



Sequencing QC Report  
Based upon: 20.088.878 sequences in 8 data sets  
Generated by: IMG  
Creation date: Thu Feb 18 11:14:18 CET 2016  
Software: CLC Genomics Server 7.5.1

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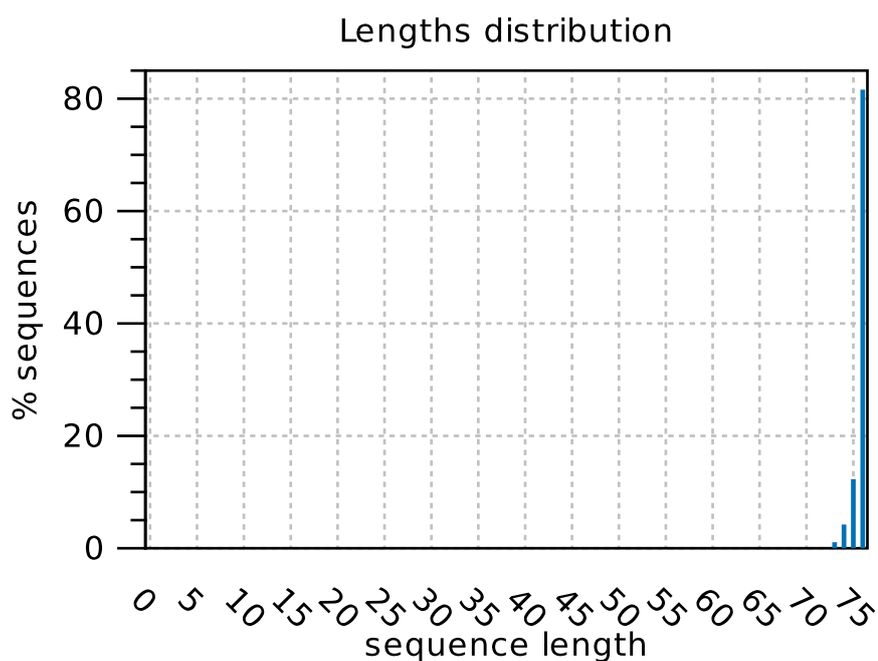
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# 1. Summary

Creation date:	Thu Feb 18 11:14:18 CET 2016
Generated by:	IMGM
Software:	CLC Genomics Server 7.5.1
Based upon:	8 data sets
NG-125-0012_S3_L001_P_run1:	1.533.830 sequences in pairs
NG-125-0012_S3_L002_P_run1:	1.493.586 sequences in pairs
NG-125-0012_S3_L003_P_run1:	1.451.974 sequences in pairs
NG-125-0012_S3_L004_P_run1:	1.531.508 sequences in pairs
NG-125-0012_S1_L001_P_run2:	3.611.860 sequences in pairs
NG-125-0012_S1_L002_P_run2:	3.532.720 sequences in pairs
NG-125-0012_S1_L003_P_run2:	3.490.402 sequences in pairs
NG-125-0012_S1_L004_P_run2:	3.442.998 sequences in pairs
Total sequences in data sets	20.088.878 sequences
Total nucleotides in data sets	1.520.260.856 nucleotides

