












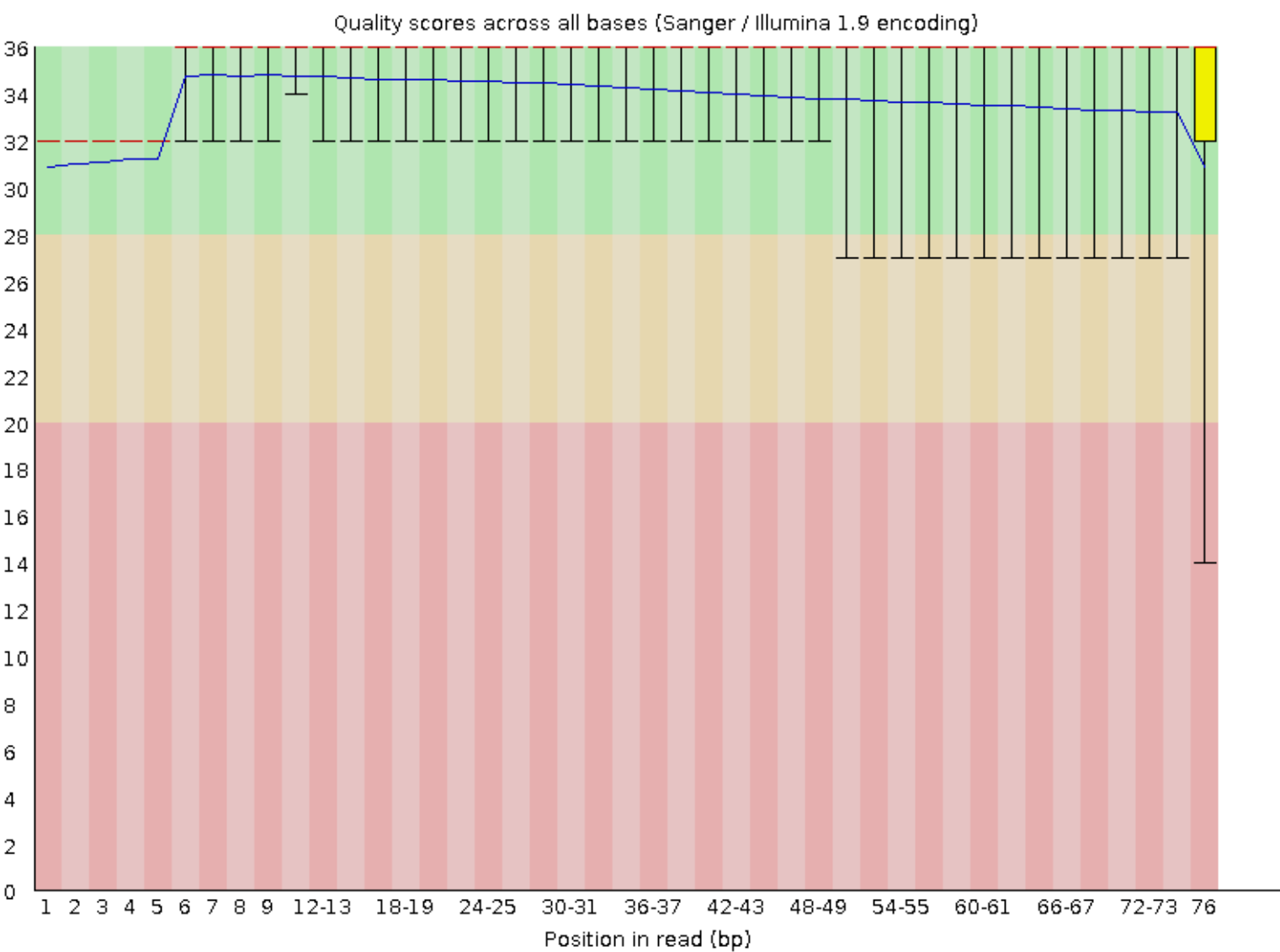
## Summary

-  [Basic Statistics](#)
-  [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](#)

