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v1.14
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MultiQC

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

Report generated on 2024-04-29, 15:08 CEST based on data in: `/home/BCG_2024_vfassi/project/676`

📢 Welcome! Not sure where to start?

[Watch a tutorial video](#)

(6:06)

don't show again ✕

General Statistics

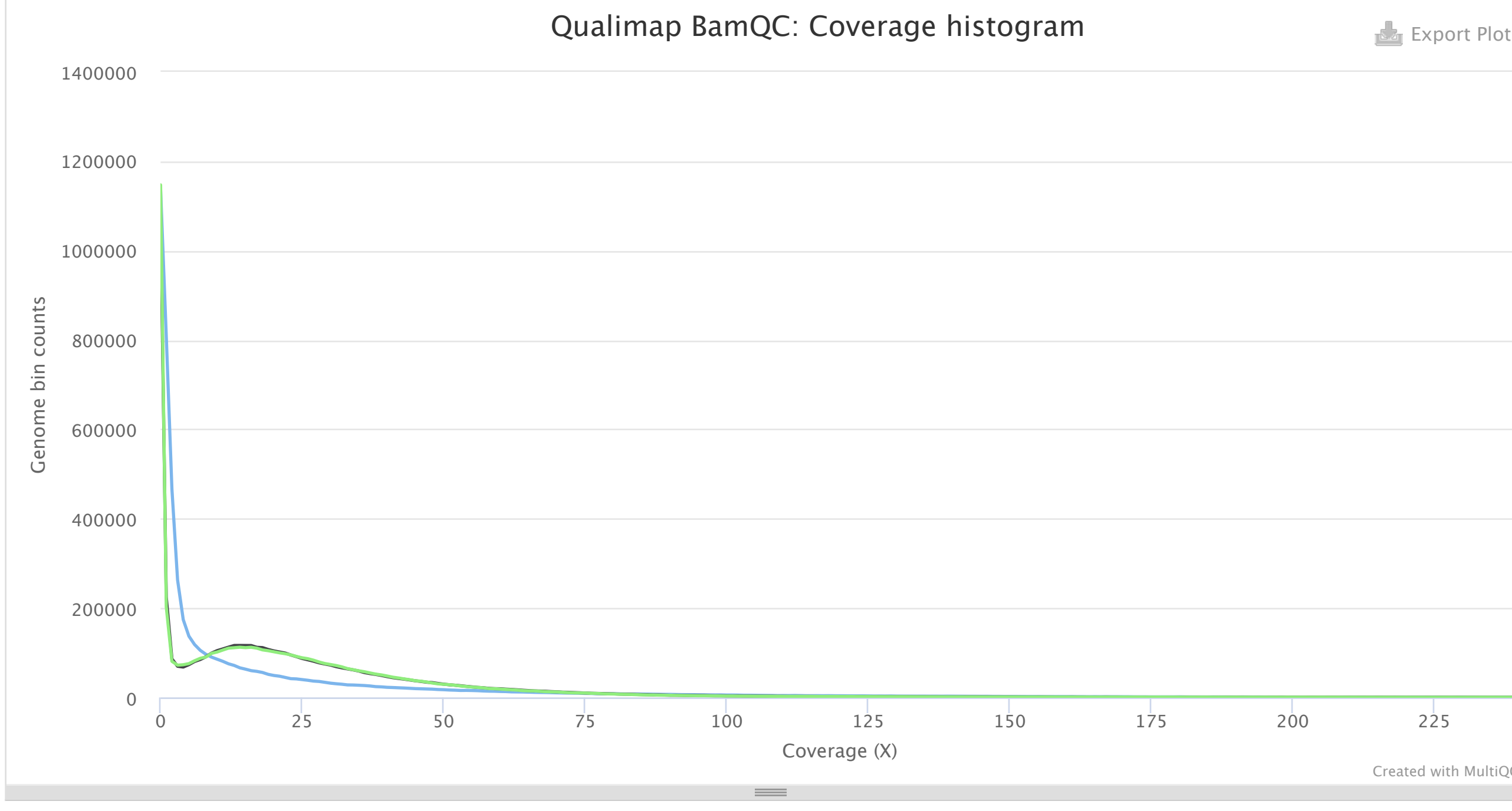
Sample Name	% GC	≥ 30X	Median cov	Mean cov	% Aligned	% Dups	% GC	M Seqs
case676_child	46%	22.8%	5.0X	24.1X	99.8%	5.4%	43%	3.0
case676_father	52%	31.2%	18.0X	27.3X	99.9%	6.1%	50%	2.2
case676_mother	52%	31.0%	18.0X	26.3X	99.8%	8.5%	50%	2.1

QualiMap

QualiMap is a platform-independent application to facilitate the quality control of alignment sequencing data and its derivatives like feature counts. DOI: [10.1093/bioinformatics/btv566](https://doi.org/10.1093/bioinformatics/btv566); [10.1093/bioinformatics/bts503](https://doi.org/10.1093/bioinformatics/bts503).