## 2. Per-sequence analysis

### 2.1 Lengths distribution

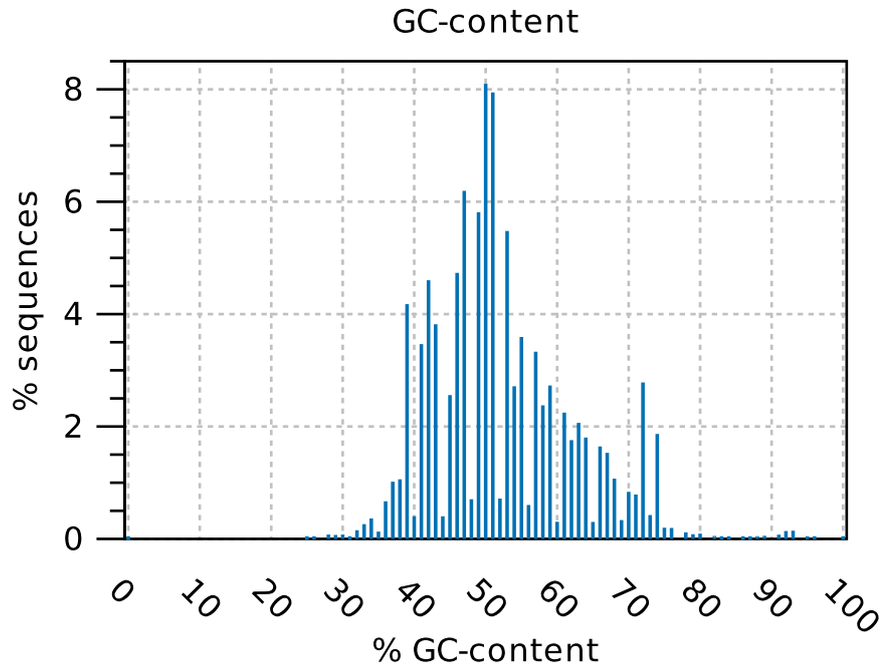


Distribution of sequence lengths. In cases of untrimmed Illumina or SOLiD reads it will just contain a single peak.

x: sequence length in base-pairs

y: number of sequences featuring a particular length normalized to the total number of sequences

## 2.2 GC-content

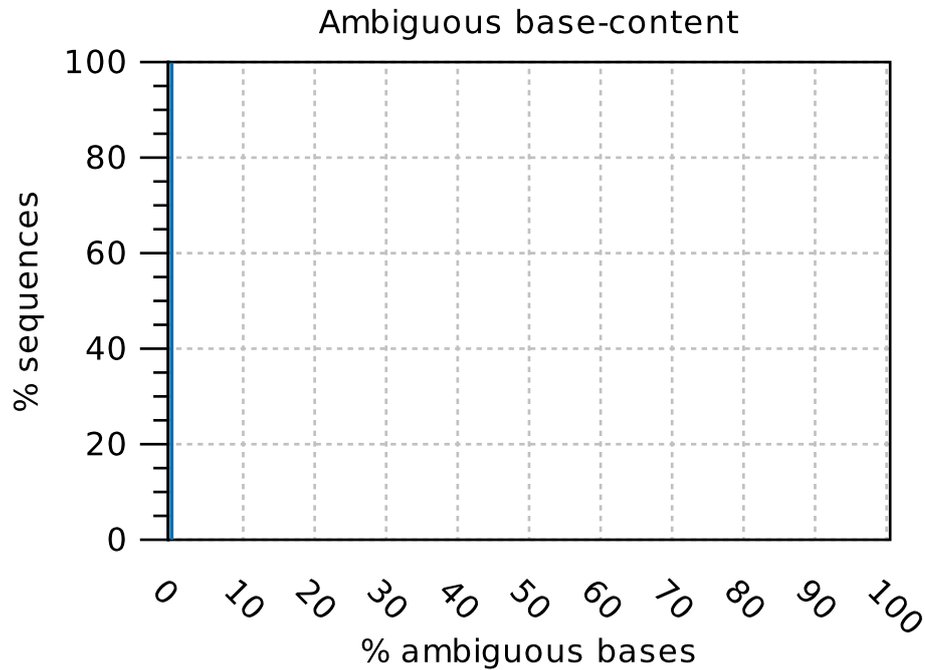


Distribution of GC-contents. The GC-content of a sequence is calculated as the number of G C-bases compared to all bases (including ambiguous bases).

x: relative GC-content of a sequence in percent

y: number of sequences featuring particular GC-percentages normalized to the total number of sequences

