-ST1136:97:HS041:7:1101:1681:2104 1:N:0:ACAGTG

- Read :
The member of a pair = 1 or 2 (for paired-end reads)
- Is filtered
Y = bad quality (the cluster do not pass filter), N otherwise
Recent versions of Illumina pipeline only supply passing filter reads

Quality in FASTQ files

- Phred quality score (Sanger format)
- Encoded in ASCII characters to save space
- 1 ASCII symbol = 1 quality value
- Phred quality scores from 0 to 93 are encoded using ASCII 33 to 126 :

032 sp	048 ò	064 @	080 P	096 `	112 p
033 !	049 1	065 A	081 Q	097 a	113 q
034 " ' "	050 2	066 B	082 R	098 b	114 r
035 #	051 3	067 C	083 S	099 c	115 s
036 \$	052 4	068 D	084 T	100 d	116 t
037 %	053 5	069 E	085 U	101 e	117 u
038 &	054 6	070 F	086 V	102 f	118 v
039 *	055 7	071 G	087 W	103 g	119 w
040 (056 8	072 H	088 X	104 h	120 x
041)	057 9	073 I	089 Y	105 i	121 y
042 *	058 :	074 J	090 Z	106 j	122 z
043 +	059 ;	075 K	091 [107 k	123 {
044 ,	060 <	076 L	092 \	108 l	124
045 -	061 =	077 M	093]	109 m	125 }
046 .	062 >	078 N	094 ^	110 n	126 ~
047 /	063 ?	079 O	095 _	111 o	127 ò

- Binned in order to save space in the last versions of Illumina software, e.g.
 - $2 < \text{real Q-score} < 9 \rightarrow \text{binned Q-score} = 6$
 - $10 < \text{real Q-score} < 19 \rightarrow \text{binned Q-score} = 15$
 - ...
 - $\text{real Q-score} \geq 40 \rightarrow \text{binned Q-score} = 40$

Paired-end FASTQ files

- 2 FASTQ files per sample



XXXX.R1.fastq.gz

XXXX.R2.fastq.gz

```
@HWI-ST1136:163:HS087:7:2310:17264:70630 1 N:0:ATCACG
GTTAGAGGCCAAGGTACAGTGGCCTGTCTTTGTAATGTGCCTTATGT
+
CCCCFFFFHHHHHJFHIIJHIJIIJIIJIIJIIJIIHIIJIIIGIIJIIJIIJII
@HWI-ST1136:163:HS087:7:2310:17415:70636 1:N:0:ATCACG
TGGAGCCTTGGTAACTTTTTGTAGTTTGTATGCGTTTTTGTGGTCTC
+
BCCFFFFHHHHHIIJJJJJJJHIJJJJJIIJJJJIIJJJJIIJJJJFGIJ
@HWI-ST1136:163:HS087:7:2310:17337:70637 1:N:0:ATCACG
CTGTTACCCCTCATTACAGGGTATGAAGAAGGGCTTCACCTGTAGTTC
+
@CCFFFFFHIIIIJIIJJJJJBFHIIJIIJIIIIJIIJIIJIIHGHIIJJ
```

```
@HWI-ST1136:163:HS087:7:2310:17264:70630 2 N:0:ATCACG
TAATTTTTGCATCCTGAAAAGTGTGGAAGTTGGGTTTTTCATAGCAA
+
CCCCFFFFHHHHHJJIIJJJJICHGIIJIIJIIJIIJIIJFHIIJHIJJIIJJ
@HWI-ST1136:163:HS087:7:2310:17415:70636 2:N:0:ATCACG
TGTTTCATATGTATGAGATAGATTTGAAAAATCTACTAATTTTTAAAATC
+
CCCCFFFFHHHHHJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJ
@HWI-ST1136:163:HS087:7:2310:17337:70637 2:N:0:ATCACG
TCCTGACATCAAGCACACTGCTTCTGCATCTATGTGGCACCTAAAACAA
+
CCCCFFFFHHHHHJIIJJJJJJJJFIIJJJJJJIIJIIJIIJIIJJJJJJJJ
```

Quality control of Illumina data

- Primary analysis
- Quality control
- Data pre-processing

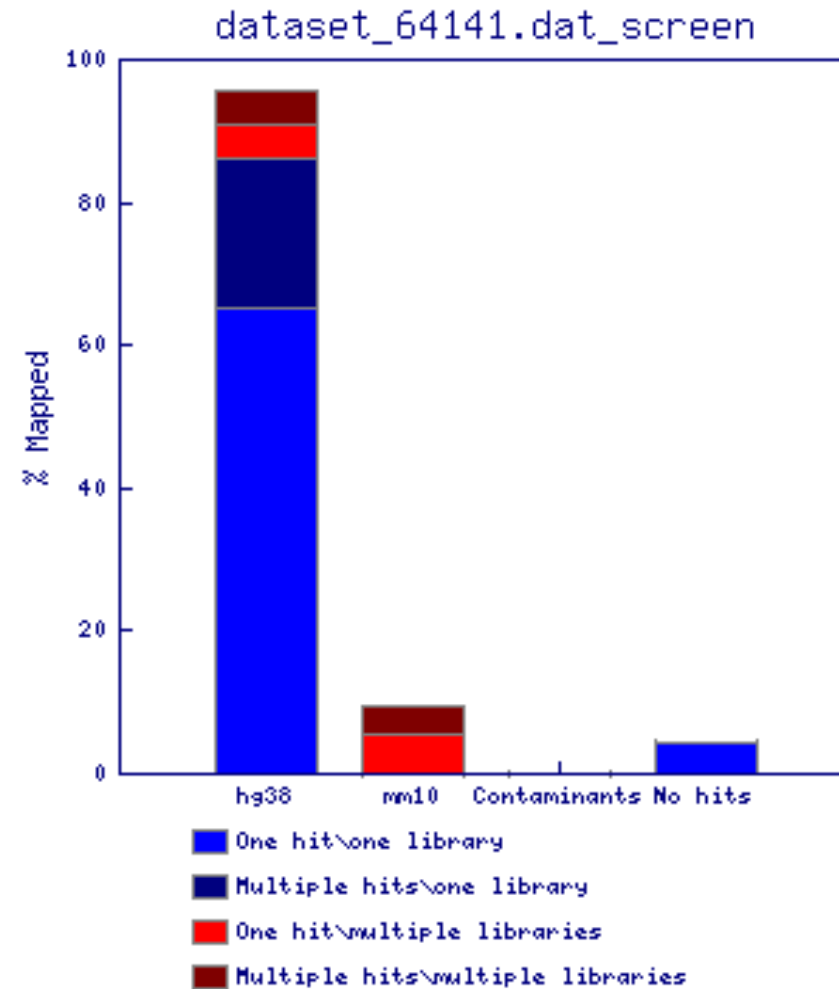
Quality control

- Why ?
 - Are the data consistent with what is expected ?
 - Identify any problems of which you should be aware before doing any further analysis
- What to look for ?
 - Number of reads
 - Base qualities
 - Sequence duplication
 - Presence of adapters
 - Contaminations

