

# QAA\_Lab\_Report

Julia Jones

2024-09-08

## File assignments

Under the path: `projects/bgmp/shared/2017_sequencing/demultiplexed/`

File	Alias
34_4H_both_S24_L008_R1_001.fastq.gz	34_4H_R1
34_4H_both_S24_L008_R2_001.fastq.gz	34_4H_R2
21_3G_both_S15_L008_R1_001.fastq.gz	21_3G_R1
21_3G_both_S15_L008_R1_001.fastq.gz	21_3G_R2

Table 1: My file assignments

File	Size	Read Count	Read Length
21_3G_R1	480M	9,237,299	101
21_3G_R2	535M	9,237,299	101
34_4H_R1	469M	9,040,597	101
34_4H_R2	525M	9,040,597	101

Table 2: Relevant information about my files

## FastQC

### 21\_3G

Score	On	Original File
PASS	Per base sequence quality	21_3G_R1
PASS	Per base N content	21_3G_R1

Table 3: FASTQC 21\_3G\_R1 output overview

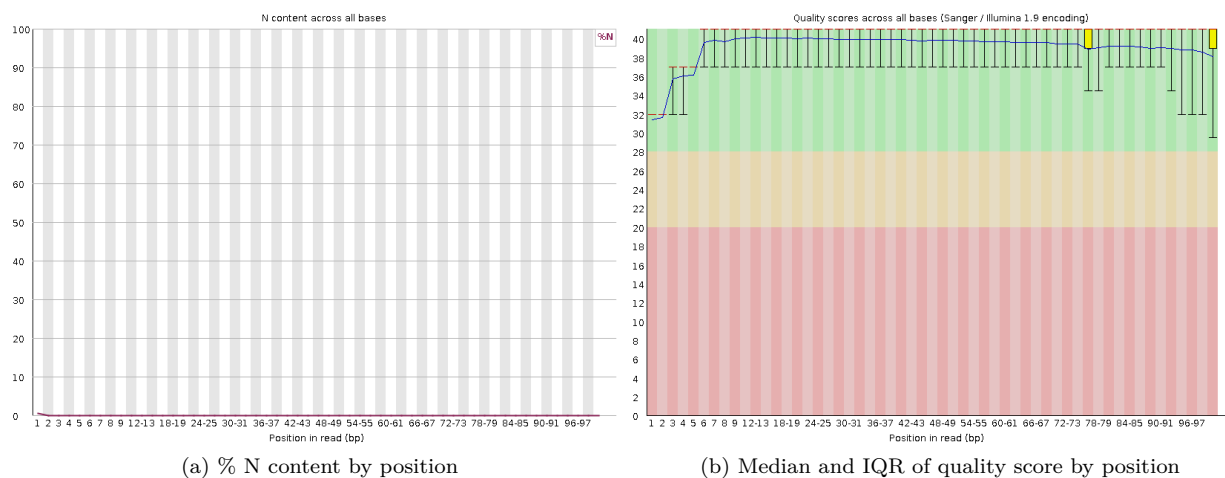


Figure 1: FASTQC output for 21\_3G\_R1

N-content stays extremely low throughout the reads, even at the susceptible early positions. As there are no quality data outliers in the red zone of the quality score distribution plot, this lack of N-content makes sense.

Quality is worse at the lower positions, while quality score variation increases at the highest positions. Despite this all qualities lie within the acceptable region.

Score	On	Original File
PASS	Per base sequence quality	21_3G_R2
PASS	Per base N content	21_3G_R2

Table 4: FASTQC 21\_3G\_R2 output overview

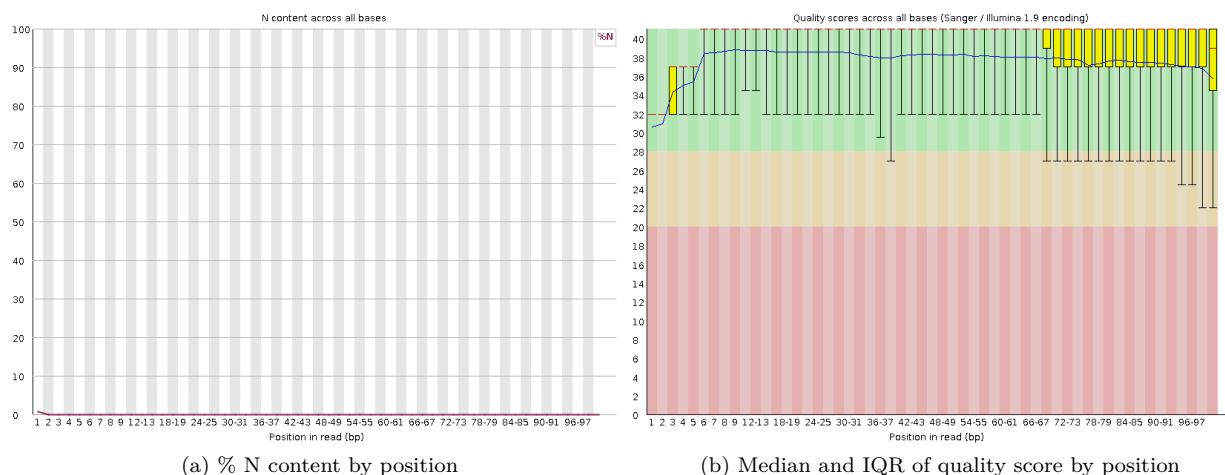


Figure 2: FASTQC output for 21\_3G\_R2

N-content stays extremely low throughout the reads, even at the susceptible early positions. As there are no quality data outliers in the red zone of the quality score distribution plot, this lack of N-content makes sense.

Average quality is worse at the lower positions, while quality score variation increases at the highest positions.

Despite this all median qualities lie within the acceptable region. The outlier warning qualities at positions 66-97 are something to look out for when examining other metrics.

