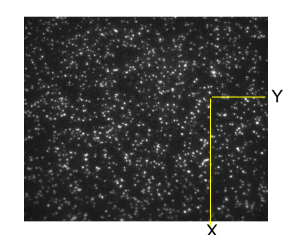
- Primary analysis
- Quality control
- Data pre-processing

- Primary analysis
- Quality control
- Data pre-processing

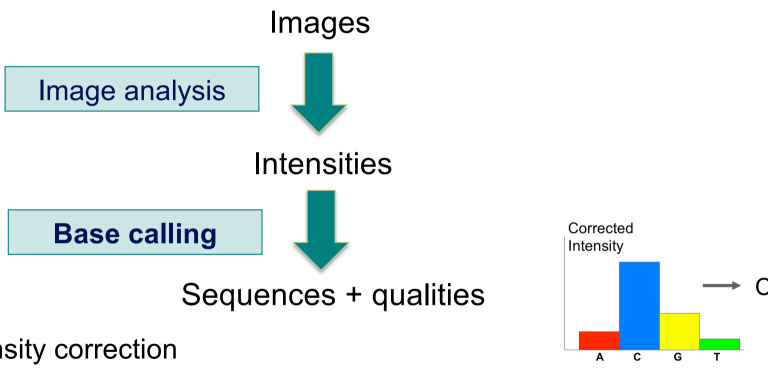
Primary analysis

Images
Image analysis
Intensities



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

Primary analysis



- Intensity correction
 - Take into account ≠ 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

