# **Report**Summary

Mon 24 Apr 2023 gc148-B\_trimmed\_R1.fastq.gz



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

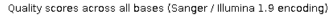
Overrepresented sequences

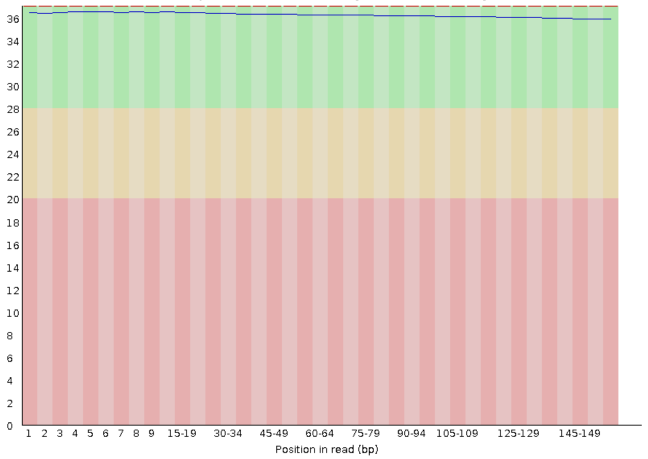
Adapter Content

#### Basic Statistics

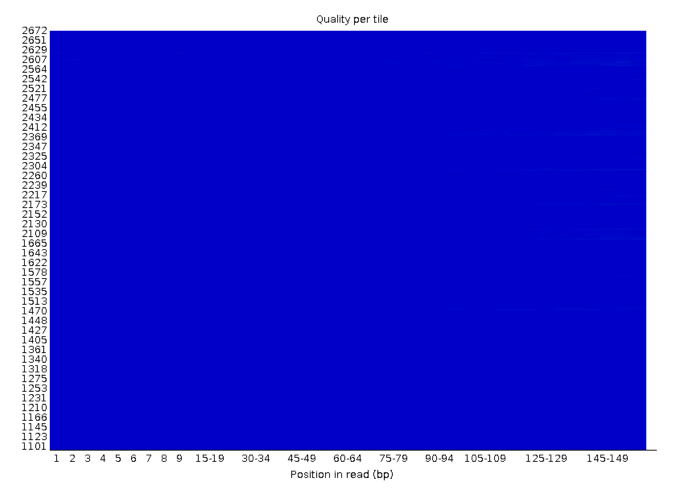
Measure	Value
Filename	<pre>gc148- Btrimmed_R1.fastq.gz</pre>
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	23668708
Sequences flagged as poor quality	0
Sequence length	35-159
%GC	50