### 34\_4H

Score	On	Original File
PASS	Per base sequence quality	34_4H_R1
PASS	Per base N content	34_4H_R1

Table 5: FASTQC 34\_4H\_R1 output overview

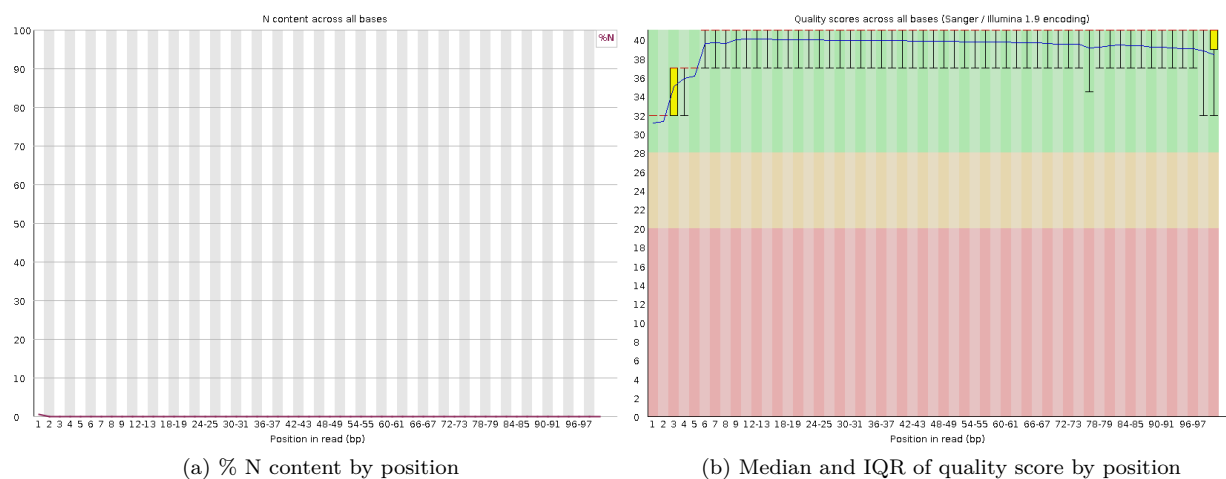


Figure 3: FASTQC output for 34\_4H\_R1

N-content stays extremely low throughout the reads, even at the susceptible early positions. As there are no quality data outliers in the red zone of the quality score distribution plot, this lack of N-content makes sense.

Quality is worse at the lower positions, while quality score variation increases at the highest positions. Despite this all qualities lie within the acceptable region.

Score	On	Original File
PASS	Per base sequence quality	34_4H_R2
PASS	Per base N content	34_4H_R2

Table 6: FASTQC 34\_4H\_R2 output overview

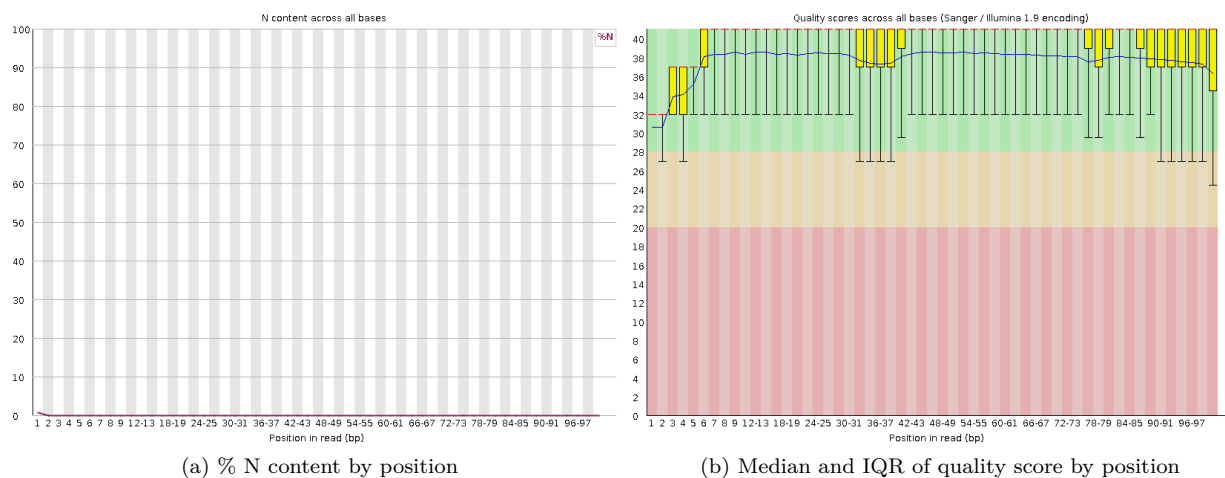


Figure 4: FASTQC output for 34\_4H\_R2

N-content stays extremely low throughout the reads, even at the susceptible early positions. As there is no quality data outliers in the red zone, this lack of N-content makes sense

Average quality is worse at the lower positions, while quality score variation increases at 31-32 and 72-101. All median qualities lie within the acceptable region. The outlier qualities at positions 31-32 and 72-101 are something to look out for when examining other metrics.

Because of how low our N content is,

## My Quality Score Script

### 21\_3G

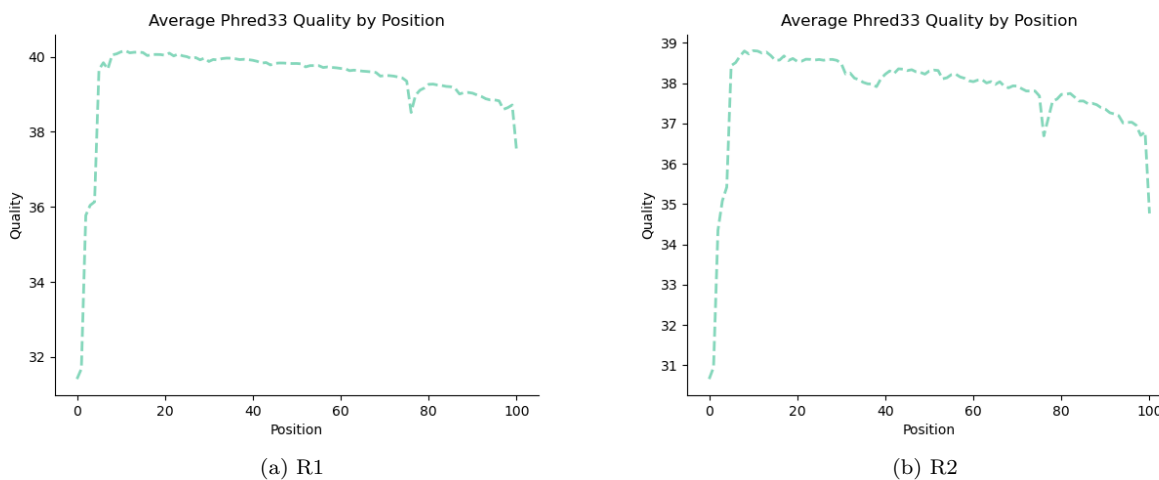


Figure 5: Average quality score by position for 21\_3G

R2 means seems to be consistently lower than R1 means by 1 point. This slight difference is probably insignificant.