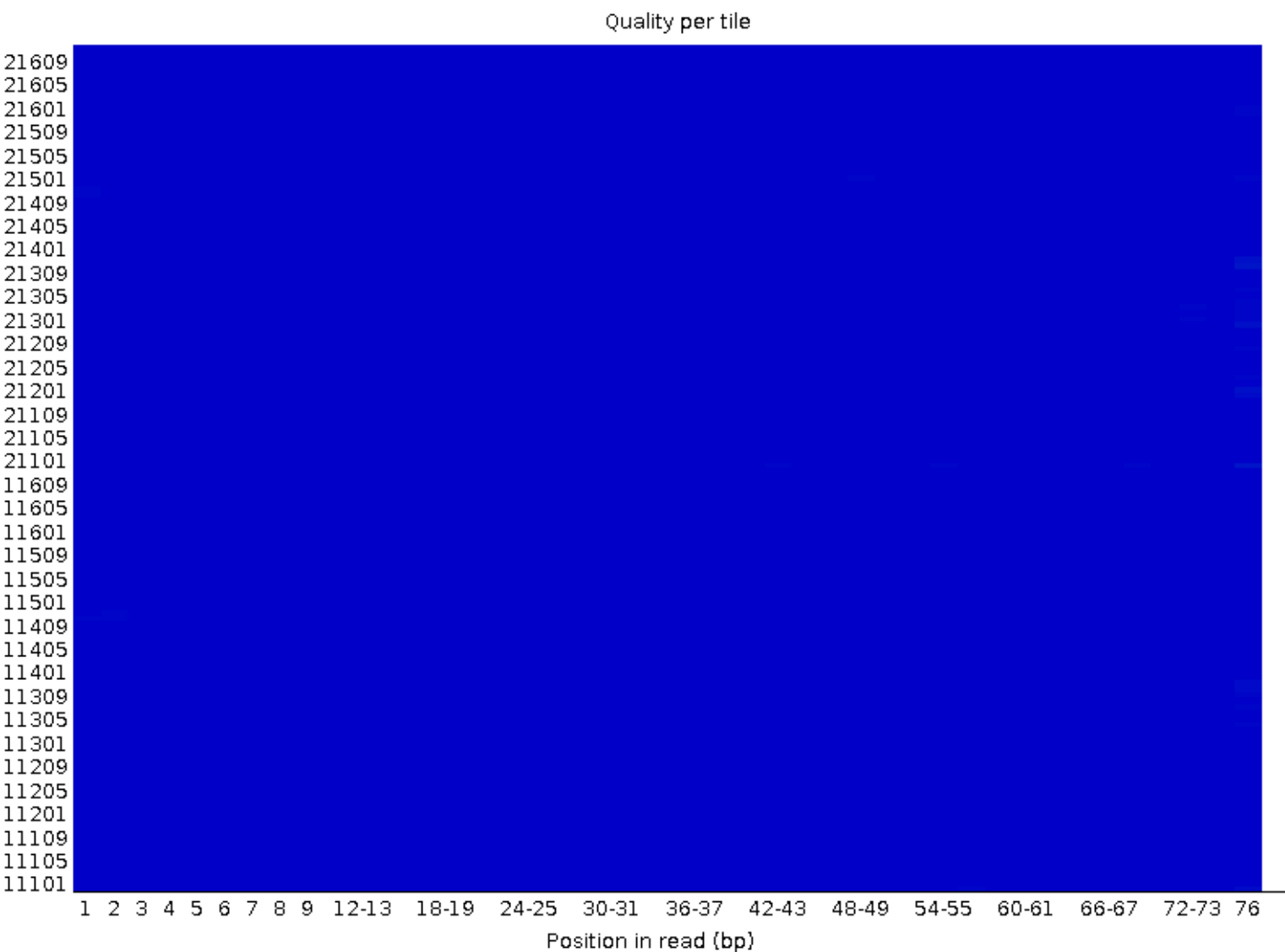
## Basic Statistics

Measure	Value
Filename	ERR13985875_2.fastq
File type	Conventional base calls
Encoding	Sanger / Illumina 1.9
Total Sequences	10689040
Sequences flagged as poor quality	0
Sequence length	76
%GC	47

