# RNA-Seq Quality Assessment

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## Part 1: Read quality score distributions

Plots of per-base quality scores and of per-base N content were generated using FASTQC version 0.11.5 as a module on Talapas, which was run by submitting the following script to Talapas' queuing system:

## #!/bin/bash

```
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00
```

module load fastqc/0.11.5

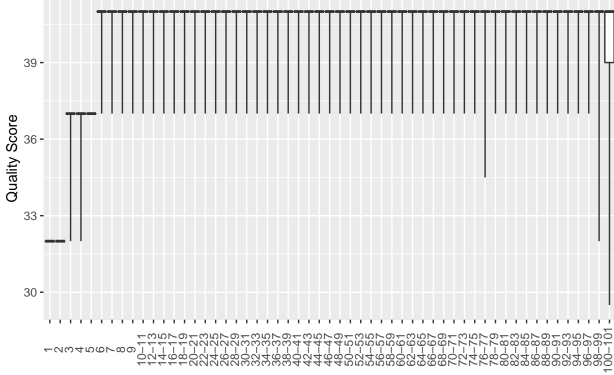
input\_dir=/projects/bgmp/shared/2017\_sequencing/demultiplexed
output\_dir=/projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/fastqc\_output/

exit

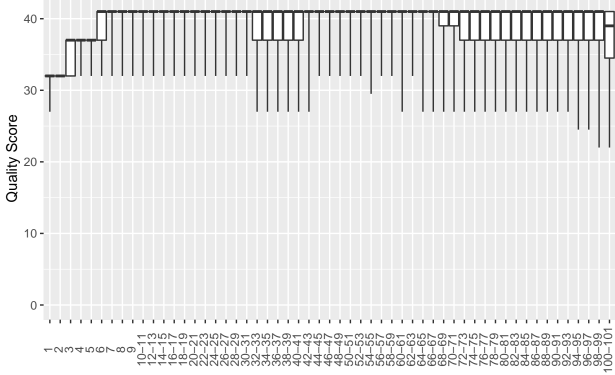
Plots from FASTQC have been re-plotted in R using the data points from FASTQC, for formatting purposes. Please see GitHub repo for original plots from FASTQC.

Per-base quality scores:

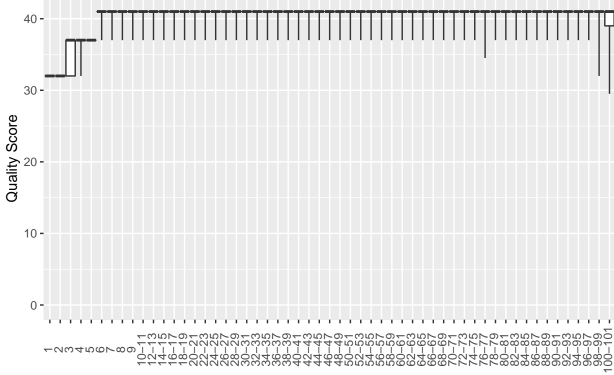
Per-Base Quality Scores in 27\_4C\_mbnl\_S19\_L008\_R1\_001.fastq



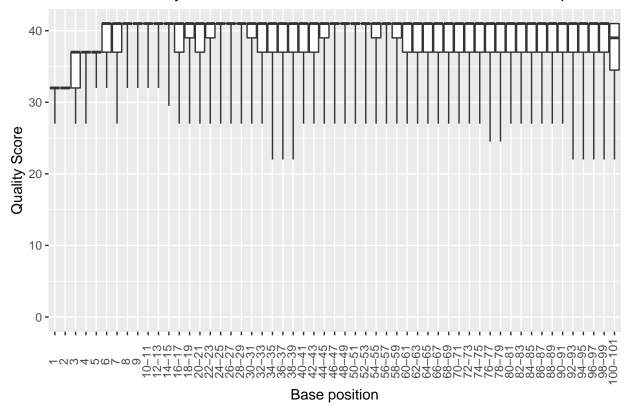
Per-Base Quality Scores in 27\_4C\_mbnl\_S19\_L008\_R2\_001.fastq