### 34\_4H

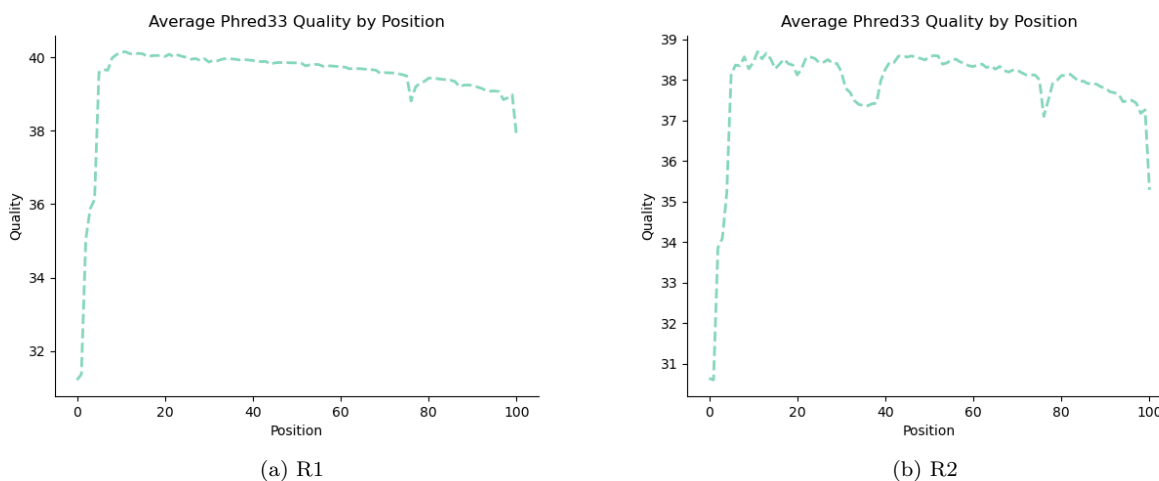


Figure 6: Average quality score by position for 34\_4H

R2 means seems to be consistently lower than R1 means by 1 point (plus a small region at 35-40 that is lower by 3-4 points). This slight difference should be insignificant.

### My script vs. FASTQC

#### Quality of quality score distribution plots

The FASTQC data plots impart much more information than my own. My plots solely map the mean quality distribution by position. Thus, I cannot see when my quality scores are lowered due to outliers or from low quality as a whole. In contrast, FASTQC calculates the mean, median, and IQR. And while my mean line and FASTQC's mean line look identical, the FASTQC's multi-metric methodology handles outliers much better.

FASTQC also incorporates great visual cues into their plots: a hard ylim of 40 and quality color coding in the figure's background. These additions make interpreting quality scores much simpler at a glance. In my code, I did not set a hard ylim. So, if a whole run is relatively worse than normal, my y-axis maximum will be lower than 40 (i.e., Figure 6b) this coupled with the lack of background color coding can mask the low-quality nature of the run at first glance.

#### Runtime difference

File	Script	Runtime (seconds)
21_3G_R1	mine	304.88
21_3G_R1	FASTQC	38.27*
21_3G_R2	mine	308.32
21_3G_R2	FASTQC	38.27*
34_4H_R1	mine	307.19

File	Script	Runtime (seconds)
34_4H_R1	FASTQC	38.27*
34_4H_R2	mine	317.67
34_4H_R2	FASTQC	38.27*
average	mine	309.52
average	FASTQC	38.27

Table 7: FASTQC vs. my run time

\*Ran all files at once and divided total run time by 4.

On average, FASTQC is  $(309.52/38.27)=8.1$  times **faster** than my code!

This large disparity in speed makes sense. I wrote my De-multiplexing code in one week in Python, while FASTQC was built by a team in 2010 in Java (a generally faster language) and continually improved over the past decade. Even in the one week I spent on De-multiplexing I was able to greatly speed up my code. This is a case of disparity in time and resources.

### Memory/CPU usage

File	Script	% CPU used
21_3G_R1	mine	99
21_3G_R1	FASTQC	97
21_3G_R2	mine	99
21_3G_R2	FASTQC	97
34_4H_R1	mine	99
34_4H_R1	FASTQC	99
34_4H_R2	mine	99
34_4H_R2	FASTQC	97

Table 8: FASTQC vs. my data usage

CPU usage was consistent across runs, with FASTQC using slightly less CPU than my own code. This difference is slight enough to be insignificant.

### Comment on overall data quality

**Comment on the overall data quality of your two libraries. Go beyond per-base qscore distributions. Make and justify a recommendation on whether these data are of high enough quality to use for further analysis.**

### 21\_3G Overall quality

Recommendation	Metric	File
PASS	Per base sequence quality	R1, R2
PASS	Per base N content	R1, R2
PASS	Sequence Length Distribution	R1, R2
WARN	Sequence Duplication Levels	R1, R2
FAIL	Per tile sequence quality	R1, R2

Recommendation	Metric	File
FAIL	Per base sequence content	R1
WARN	Per base sequence content	R2
PASS	Per sequence GC content	R1, R2
WARN	Overrepresented sequences	R1, R2
PASS	Adapter Content	R1, R2

Table 9: FASTQC Calls on all metrics

### Per base sequence quality, Per base N content

This metric is not a concern with this data. See FASTQC section above for more information.

### Sequence Length Distribution

All sequences were length 101. See File assignments section above for more information.

### Sequence duplication levels

Because this is an RNA-seq experiment, we don't care about sequence duplication levels. In RNA-seq, sequence duplication imparts differential expression.

### Per tile sequence quality

Both R1 and R2 failed this metric, lets examine the tile results in more detail.

