Quality control and artefact removal

FastQC, Cutadapt, Trimmomatic, Fastx toolkit

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Overview

- Quality control
 - FastQC
- Artefact removal
 - Cutadapt/TrimGalore, Trimmomatic

Why do we need quality control?

- NGS sequencing generates highly accurate data, but it can have certain types of errors:
 - Contamination with adapters
 - Technical duplication in the library
 - Failure at specific parts of the flowcell
 - PCR duplicates
 - Etc.
- This is why it is important to check the data quality before alignment
- FastQC:
 - <u>http://www.bioinformatics.babraham.ac.uk/projects/fastqc/</u>
 - Reads in fastq files and generates reports based on the quality information that the sequencer provided
 - Command line and interactive mode
 - Outputs an html report and a .zip file with the raw quality data
- MultiQC:
 - <u>http://multiqc.info/</u>
 - Aggregates FastQC results of multiple analyses into a single report

FastQC - basic statistics

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	



Measure	Value		
Filename	bad_sequence.txt		
File type	Conventional base calls		
Encoding	Illumina 1.5		
Total Sequences	395288		
Sequences flagged as poor quality	0		
Sequence length	40		
%GC	47		

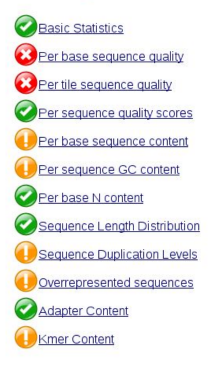
http://www.bioinformatics.babraham.ac.uk/projects/fastqc/ good_sequence_short_fastqc.html http://www.bioinformatics.babraham.ac.uk/projects/fastqc/ bad_sequence_fastqc.html