## 2.3 Ambiguous base-content

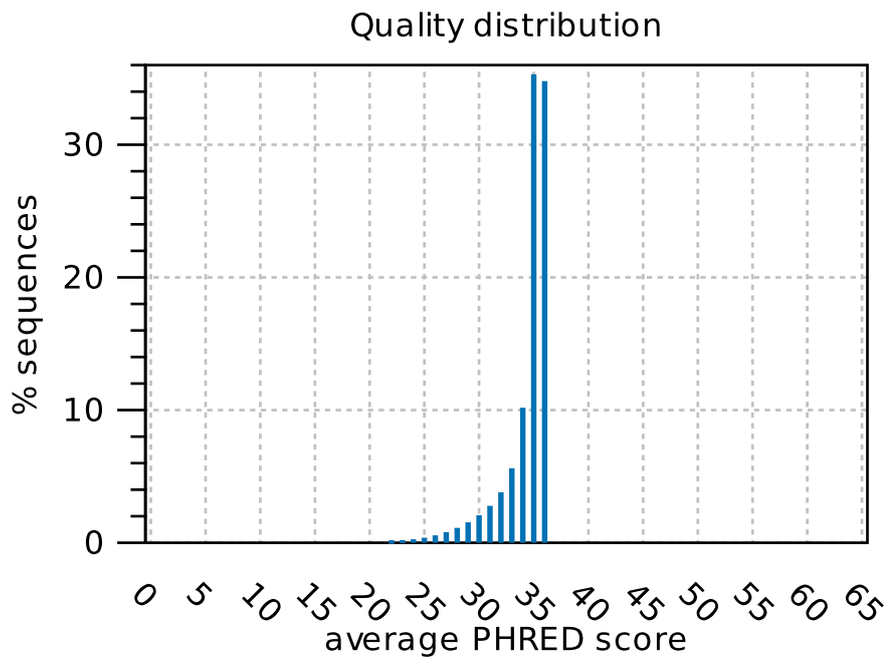


Distribution of N-contents. The N-content of a sequence is calculated as the number of ambiguous bases compared to all bases.

x: relative N-content of a sequence in percent

y: number of sequences featuring particular N-percentages normalized to the total number of sequences

## 2.4 Quality distribution



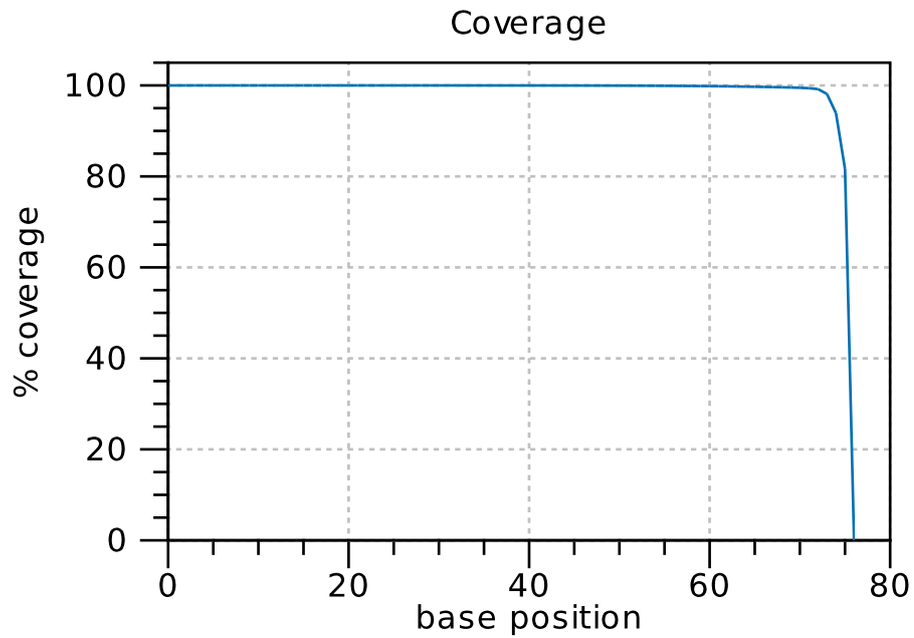
Distribution of average sequence quality scores. The quality of a sequence is calculated as the arithmetic mean of its base qualities.

x: PHRED-score

y: number of sequences observed at that qual. score normalized to the total number of sequences

