ReportSummary



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

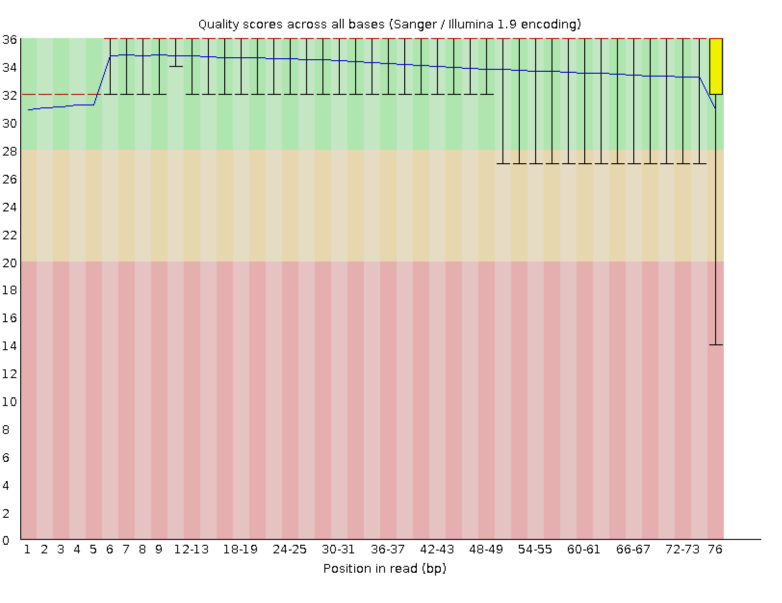
Overrepresented sequences

Adapter Content

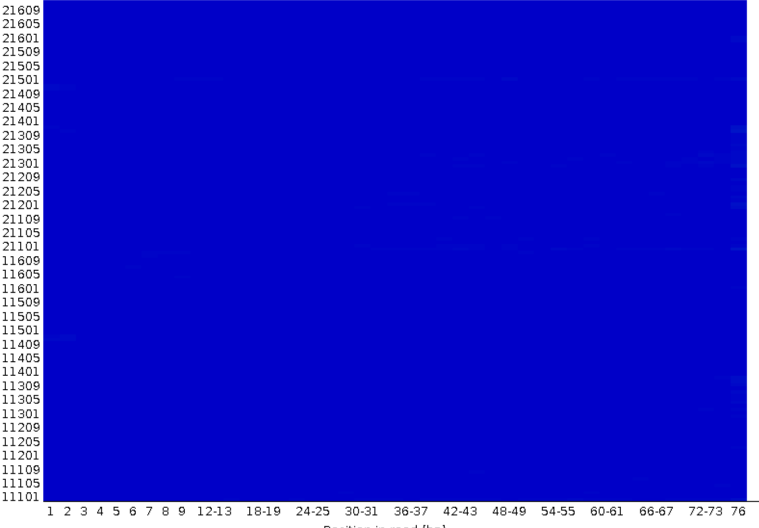
Basic Statistics

Measure	Value
Filename	ERR13985875_2.fastq
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	10689040
Sequences flagged as poor quality	0
Sequence length	76
%GC	47

