



Quality control and artefact removal

FastQC, Cutadapt, Trimmomatic, Fastx toolkit

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CRUK CI Bioinformatics Summer School July 2019

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
 - o Etc.
- This is why it is important to check the data quality before alignment
- FastQC:
 - http://www.bioinformatics.babraham.ac.uk/projects/fastqc/
 - Reads in .fastq (or .bam) files and generates reports based on the quality information that the sequencer provided (or alignment generated)
 - Command line and interactive mode
 - Outputs an html report and a .zip file with the raw quality data
- MultiQC:
 - http://multigc.info/
 - 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 | |

Basic Statistics

| 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

- 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
- Kmer Content

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

FastQC - per base sequence quality

median value

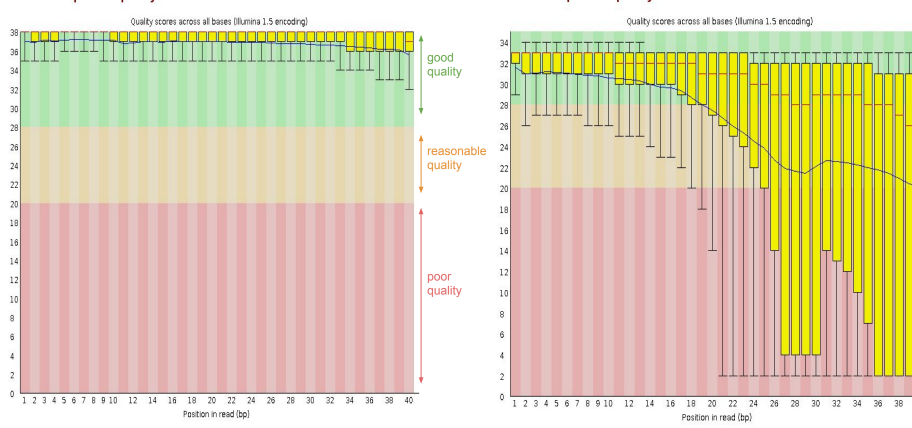
mean value

inter-quartile range (25-75%)