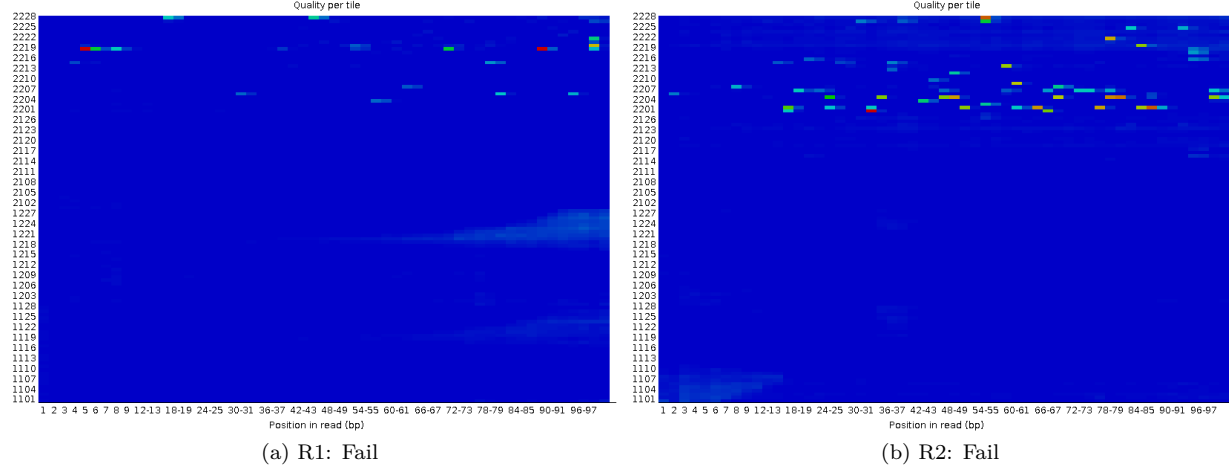


Figure 7: Per tile sequence quality for 21\_3G

An ideal per-tile quality plot should be completely blue (high quality). Here our R1 file has some low quality hotspots from the 3' and 5' end of reads constructed in tile 2219. This cell constructed a small proportion of our total R1 reads. Though R2 has more middling quality tiles, its number of consistently low-quality tiles is small (2201, 2204). Thus, I would not consider tile-quality to be a concern for our data.

## Per base sequence content

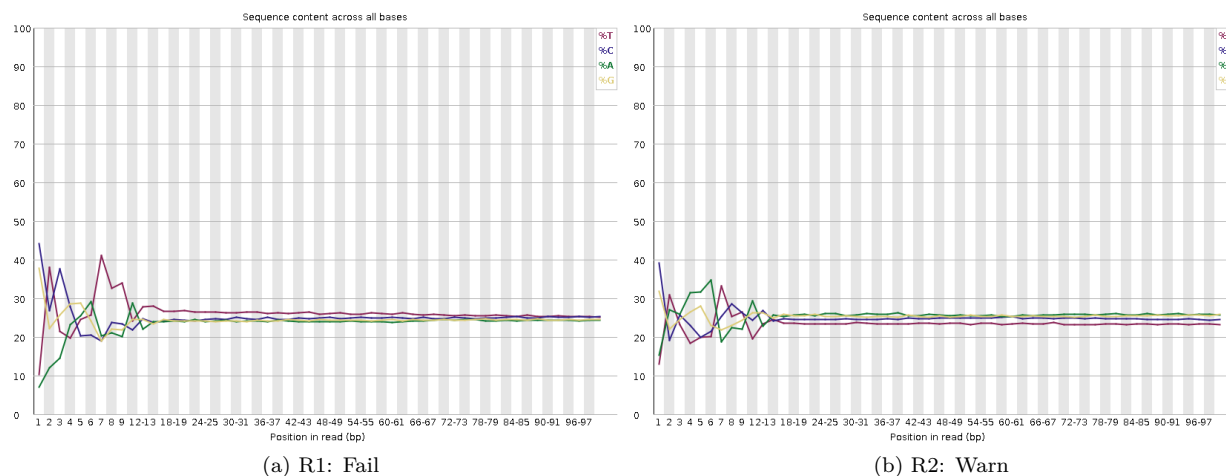


Figure 8: Per base sequence content for 21\_3G

In an ideal per-base content graphs the A, T, C, and G y-values would remain consistent as you move along the x-axis. In our data, the y-values (sequence contents) spike and drop until position 13 where they level out. Thus, R1 is called a fail, and R2 is a warning. R1 fails because its T sequence content spike from position 5-9 is higher than anything in our R2 plots. But, because this y-value volatility is localized to the start of the reads, where we have low-quality data to be trimmed downstream by **Trimmomatic**, I would not be concerned by R1's or R2's per-base sequence content.

## Per base GC content

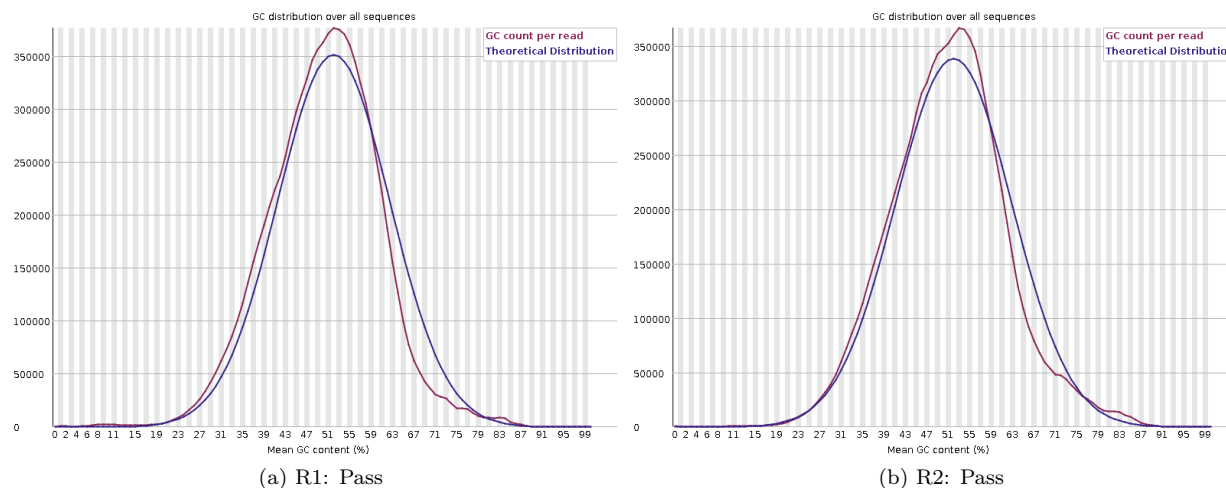


Figure 9: Per base GC content for 21\_3G

For both R1 and R2, our GC content closely follows the theoretical distribution provided by FASTQC. The center of our distribution is a bit higher than the theoretical distribution, but as this is the value where our



nucleotides are balanced (50%) this is not problematic. Thus, GC content is likely not an area of concern here.