Per-Base Quality Scores in 28\_4D\_mbnl\_S20\_L008\_R1\_001.fastq

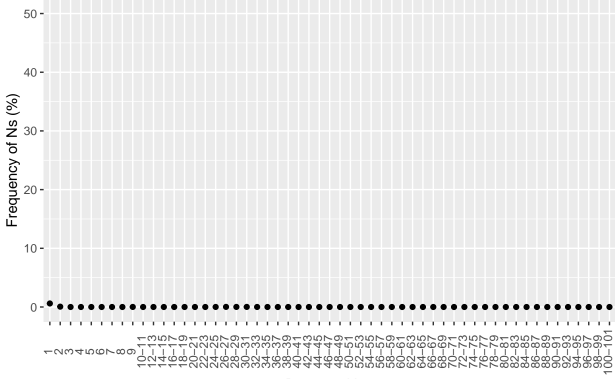


Per-Base Quality Scores in 28\_4D\_mbnl\_S20\_L008\_R2\_001.fastq

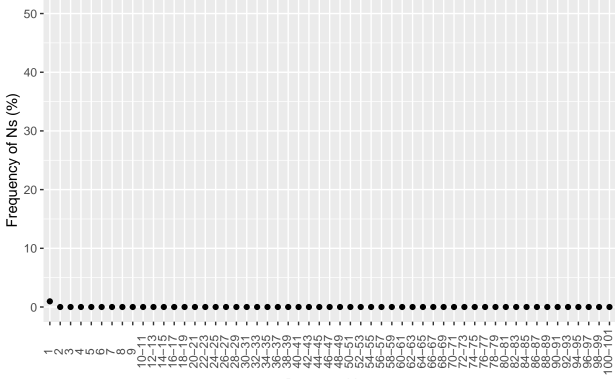


Per-base N content:

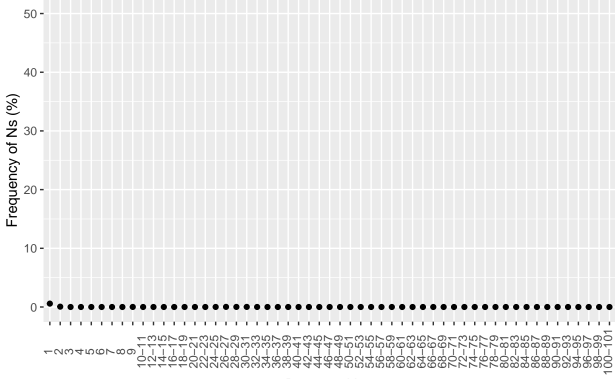
Per-Base N Content in 27\_4C\_mbnl\_S19\_L008\_R1\_001.fastq

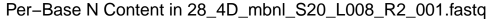


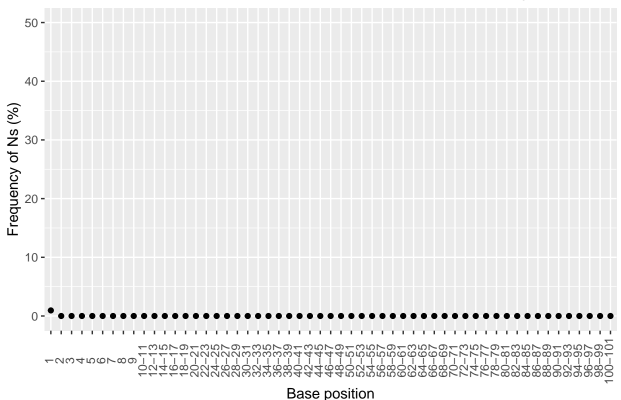
Per-Base N Content in 27\_4C\_mbnl\_S19\_L008\_R2\_001.fastq



Per-Base N Content in 28\_4D\_mbnl\_S20\_L008\_R1\_001.fastq







For all 4 reads, the per-base N content is consistent with the per-base quality scores. Each quality score plot shows that the quality score starts lower and rises over the first 5 bases or so, after which it stabilizes. Each N content plots shows that the N content starts a little higher and then quickly drops down to around 0. So, both plots show that the N content and quality scores are slightly worse at the very beginning of the read. Also, the N content is very low for all 4 reads, and all 4 reads have decent quality scores (well above 35, and closer to 40 for the read 1 reads).