### 3. Per-base analysis

#### 3.1 Coverage

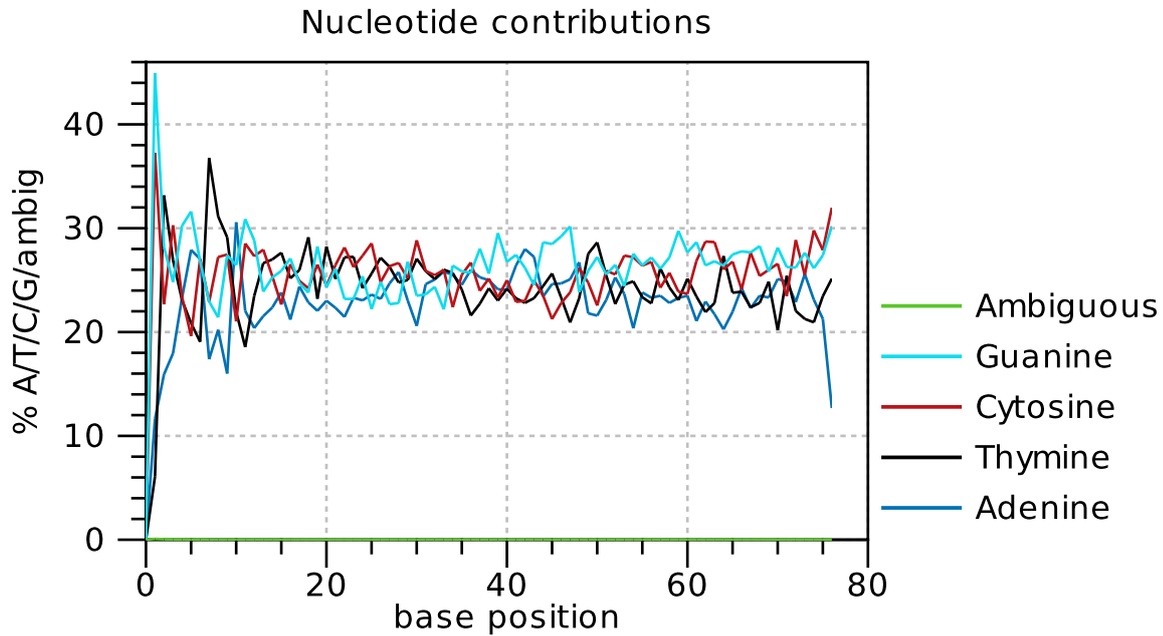


The number of sequences that support (cover) the individual base positions. In cases of un trimmed Illumina or SOLiD reads it will just contain a rectangle.

x: base position

y: number of sequences covering individual base positions normalized to the total number of sequences

### 3.2 Nucleotide contributions

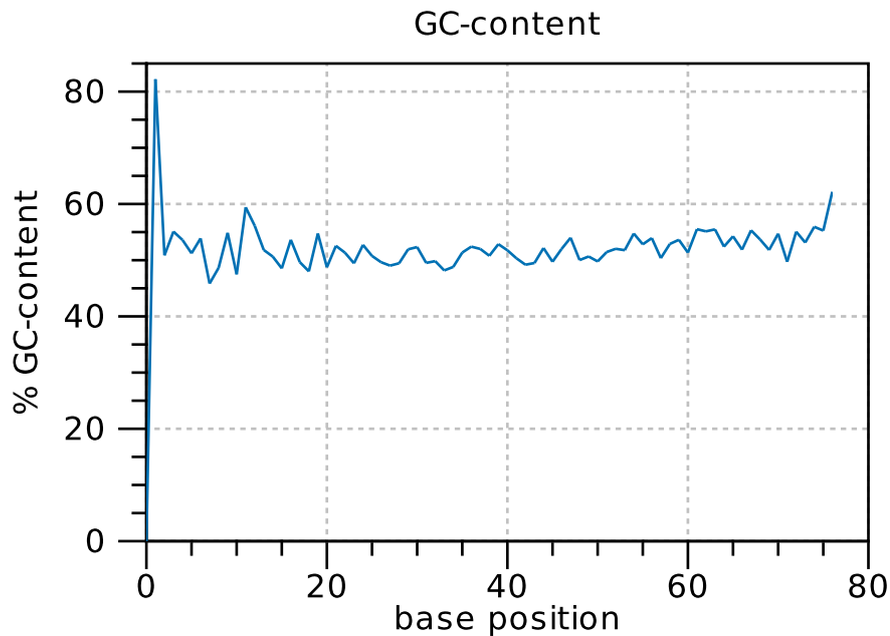


Coverages for the four DNA nucleotides and ambiguous bases.

x: base position

y: number of nucleotides observed per type normalized to the total number of nucleotides observed at that position