Some quality control tools

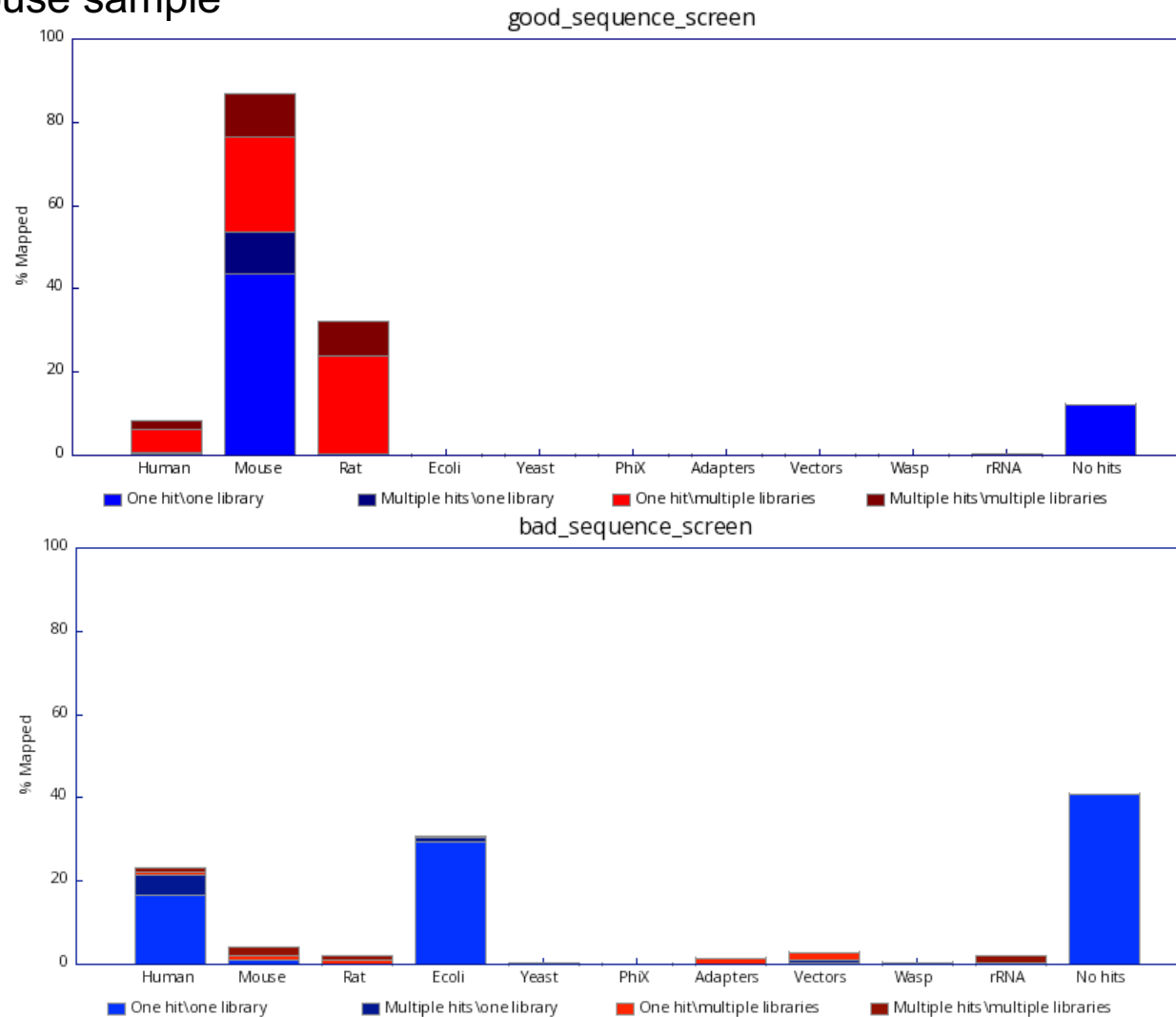
- FastQC
 - <https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>
- RSeQC – quality controls specific to RNAseq data
 - <http://rseqc.sourceforge.net/>
- FastQ Screen – to verify the composition of a library and search for possible contaminations
 - https://www.bioinformatics.babraham.ac.uk/projects/fastq_screen/

FastQ Screen result on siLuc3_S12040.fastq



FastQ Screen result examples

On a mouse sample



FastQC

- Allows quality control of NGS data
- Input
 - FASTQ or SAM/BAM alignment files
- Can be used *via* a graphical interface, in command-line or in Galaxy
- Generates graphs and tables with several quality control analyses
 - ➔ Allows a global quality assessment of NGS data and rapid identification of possible problems

Exercise : quality analysis

- Analyse the quality of siLuc3_S12040.fastq file
 - How many reads have been sequenced in this sample ?
 - What do you think about the quality of this sample ?
 - Do you identify bias in these data ?

FastQC results

FastQC Report

Wed 5 Apr 2023
siLuc3_S12040_fastq_gz.gz

History

Rechercher des données

RNA-seq data analysis

2.11 GB

- 3 : FastQC on data 1: RawData
- 2 : FastQC on data 1: Webpage
- 1 : siLuc3_S12040.fastq.gz

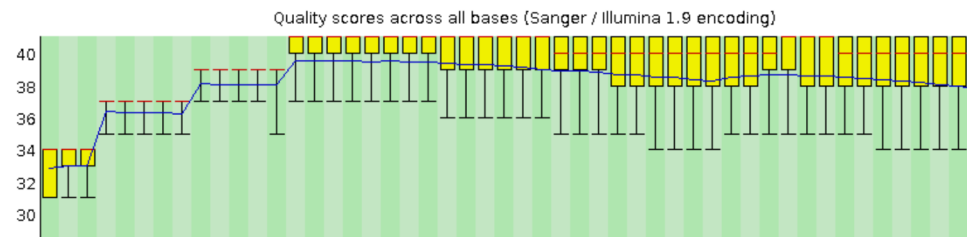
Summary

- Basic Statistics
- Per base sequence quality
- Per tile sequence quality
- Per sequence quality scores
- Per base sequence content
- Per sequence GC content
- Per base N content
- Sequence Length Distribution
- Sequence Duplication Levels
- Overrepresented sequences
- Adapter Content

Basic Statistics

Measure	Value
Filename	siLuc3_S12040_fastq_gz.gz
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	50079515
Sequences flagged as poor quality	0
Sequence length	50
%GC	49

Per base sequence quality

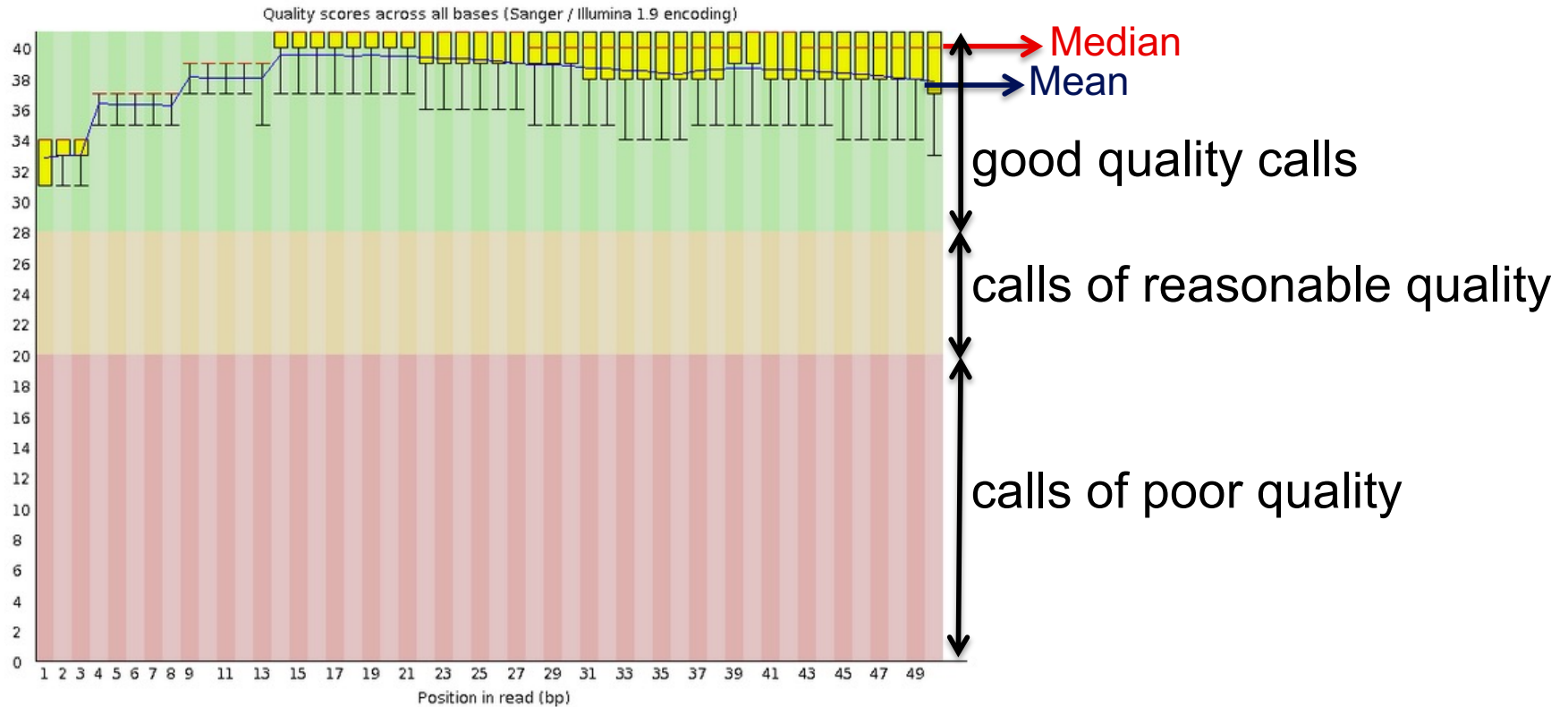


Basic Statistics

Measure	Value
Filename	siLuc3_S12040_fastq_gz.gz
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	50079515
Sequences flagged as poor quality	0
Sequence length	50
%GC	49

- **File type** : Base calls or colorspace data
- **Encoding** : Which ASCII encoding of quality values was found in this file
- **Total Sequences**: A count of the total number of sequences in the file
- **Sequences flagged as poor quality** : Sequences flagged will be removed from all analyses. The total sequences count above will not include these filtered sequences
- **Sequence length**: Length of the shortest and longest sequence
If all sequences have the same length only one value is reported
- **%GC**: The overall %GC of all bases in all sequences