### Overrepresented sequences

File	Source	Count	Percentage
R1	TruSeq Adapter, Index 18	29130	0.32
R2	Illumina Single End Sequencing Primer	30251	0.33

Though we got a warning for R1 and R2, I would not call this data problematic (even though 21\_3G has more overrepresented sequences than 34\_4H). Both of our over represented sequences are known adapters, and thus will be captured and trimmed downstream with cutadapt. Thus over represented sequences are not an area of concern for our data.

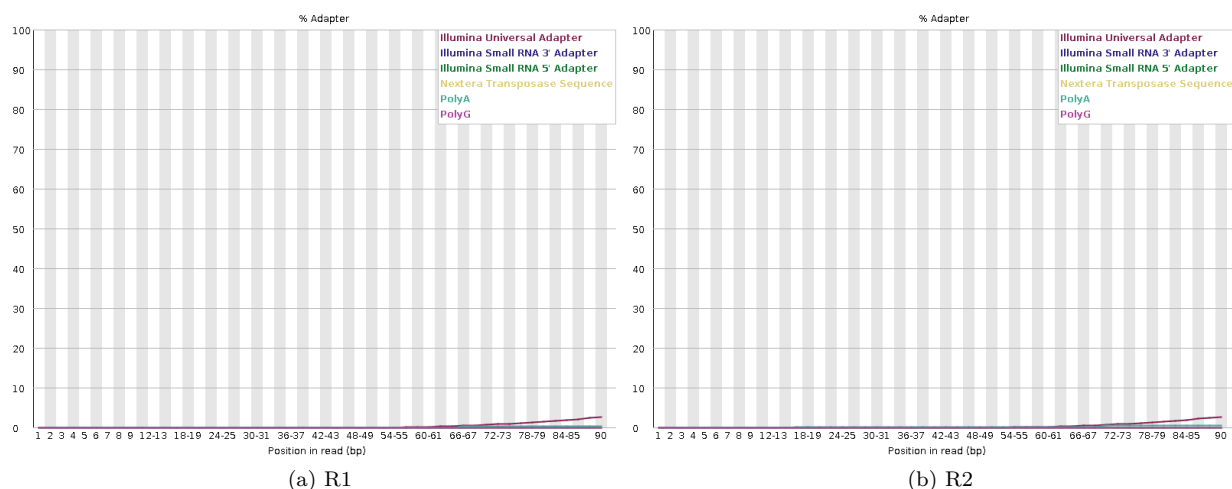


Figure 10: Adapter content 21\_3G

**Adapter Content** As you approach the 3' end of our read, adapter content does increase. In both our R1 and R2 files this difference is slight. Furthermore, the vast majority of this adapter content is the Illumina universal adapter which will be trimmed with cutadapt. Thus, adapter content is not a concern for this dataset.

### 21\_3G Final recommendation

With the metrics discussed above in mind, it's safe to say that both 21\_3G files are high enough quality to proceed with further analysis. Most metrics passed, and those that produced a warning or failure make sense (sequence duplication) or can be fixed downstream (per-base sequence content).

### 34\_4H Overall quality

Recommendation	Metric	File
PASS	Per base sequence quality	R1, R2

Recommendation	Metric	File
PASS	Per base N content	R1, R2
PASS	Sequence Length Distribution	R1, R2
WARN	Sequence Duplication Levels	R1, R2
FAIL	Per tile sequence quality	R1, R2
FAIL	Per base sequence content	R1
WARN	Per base sequence content	R2
PASS	Per sequence GC content	R1, R2
WARN	Overrepresented sequences	R1, R2
PASS	Adapter Content	R1, R2

Table 9: FASTQC Calls on all metrics

### Per base sequence quality, Per base N content

Overall, this metric is not a concern with this data. See FASTQC section above for more information.

### Sequence Length Distribution

All sequences were length 101. See File assignments section above for more information.

### Sequence duplication levels

Because this is an RNA-seq experiment, we don't care about sequence duplication levels. In RNA-seq, sequence duplication imparts differential expression.

### Per tile sequence quality

Both R1 and R2 failed this metric, lets examine the tile results in more detail.

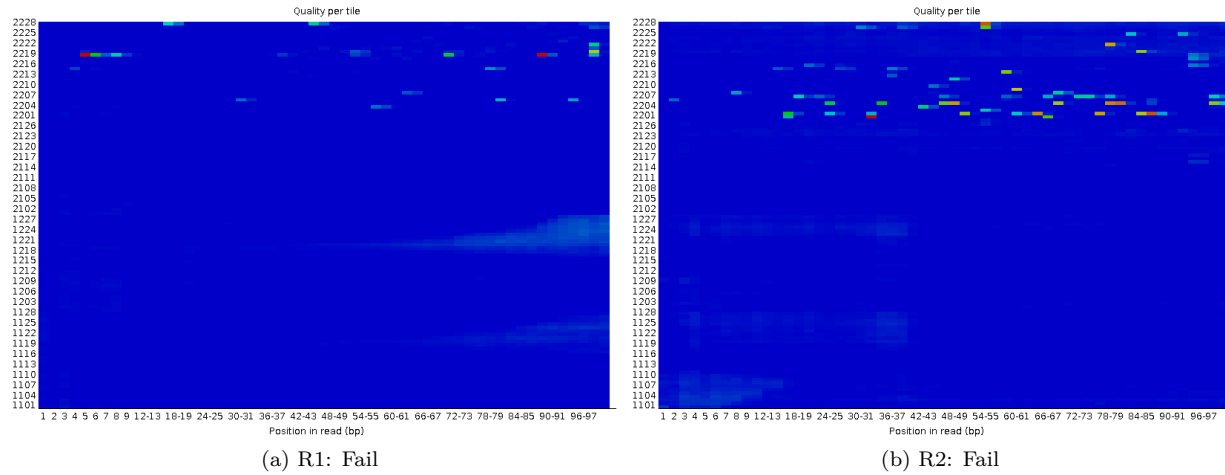


Figure 11: Per tile sequence quality for 34\_4H

An ideal per-tile quality plot should be completely blue (high quality). Here our R1 file has some low quality hotspots from the 3' and 5' end of reads constructed in tile 2219 and 2222. These cells constructed a small proportion of our total R1 reads. Though R2 has more middling quality tiles, its number of low-quality tiles is small. Thus, I would not consider tile-quality to be a concern for our data here.