Per base sequence quality

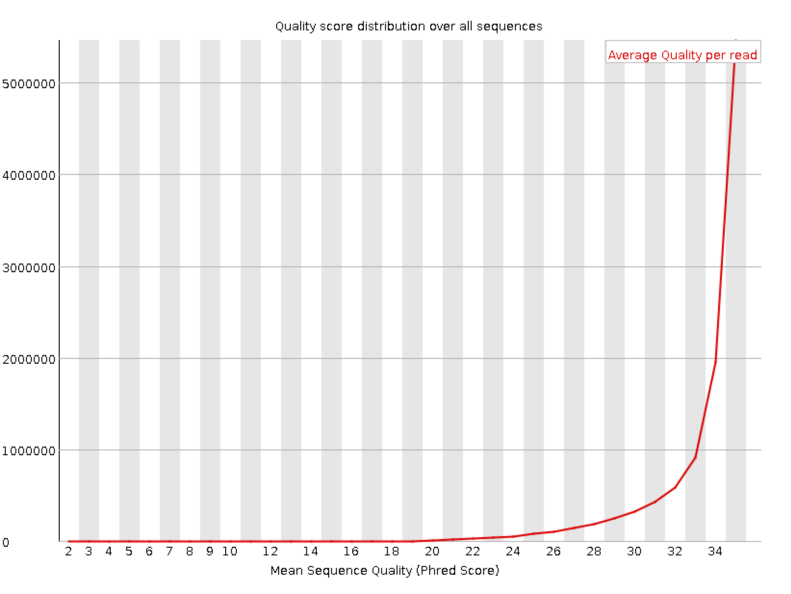




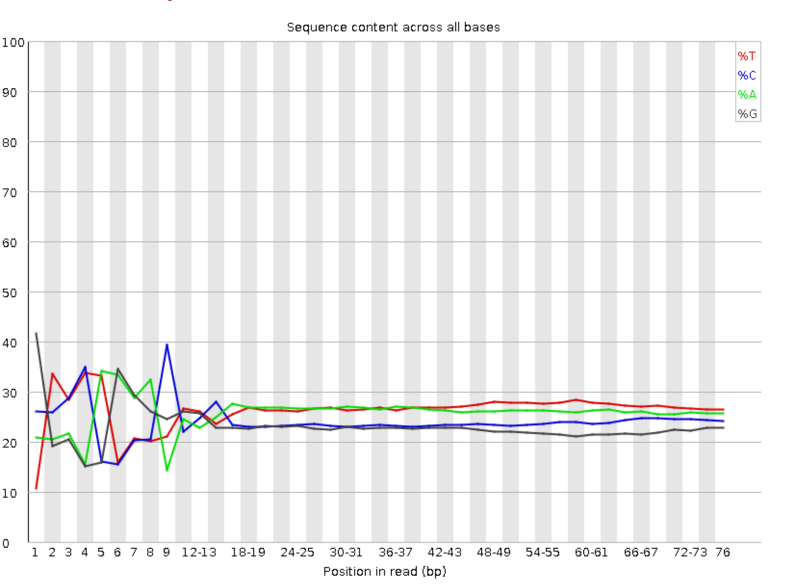


Position in read (bp)