### 3.3 GC-content

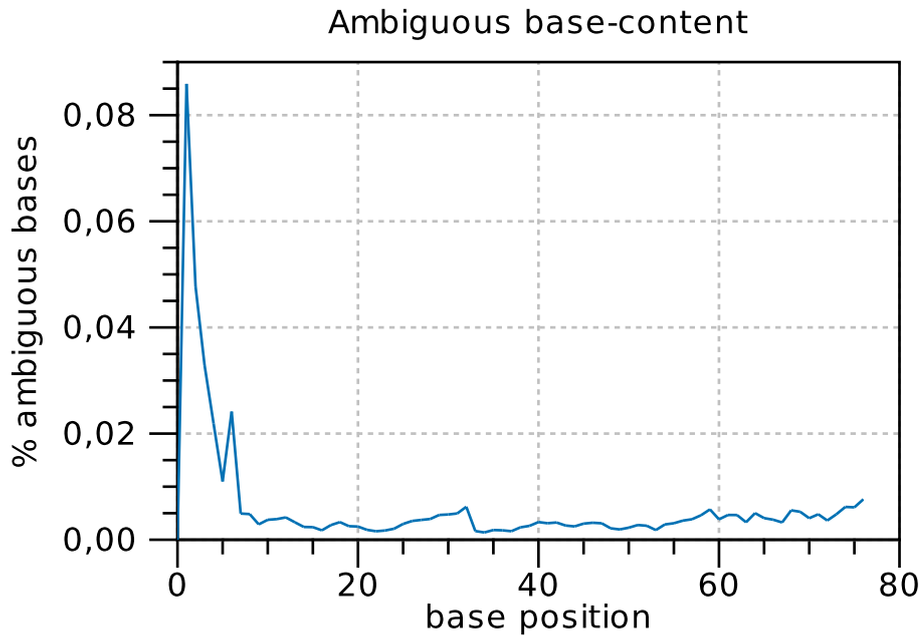


Combined coverage of G- and C-bases.

x: base position

y: number of G- and C-bases observed at current position normalized to the total number of bases observed at that position

### 3.4 Ambiguous base-content

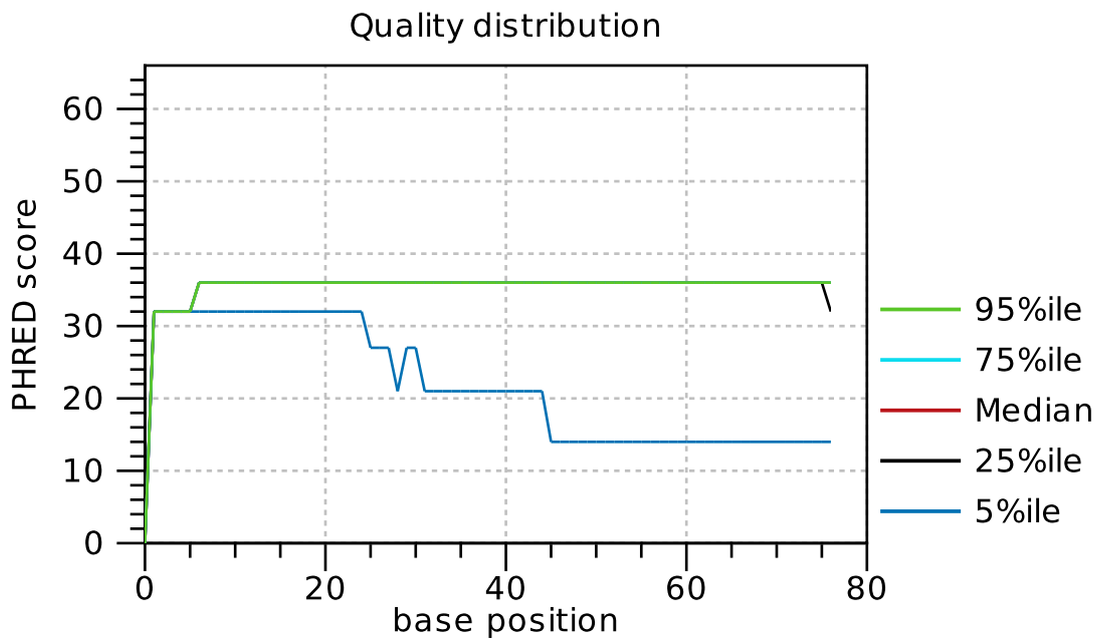


Combined coverage of ambiguous bases.

x: base position

y: number of ambiguous bases observed at current position normalized to the total number of bases observed at that position

