

A modular tool to aggregate results from bioinformatics analyses across many samples into a single report.

Report generated on 2022-07-01, 14:04 based on data in: /data/jwd/main/047/988/47988471/working/multiqc\_WDir

Copy table

Configure Columns

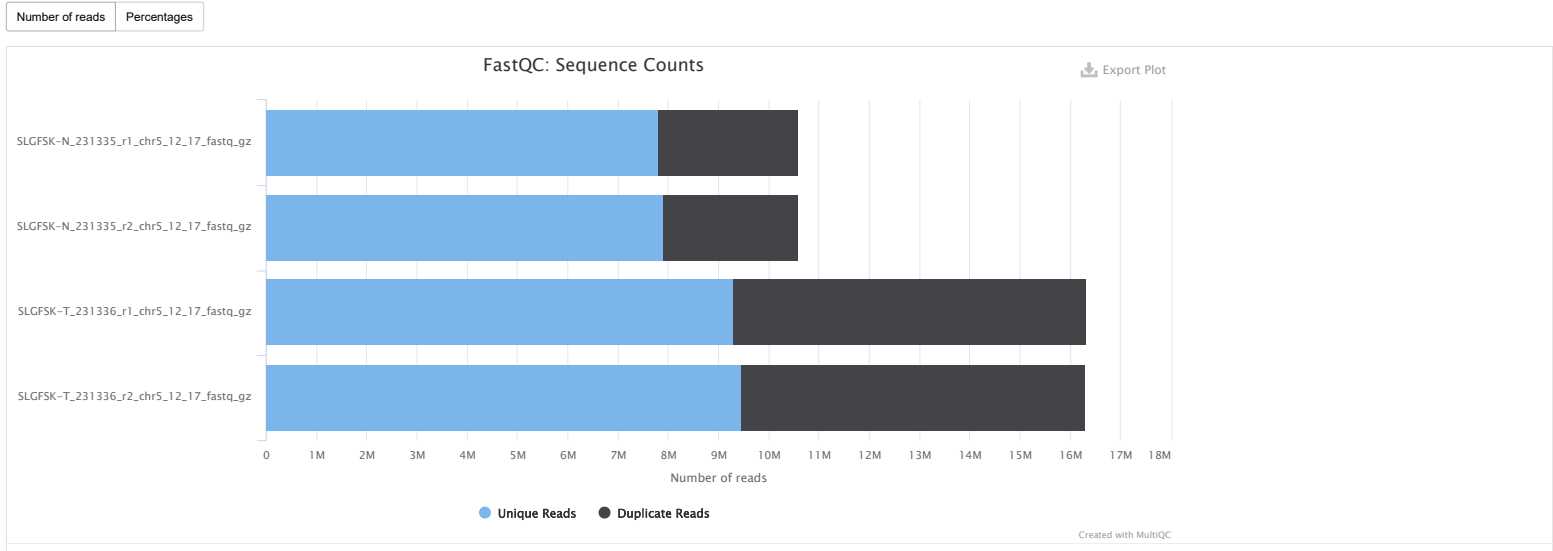
Plot

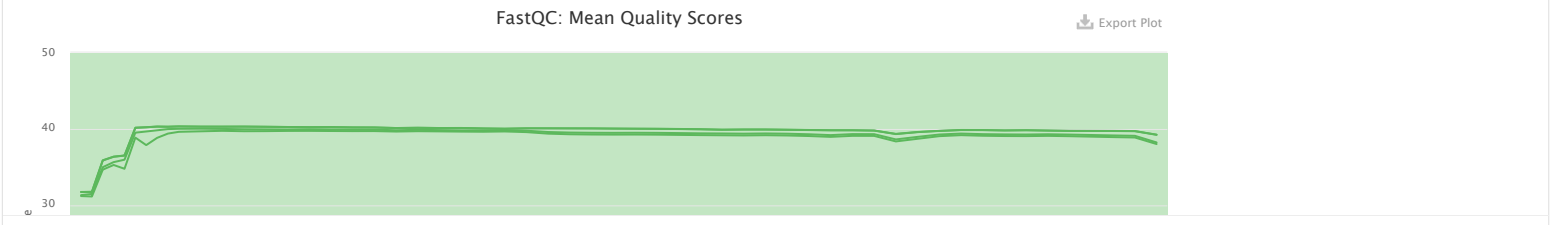
Showing 4/4 rows and 3/5 columns.