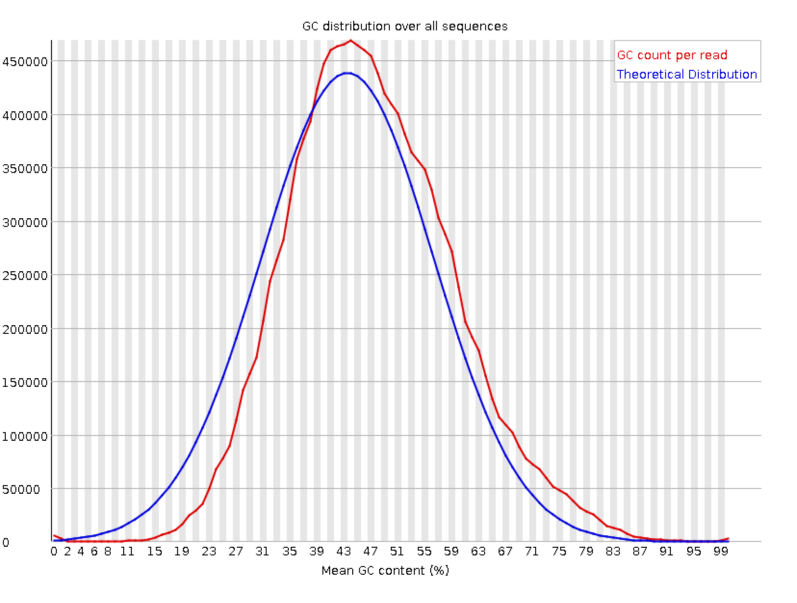
Per sequence quality scores



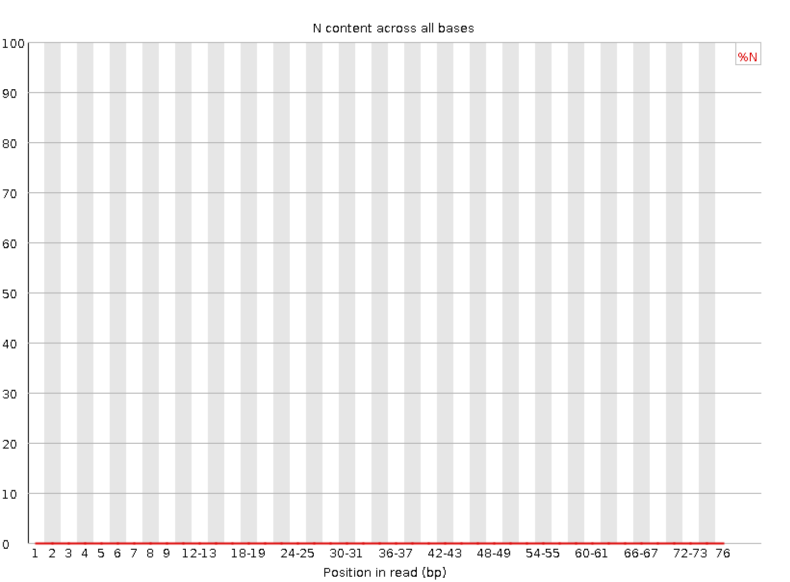
Per base sequence content



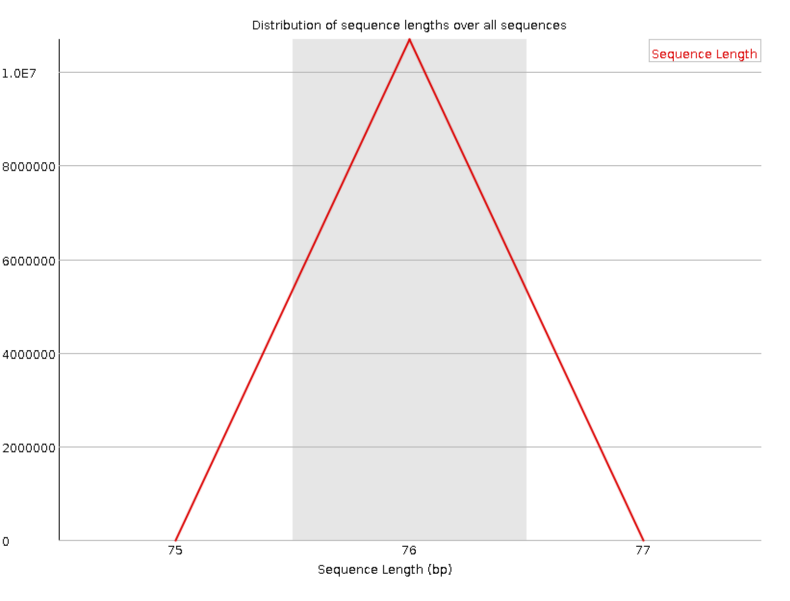
Per sequence GC content



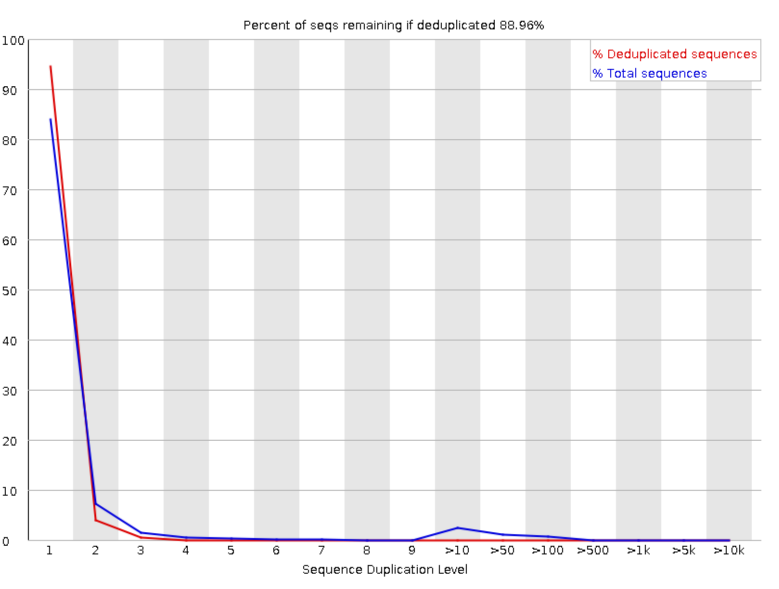
Per base N content



Sequence Length Distribution

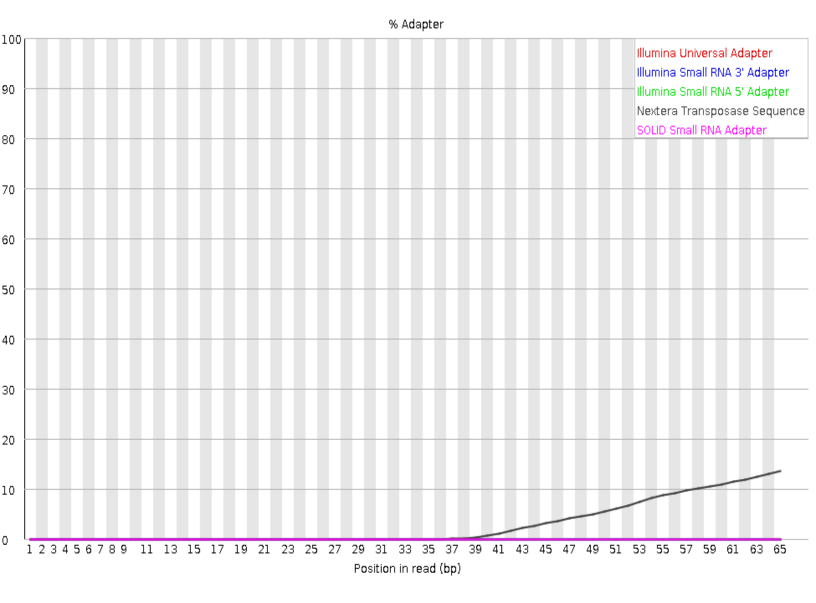


Sequence Duplication Levels









Produced by FastQC (version 0.11.7)