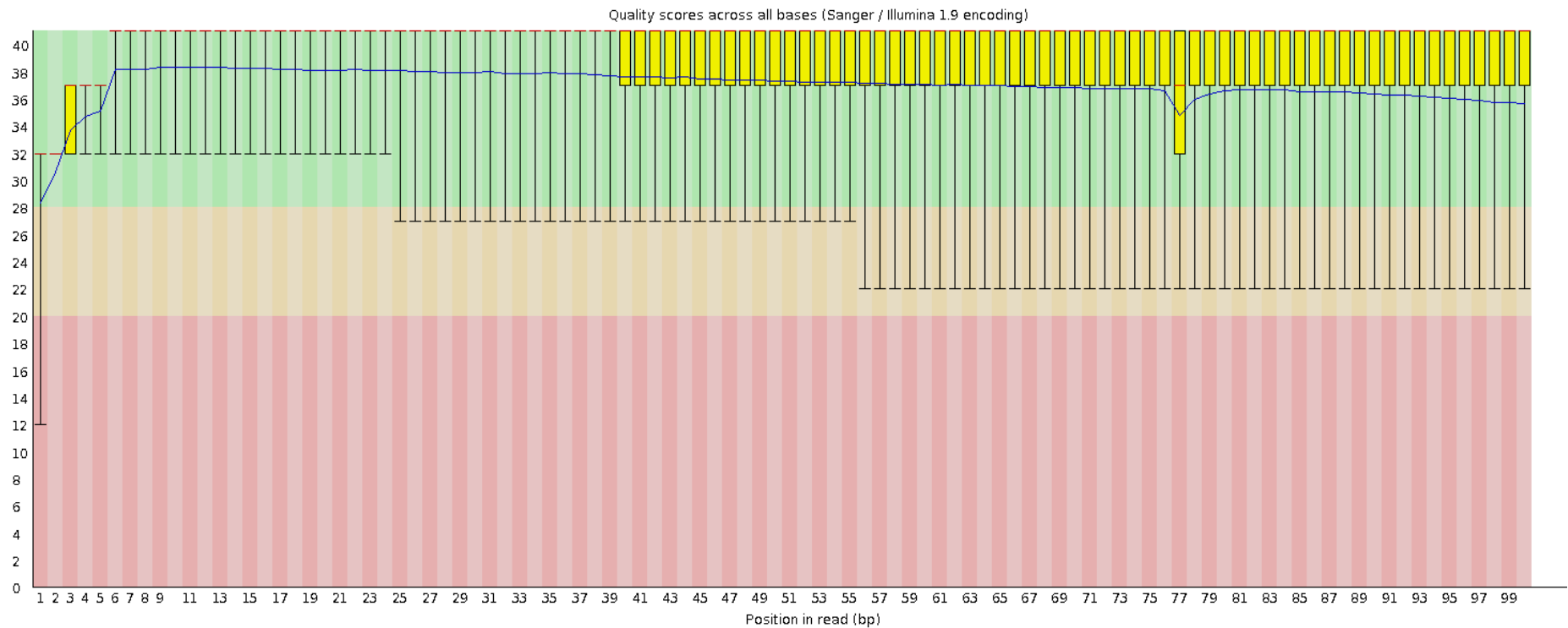
Per base sequence quality



- Yellow boxes : inter-quartile range (25-75%)
 - Lower and upper whiskers : 10% and 90%
- ➔ **Sample of good quality**

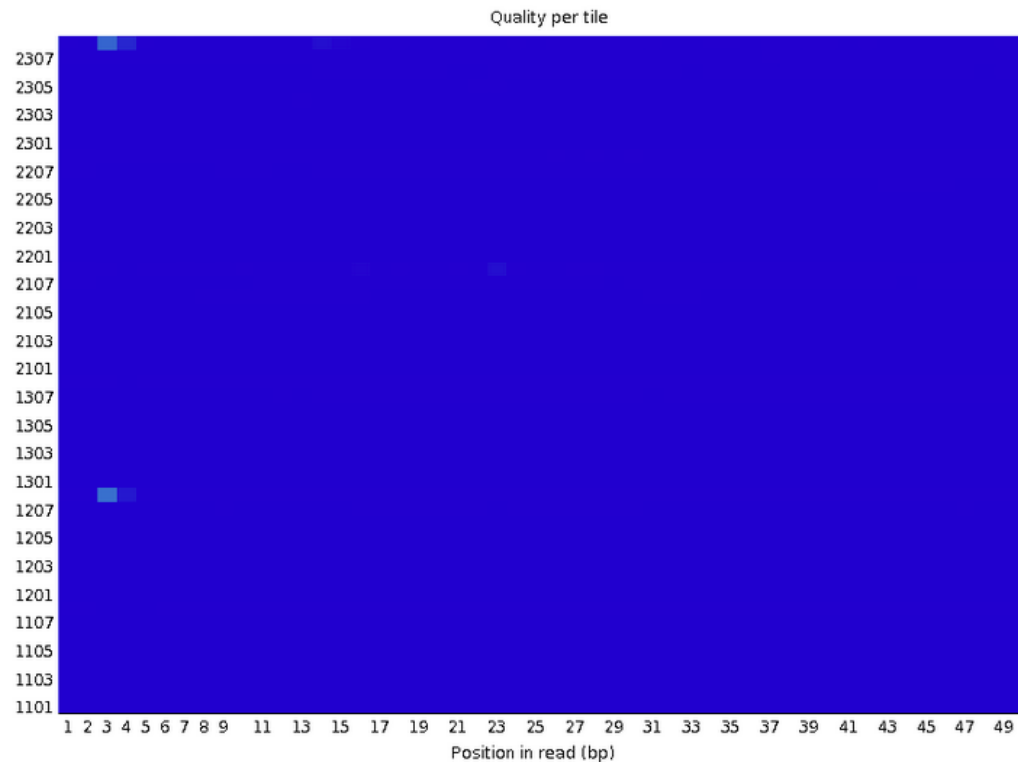
Per base sequence quality on another sample

- The quality of calls decreases as the run progress
e.g. 2nd read of a 2x100bp run :



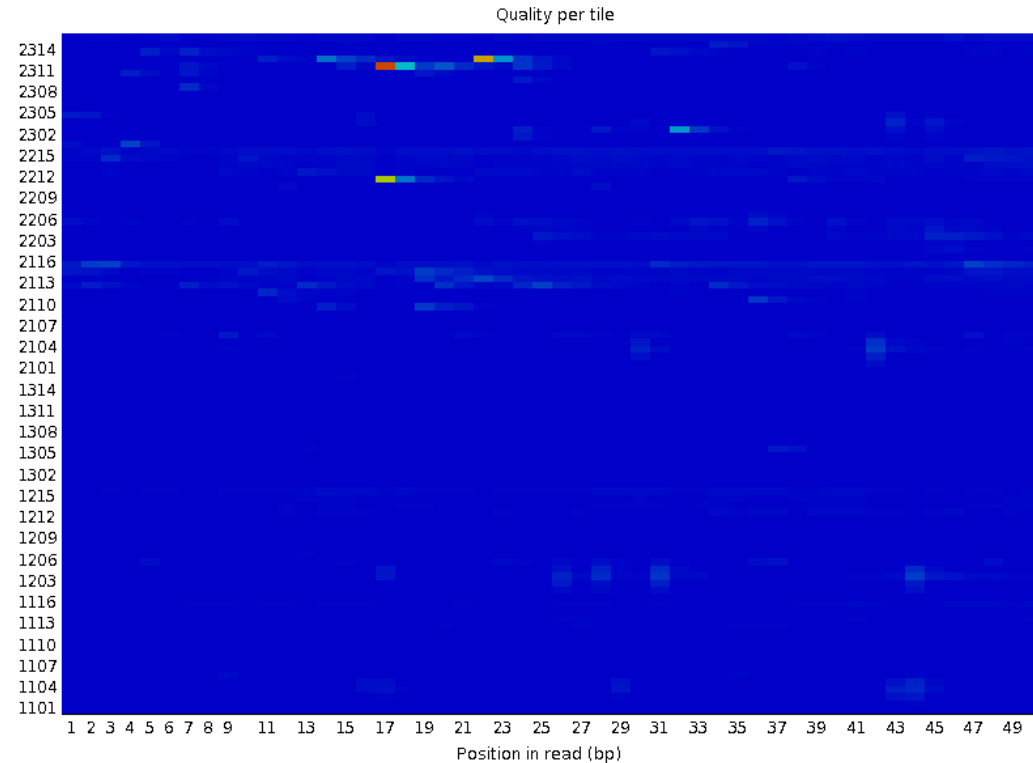
Per tile sequence quality

Quality scores from each tile across all bases :
show the deviation from the average quality for each tile