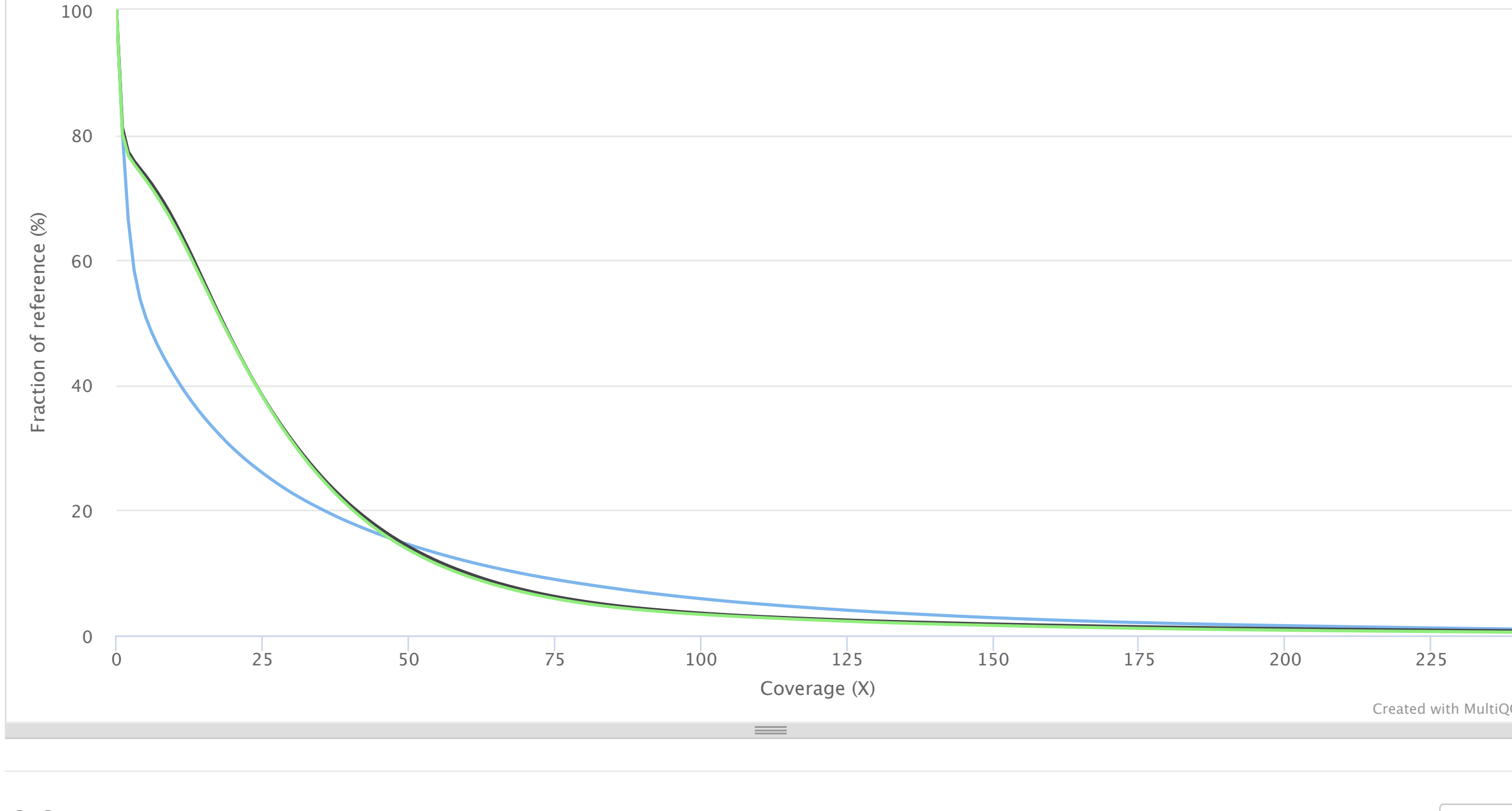
Coverage histogram

Distribution of the number of locations in the reference genome with a given depth of coverage.