Sample Name	% Dups	% GC	M Seqs
SLGFSK-N_231335_r1_chr5_12_17_fastq_gz			
SLGFSK-N_231335_r2_chr5_12_17_fastq_gz			
SLGFSK-T_231336_r1_chr5_12_17_fastq_gz			
SLGFSK-T_231336_r2_chr5_12_17_fastq_gz			

## Sequence Counts Help

Sequence counts for each sample. Duplicate read counts are an estimate only.





Per Sequence Quality Scores

4

Help

The number of reads with average quality scores. Shows if a subset of reads has poor quality.

Y-Limits: on



Per Base Sequence Content

4

Help

The proportion of each base position for which each of the four normal DNA bases has been called.

Click a sample row to see a line plot for that dataset.

Info

Rollover for sample name

Position: -

%T: -

%C: -

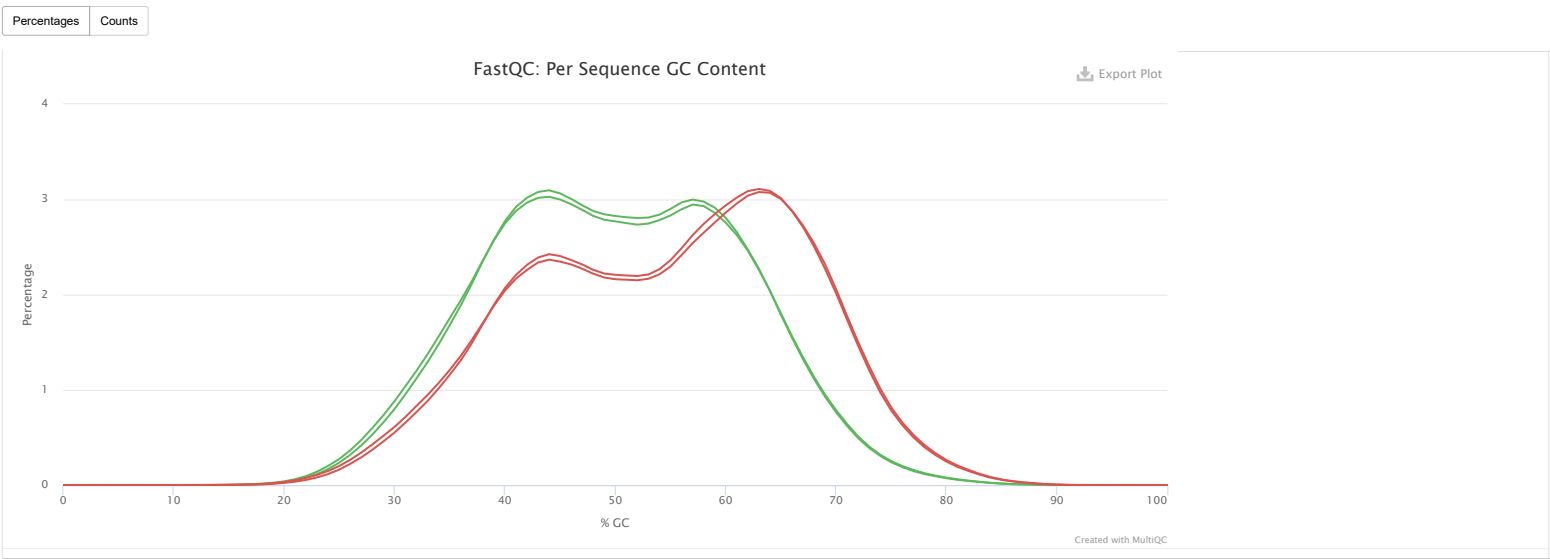
%A: -

%G: -

Export Plot

The average GC content of reads. Normal random library typically have a roughly normal distribution of GC content.