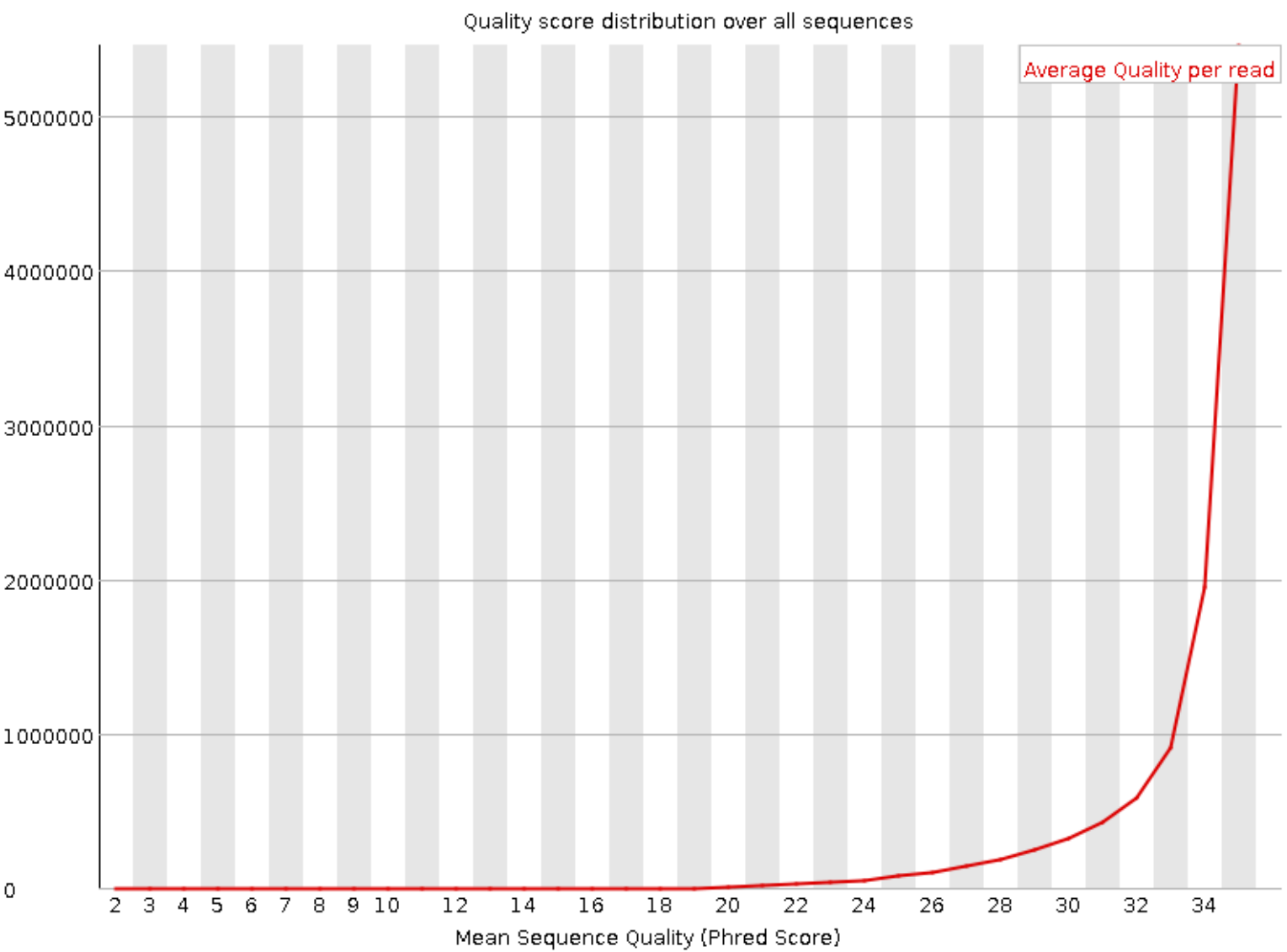
## ✔ Per base sequence quality



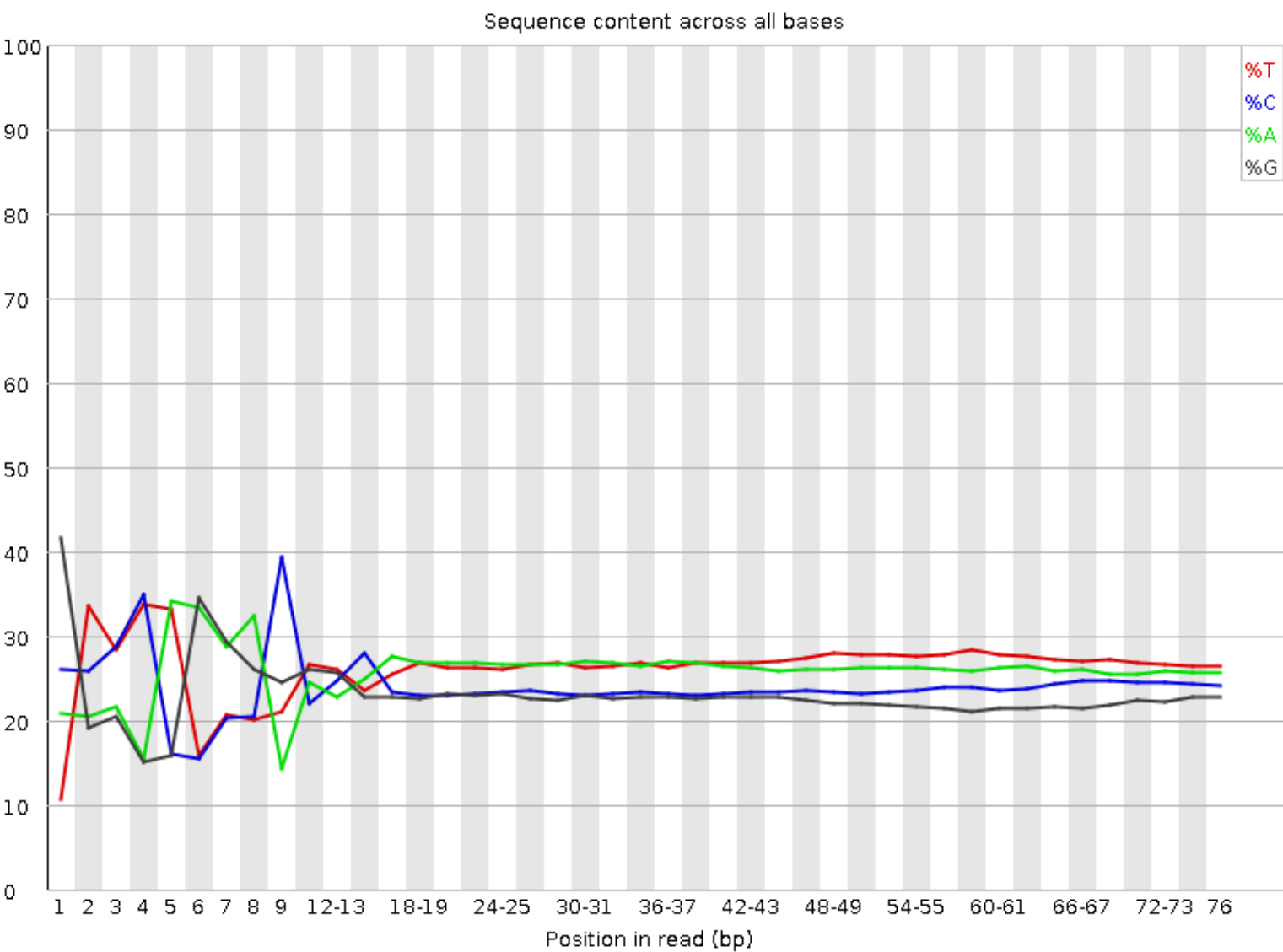