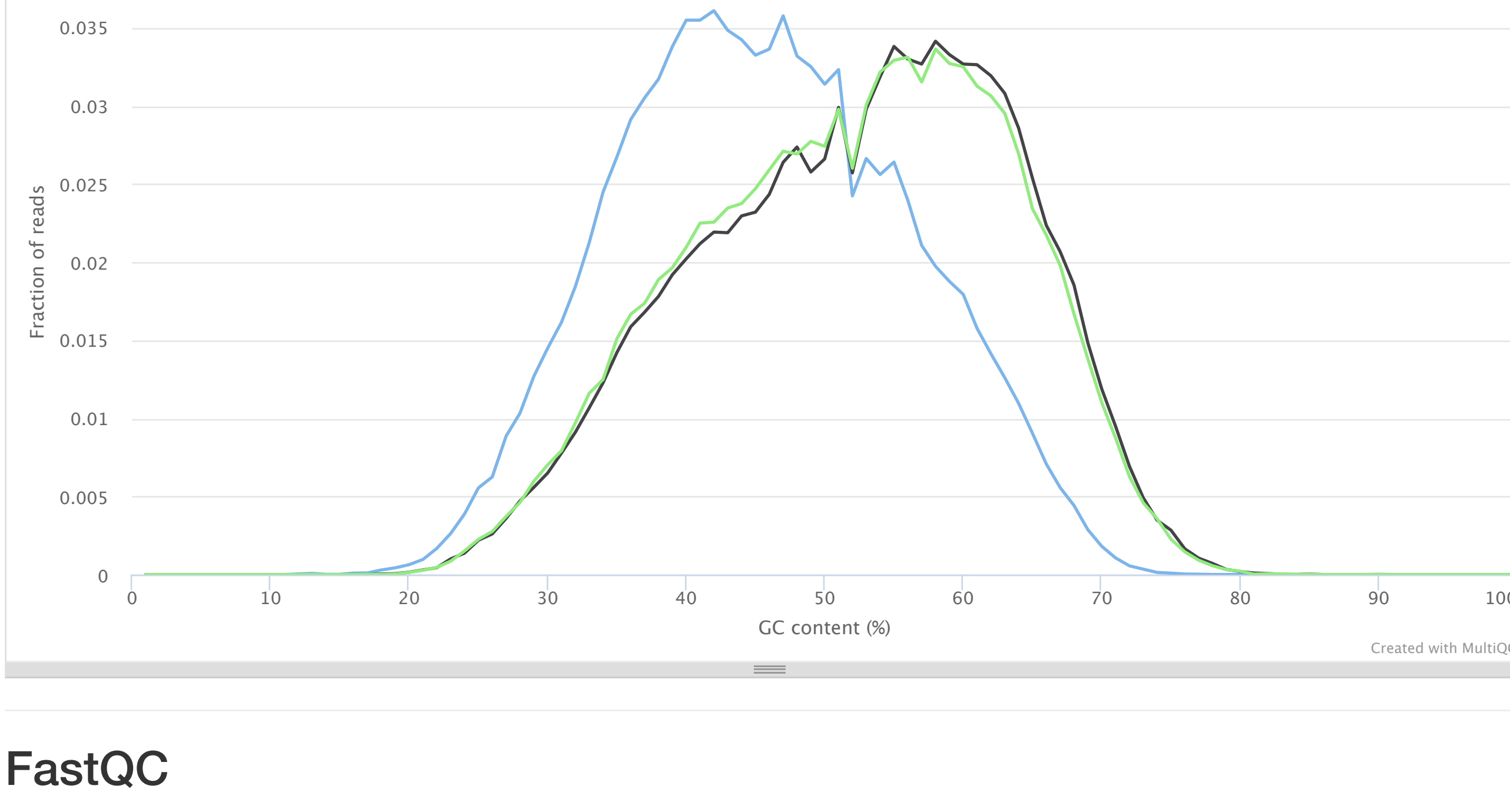
Cumulative genome coverage

Percentage of the reference genome with at least the given depth of coverage.



GC content distribution

Each solid line represents the distribution of GC content of mapped reads for a given sample.