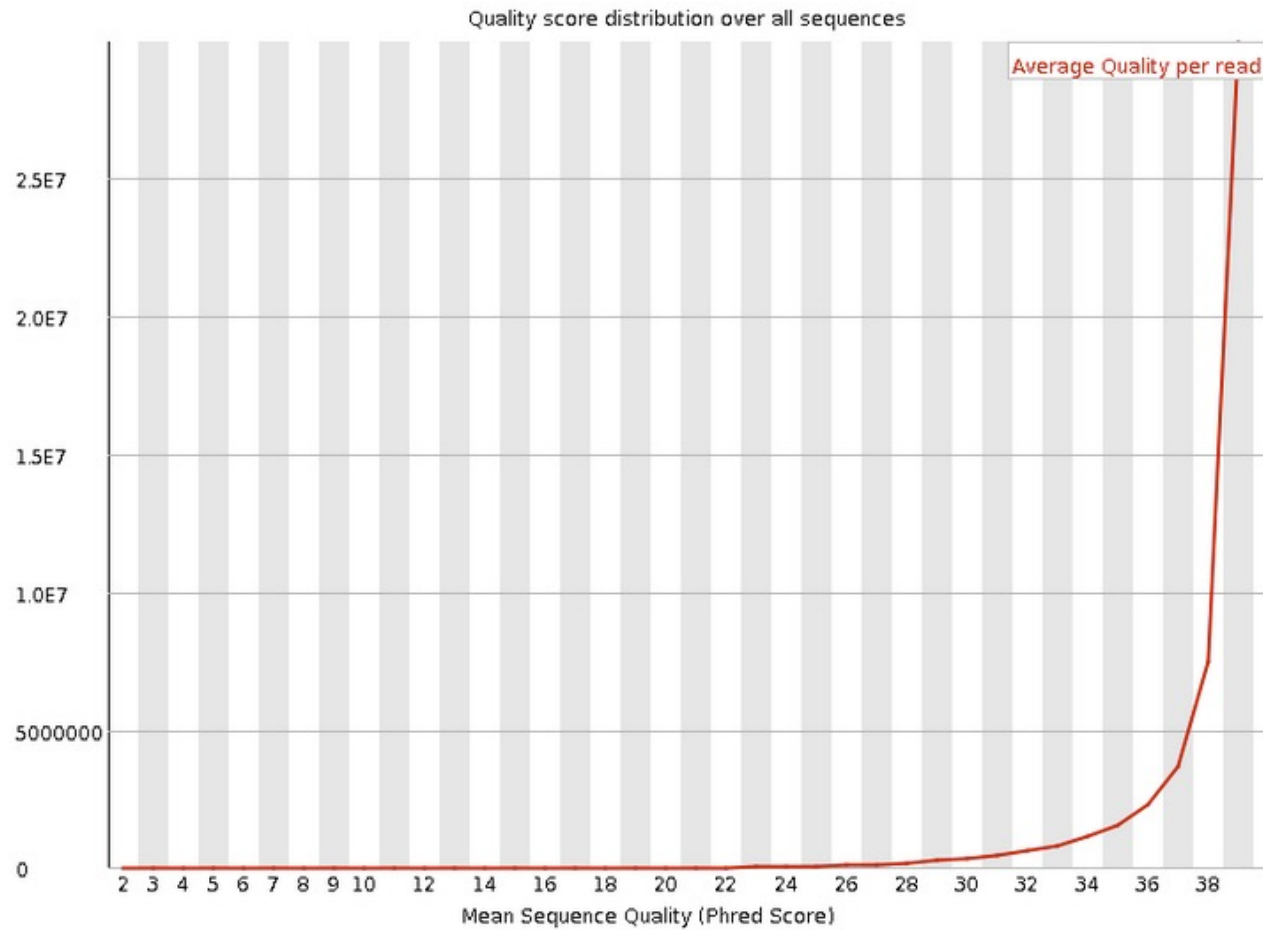
- To see if there was a loss in quality associated with only one part of the flowcell
- No poor quality tile for this sample

Per tile sequence quality on another sample



- Colours on a cold to hot scale
 - Cold colours : positions where the quality was at or above the average for that base in the run
 - Hotter colours : a tile had worse qualities than other tiles for that base
- A good plot should be blue all over

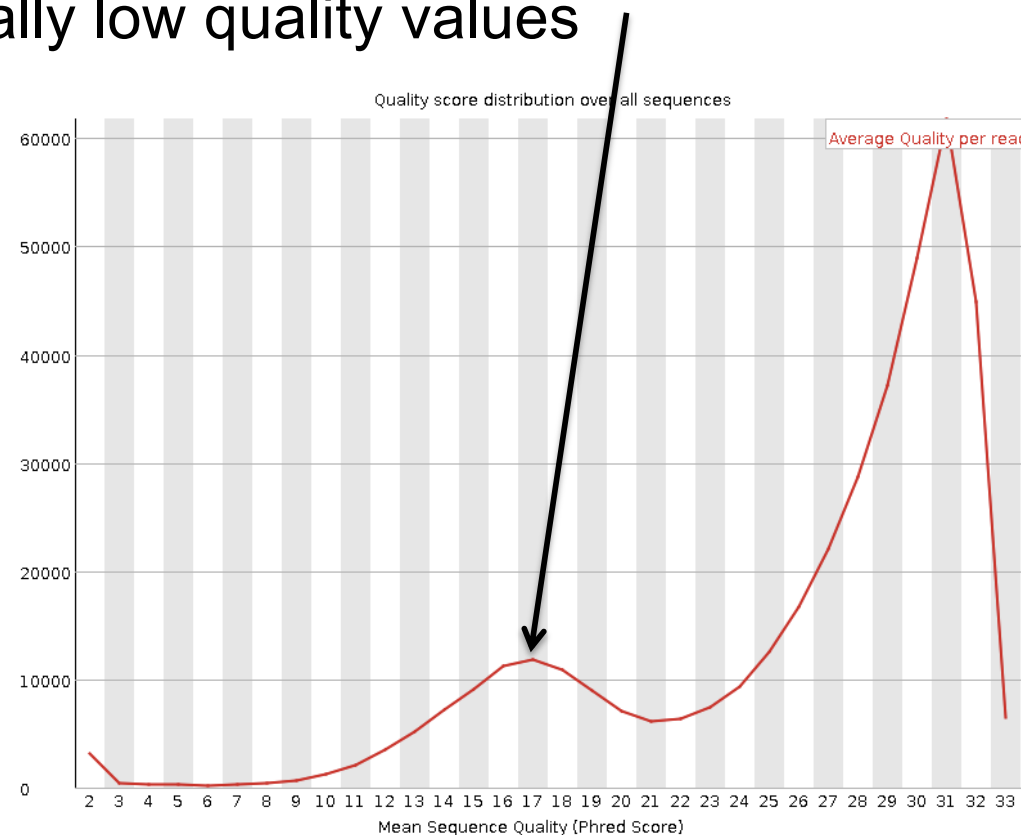
Per sequence quality scores



→ Good quality of all sequences

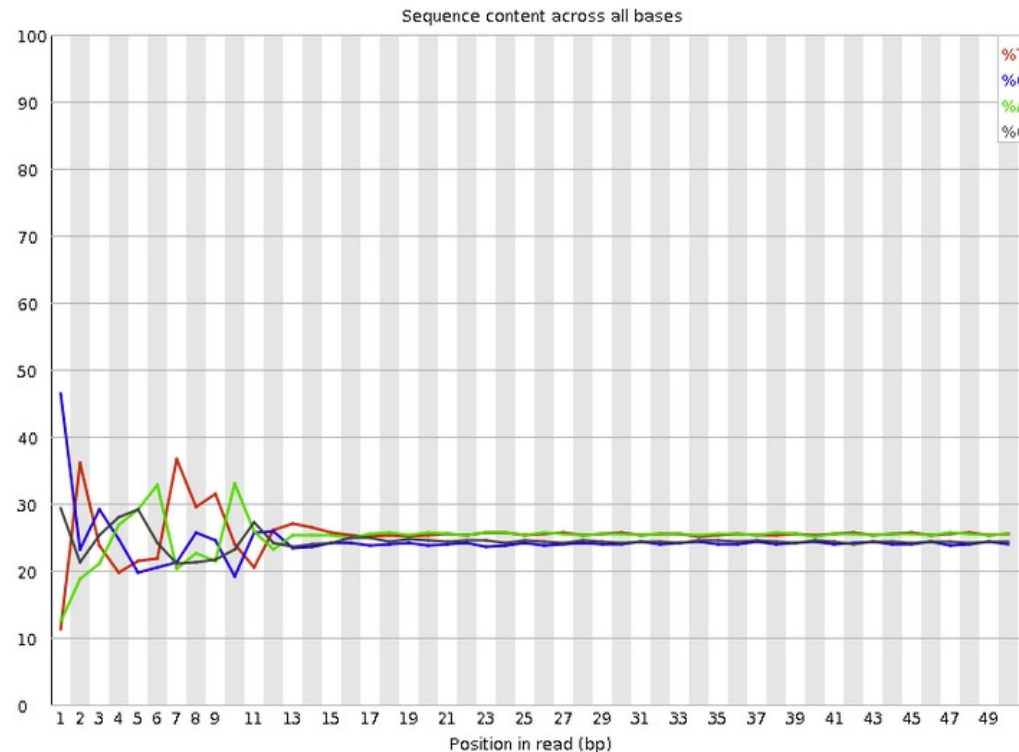
Per sequence quality score on another sample

- Allows you to see if a subset of your sequences have universally low quality values