# Per tile sequence quality



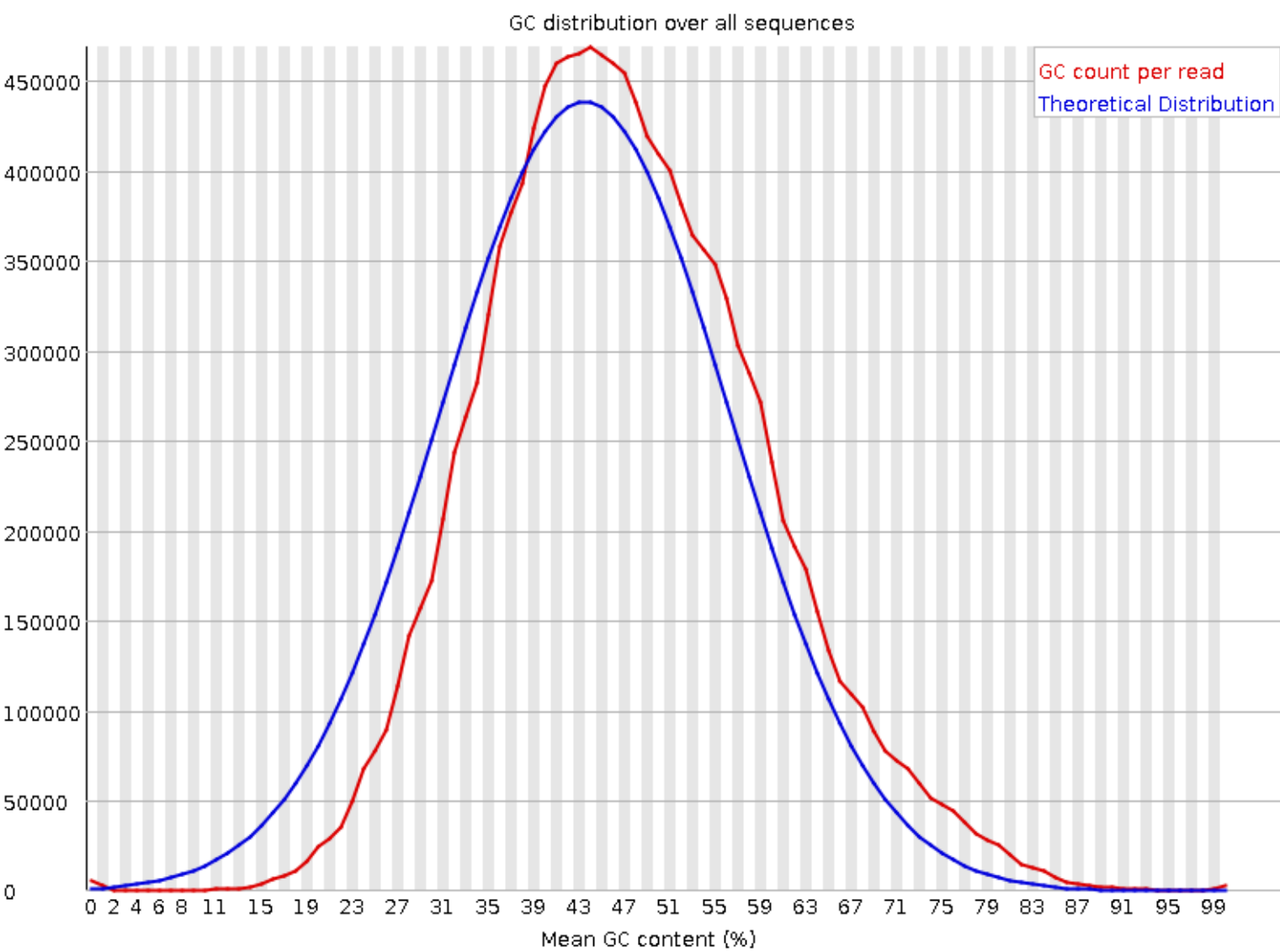