I also produced per-base quality score plots directly from the FASTQs using my own Python script. The Python script, followed by its dependent module Bioinfo.py and the bash scripts used to run the program, are shown here:

Bioinfo.py module:

```
#!/usr/bin/env python
```

```
def validate_base_seq(seq: str,RNAflag: bool =False) -> bool:
    '''This function takes a string. Returns True if string is composed
    of only As, Ts (or Us if RNAflag), Gs, Cs. False otherwise. Case insensitive.'''
    DNAbases = set('ATGCatcg')
    RNAbases = set('AUGCaucg')
    return set(seq)<=(RNAbases if RNAflag else DNAbases)

if __name__ == '__main__':
    assert validate_base_seq("AATAGAT") == True, "Validate base seq does not work on DNA"
    assert validate_base_seq("AAUAGAU", True) == True, "Validate base seq does not work on RNA"
    assert validate_base_seq("Hi there!") == False, "Validate base seq fails to recognize nonDNA"
    assert validate_base_seq("Hi there!", True) == False, "Validate base seq fails to recognize nonDNA"</pre>
```

```
print("Passed DNA and RNA tests")
def convert phred(letter: str) -> int:
    """Converts a single character into a phred score for quality scores encoded with Phred+33"""
   dec: int = ord(letter)
   phred: int = dec - 33
   return phred
if __name__ == '__main__':
    assert convert_phred('I') == 40, 'Wrong phred score'
   print('Converted character into phred score correctly')
def gc_content(DNA: str) -> int:
    '''Takes DNA (or RNA) sequence and returns GC content of the sequence in decimal format as a fracti
   GC_count = 0
    if validate_base_seq(DNA):
       DNA = DNA.upper()
        for letter in DNA:
            if letter == 'G' or letter == 'C':
                GC count += 1
    else:
       print('Not a valid sequence')
   return GC_count/len(DNA)
if __name__ == '__main__':
    assert gc_content('ATGCGA') == 0.5, 'gc_content does not find correct GC content'
   print('Calculated gc content correctly')
def qual_score(phred_score: str) -> int:
    '''Calculates average quality score of a read, given a quality score string in Phred+33 notation'''
    sum: int = 0
   for letter in phred_score:
        sum += convert_phred(letter)
   return sum/len(phred_score)
if __name__ == '__main__':
    qual_score('JJII') == 40.5, 'Incorrect average phred quality score'
   print('Correct average phred quality score')
def fasta_fixer(fasta: str):
    '''Takes FASTA filename as input and outputs a FASTA file that has exactly 2 lines per record (one
   output = fasta[0:-2] + 'fixed.fa'
    seq = "
   with open(fasta, 'r') as fh, open(output, 'x') as out:
        for line in fh:
            line = line.strip()
            if line[0] == '>' and seq != '':
                out.write(header+'\n')
                out.write(seq+'\n')
                seq = ''
            if line[0] == '>':
               header = line
            else:
                seq += line
```

```
out.write(header+'\n')
        out.write(seq+'\n')
DNA_bases = 'ATCGNatcgn'
RNA_bases = 'AUCGNaucgn'
Python script:
#!/usr/bin/env python
import argparse
import Bioinfo
from matplotlib import pyplot as plt
import gzip
def get_args():
   parser = argparse.ArgumentParser(description='Takes a zipped fastq file and outputs a png plot of t
   parser.add_argument('-i', '--input',help='Path to input zipped fastq file',required=True)
   parser.add_argument('-o','--output',help='Desired name (not including .png and not including any di
   parser.add_argument('-1','--length',help='Read length', required=True, type=int)
   return parser.parse_args()
args = get_args()
def init_list(lst: list, read_length: int, value: float=0.0) -> list:
    '''This function takes an empty list and will populate it with
   the value passed in "value". If no value is passed, initializes list
   with 101 values of 0.0.'''
   for i in range(read_length):
        lst.append(value)
   return 1st
def populate_list(file, read_length):
   """Takes a FASTQ file and returns a list in which each item is the total sum of Q scores across all
   qscore_list = []
   qscore_list = init_list(qscore_list, read_length)
   with gzip.open(file, 'rt') as fh:
       num_lines = 0
        for line in fh:
           line = line.strip()
            num_lines += 1
            if num_lines % 4 == 0:
                i = 0
                for score in line:
                    qscore_list[i] += Bioinfo.convert_phred(score)
                    i += 1
        i = 0
        for score in qscore_list:
            qscore_list[i] = score/(num_lines/4)
            i +=1
   return qscore_list
```

```
read_list = populate_list(args.input, args.length)
def make_hist(some_list:list, name:str):
   plt.bar(range(len(some_list)), some_list)
   plt.xlabel('# Base Pair')
   plt.ylabel('Mean quality score')
   plt.title(f'Quality Score Distribution in {name}')
   plt.savefig(f'{name}_qscore_distribution.png')
   plt.close()
make_hist(read_list,args.output)
Scripts submitted to Talapas queue:
#!/bin/bash
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00
conda activate bgmp_py39
/usr/bin/time -v ./qual_scores.py -i /projects/bgmp/shared/2017_sequencing/demultiplexed/27_4C_mbnl_S19
exit
#!/bin/bash
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00
conda activate bgmp_py39
/usr/bin/time -v ./qual_scores.py -i /projects/bgmp/shared/2017_sequencing/demultiplexed/27_4C_mbnl_S19
exit
#!/bin/bash
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00
conda activate bgmp_py39
/usr/bin/time -v ./qual_scores.py -i /projects/bgmp/shared/2017_sequencing/demultiplexed/28_4D_mbnl_S20
exit
```