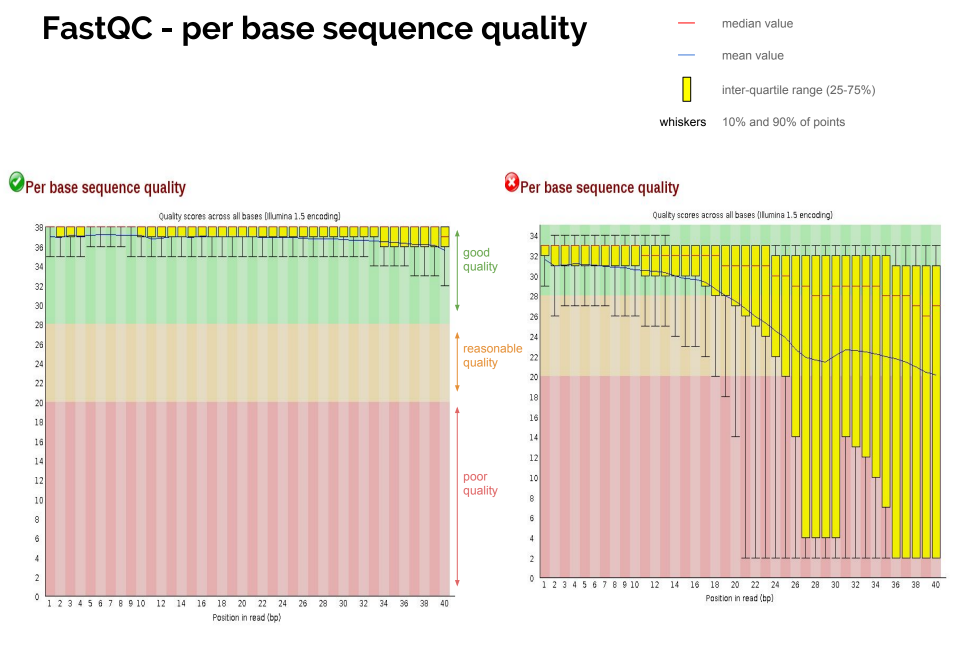
FastQC - summary

Summary



Summary

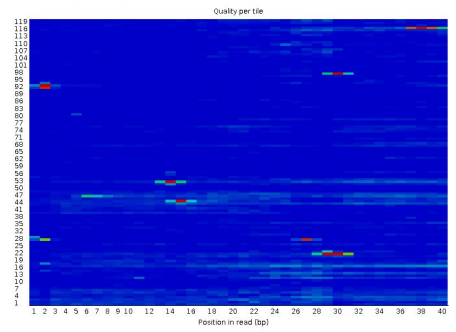




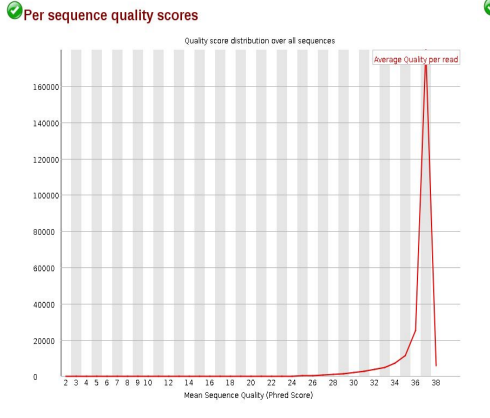
FastQC - per tile sequence quality

Per tile sequence quality Quality per tile 1 2 3 4 5 6 7 8 9 10 12 14 Position in read (bp)