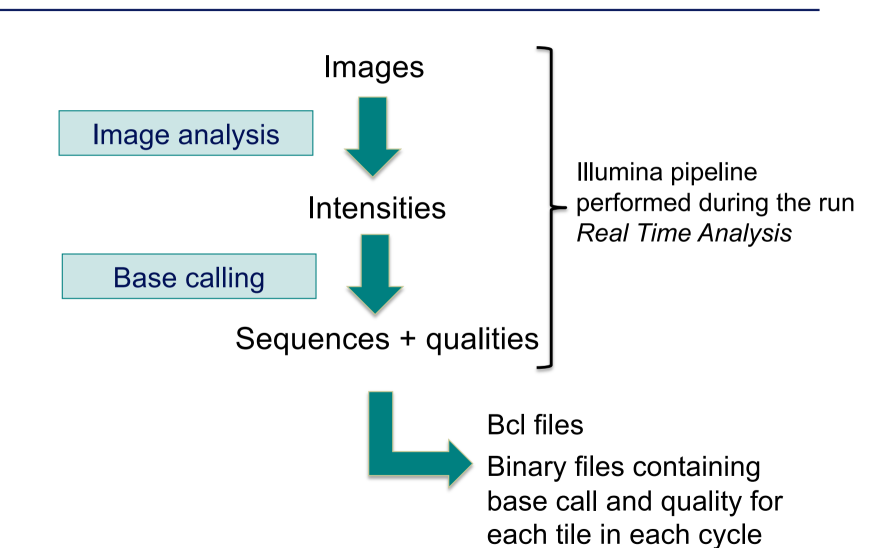
$$Q = -10 \, \log_{10} P$$

 $Q: quality\ score$

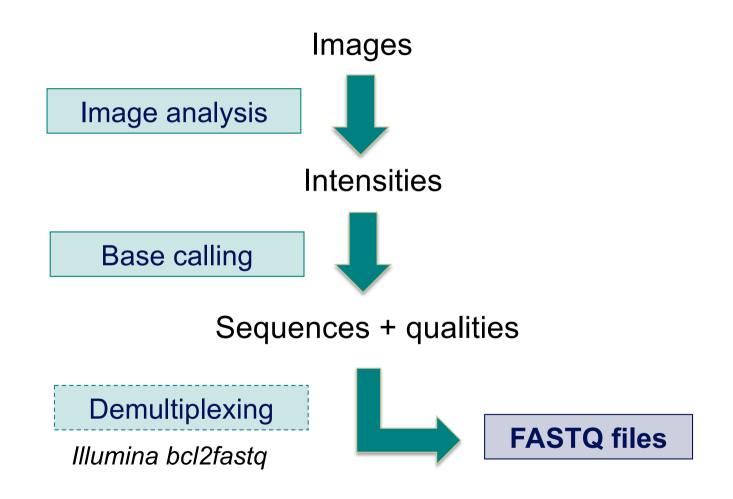
 $P: error\ probability$

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

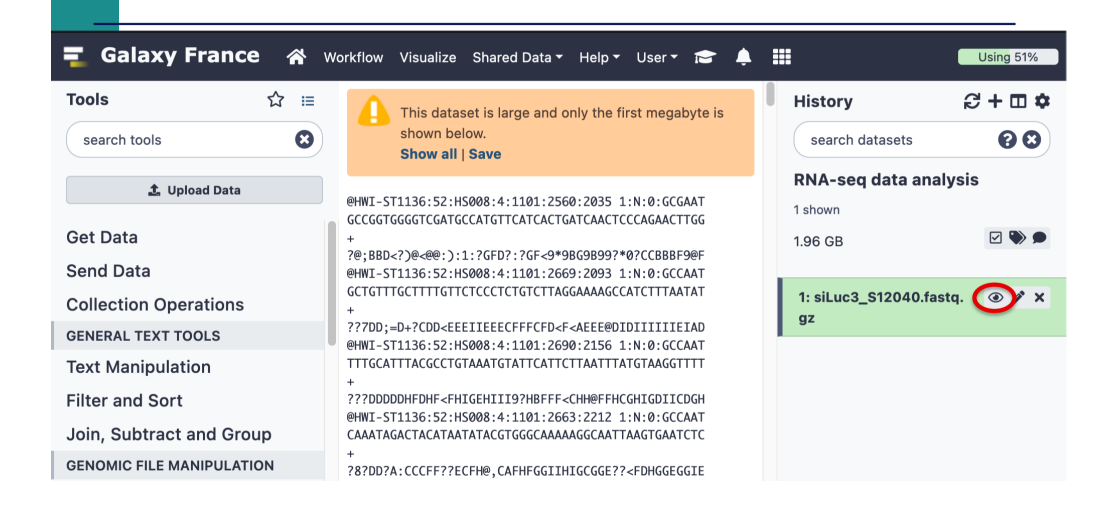
- Text file containing
 - Sequences
 - Qualities

Probability that the corresponding base call is incorrect

4 lines per sequence:

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

Beginning of siLuc3_S12040.fastq file



Sequence identifier in FASTQ files

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

@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 0
                 064 @
                         080 P
033 !
         049 1
                 065 A
                        081 Q
                                 097 a
                                         113 g
034 "
         050 2
                 066 B
                        082 R
                                 098 b
         051 3
                 067 C
                        083 S
                                 099 c
036 $
                 068 D
                        084 T
                                 100 d
037 %
         053 5
                 069 E
                        085 U
                                 101 e
                                         117 u
038 &
                070 F
        054 6
                        086 V
                                 102 f
                                         118 v
039 *
         055 7
                071 G
                                 103 q
                         087 W
040 (
         056 8
                 072 H
                        088 X
                                 104 h
                                         120 x
        057 9
041 )
               073 I
                        089 Y
                                 105 i
                                         121 y
042 *
         058 :
                074 J
                         090 Z
                                 106 i
043 +
         059 ;
                 075 K
                         091 Г
                                 107 k
044 ,
         060 < 076 L
                        092 \
                                 108 1
045 -
         061 =
                 077 M
                         093 1
                                 109 m
                         094 ^
046 .
         062 >
                 078 N
                                 110 n
047 /
        063 ?
                 079 0
                         095
                                 111 o
                                         127 🗅
```

- Binned in order to save space in the last versions of Illumina software, e.g.
 - 2 < real Q-score < 9 → binned Q-score = 6
 - 10 < real Q-score < 19 → binned Q-score = 15
 - **...**
 - real Q-score ≥ 40 → binned Q-score = 40

Paired-end FASTQ files

2 FASTQ files per sample



XXXX.R1.fastq.gz

XXXX.R2.fastq.gz

- Primary analysis
- Quality control
- Data pre-processing

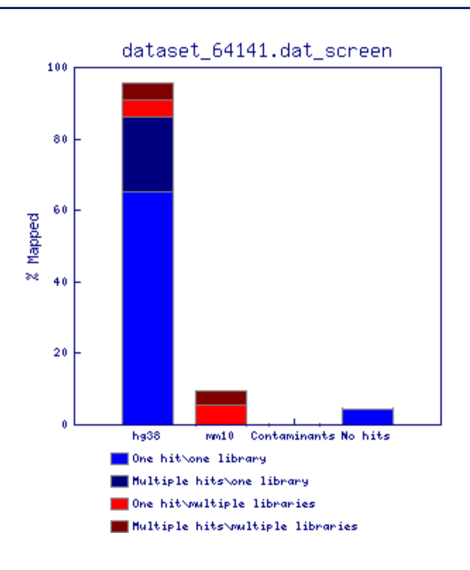
Quality control

- 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

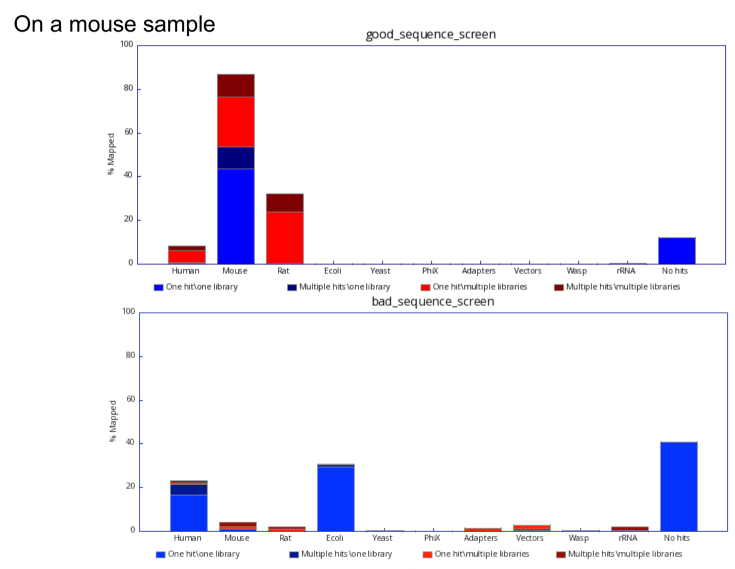
Some quality control tools

- 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



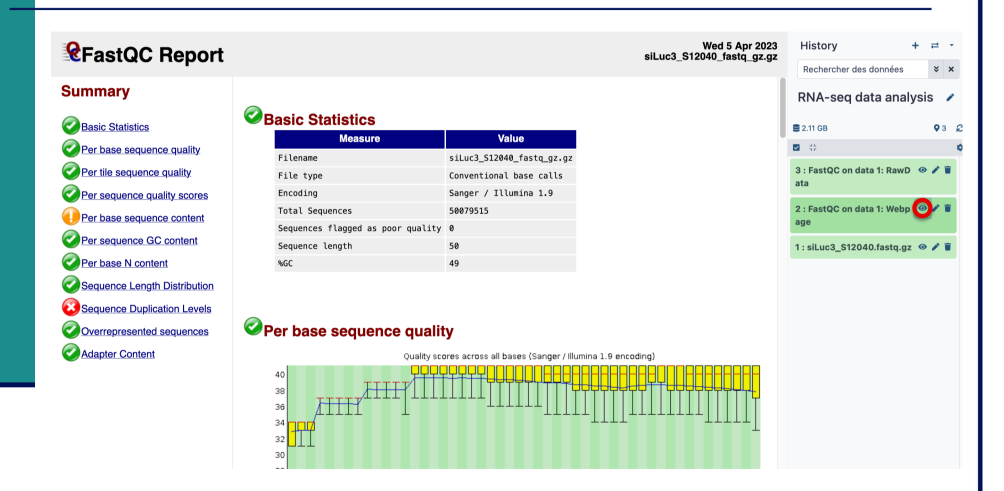
FastQC

- 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