➔ these should represent only a small percentage of the total sequences

Per base sequence content

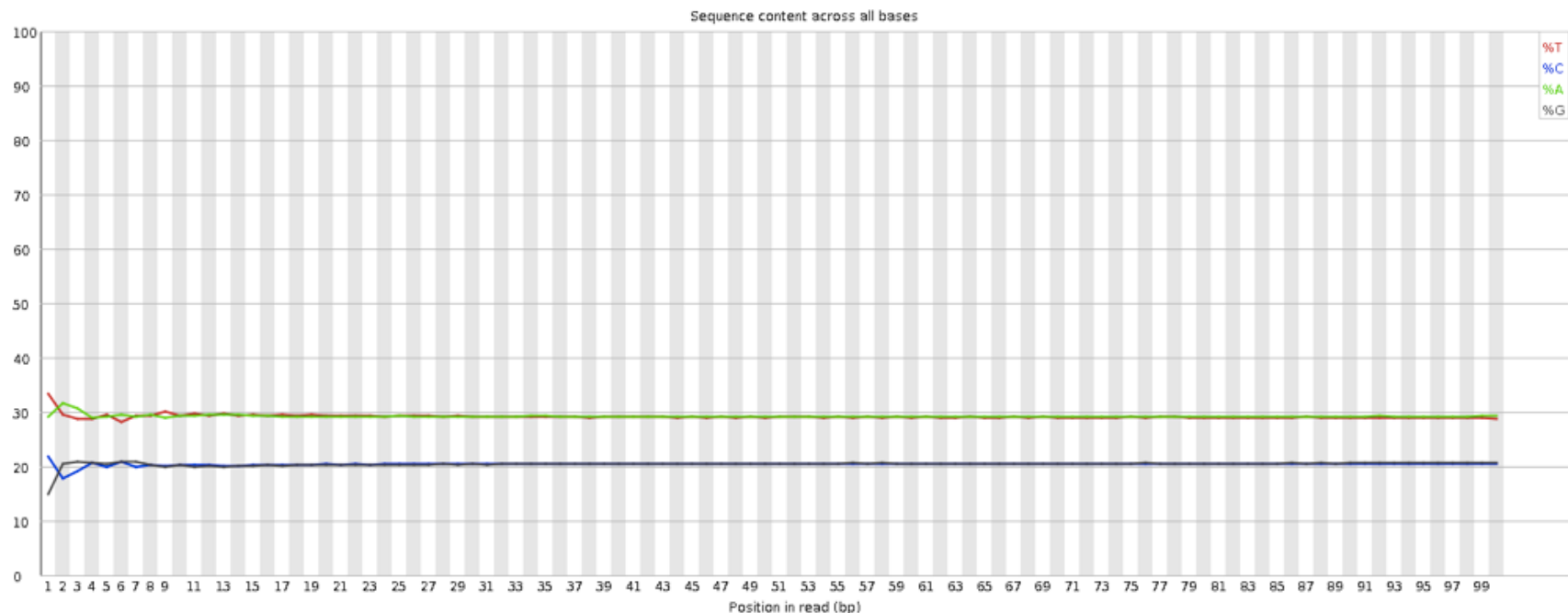


- **Known bias in the repartition of the first nt in RNA-seq libraries**

- Because random primers used during RT are “not so random”
- “Reproducible bias” → Comparative analyses OK
- c.f. Hansen et al. 2010;38(12):e131.
Li et al. Genome Biology 2010;11(5):R50.

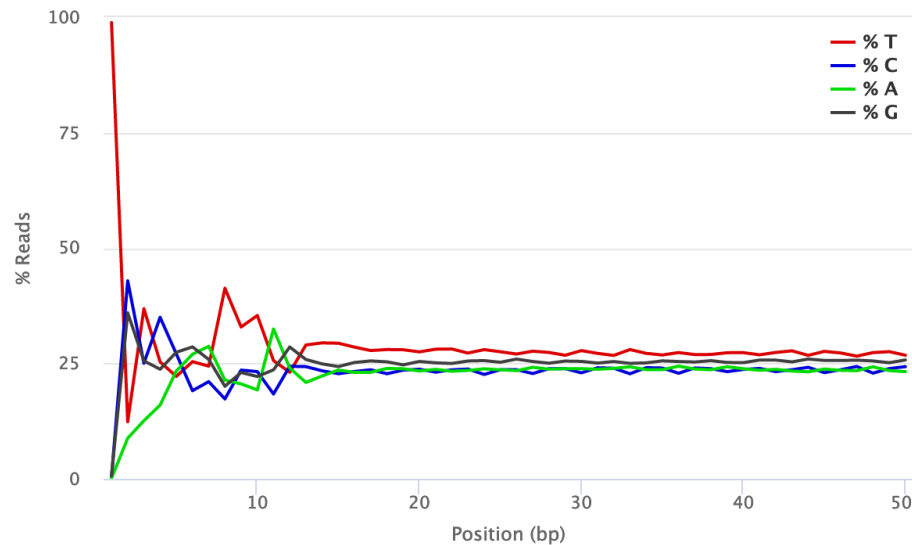
Per base sequence content on other samples

- The lines in this plot should run parallel with each other
- The relative amount of each base should reflect the overall amount of these bases in your genome
- Example for a DNaseq sample :



Per base sequence content on other samples

- Strong biases which change in different bases
 - Usually indicates an overrepresented sequence

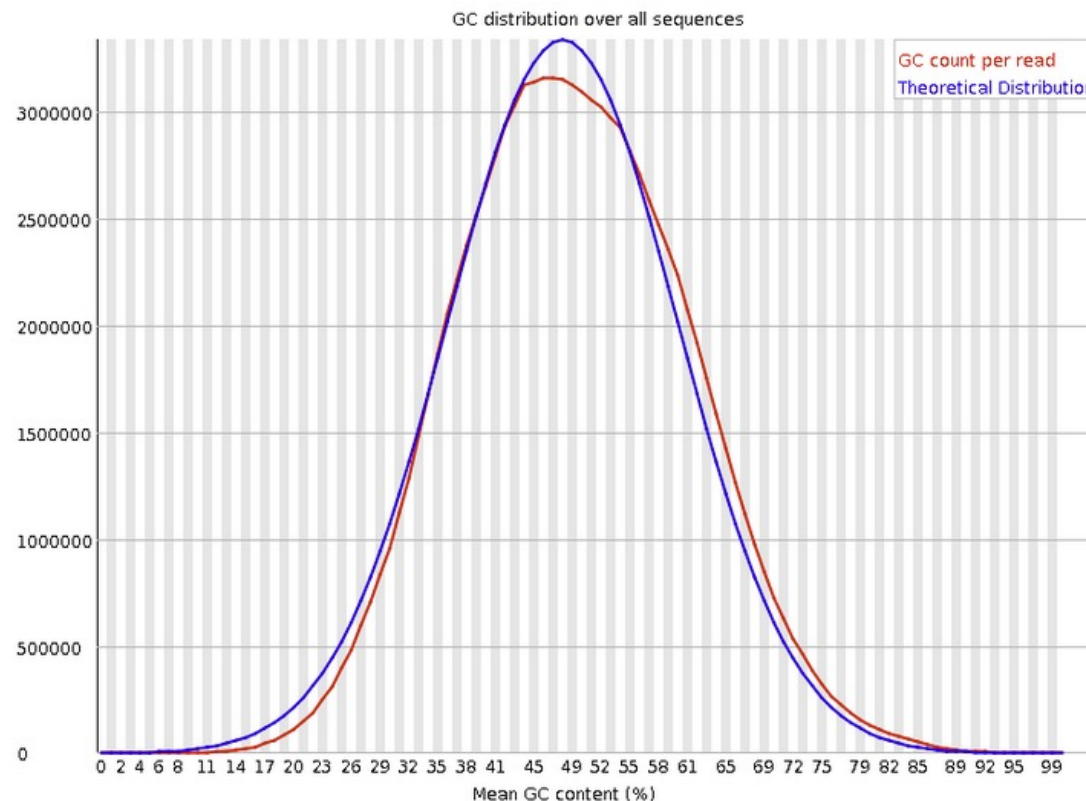


Library prepared with the Illumina Stranded mRNA Prep, Ligation protocol

- Bias which is consistent across all bases
 - indicates that the original library was sequence biased (e.g. adapter)
 - or that there was a systematic problem during sequencing