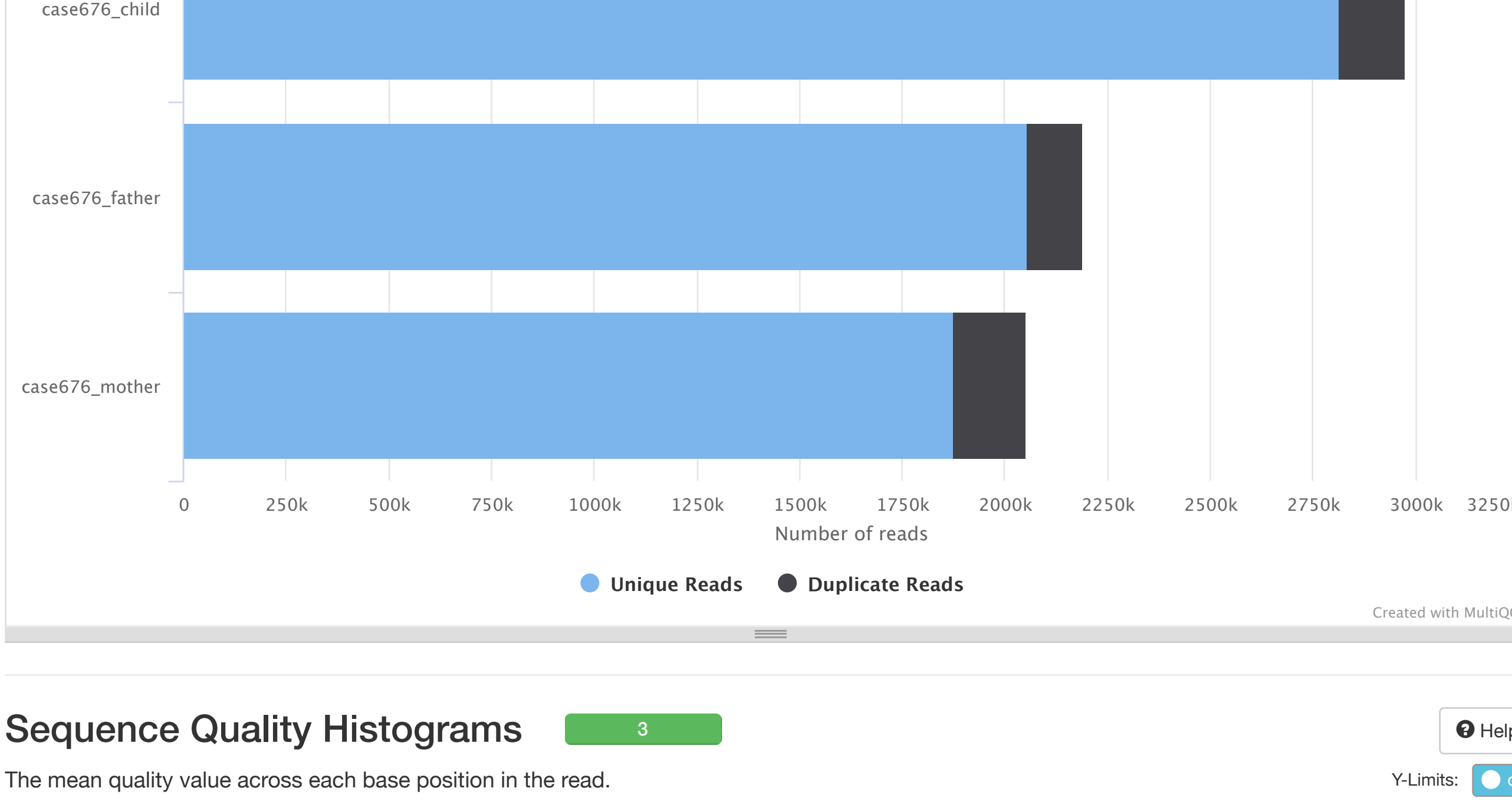


FastQC

FastQC is a quality control tool for high throughput sequence data, written by Simon Andrews at the Babraham Institute in Cambridge.