## ✔ Per sequence quality scores



## 🚨 Per base sequence content

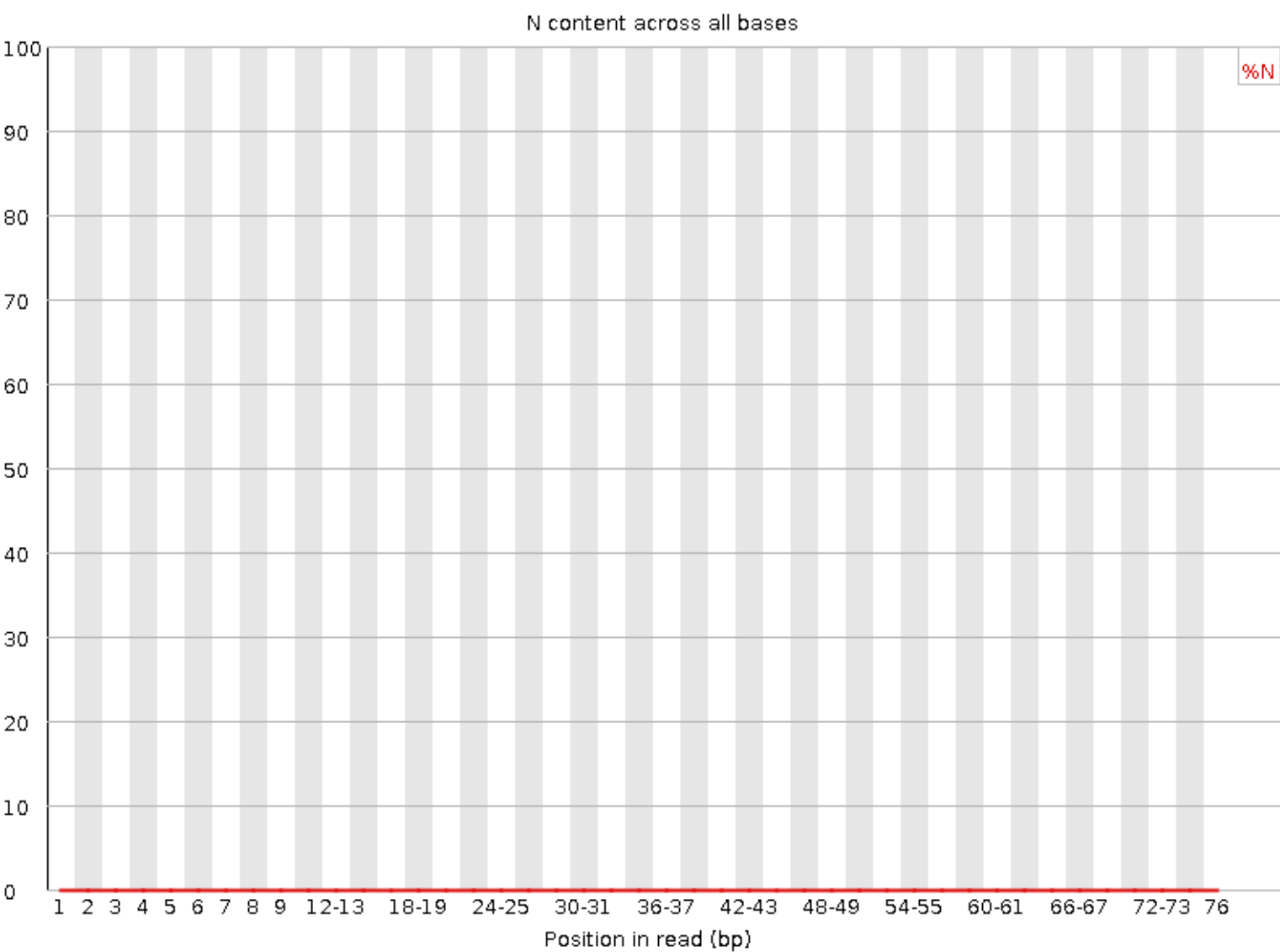