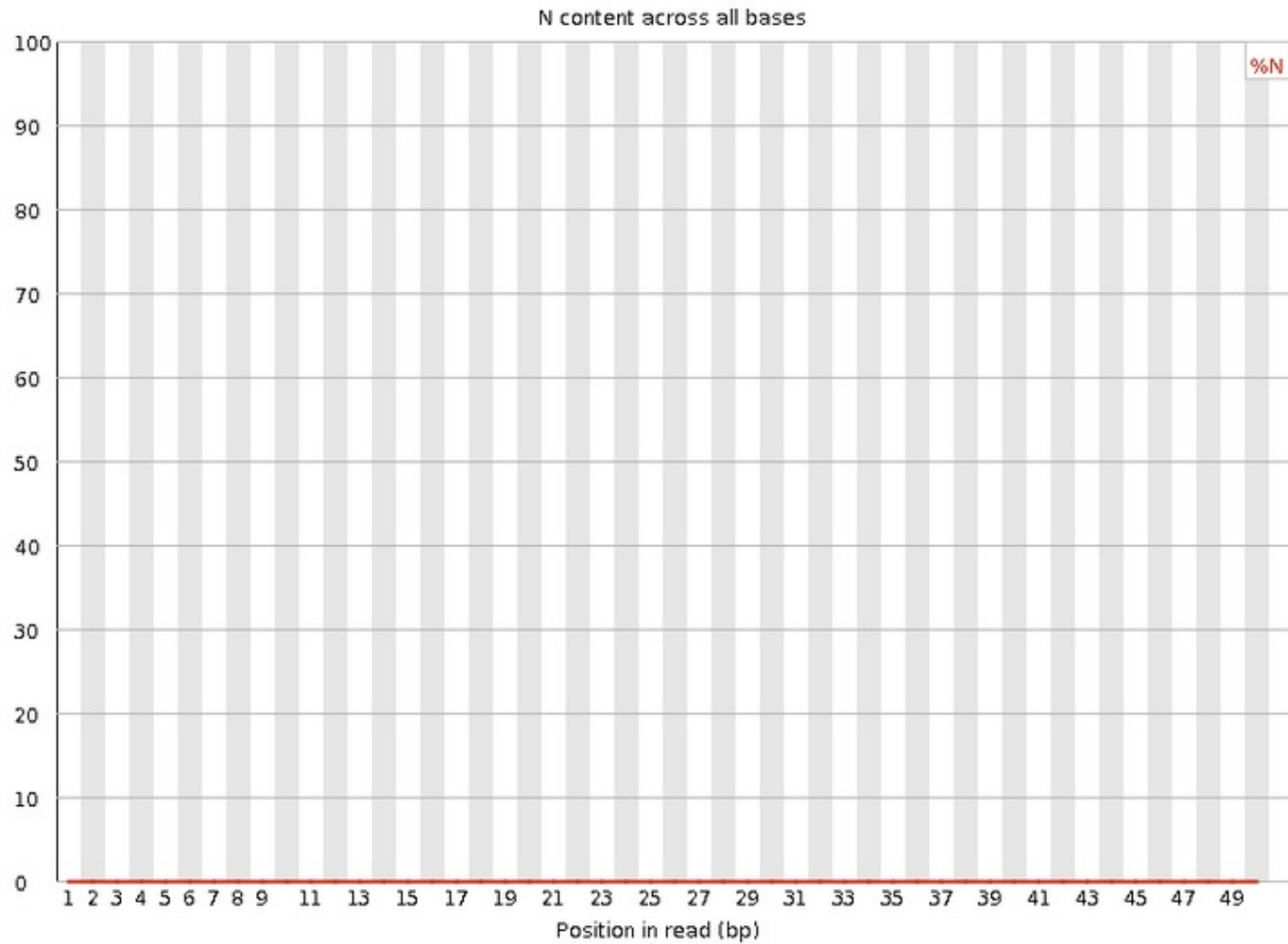
Per sequence GC content

- Compares GC content of all sequences to a modelled normal distribution of GC content (mode calculated from the data and used to build the reference distribution)



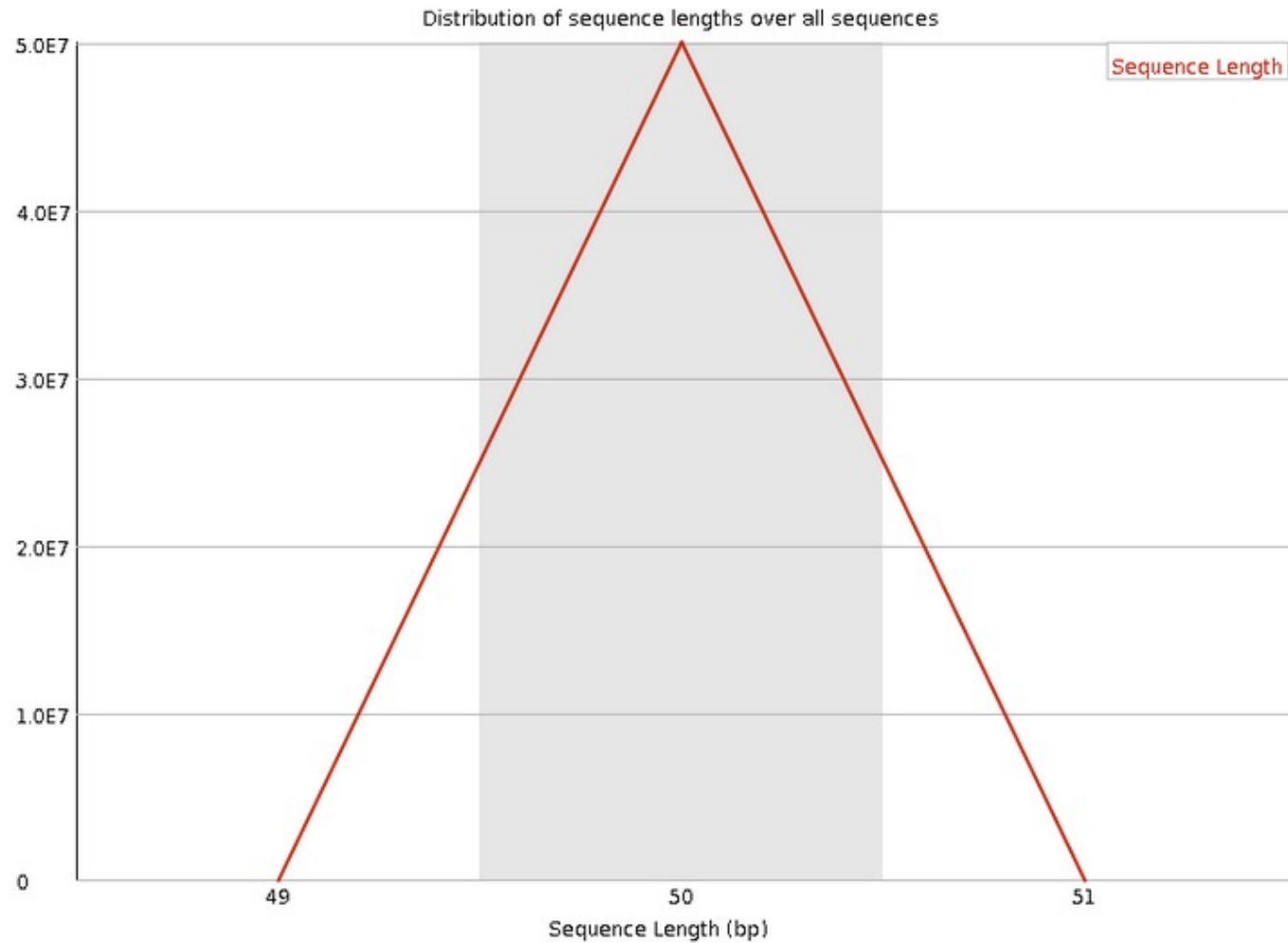
➔ Observed GC distribution similar to the theoretical one