## Per base sequence content

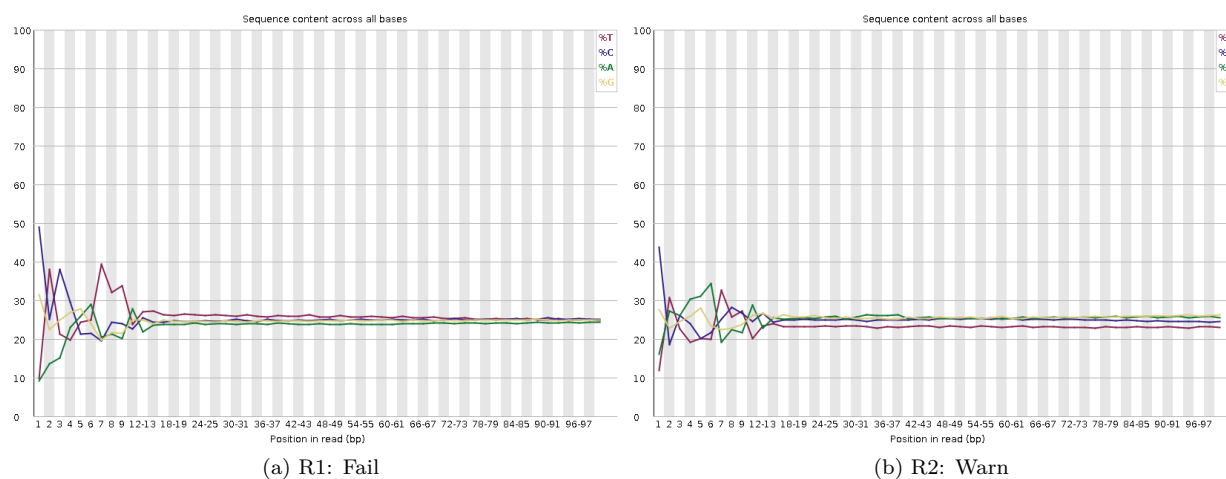


Figure 12: Per base sequence content for 34\_4H

These R1 and R2 graphs are almost identical to the 21\_3G graphs, with slightly more volatility around the first base. Recall, in an ideal per-base content graphs the A, T, C, and G y-values would remain consistent as you move along the x-axis. In our data, the y-values (sequence contents) spike and drop until position 13 where they level out. Thus, R1 is called a fail, and R2 is a warning. R1 fails because its T sequence content spike from position 6-12 is higher than any spike in our R2 plots. But, because y-value volatility is localized to the start of the reads, where our low-quality data will be trimmed downstream, I would not be concerned by R1's or R2's per-base sequence content.

## Per base GC content

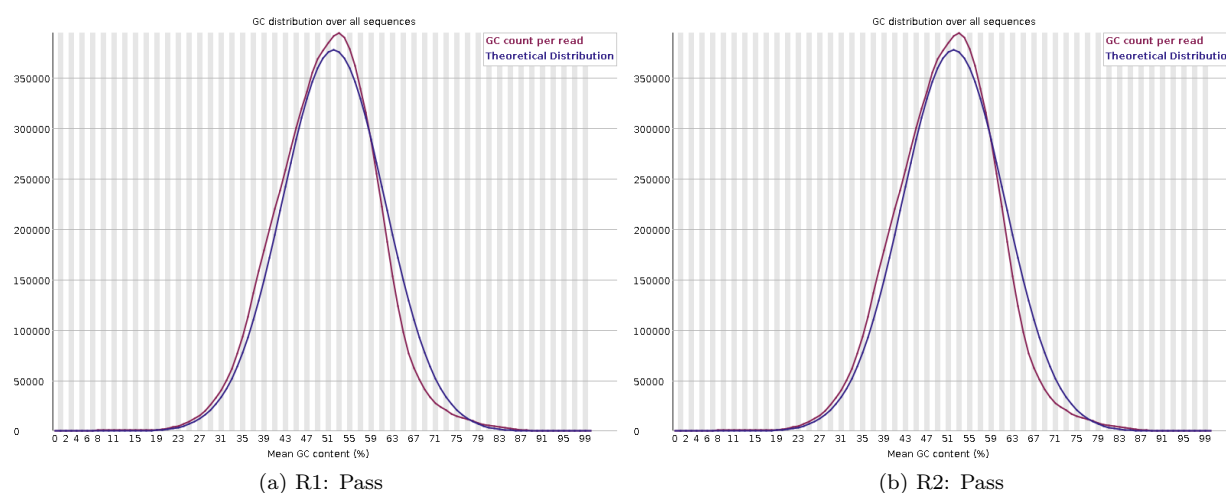


Figure 13: Per base GC content for 34\_4H

For both R1 and R2, our GC content closely follows the theoretical distribution provided by FASTQC. Thus, GC content is likely not an area of concern here.

### Overrepresented sequences

File	Source	Count	Percentage
R1	TruSeq Adapter, Index 15	9502	0.11
R2	Illumina Single End Sequencing Primer	9611	0.11

Though we got a warning for R1 and R2, I would not call this data problematic. Both of our over represented sequences are known adapters, and will be captured and trimmed downstream with `cutadapt`. Thus overrepresented sequences are not an area of concern for our data.

### Adapter Content

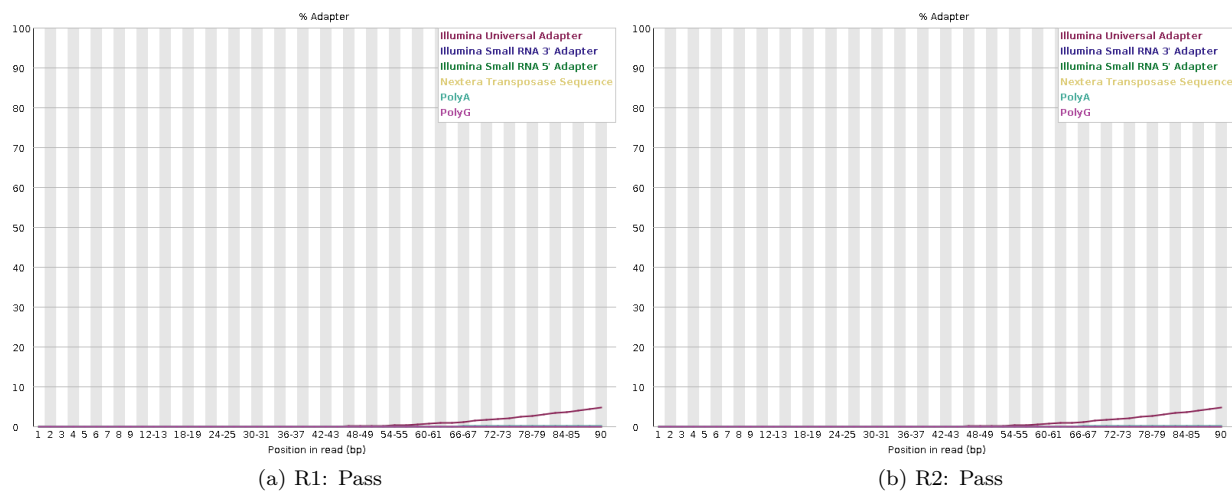


Figure 14: Adapter content 34\_4H

Adapter content appears to be identical between R1 and R2. As you approach the 3' end of our read, adapter content does increase. Adapter content at the 3' end is higher in 34\_4H than 21\_3G, though for both replicates the level of adapter content is not a concern. This is because the vast majority of the adapter content is the Illumina universal adapter which will be trimmed with `cutadapt`. Thus, adapter content is not a concern for this dataset.