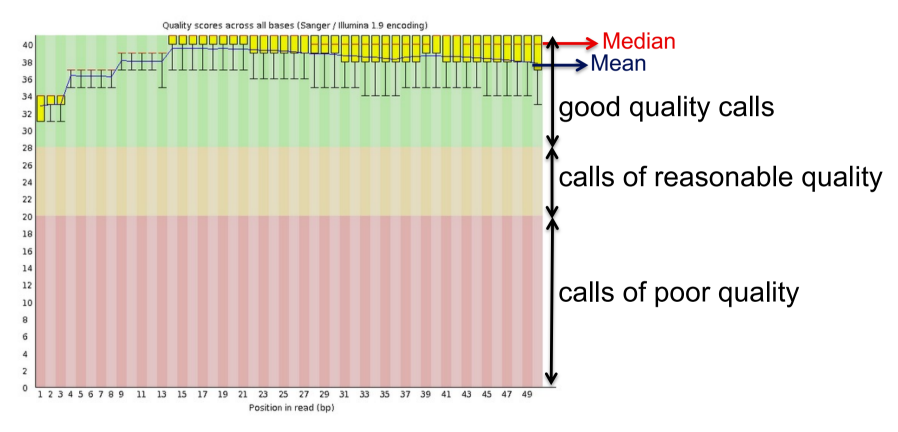


Basic Statistics

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

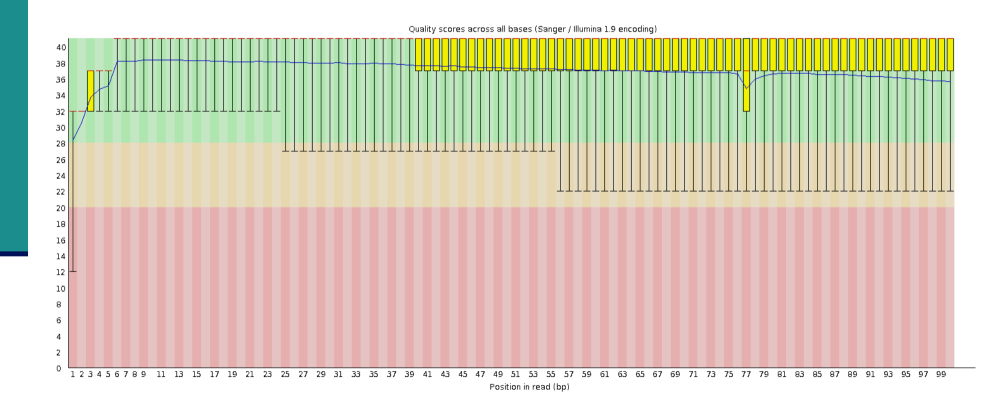
Per base sequence quality



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

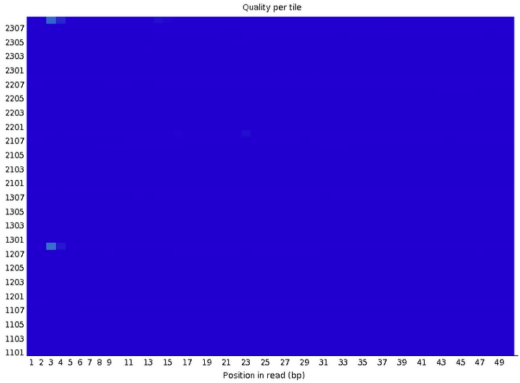
Per base sequence quality on another sample

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



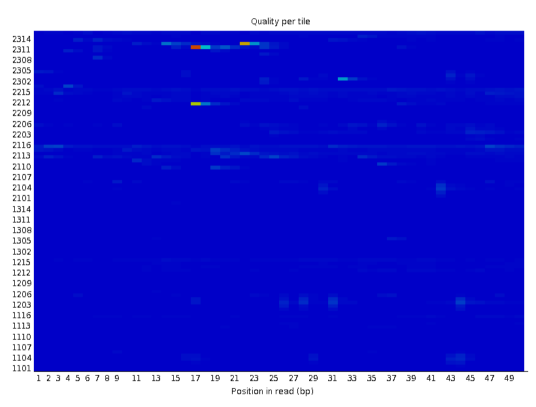
Per tile sequence quality

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



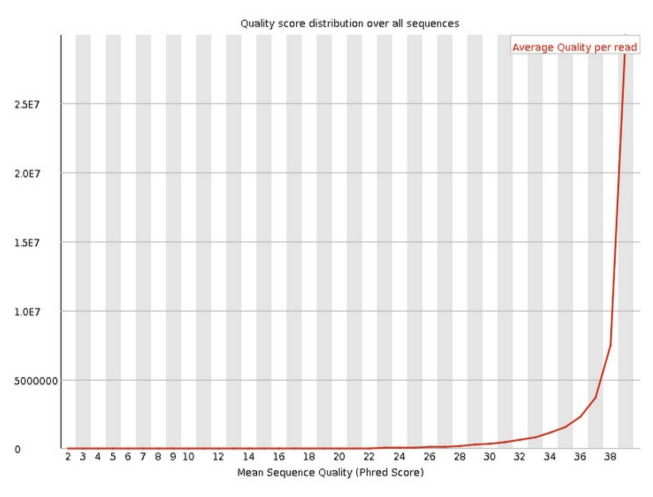
- → To see if there was a loss in quality associated with only one part of the flowcell
- → No poor quality tile for this sample

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

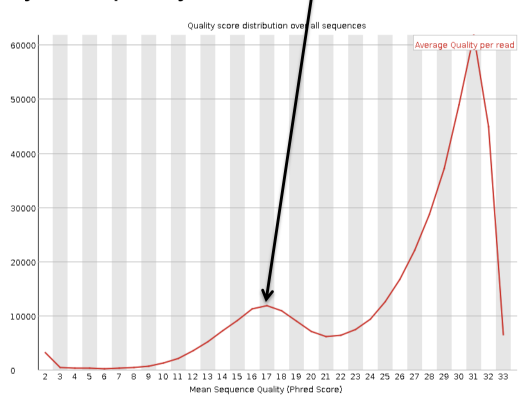
Per sequence quality scores



→ Good quality of all sequences

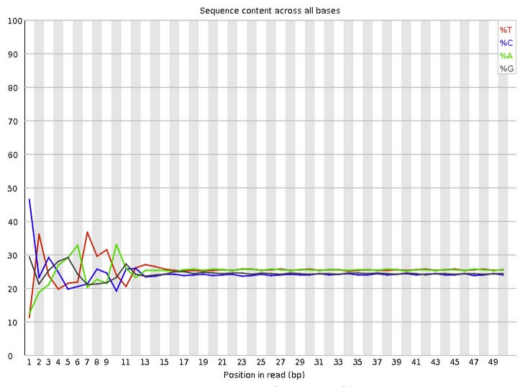
Per sequence quality score on another sample