Per base N content



→ Very low N content

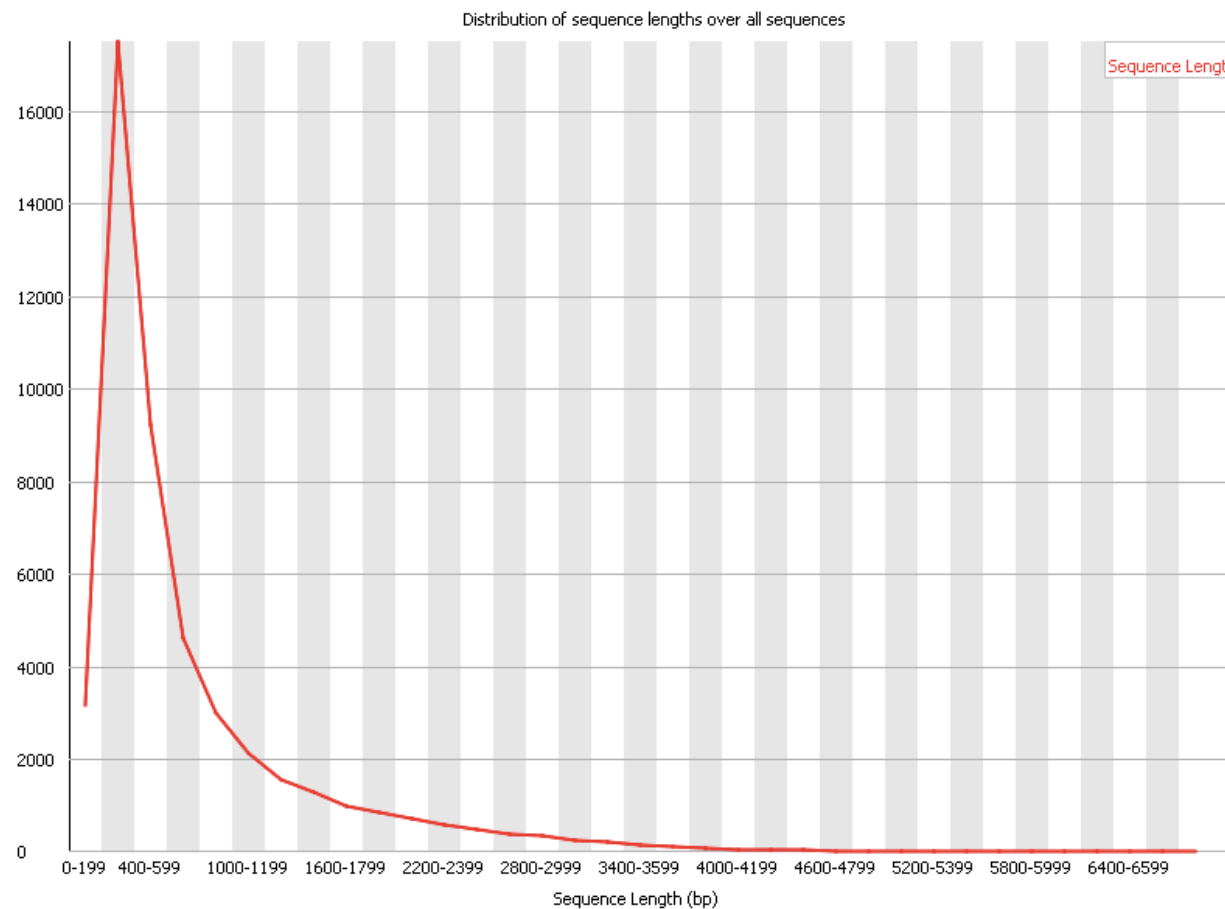
Sequence length distribution



→ All sequences = 50bp reads

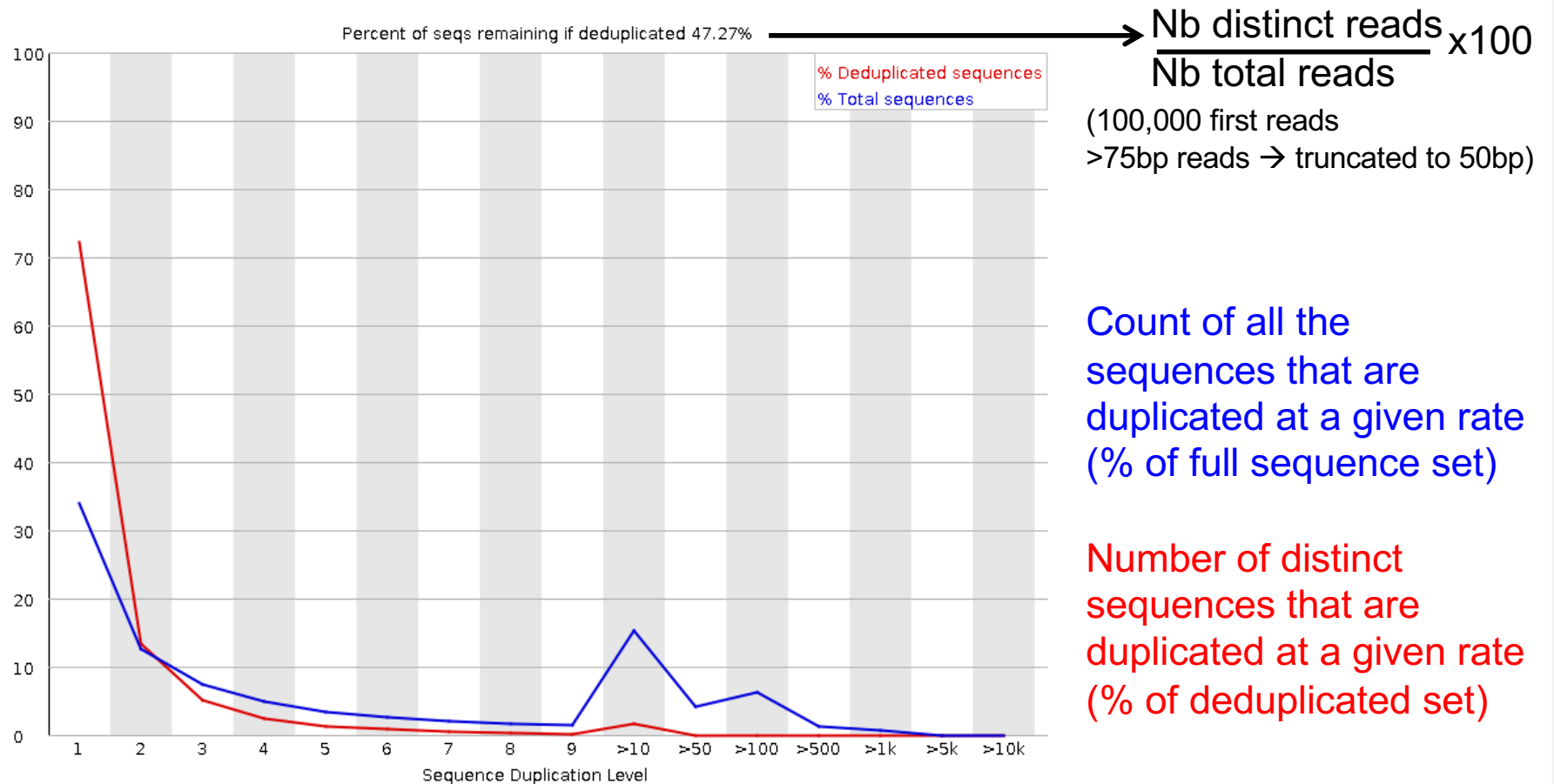
Sequence length distribution on another sample

- Useful when different sequence lengths in the file



Sequence duplication levels

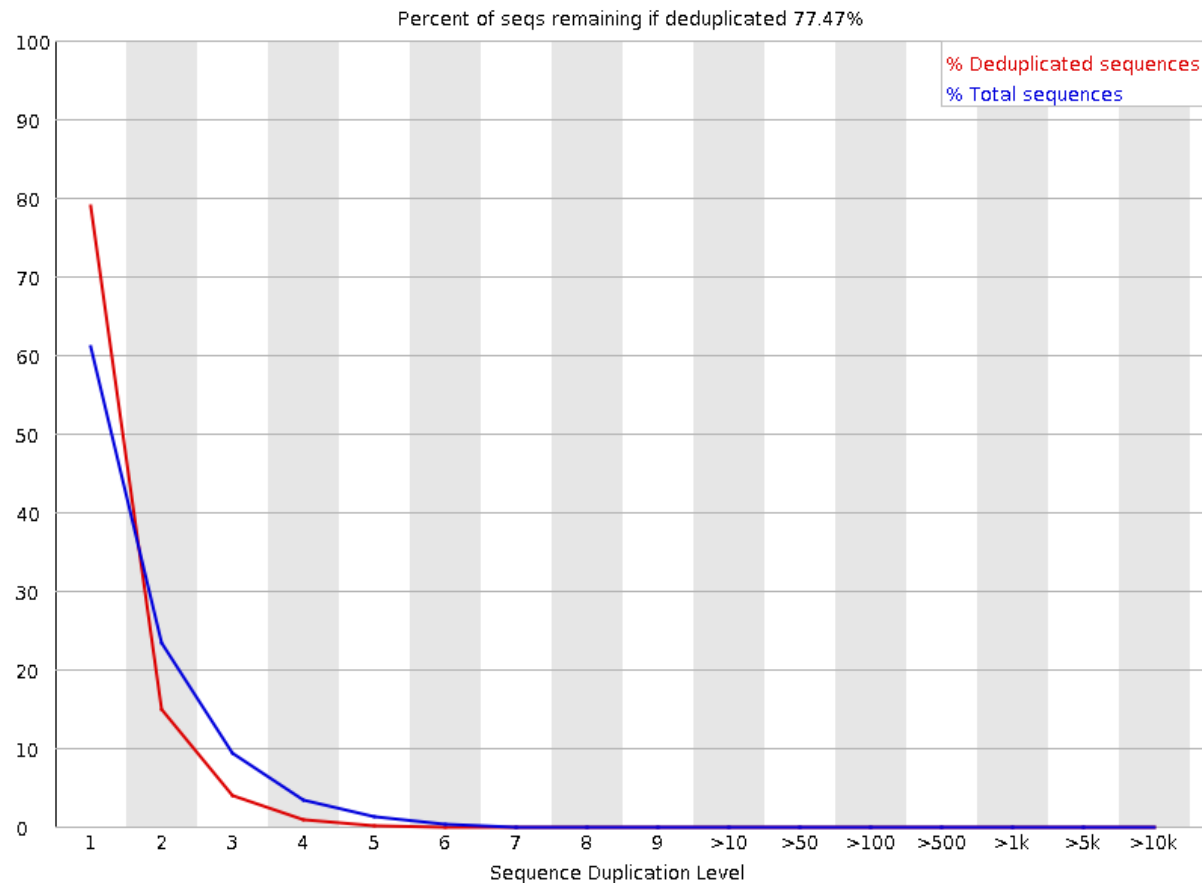
■ Relative number of sequences with different degrees of duplication



➔ OK for an RNA-seq sample :
Abundant mRNAs could lead to duplicated sequences

Sequence duplication levels on other samples

■ Example for a DNA-seq sample



- A high level of duplication may indicate an enrichment bias, e.g. PCR over amplification