## 🚨 Per sequence GC content

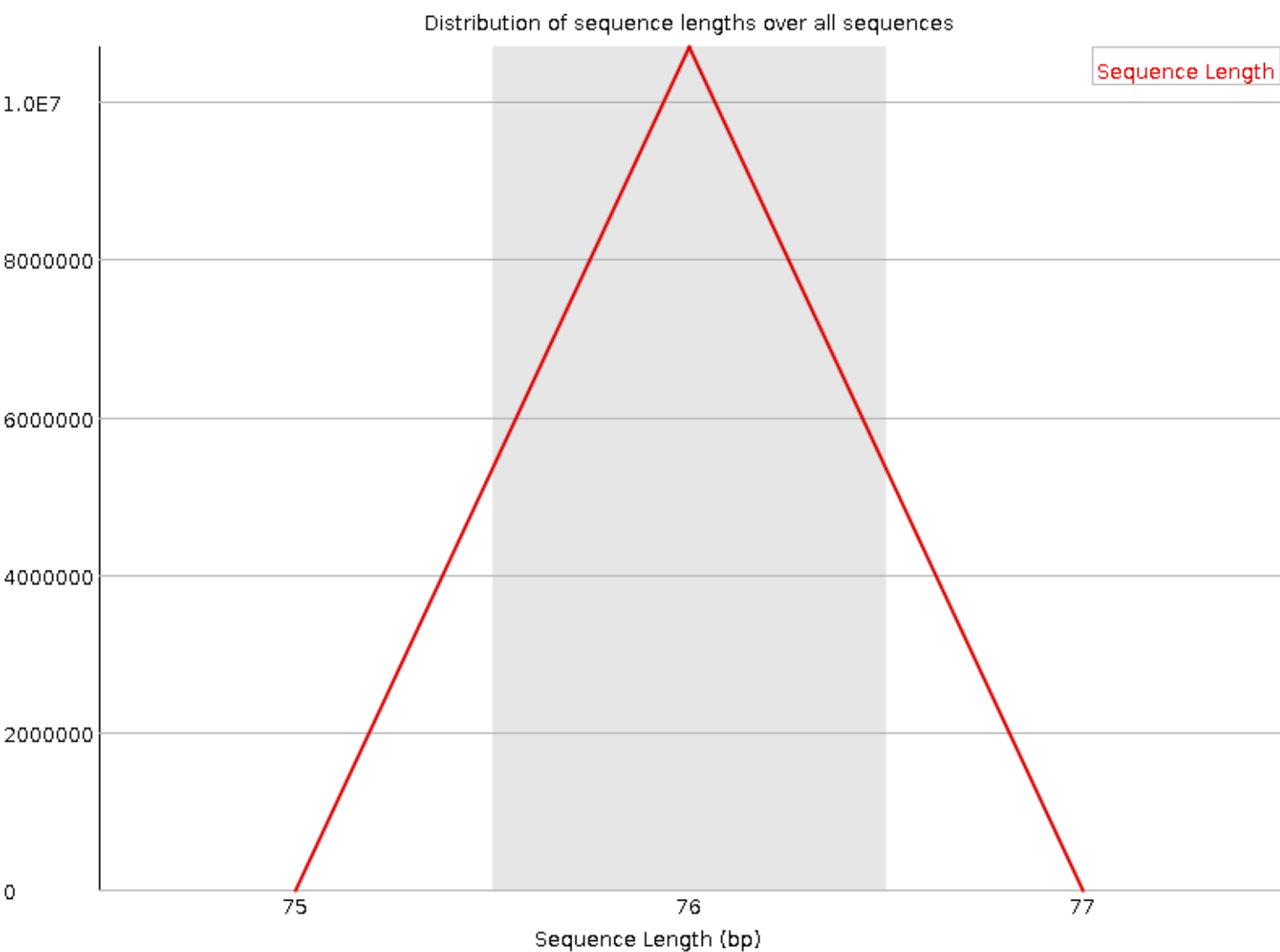




## Per base N content