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

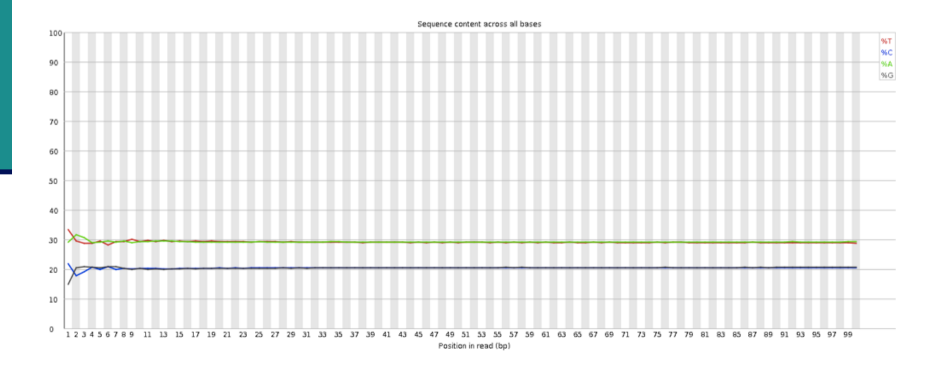


■ Known bias in the repartition of the first nt in RNA-seq libraries

- 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.

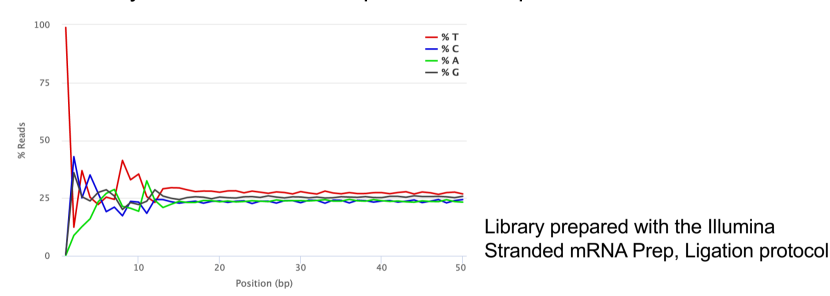
Per base sequence content on other samples

- 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

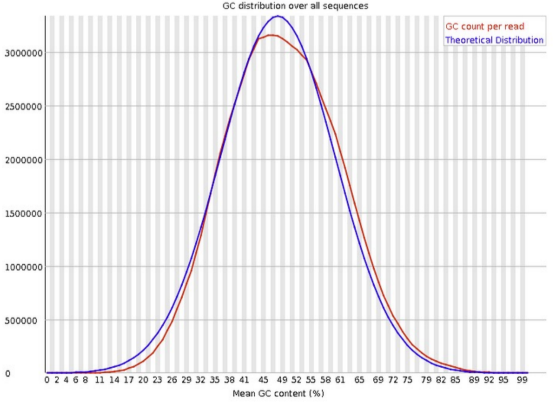
- Strong biases which change in different bases
 - Usually indicates an overrepresented sequence



- 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

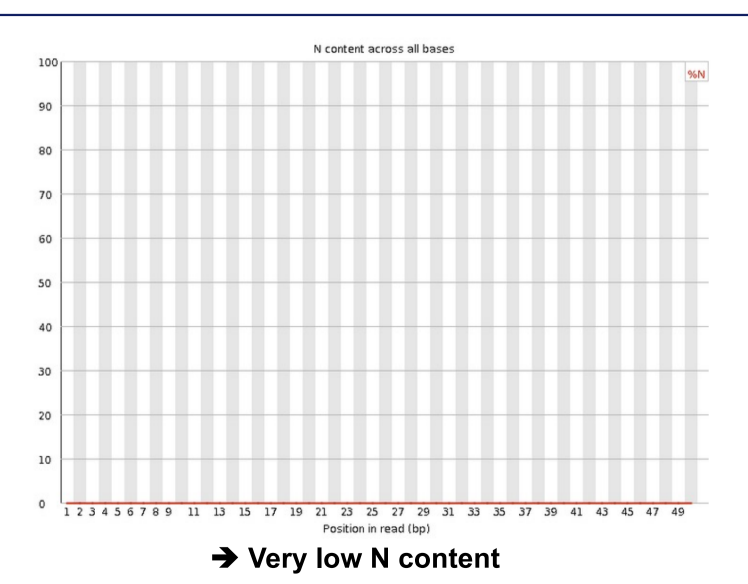
Per sequence GC content

 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)

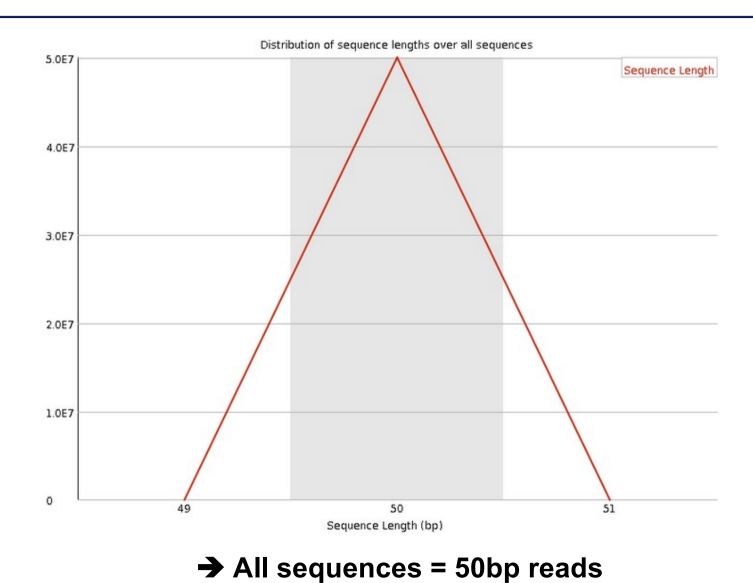


→ Observed GC distribution similar to the theoretical one

Per base N content

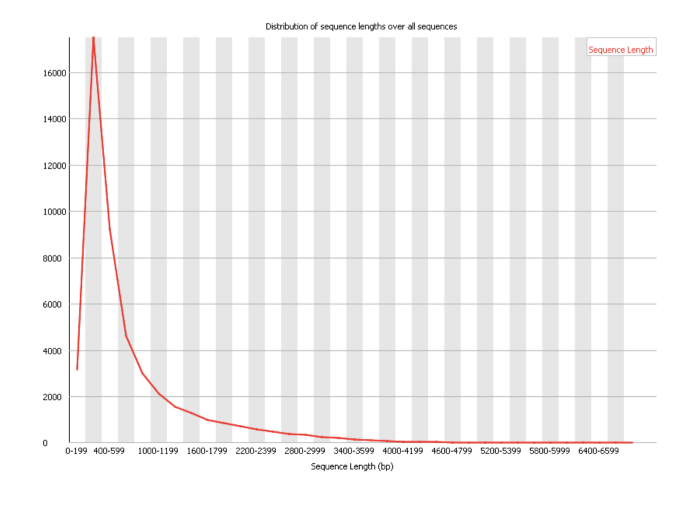


Sequence length distribution



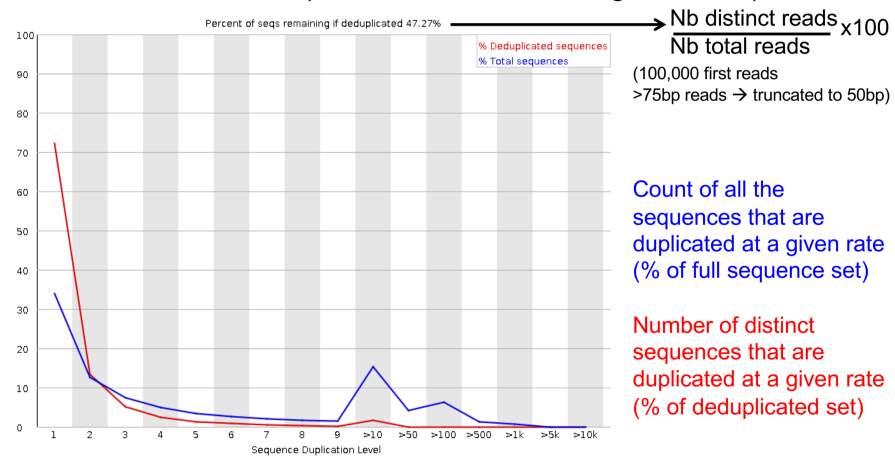
Sequence length distribution on another sample