### Per base sequence quality

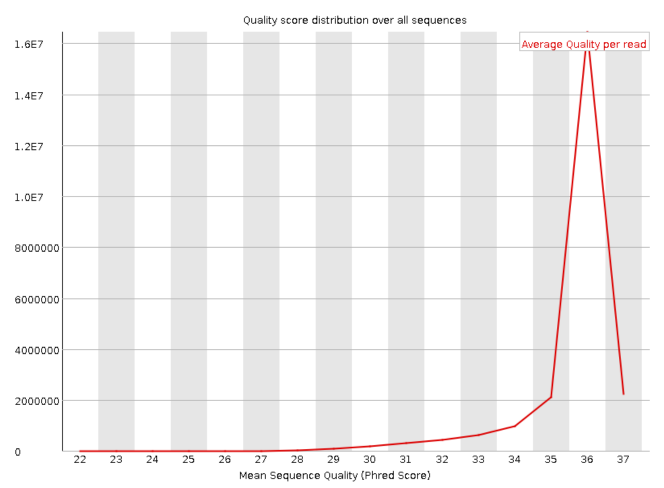




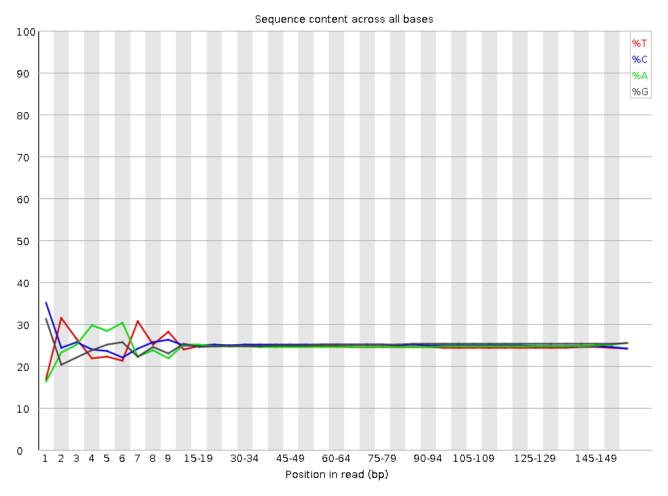
# Per tile sequence quality



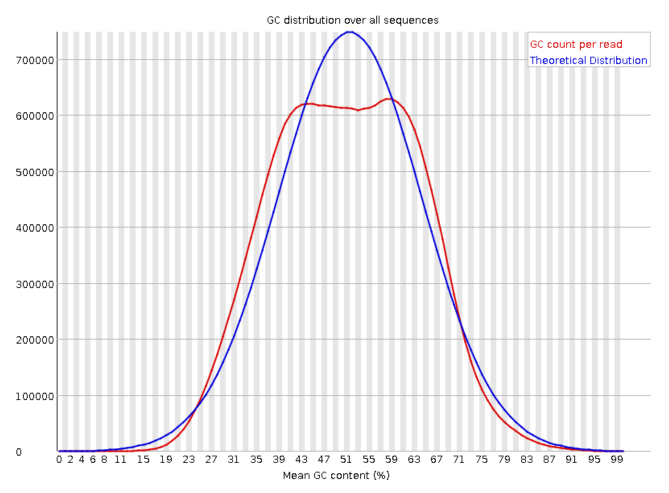
# **Per sequence quality scores**



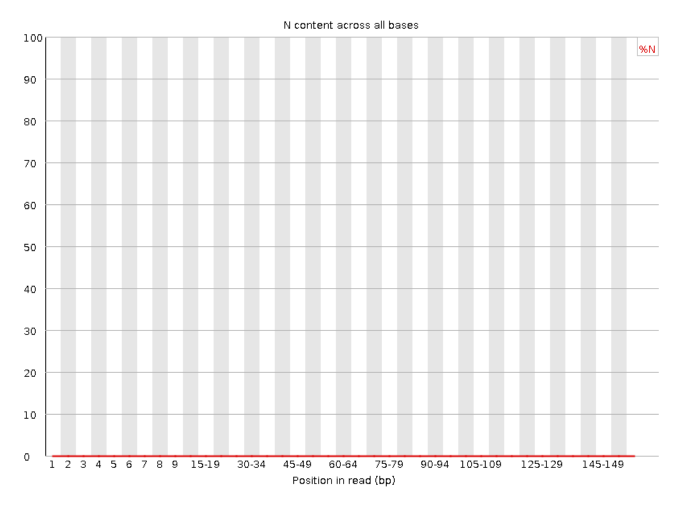
# Per base sequence content



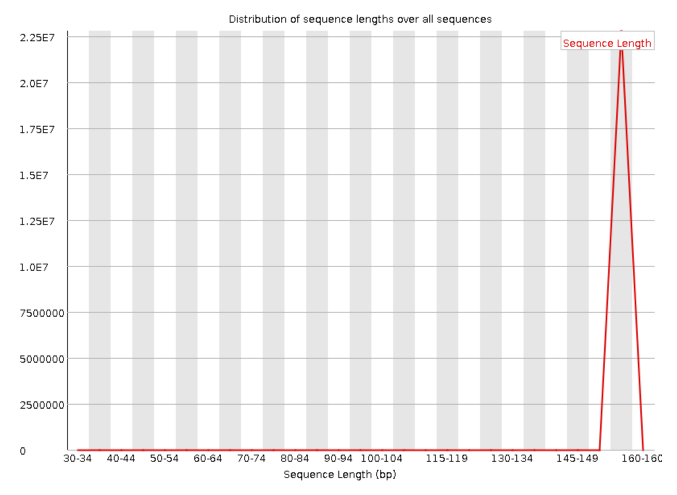
# Per sequence GC content



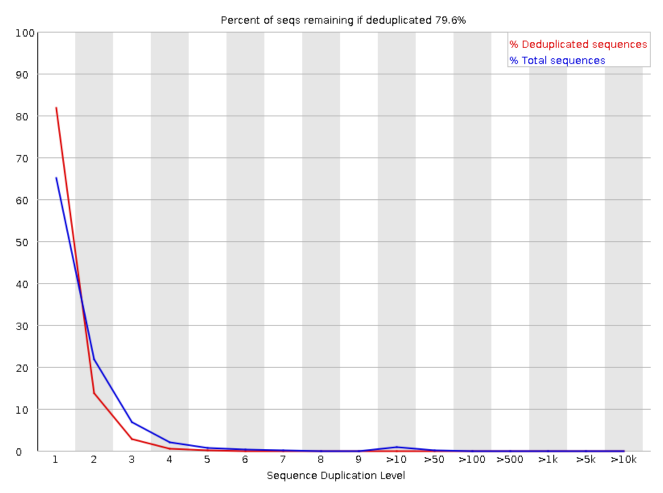
## Per base N content



### Sequence Length Distribution

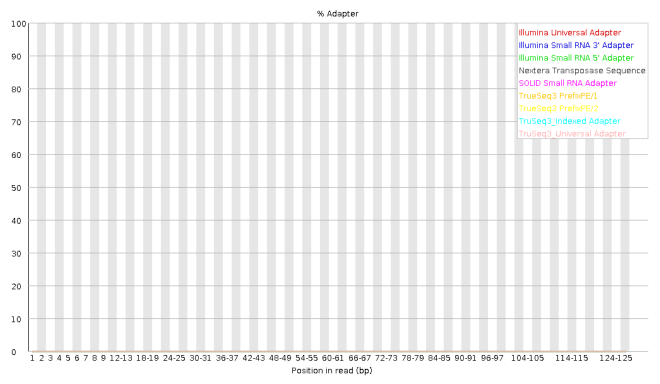


# Sequence Duplication Levels









Produced by FastQC (version 0.11.9)