### 34\_4H final quality determination

With the metrics discussed above in mind, it's safe to say that both 34\_4H files are high enough quality to proceed with further downstream analysis. Though my 21\_3G data is higher quality than my 34\_4H most metrics passed, and those that produced a warning or failure make sense (sequence duplication) or can be fixed downstream (per-base sequence content).

## Part 2

### Cutadapt

#### Determine adapters

Mechanistically in one run, adapters won't be present in every sequence. They may not even be the top hit. But because Illumina uses universal adapters, we can compare across disparate runs (34\_4H\_RN and 21\_3G\_RN). Comparing across files, the universal adapter is the unifying sequence.

File name	Adapter
21_3G_R1	AGATCGGAAGAGCACACGTCTGAACTCCAGTCA
21_3G_R2	AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT
34_4H_R1	AGATCGGAAGAGCACACGTCTGAACTCCAGTCA
34_4H_R2	AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT

*Table 9: Universal adapter by file*

With my adapters determined, I could proceed to actually running `cutadapt`

#### Cutadapt results

File name	Percent adapter trimmed
21_3G_R1	6.6%
21_3G_R2	7.4%
34_4H_R1	9.1%
34_4H_R2	9.8%

*Table 10: Percentage of file requiring adapter trimming*

#### Cutadapt interpretation

As discussed in the 34\_4H Overall quality section, my 34\_4H data had higher adapter content (and were trimmed more) than my 21\_3G data. Thus, it is not surprising that my 34\_4H files were more adapter trimmed than my 21\_3G files. Nevertheless, in all my assigned files, the percent that needed to be adapter trimmed were relatively low. In both 21\_3G and 34\_4H our R2 files were trimmed more than our R1 files. This makes sense because R2 runs last on the sequencer and is typically lower quality than R1. For more information on why R2 files were more trimmed than R1 files see the section **R1 and R2 adapter trimmed at different rates?** below.

#### Trimmomatic

With our adapters removed, our data is now ready for Trimmomatic

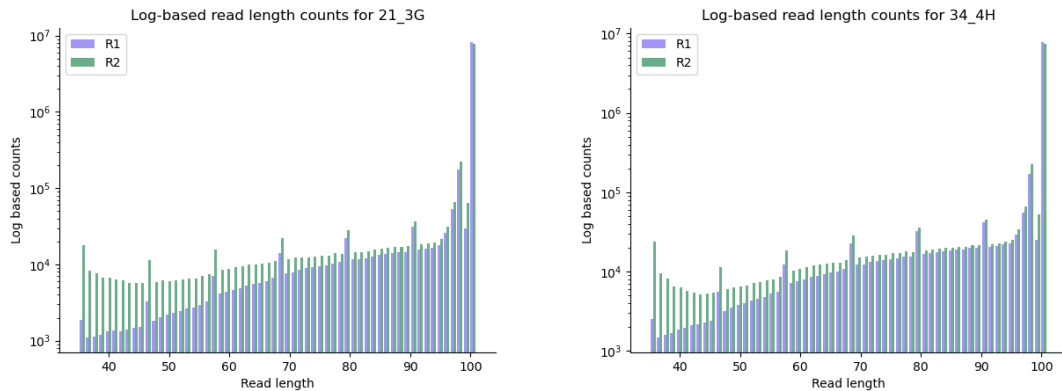


Figure 15: Read length distribution by file

### R1 and R2 adapter trimmed at different rates?

R2 files should be adapter trimmed at higher rates than R1 files because R2 files are completed at the end of the run. As the run continues reagents and the signal degrade. Signal degradation causes adapter bases to be called over the insert bases, causing adapters to be overrepresented. This contributes to the overall lowering of run quality, which we can see from our FASTQ output above.

## Part 3

Genome generate :angel:

Align reads

Replicate	Mapped reads	Unmapped reads	% mapped
21_3G	15,125,868	2,580,756	85.4
34_4H	15,631,049	1,675,233	90.3

Table 11: Percentage of mapped reads by replicate

### Determining strandedness

I propose that both my 21\_3G and 34\_4H libraries are stranded. The reasoning behind this determination is clear once one examines the htseq-count output in detail.

## 21\_3G

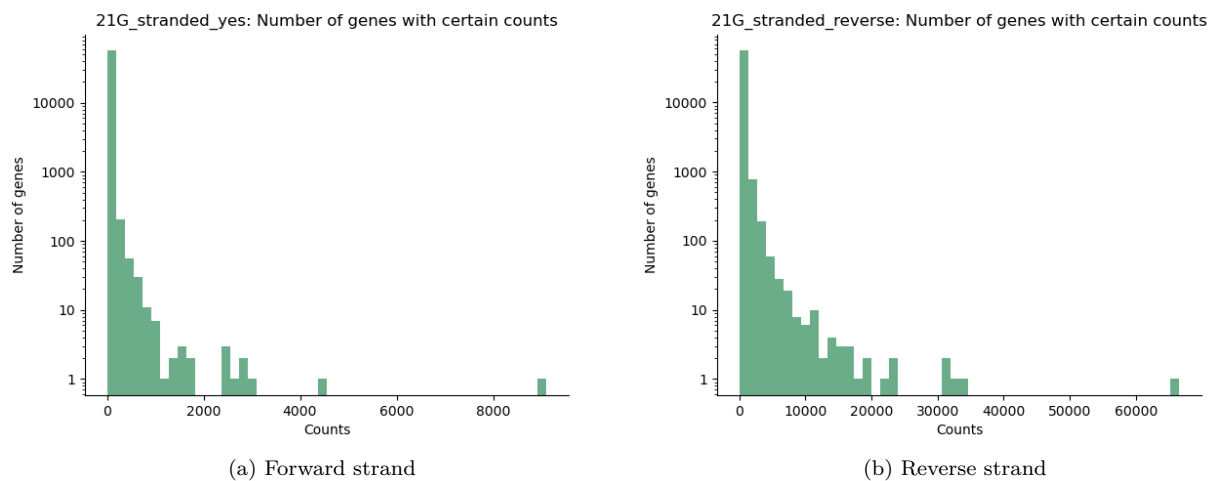


Figure 16: Number of genes mapped X times by strand

If our data was unstranded, the number of genes mapped to specific runs would be the same whether the strand direction was set to forward or reverse. This is not the case: visually the reverse strand plot has much more data mapped than the forward strand plot. Numerically 303,726 reads mapped on the forward strand and 6,636,009 reads mapped on the reverse strand. This 2084.87% increase