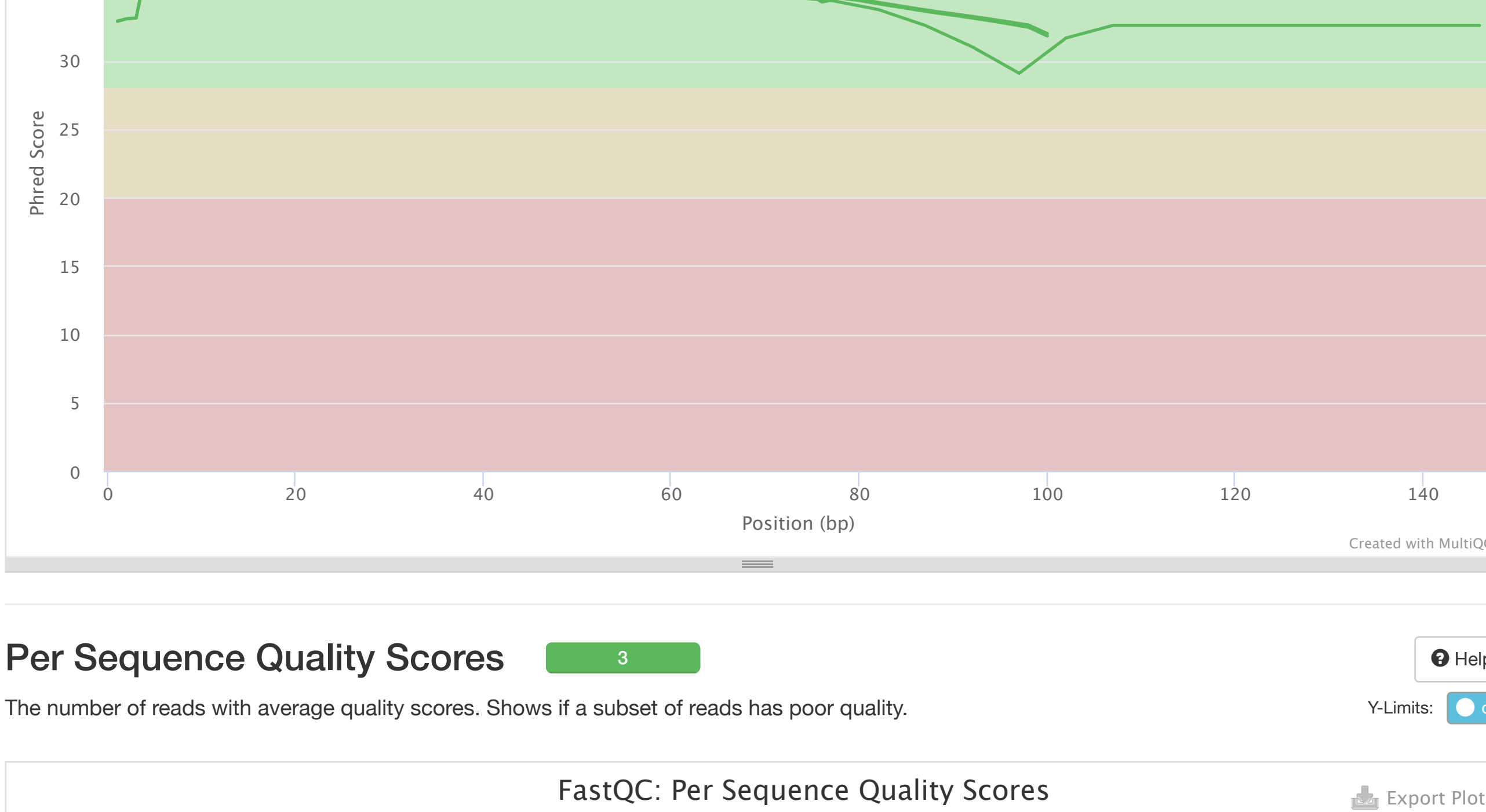
Sequence Counts

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



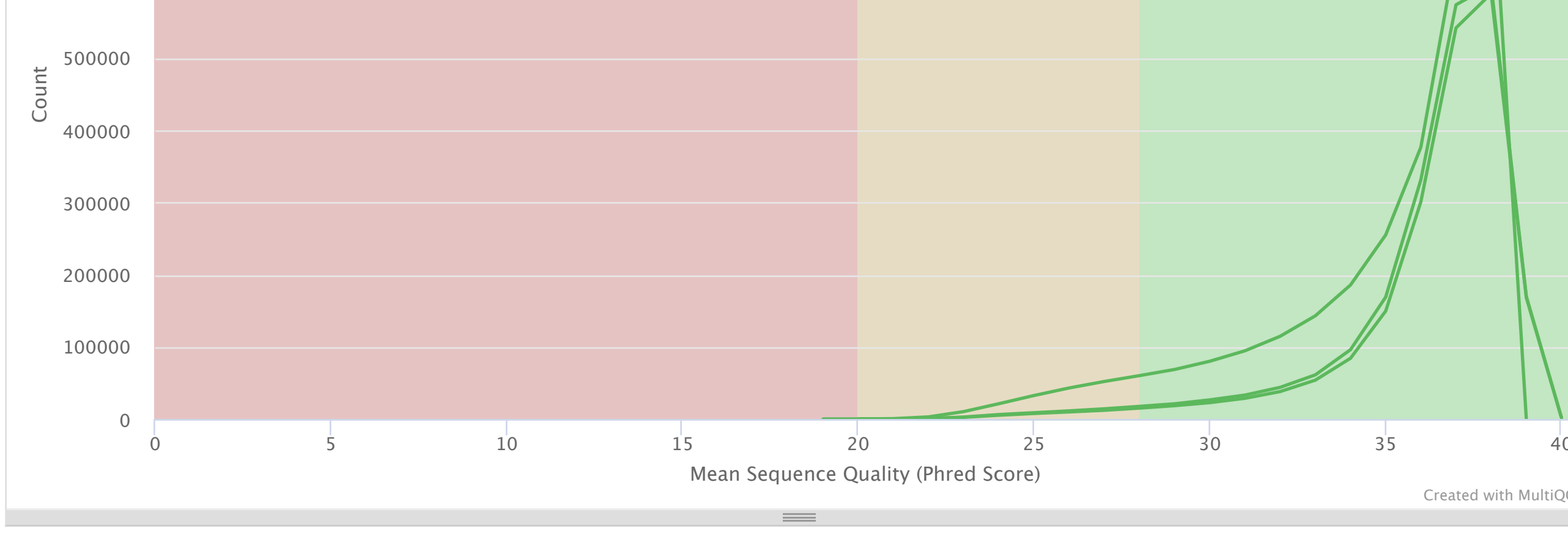
Sequence Quality Histograms

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



Per Sequence Quality Scores

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



