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Quality Histograms	
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## MultiQC

(http://multiqc.info)

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

## **General Statistics**

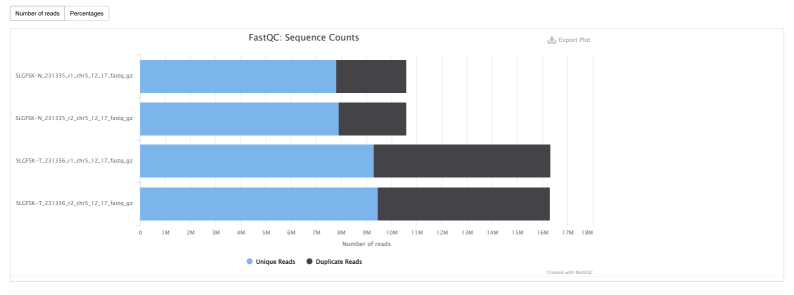
☐ Showing 4/4 rows and 3/5 columns. ☐ ♣ Plot Showing 4/4 rows and 3/5 columns.			
Sample Name	% Dups	% GC	M Seqs
SLGFSK-N_231335_r1_chr5_12_17_fastq_gz			
SLGFSK-N_231335_r2_chr5_12_17_fastq_gz			
SLGFSK-T_231336_r1_chr5_12_17_fastq_gz			
SLGFSK-T_231336_r2_chr5_12_17_fastq_gz			

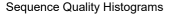
## FastQC

FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) is a quality control tool for high throughput sequence data, written by Simon Andrews at the Babraham Institute in Cambridge.

## Sequence Counts

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



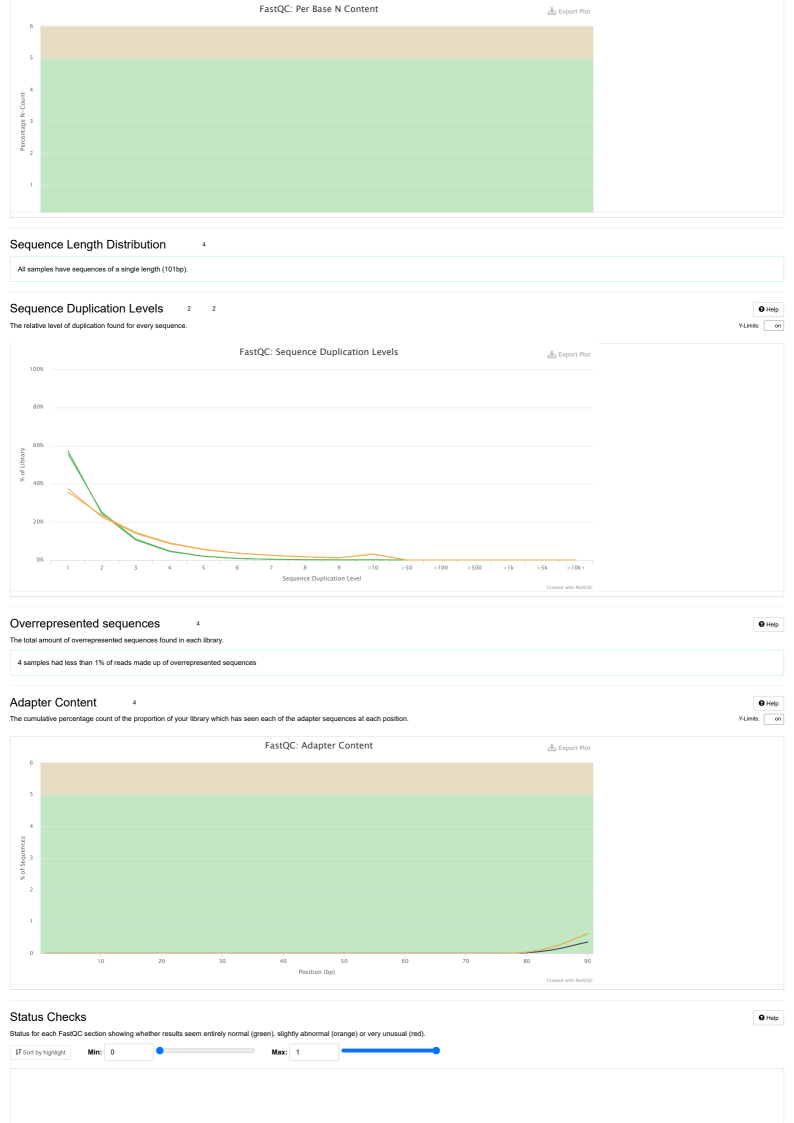


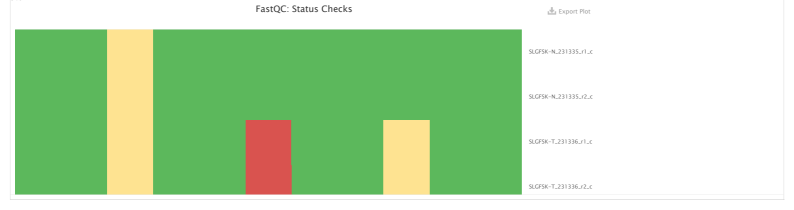
The mean quality value across each base position in the read.



**9** Help







MultiQC v1.11 (http://multiqc.info) - Written by Phil Ewels (http://phil.ewels.co.uk), available on GitHub (https://github.com/ewels/MultiQC).

This report uses HighCharts (http://www.highcharts.com/), |Query UI (https://giuery.com/), |Query UI (https://github.com/ewels/MultiQC).

Clipboard is (https://clipboardjs.com/).

FileSaver is (https://github.com/eligrey/FileSaver.js) and clipboard is (https://clipboardjs.com/).