### 3.5 Quality distribution



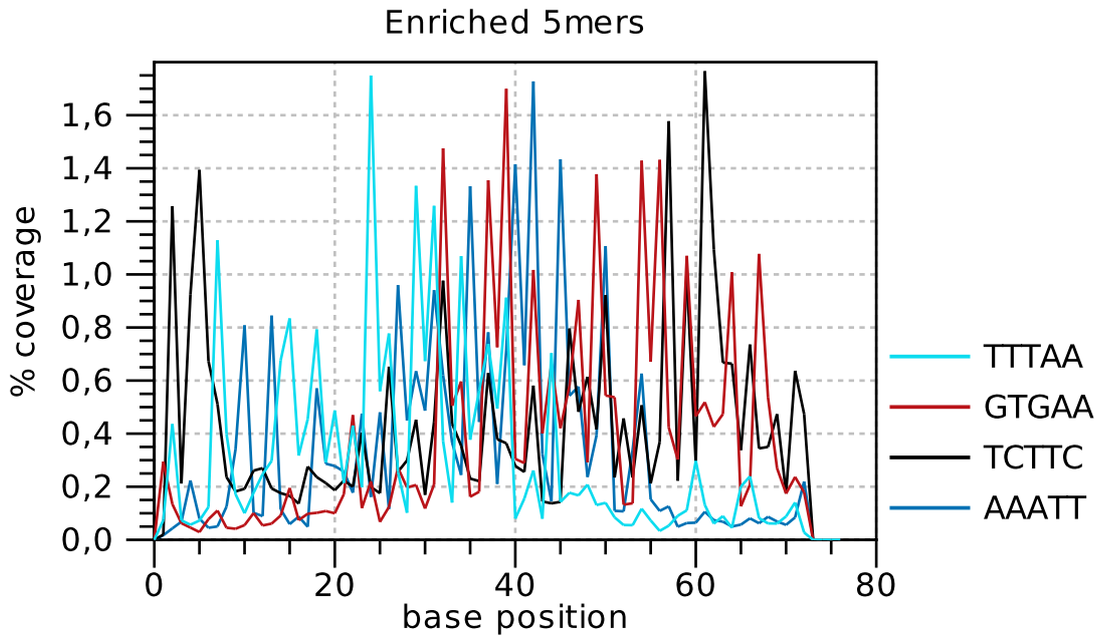
Base-quality distribution along the base positions.

x: base position

y: median & percentiles of quality scores observed at that base position

## 4. Over-representation analyses

### 4.1 Enriched 5mers

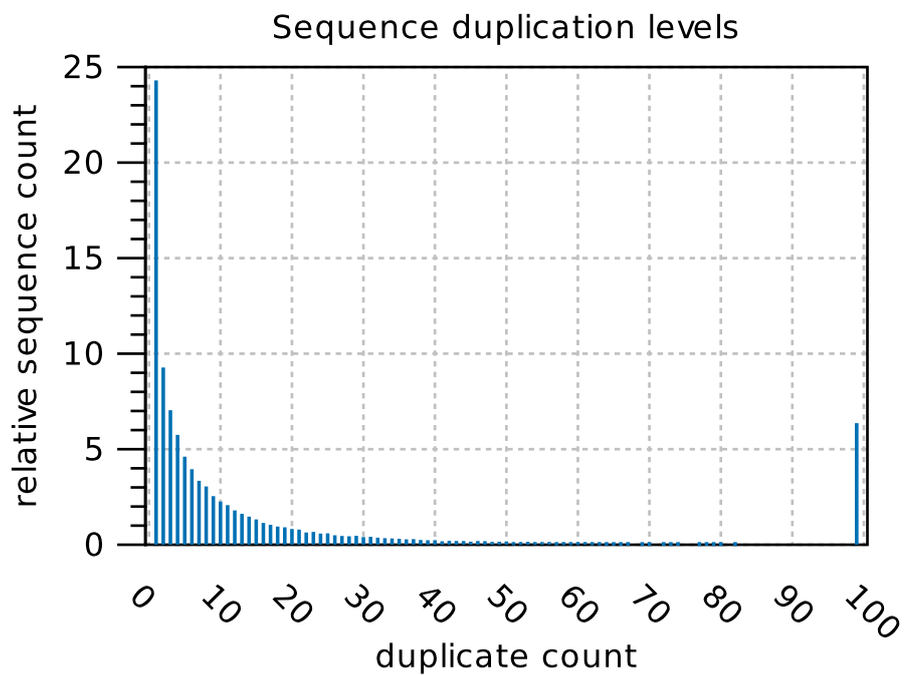


The five most-overrepresented 5mers. The over-representation of a 5mer is calculated as the ratio of the observed and expected 5mer frequency. The expected frequency is calculated as product of the empirical nucleotide probabilities that make up the 5mer. (5mers that contain ambiguous bases are ignored)

x: base position

y: number of times a 5mer has been observed normalized to all 5mers observed at that position

## 4.2 Sequence duplication levels



Duplication level distribution. Duplication levels are simply the count of how often a particular sequence has been found.

x: duplicate count

y: number of sequences that have been found that many times normalized to the number of unique sequences

## 4.3 Duplicated sequences

A table of over-represented sequences is given in the supplementary report