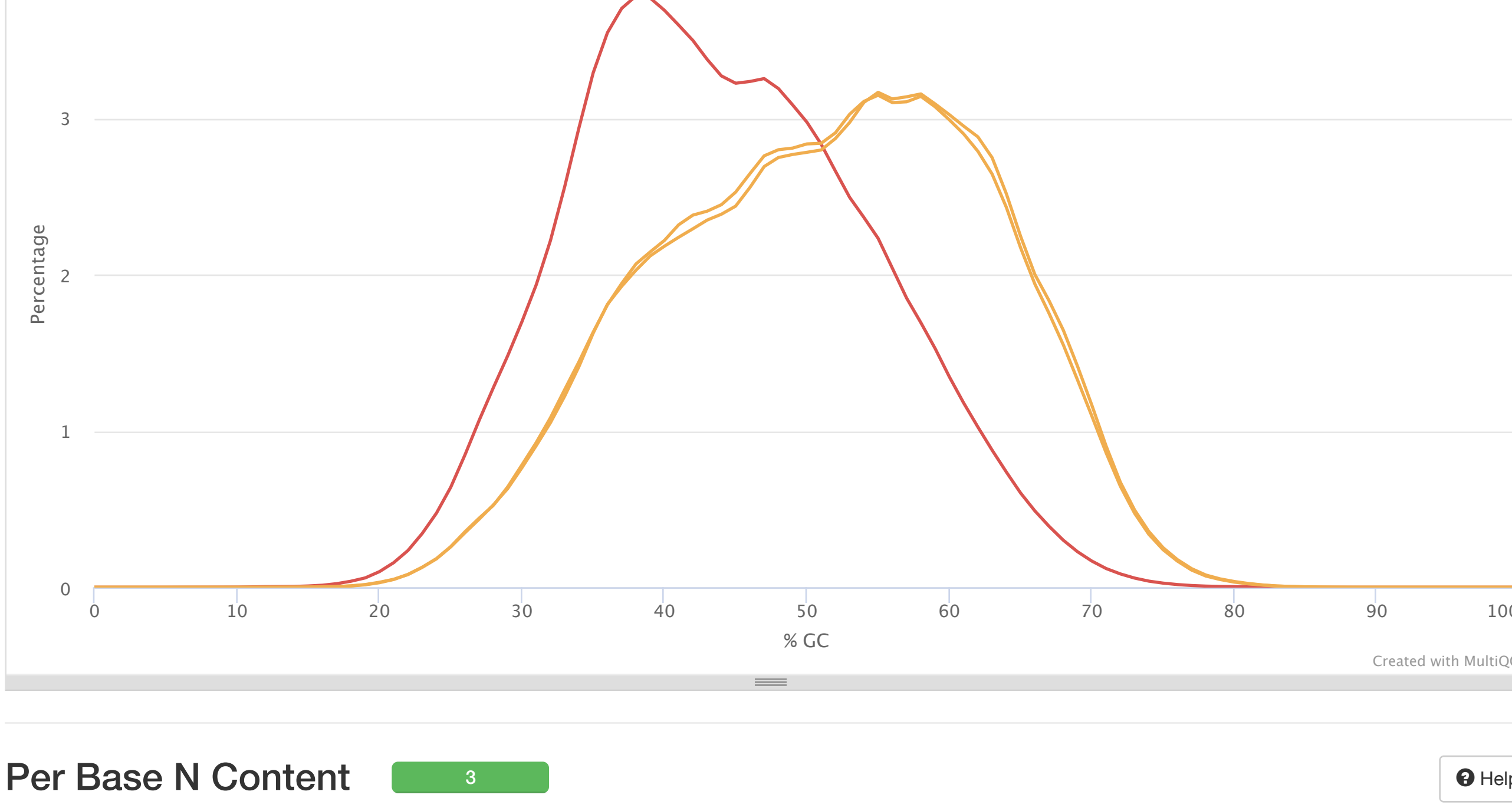
Per Base Sequence Content

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



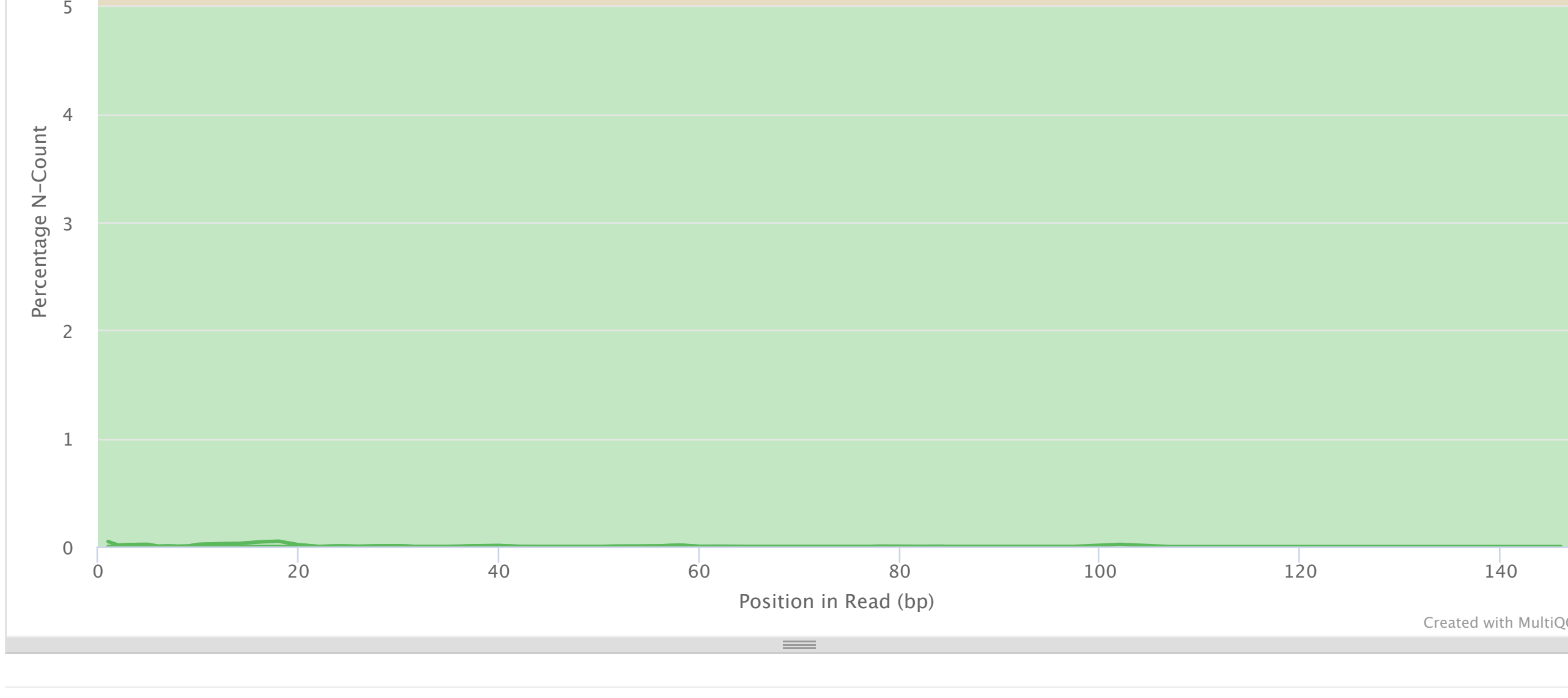
Per Sequence GC Content

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



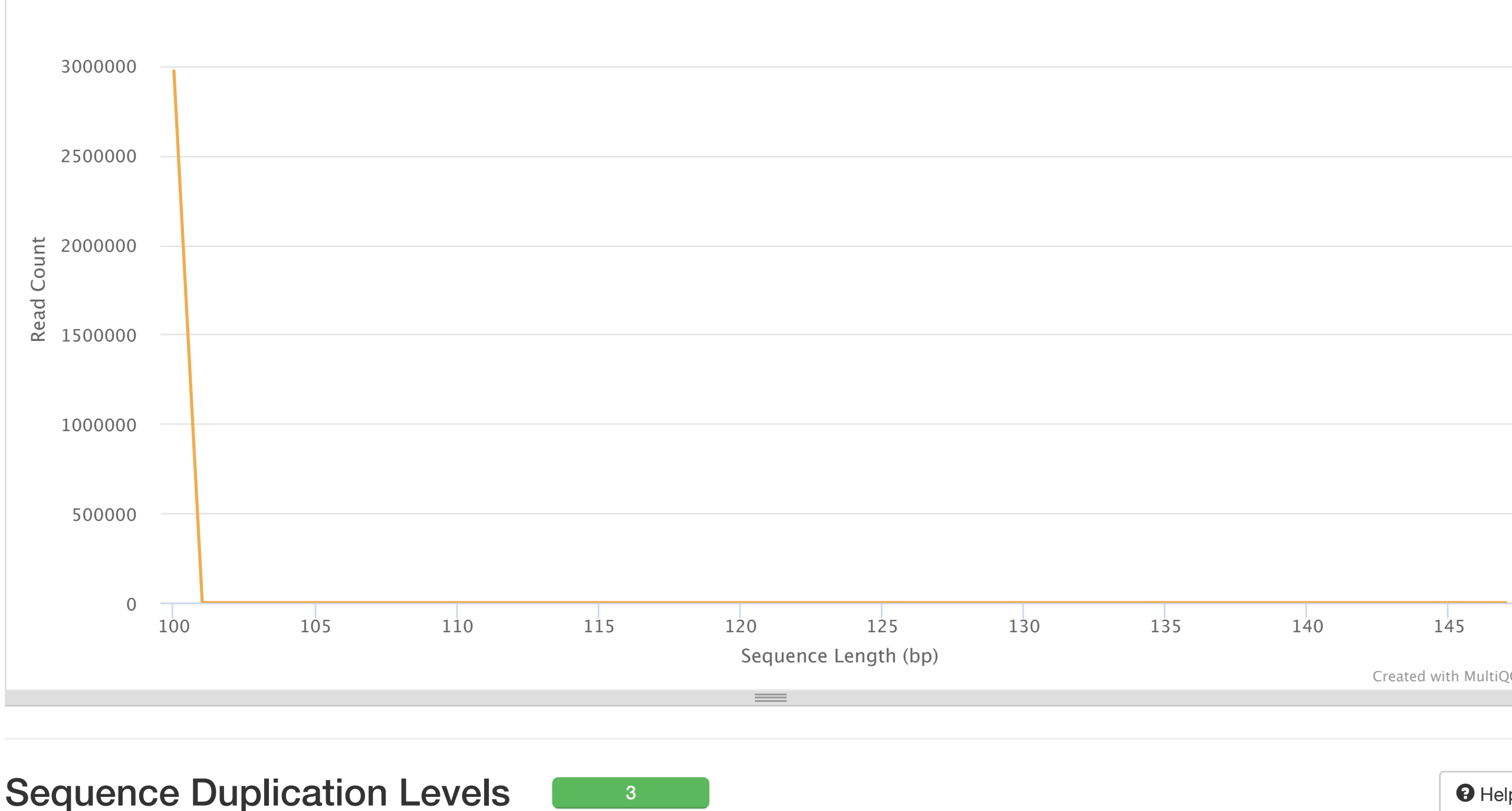
Per Base N Content