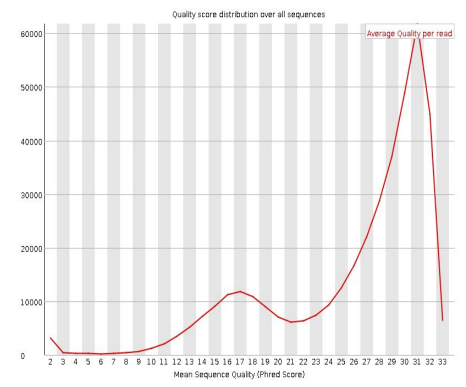
Per tile sequence quality



FastQC - per sequence quality scores

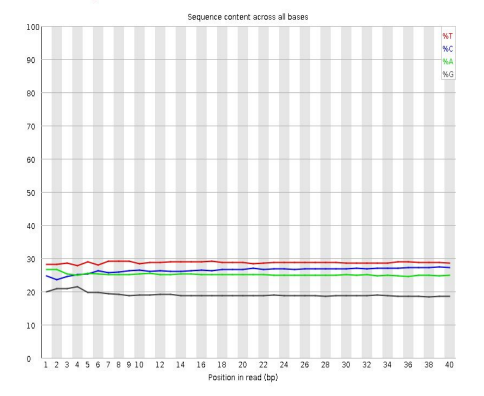


Per sequence quality scores

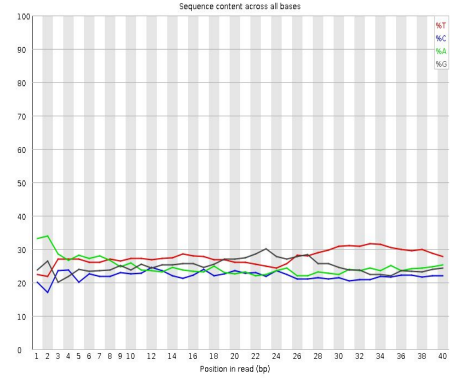


FastQC - per base sequence content

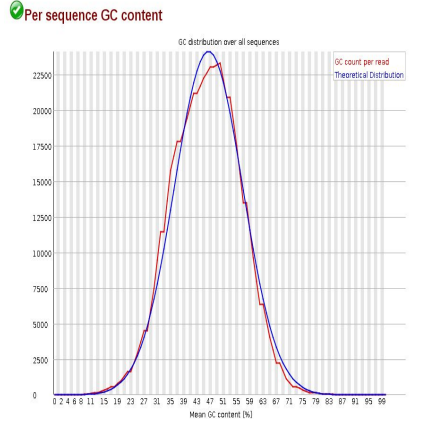
Per base sequence content



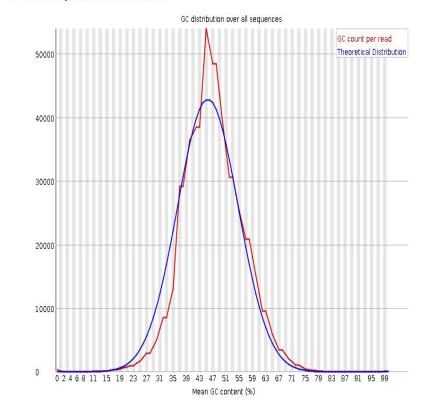
Per base sequence content



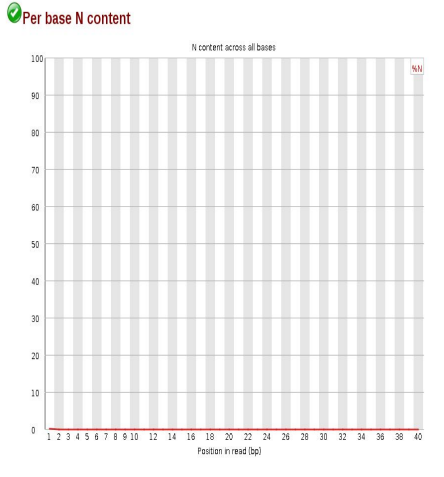
FastQC - per sequence GC content



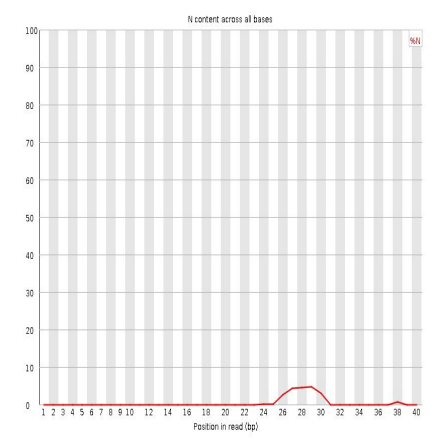
UPer sequence GC content



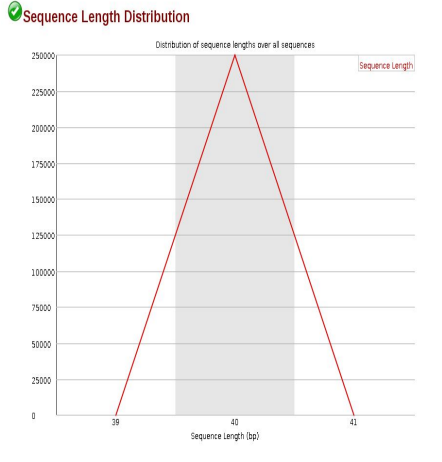
FastQC - per base N content



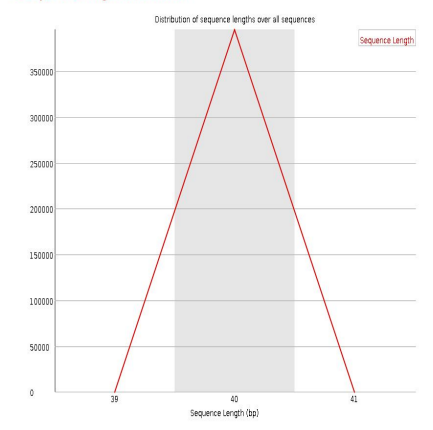
Per base N content



FastQC - sequence length distribution

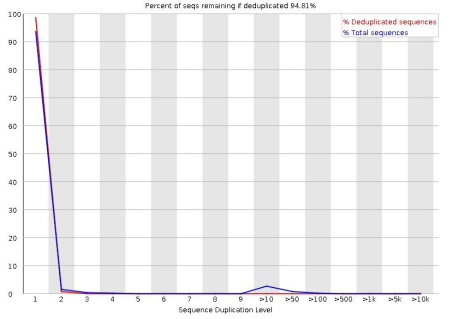


Sequence Length Distribution

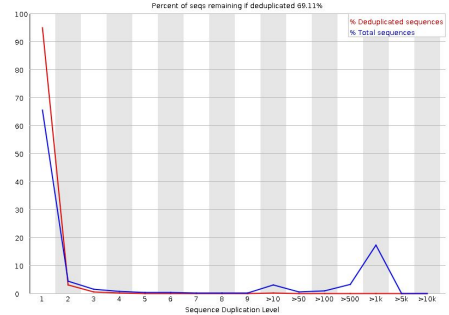


FastQC - sequence duplication level

Sequence Duplication Levels



Sequence Duplication Levels



FastQC - overrepresented sequences

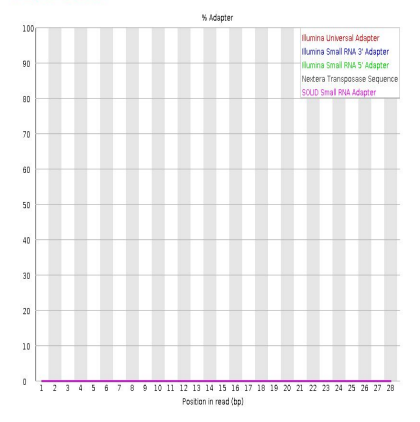
Overrepresented sequences No overrepresented sequences

Overrepresented sequences

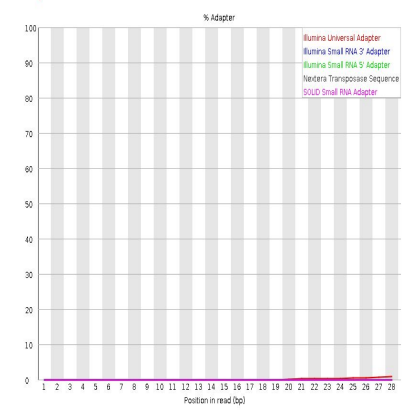
Sequence	Count	Percentage	Possible Source
AGAGTTTTATCGCTTCCATGACGCAGAAGTTAACACTTTC	2065	0.5224039181558763	No Hit
GATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCCATG	2047	0.5178502762542754	No Hit
ATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCCATGA	2014	0.5095019327680071	No Hit
CGATAAAAATGATTGGCGTATCCAACCTGCAGAGTTTTAT	1913	0.4839509420979134	No Hit
GT AT CCAACCT GCAGAGTTTT AT CGCTT CCAT GACGCAGA	1879	0.47534961850600066	No Hit