whiskers 10% and 90% of points

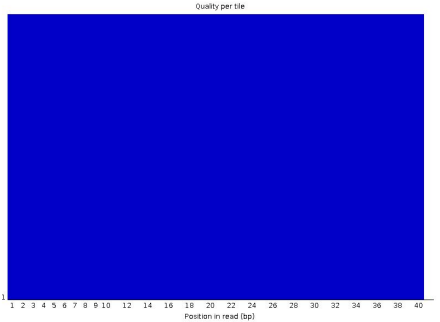


Per base sequence quality

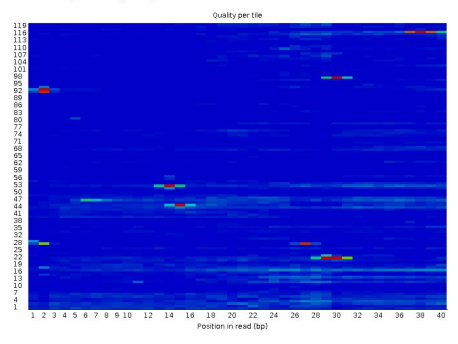


FastQC - per tile sequence quality



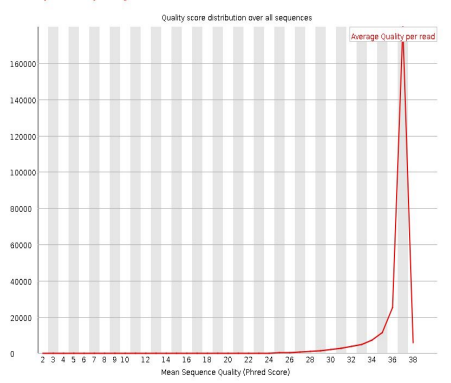


Per tile sequence quality

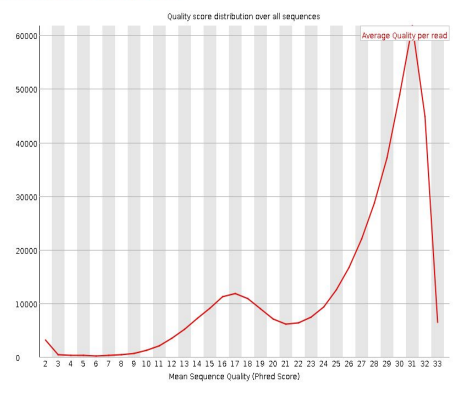


FastQC - per sequence quality scores



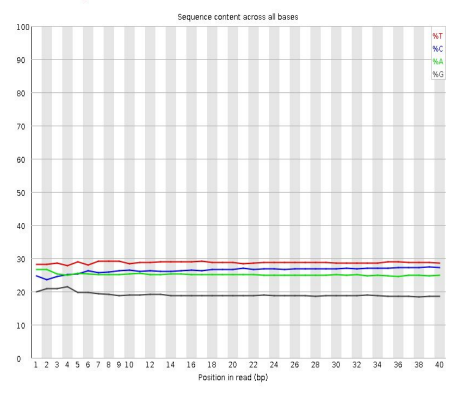


Per sequence quality scores

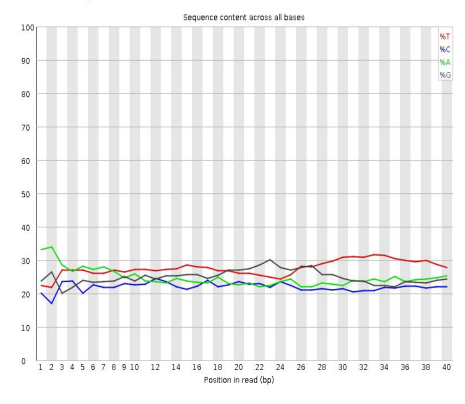


FastQC - per base sequence content



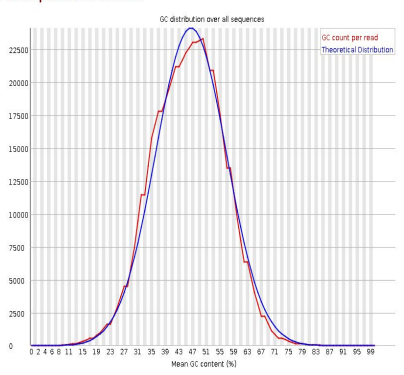


Per base sequence content

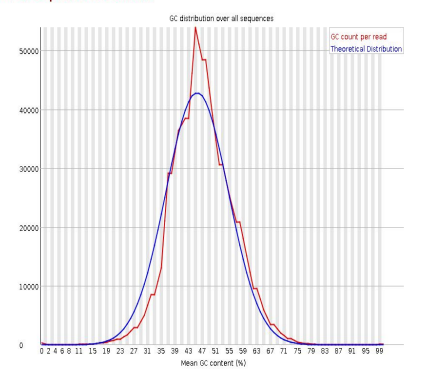


FastQC - per sequence GC content

Per sequence GC content

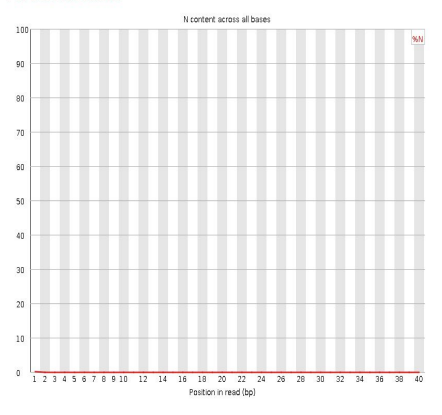


Per sequence GC content

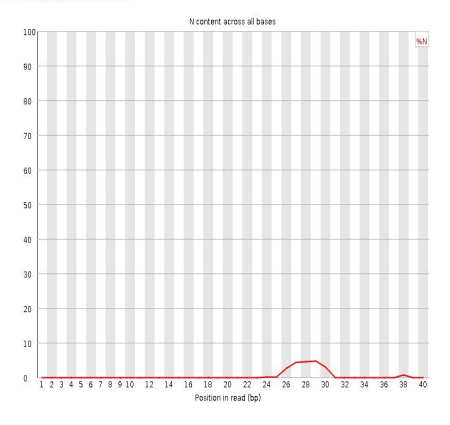


FastQC - per base N content



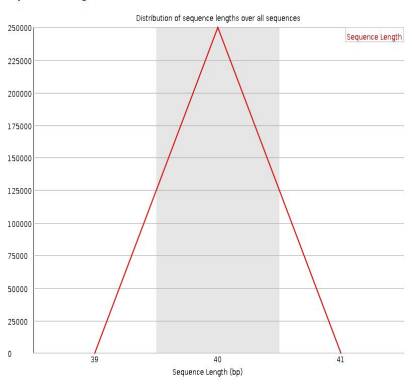


Per base N content

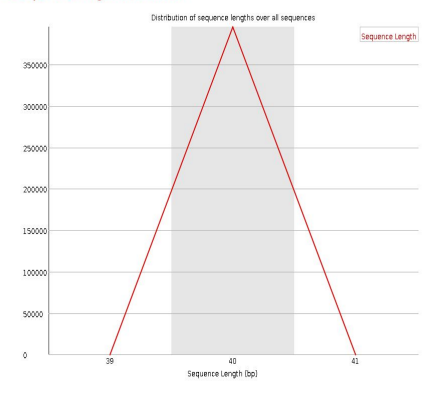


FastQC - sequence length distribution

Sequence Length Distribution

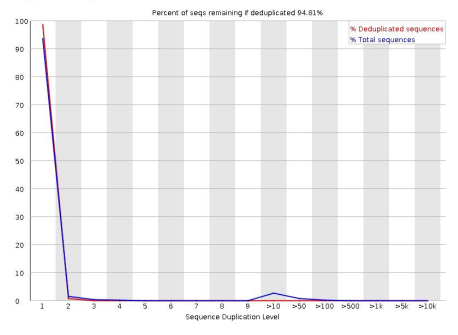


Sequence Length Distribution

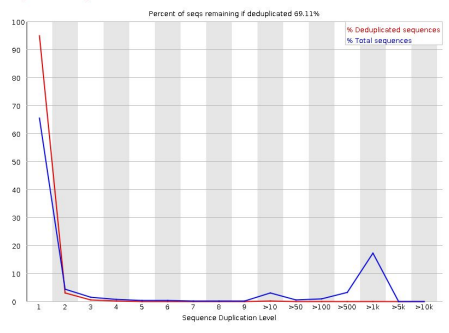


FastQC - sequence duplication level





Sequence Duplication Levels



FastQC - overrepresented sequences

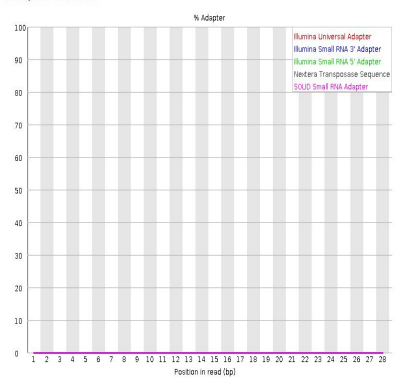


Overrepresented sequences

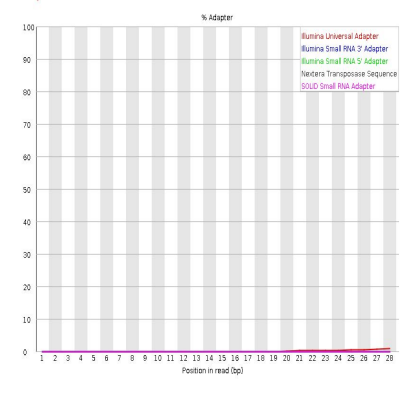
| Sequence | Count | Percentage | Possible Source |
|--|-------|---------------------|--|
| AGAGTTTTATCGCTTCCATGACGCAGAAGTTAACACTTTC | 2065 | 0.5224039181558763 | No Hit |
| GATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCCATG | 2047 | 0.5178502762542754 | No Hit |
| ATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCCATGA | 2014 | 0.5095019327680071 | No Hit |
| CGATAAAAATGATTGGCGTATCCAACCTGCAGAGTTTTAT | 1913 | 0.4839509420979134 | No Hit |
| GTATCCAACCTGCAGAGTTTTATCGCTTCCATGACGCAGA | 1879 | 0.47534961850600066 | No Hit |
| AAAAATGATTGGCGTATCCAACCTGCAGAGTTTTATCGCT | 1846 | 0.4670012750197325 | No Hit |
| TGATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCCAT | 1841 | 0.46573637449150995 | No Hit |
| AACCTGCAGAGTTTTATCGCTTCCATGACGCAGAAGTTAA | 1836 | 0.46447147396328753 | No Hit |
| GATAAAAATGATTGGCGTATCCAACCTGCAGAGTTTTATC | 1831 | 0.4632065734350651 | No Hit |
| AAATGATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTC | 1779 | 0.45005160794155147 | No Hit |
| ATGATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCCA | 1779 | 0.45005160794155147 | No Hit |
| AATGATTGGCGTATCCAACCTGCAGAGTTTTATCGCTTCC | 1760 | 0.4452449859343061 | No Hit |
| AAAATGATTGGCGTATCCAACCTGCAGAGTTTTATCGCTT | 1729 | 0.4374026026593269 | No Hit |
| CGTATCCAACCTGCAGAGTTTTATCGCTTCCATGACGCAG | 1713 | 0.43335492096901496 | No Hit |
| ATCCAACCTGCAGAGTTTTATCGCTTCCATGACGCAGAAG | 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 | 599 | 0.15153508328105078 | Illumina Paired End PCR Primer 2 (96% over 25bp) |
| TCTGCAGGTTGGATACGCCAATCATTTTTATCGAAGCGCG | 585 | 0.1479933618020279 | No Hit |