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



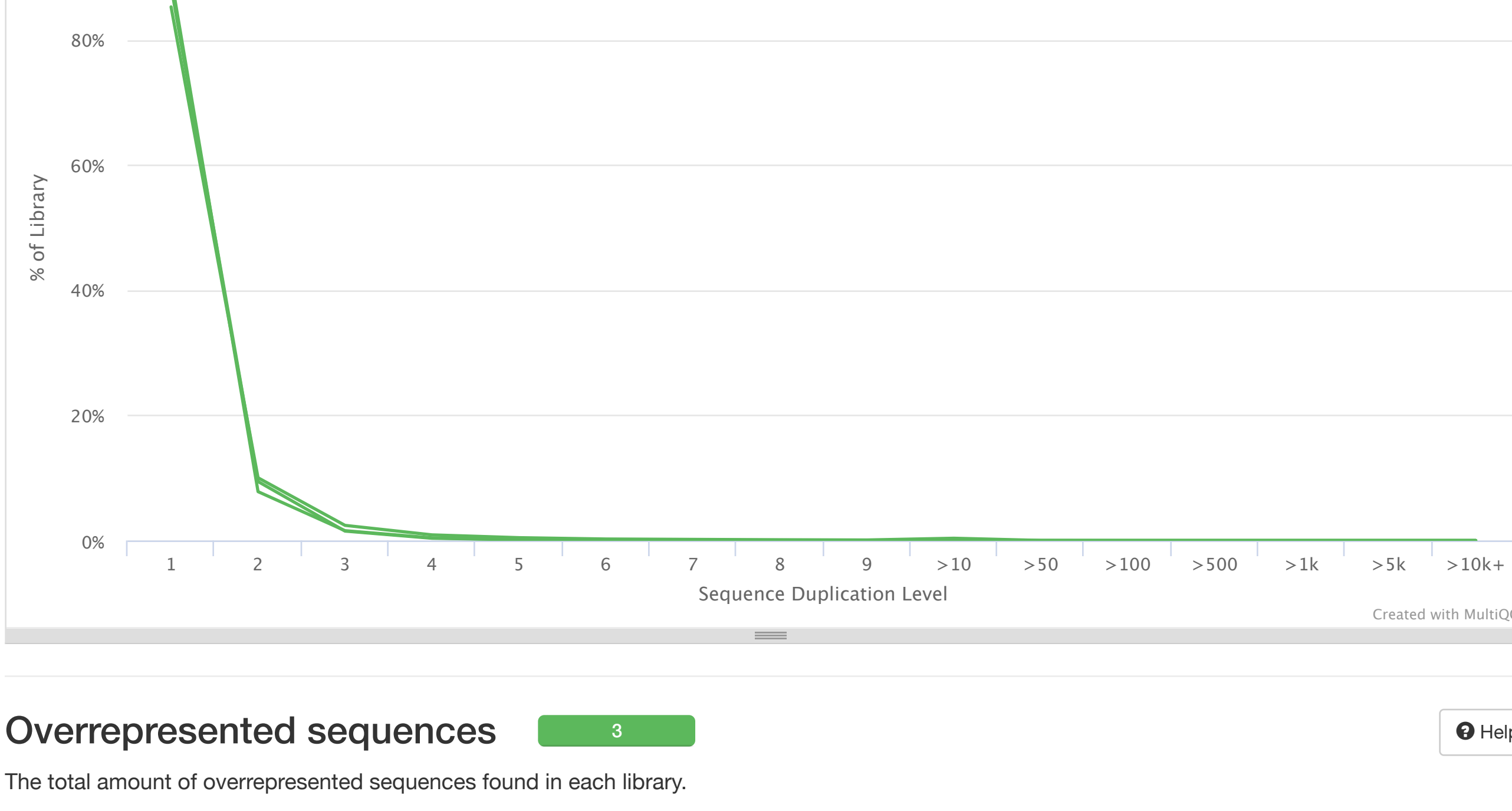
Sequence Length Distribution

The distribution of fragment sizes (read lengths) found. See the [FastQC help](#).



Sequence Duplication Levels

The relative level of duplication found for every sequence.



Overrepresented sequences

The total amount of overrepresented sequences found in each library.

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

Adapter Content

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

No samples found with any adapter contamination > 0.1%

Status Checks

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