Overrepresented sequences

- Lists all sequences representing more than 0.1% of the total

No overrepresented sequences

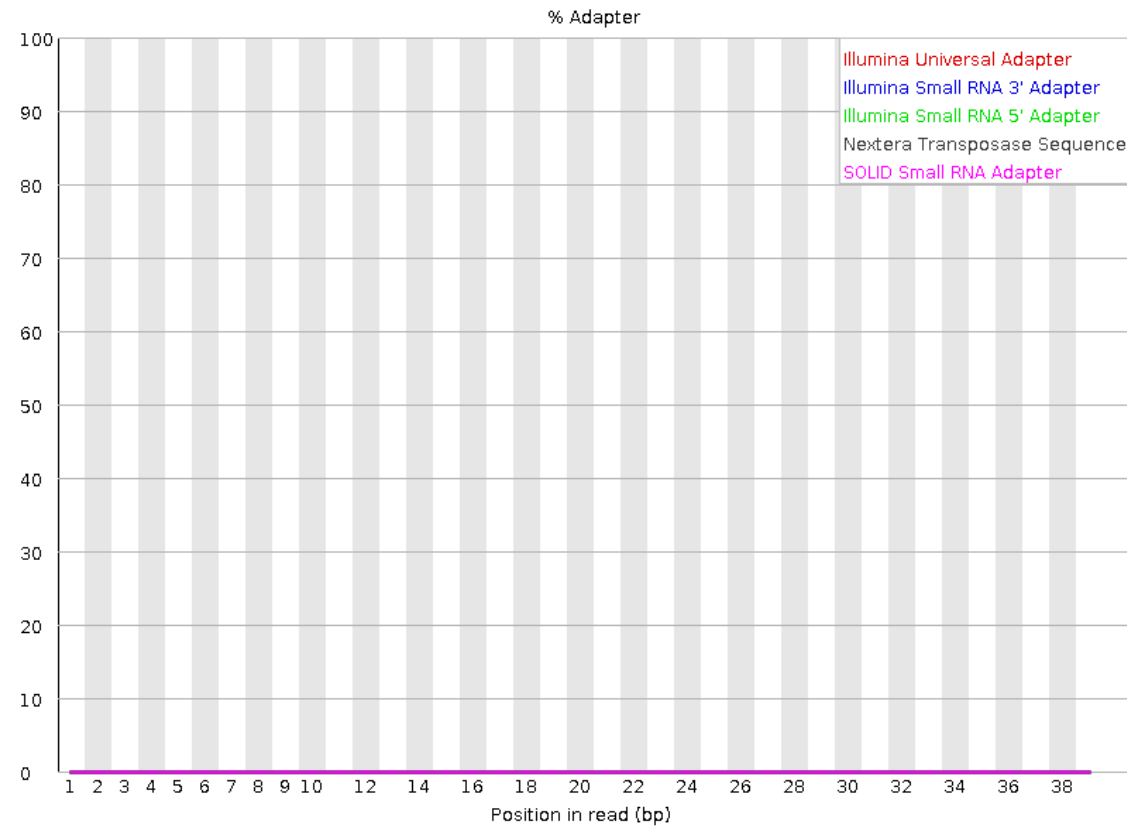
→ No sequences representing > 0.1% of the total

Overrepresented sequences on another sample

- For each overrepresented sequence, FastQC will look for matches in a database of common contaminants
 - ➔ report the best hit, e.g. :

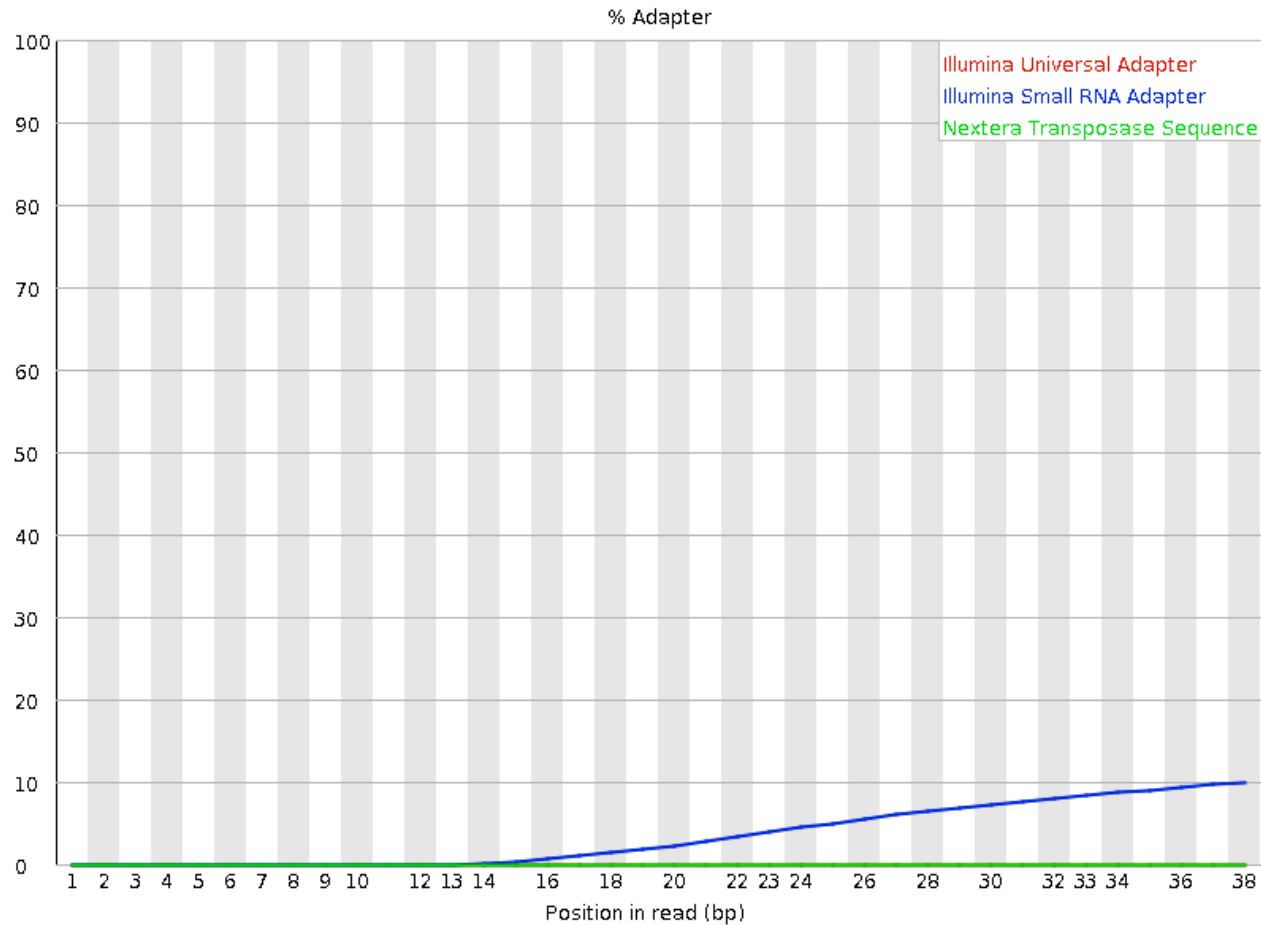
Sequence	Count	Percentage	Possible Source
AGATCGGAAGAGCACACTTCTGAACTCCAGTCACCGATGTATCTCGTATG	113163	0.614990735439532	TruSeq Adapter, Index 2 (97% over 49bp)
AGATCGGAAGAGCACACGTCTGAACTCAAGTCACCGATGTATCTCGTATG	41889	0.22764814397662272	TruSeq Adapter, Index 2 (97% over 49bp)
AGATCGGAAGAGCACACCTCTGAACTCCAGTCACCGATGTATCTCGTATG	39078	0.21237160520228368	TruSeq Adapter, Index 2 (97% over 49bp)

Adapter content



➔ No adapters

Adapter content on another sample



➔ Reads have to be trimmed before analysis

Quality control of Illumina data

- Primary analysis
- Quality control
- Data pre-processing

Data pre-processing

- Why ?
 - Remove bad quality/contaminant data
 - Improve confidence of downstream analysis
- Needed ?
 - Depend on what type of data and what type of analysis you want to perform on your data
 - e.g. small RNA-seq : adapters removal required
 - e.g. assembly : cleaned data required
 - e.g. variant calling : has to be performed only on good quality reads / part of reads
- Example of tools
 - Cutadapt
 - Trimmomatic

Data pre-processing

- Trimming
 - Remove low quality bases from the sequence end
- Filtering low quality reads
 - Keep only reads with a sufficient quality
- Removing/clipping adapter sequences
 - e.g. small RNA-seq library
 - Remove adapter sequences
 - Remove too-short sequences
 - Remove too-long sequences
 - Clip adapters
- Removing contaminants
 - e.g. sequences used during library preparation (e.g. spikes), from other organisms (e.g. xenografts), rRNA sequences

smallRNA sequence