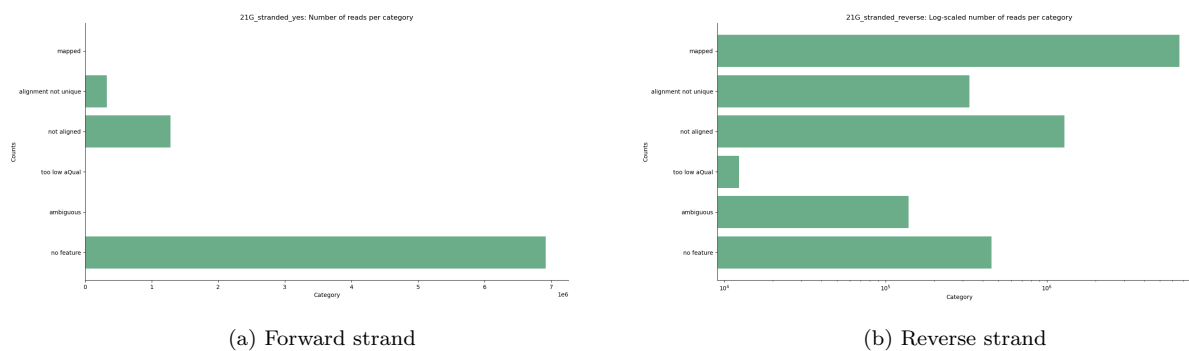


Figure 17: Number of reads within each category

In the forward strand data, “no feature” dominates our read categories. In the reverse strand data, “mapped” dominates our read categories. This indicates that our data is stranded in the **reverse** direction.

## 34\_4H

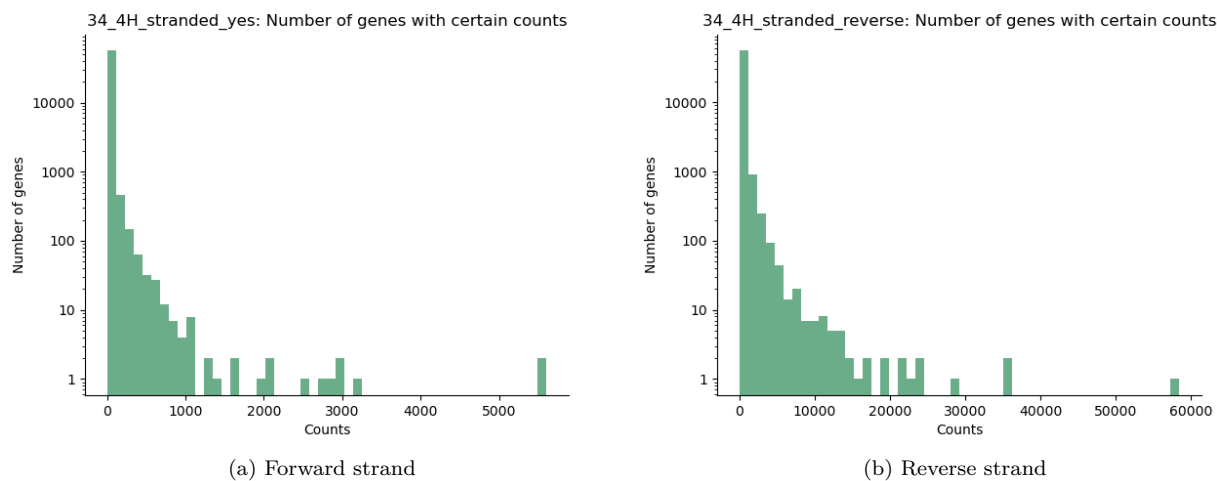


Figure 18: Number of genes mapped X times by strand

The same as was mentioned for 21\_3G holds true for 34\_4H. If our data was unstranded, the number of genes mapped to specific runs would be the same whether the strand direction was set to forward or reverse. This is not the case: visually the reverse strand plot has much more data mapped than the forward strand plot. Numerically,

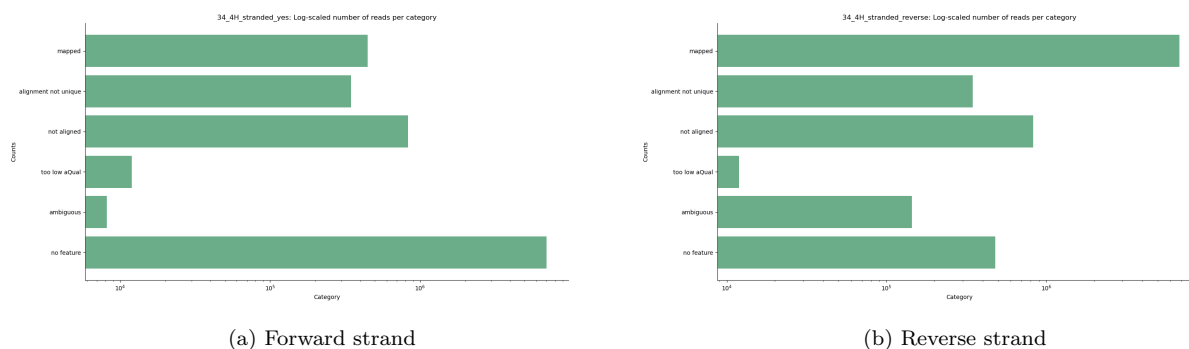


Figure 19: Number of reads within each category

As mentioned in 21\_3G, in the forward strand data “no feature” dominates our read categories. In the reverse strand data, “mapped” dominates our read categories. This indicates that our data is stranded in the **reverse** direction.