AAAAAT GATT GGCGT AT CCAACCT GCAGAGTTTT AT CGCT	1846	0.4670012750197325	No Hit
TGATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCCAT	1841	0.46573637449150995	No Hit
AACCTGCAGAGTTTTATCGCTTCCATGACGCAGAAGTTAA	1836	0.46447147396328753	No Hit
GAT AAAAAT GATT GGCGT AT CCAACCT GCAGAGTTTT AT C	1831	0.4632065734350651	No Hit
AAAT GATT GGCGT AT CCAACCT GCAGAGTTTT AT CGCTT C	1779	0.45005160794155147	No Hit
AT GATT GGCGT AT CCAACCT GCAGAGTTTT AT CGCTT CCA	1779	0.45005160794155147	No Hit
AATGATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCC	1760	0.4452449859343061	No Hit
AAAATGATTGGCGTATCCAACCTGCAGAGTTTTATCGCTT	1729	0.4374026026593269	No Hit
CGTATCCAACCTGCAGAGTTTTATCGCTTCCATGACGCAG	1713	0.43335492096901496	No Hit
AT CCAACCT GCAGAGTTTT AT CGCTT CCAT GACGCAGAAG	1708	0.43209002044079253	No Hit
CAGAGTTTTATCGCTTCCATGACGCAGAAGTTAACACTTT	1684	0.42601849790532476	No Hit
TGCAGAGTTTTATCGCTTCCATGACGCAGAAGTTAACACT	1668	0.4219708162150128	No Hit
CAACCTGCAGAGTTTTATCGCTTCCATGACGCAGAAGTTA	1668	0.4219708162150128	No Hit
TATCCAACCTGCAGAGTTTTATCGCTTCCATGACGCAGAA	1630	0.4123575722005221	No Hit
CGGTTCAGCAGGAATGCCGAGATCGGAAGAGCGGTTCAGC			Illumina Paired End PCR Primer 2 (96% over 25b)
TCT GCAGGTT GGAT ACGCCAAT CATTTTT AT CGAAGCGCG		0.1479933618020279	
CGCTT AAAGCT ACCAGTT AT AT GGCT GGGGGGTTTTTTT	552	0.13964501831575965	No Hit
CT CT GCAGGTT GGAT ACGCCAAT CATTTTT AT CGAAGCGC	532	0.1345854162028698	No Hit
CT GCGT CAT GGAAGCGAT AAAACT CT GCAGGT T GGAT ACG	515	0.13028475440691342	No Hit
CTGCAGGTTGGATACGCCAATCATTTTTATCGAAGCGCGC	505	0.12775495335046852	No Hit
GCTT AAAGCT ACCAGTT AT AT GGCT GGGGGGTTTTTTTT G	411	0.10397482341988626	No Hit