| CGCTTAAAGCTACCAGTTATATGGCTGGGGGGTTTTTTTT | 552 | 0.13964501831575965 | No Hit |
| CTCTGCAGGTTGGATACGCCAATCATTTTTATCGAAGCGC | 532 | 0.1345854162028698 | No Hit |
| ${\tt CTGCGTCATGGAAGCGATAAAACTCTGCAGGTTGGATACG}$ | 515 | 0.13028475440691342 | No Hit |
| $\tt CTGCAGGTTGGATACGCCAATCATTTTTATCGAAGCGCGC$ | 505 | 0.12775495335046852 | No Hit |
| ${\tt GCTTAAAGCTACCAGTTATATGGCTGGGGGGTTTTTTTTG}$ | 411 | 0.10397482341988626 | No Hit |

FastQC - adapter content

Adapter Content

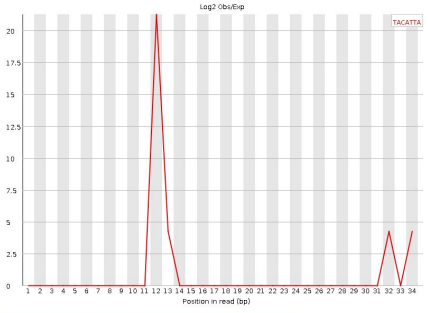


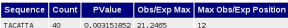
Adapter Content



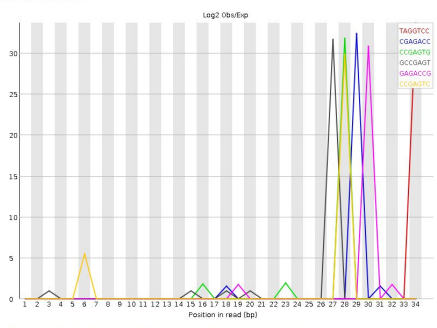
FastQC - kmer content







Windows 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

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 - Cutadapt/TrimGalore, Trimmomatic

Artefact removal

- Important when the quality needs to be increased
- Adapter trimming
 - Based on "Overrepresented Sequences", "Adapter Content" 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 ("Per base sequence quality")
 - 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
 - http://cutadapt.readthedocs.io/en/stable/index.html
 - o TrimGalore: wrapper around Cutadapt
- Trimmomatic
 - http://www.usadellab.org/cms/?page=trimmomatic
- Fastx toolkit
 - http://hannonlab.cshl.edu/fastx_toolkit/
 - short read pre-processing tool
 - fastx_trimmer: fixed length trimmer
 - fastq_quality_filter: quality based trimmer
 - fastx_artifacts_filter: artefact remover