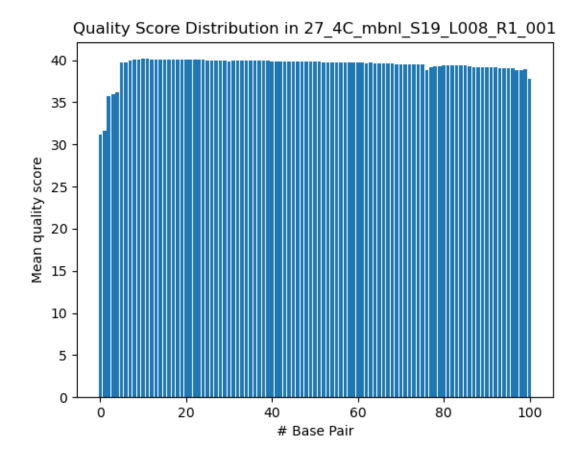
## #!/bin/bash

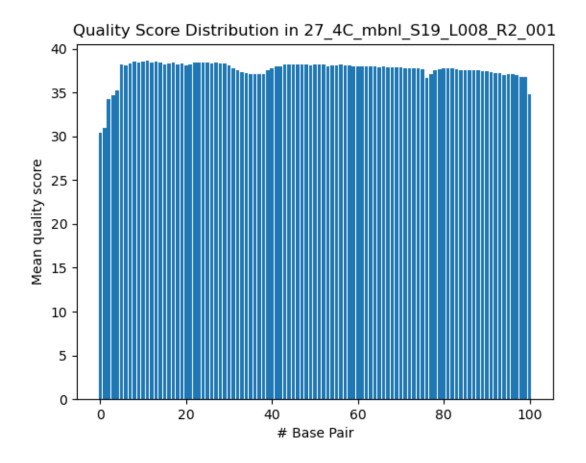
```
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00
```

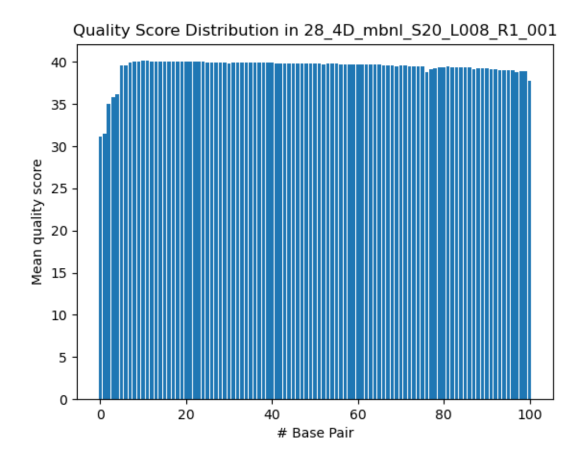
conda activate bgmp\_py39

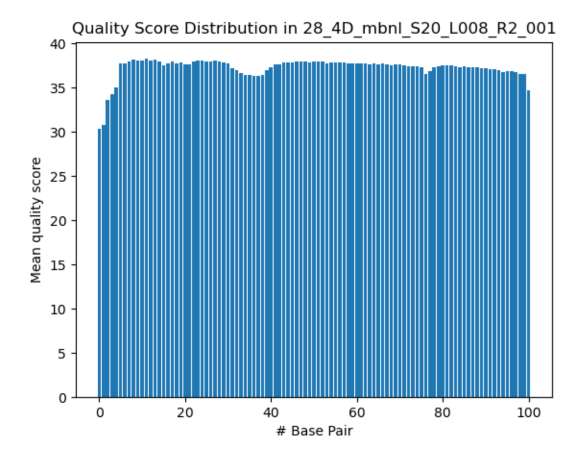
/usr/bin/time -v ./qual\_scores.py -i /projects/bgmp/shared/2017\_sequencing/demultiplexed/28\_4D\_mbnl\_S20 exit

My per-base quality score plots:









My quality score plots appear to match the quality score plots made with data produced by FASTQC. The only difference I can see is that my plots have more data points, because I calculated the mean quality score at each base position, while FASTQC calculated the mean across two bases at a time. This does not affect the overall shape of the quality score distributions.