Useful when different sequence lengths in the file



Sequence duplication levels

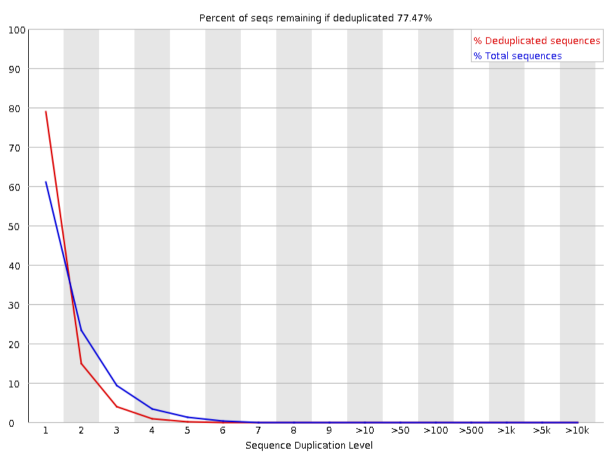
Relative number of sequences with different degrees of duplication



→ 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

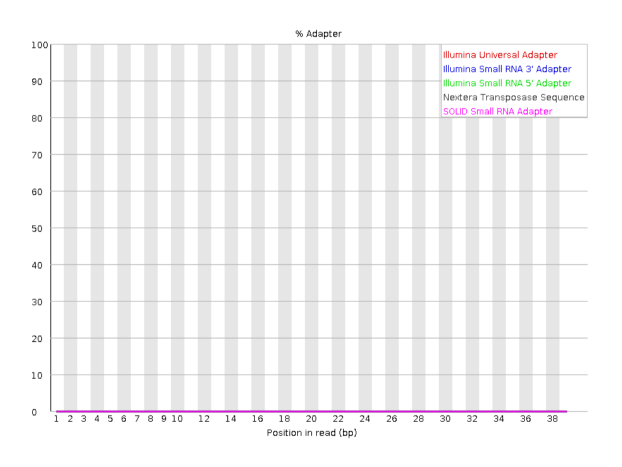
→ No sequences representing > 0.1% of the total

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.:

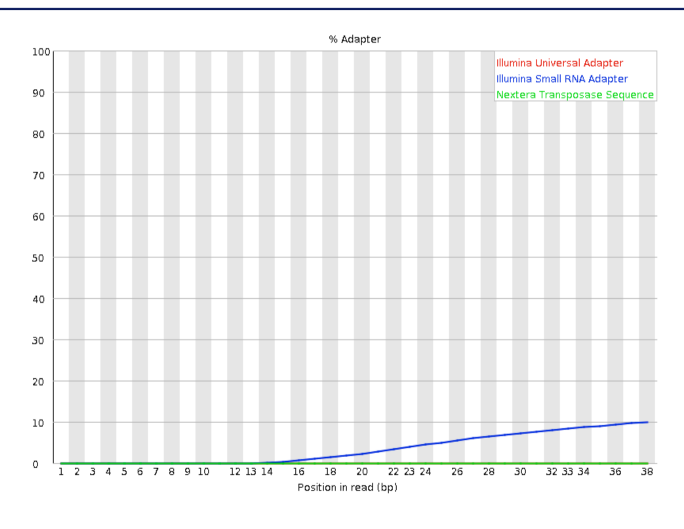
Sequence	Count	Percentage	Possible Source
${\tt AGATCGGAAGAGCACACTTCTGAACTCCAGTCACCGATGTATCTCGTATG}$	113163	0.614990735439532	TruSeq Adapter, Index 2 (97% over 49bp)
${\tt AGATCGGAAGAGCACACGTCTGAACTCAAGTCACCGATGTATCTCGTATG}$	41889	0.22764814397662272	TruSeq Adapter, Index 2 (97% over 49bp)
AGATCGGAAGAGCACCTCTGAACTCCAGTCACCGATGTATCTCGTATG	39078	0.21237160520228368	TruSeq Adapter, Index 2 (97% over 49bp)

Adapter content





Adapter content on another sample



→ Reads have to be trimmed before analysis

- 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