Y-Limits: on



Per Base N Content

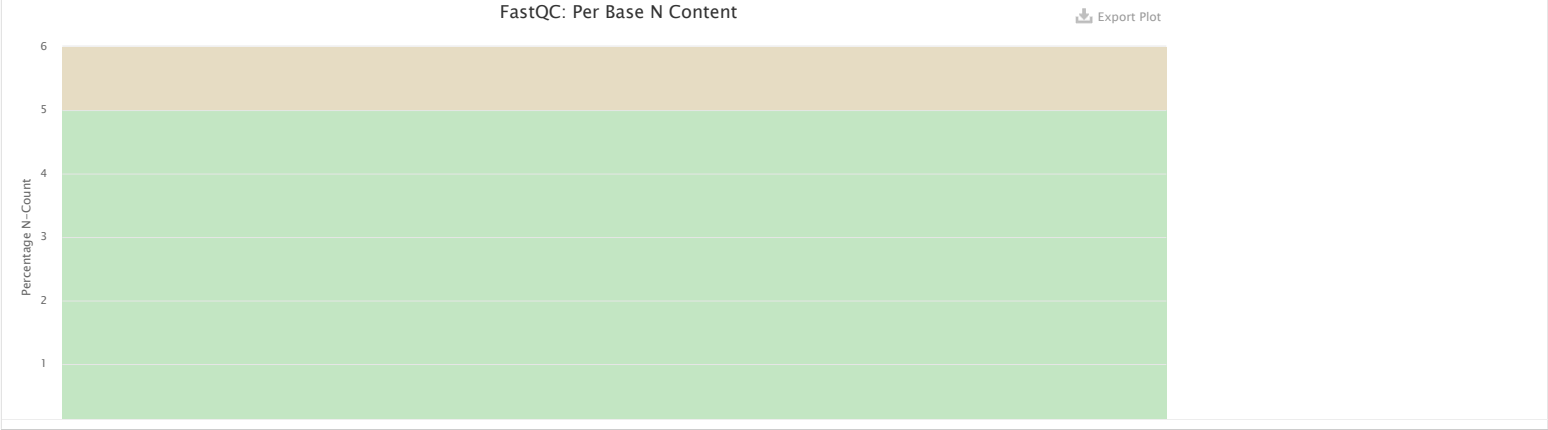
4

Help

The percentage of base calls at each position for which an N was called.

Y-Limits: on





## Sequence Length Distribution

4

All samples have sequences of a single length (101bp).