Overall, the data quality of both libraries looks sufficient for downstream analysis. N-content is low across both reads in both libraries, with only slightly raised values at the beginning of each read. Read 1 quality scores are near 40 across the bulk of the read for both libraries. Read 2 quality scores are slightly lower, closer to 38, but this is expected for libraries sequenced with Illumina, because the sequencing molecule degrades as it sits on the flow cell and is repeatedly washed with harsh chemicals during read 1. FASTQC flags k-mer content and duplication levels as being potential problems for these libraries, but I know that these are RNA-seq libraries and are therefore expected to have differing levels of expression across the genome and to have exact copies of the same transcript due to the overexpression of some transcripts.

### Part 2: Adapter trimming comparison

Adapter sequences were trimmed using cutadapt version 3.4. The adapter sequences were identified by searching for Illumina TruSeq adapter sequences on Illumina's website, and adapter presence and orientation was confirmed using the following bash commands in an interactive session on Talapas:

```
# check that read 1 adapter sequence is in read 1 file
(base) [sgrindst@n225 demultiplexed]$ zcat 27_4C_mbnl_S19_L008_R1_001.fastq.gz | grep -c
AGATCGGAAGAGCACACGTCTGAACTCCAGTCA
132794
# check that read 2 adapter sequence is not in read 1 file
(base) [sgrindst@n225 demultiplexed]$ zcat 27_4C_mbnl_S19_L008_R1_001.fastq.gz | grep -c
```

```
AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT
# check that read 1 adapter sequence is not in read 2 file
(base) [sgrindst@n225 demultiplexed] $ zcat 27 4C mbnl S19 L008 R2 001.fastq.gz | grep -c
AGATCGGAAGAGCACACGTCTGAACTCCAGTCA
# check that read 2 adapter sequence is in read 2 file
(base) [sgrindst@n225 demultiplexed] $ zcat 27_4C_mbnl_S19_L008_R2_001.fastq.gz | grep -c
AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT
135250
# check that reverse complement of read 1 adapter sequence is not in read 1 file
(base) [sgrindst@n225 demultiplexed] $ zcat 27_4C_mbnl_S19_L008_R1_001.fastq.gz | grep -c
TGACTGGAGTTCAGACGTGTGCTCTTCCGATCT
# check that reverse complement of read 2 adapter sequence is not in read 1 file
(base) [sgrindst@n225 demultiplexed] $ zcat 27_4C_mbnl_S19_L008_R1_001.fastq.gz | grep -c
ACACTCTTTCCCTACACGACGCTCTTCCGATCT
# check that reverse complement of read 1 adapter sequence is not in read 1 file
(base) [sgrindst@n225 demultiplexed] $ zcat 27_4C_mbnl_S19_L008_R2_001.fastq.gz | grep -c
TGACTGGAGTTCAGACGTGTGCTCTTCCGATCT
# check that reverse complement of read 2 adapter sequence is not in read 2 file
(base) [sgrindst@n225 demultiplexed] $ zcat 27_4C_mbnl_S19_L008_R2_001.fastq.gz | grep -c
ACACTCTTTCCCTACACGACGCTCTTCCGATCT
# perform all the same checks as above for the other library
(base) [sgrindst@n225 demultiplexed] $ zcat 28_4D_mbnl_S20_L008_R1_001.fastq.gz | grep -c
AGATCGGAAGAGCACACGTCTGAACTCCAGTCA
56758
(base) [sgrindst@n225 demultiplexed] $ zcat 28_4D_mbnl_S20_L008_R1_001.fastq.gz | grep -c
AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT
(base) [sgrindst@n225 demultiplexed] $ zcat 28_4D_mbnl_S20_L008_R2_001.fastq.gz | grep -c
AGATCGGAAGACCACGTCTGAACTCCAGTCA
(base) [sgrindst@n225 demultiplexed] $ zcat 28_4D_mbnl_S20_L008_R2_001.fastq.gz | grep -c
AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT
57626
(base) [sgrindst@n225 demultiplexed] $ zcat 28_4D_mbnl_S20_L008_R1_001.fastq.gz | grep -c
TGACTGGAGTTCAGACGTGTGCTCTTCCGATCT
(base) [sgrindst@n225 demultiplexed] $ zcat 28_4D_mbnl_S20_L008_R1_001.fastq.gz | grep -c
ACACTCTTTCCCTACACGACGCTCTTCCGATCT
(base) [sgrindst@n225 demultiplexed] $ zcat 28_4D_mbnl_S20_L008_R2_001.fastq.gz | grep -c
TGACTGGAGTTCAGACGTGTGCTCTTCCGATCT
(base) [sgrindst@n225 demultiplexed] $ zcat 28_4D_mbnl_S20_L008_R2_001.fastq.gz | grep -c
ACACTCTTTCCCTACACGACGCTCTTCCGATCT
0
```