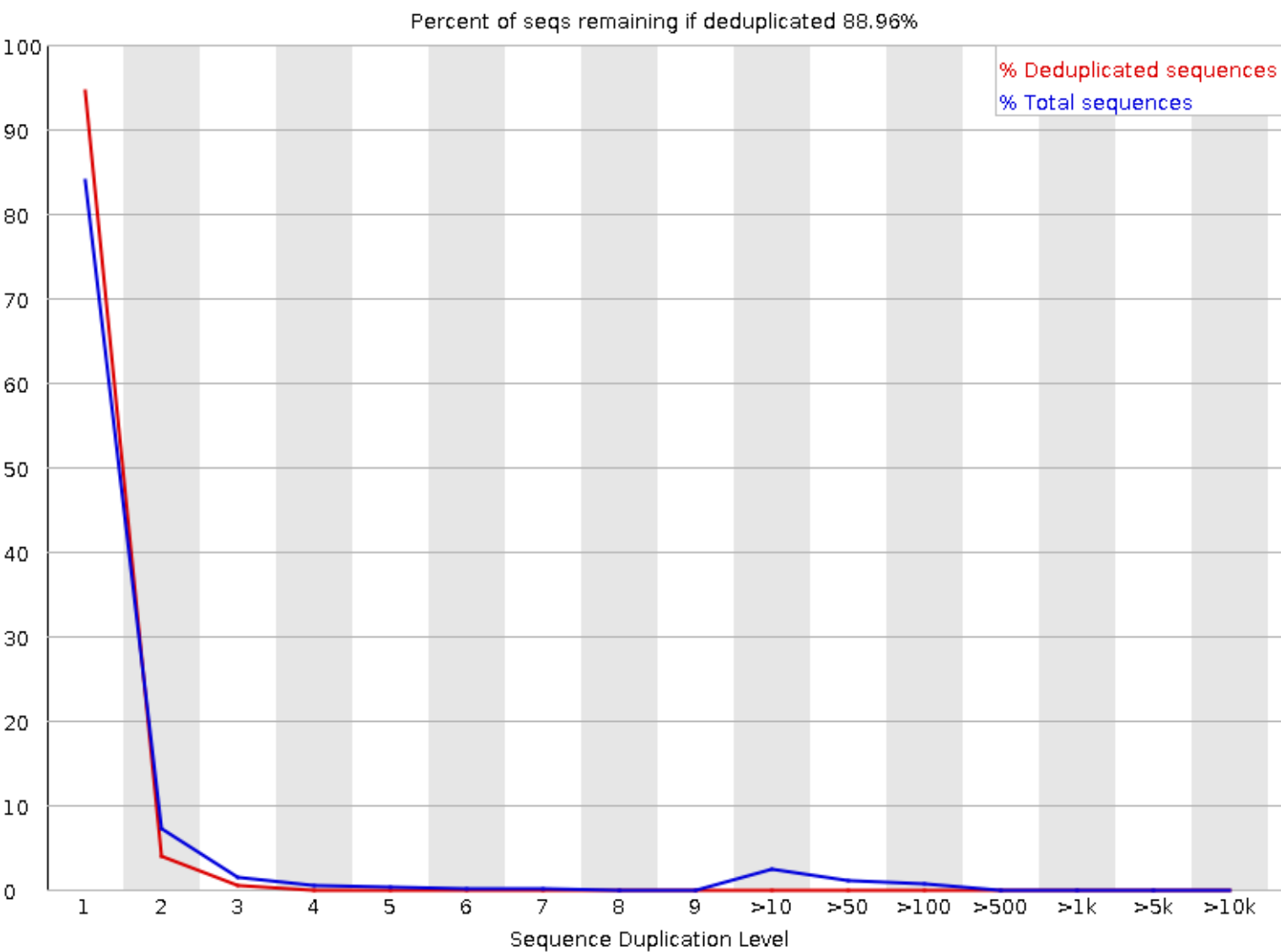


# Sequence Length Distribution





## Sequence Duplication Levels



## Overrepresented sequences

No overrepresented sequences

# Adapter Content