FastQC - adapter content

Adapter Content

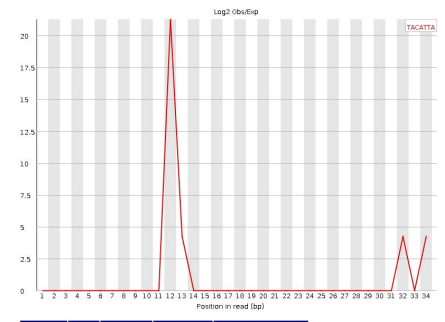


Adapter Content



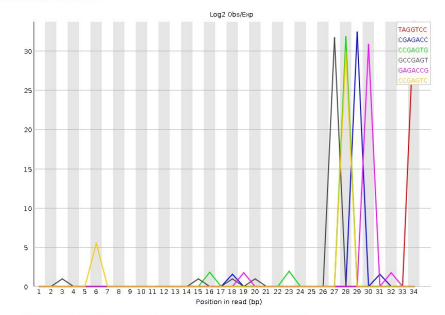
FastQC - kmer content

Kmer Content



Sequence	Count	PValue	Obs/Exp Max	Max Obs/Exp Position
TACATTA	40	0.003151852	21.2465	12

WKmer Content



Sequence	Count	PValue	Obs/Exp Max	Max Obs/Exp Position
TAGGTCC	30	1.5992917E-5	33.6211	34
CGAGACC	105	0.0	32.37975	29
CCGAGTG	90	0.0	31.803032	28
GCCGAGT	170	0.0	31.625078	27
GAGACCG	95	0.0	30.826315	30
CCGAGTC	30	4.3762376E-4	29.815344	28

Overview

- Quality control
 - FastQC
- Artefact removal
 - Cutadapt/TrimGalore, Trimmomatic

Artefact removal

- Important when the quality needs to be increased
- Adapter trimming
 - Based on "Overrepresented sequences" and/or "Kmer content" you might identify certain adapter contaminations that needs to be trimmed
 - Spikes in "Per sequence GC content" usually indicate adapter contamination
- Quality-based trimming
 - When the quality drops eg. towards the end of reads
 - When the "Per base sequence content" shows bias in sequence composition towards beginning/end
 - You can trim regions below a certain quality threshold (eg. 20)
 - You can trim *n* bases from beginning/end of all your reads

Artefact removal - paired-end data

- We want to preserve the pairs so that aligners will know which reads belong together
- We have to keep track of the pairs of those reads that are removed from one of the paired files
- Four output files will be produced, two with the trimmed paired reads and two with the unpaired ones.

Artefact removal - tools

- Cutadapt/TrimGalore
 - <u>http://cutadapt.readthedocs.io/en/stable/index.html</u>
 - TrimGalore: wrapper around Cutadapt
- Trimmomatic
 - <u>http://www.usadellab.org/cms/?page=trimmomatic</u>
- Fastx toolkit
 - <u>http://hannonlab.cshl.edu/fastx_toolkit/</u>
 - short read pre-processing tool
 - fastx_trimmer: fixed length trimmer
 - fastq_quality_filter: quality based trimmer
 - fastx_artifacts_filter: artefact remover