## Sequence Duplication Levels

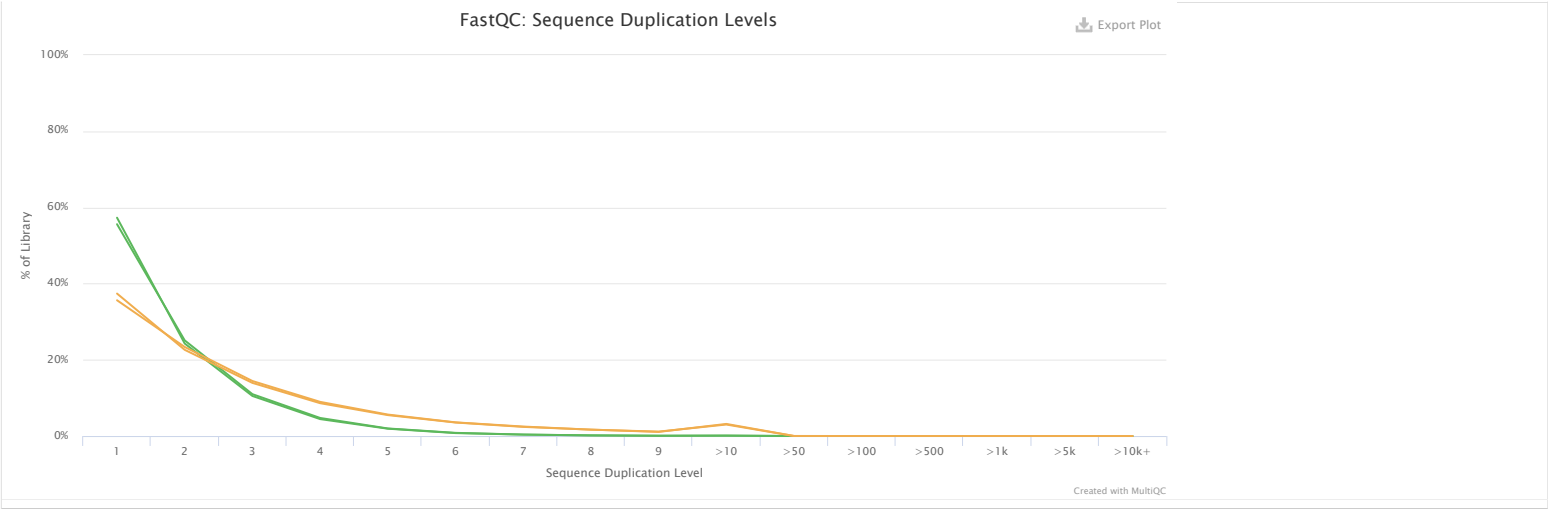
2

2

The relative level of duplication found for every sequence.

Help

Y-Limits: on



## Overrepresented sequences

4

The total amount of overrepresented sequences found in each library.

4 samples had less than 1% of reads made up of overrepresented sequences

Help

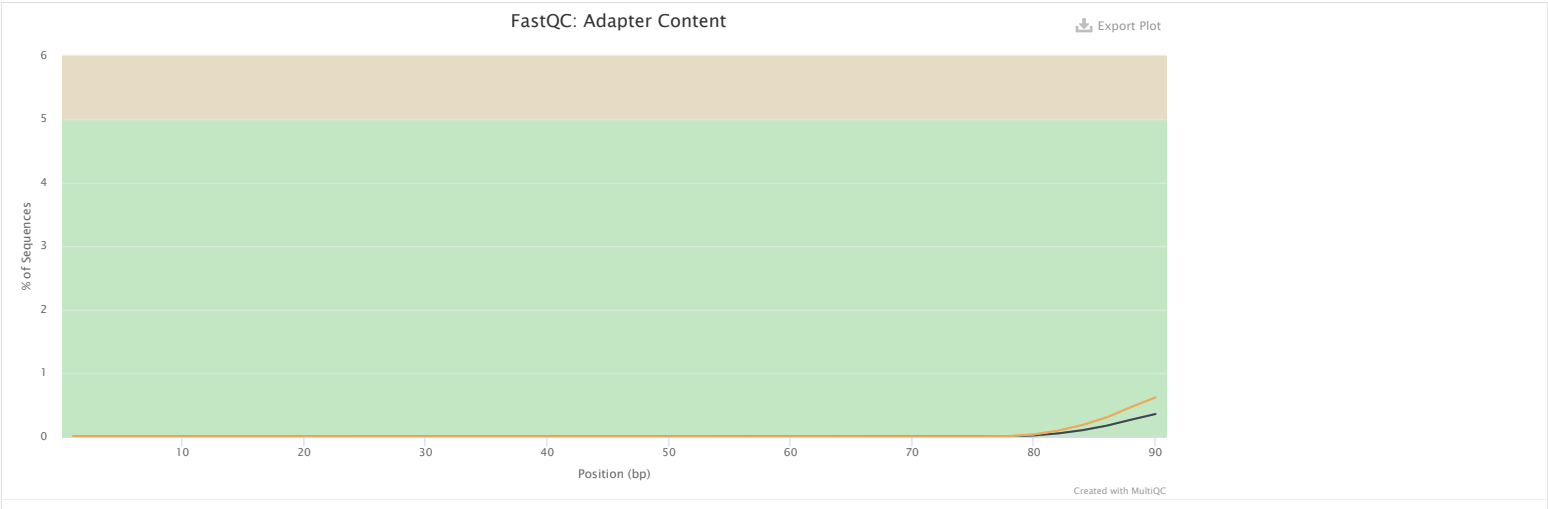
## Adapter Content

4

The cumulative percentage count of the proportion of your library which has seen each of the adapter sequences at each position.

Help

Y-Limits: on



## Status Checks

Status for each FastQC section showing whether results seem entirely normal (green), slightly abnormal (orange) or very unusual (red).

Help

Sort by highlight Min: 0 Max: 1

# FastQC: Status Checks

 Export Plot