As expected, the read 1 adapter sequence was only found in the read 1 files, and the read 2 adapter sequence was only found in the read 2 files. The reverse complements of both adapter sequences were not found in

any of the files. Therefore, the adapter sequences that I expect are present in the files that I would expect to find them in.

I then did the following commands to check that the adapter sequences were at the 3' ends of the reads as expected:

```
(base) [sgrindst@n225 demultiplexed] $ zcat 27_4C_mbnl_S19_L008_R1_001.fastq.gz | grep AGATCGGAAGAGCACGTCTGAACTCCAGTCA | head
```

The adapters are toward the 3' end of the reads, as expected. I did the same for the read 2 file with the r2 adapter sequence and got the same results.

The following bash script was submitted to Talapas to run cutadapt:

```
#SBATCH --account=bgmp
```

#!/bin/bash

#SBATCH --partition=bgmp

#SBATCH --cpus-per-task=1

#SBATCH --time=1-0:00:00

conda activate QAA

in\_dir=/projects/bgmp/shared/2017\_sequencing/demultiplexed
out\_dir=/projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/cutadapt

```
/usr/bin/time -v cutadapt -a AGATCGGAAGAGCACACGTCTGAACTCCAGTCA -A AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT \
-o $out_dir/27_4C_mbnl_S19_L008_R1_001.adaptertrimmed.fastq.gz \
-p $out_dir/27_4C_mbnl_S19_L008_R2_001.adaptertrimmed.fastq.gz \
$in dir/27 4C mbnl S19 L008 R1 001.fastq.gz $in dir/27 4C mbnl S19 L008 R2 001.fastq.gz
```

```
/usr/bin/time -v cutadapt -a AGATCGGAAGAGCACACGTCTGAACTCCAGTCA -A AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT \
-o $out_dir/28_4D_mbnl_S20_L008_R1_001.adaptertrimmed.fastq.gz \
-p $out_dir/28_4D_mbnl_S20_L008_R2_001.adaptertrimmed.fastq.gz \
```

exit

Cutadapt reports that for library  $27\_4C\_mbnl\_S19$ , 10.4% of read 1 reads were adapter-trimmed and 11.1% of read 2 reads were adapter-trimmed. For library  $28\_4D\_mbnl\_S20$ , 6.0% of read 1 reads were adapter-trimmed and 6.8% of read 2 reads were adapter-trimmed. This suggests that perhaps library  $28\_4D\_mbnl\_S20$  had a fragment distribution shifted toward longer reads before sequencing.

Trimmomatic version 0.39 was used to quality-trim the reads and was run with the following script on Talapas:

#### #!/bin/bash

```
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00
conda activate QAA
in_dir=/projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/cutadapt
out_dir=/projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/trimmed
/usr/bin/time -v trimmomatic PE -phred33 \
$in_dir/27_4C_mbnl_S19_L008_R1_001.adaptertrimmed.fastq.gz \
\sin_dir/27_4C_mbnl_S19_L008_R2_001.adaptertrimmed.fastq.gz \setminus
$out_dir/27_4C_mbnl_S19_L008_R1_001.trimmed.paired.fastq.gz \
$out_dir/27_4C_mbnl_S19_L008_R1_001.trimmed.unpaired.fastq.gz \
$out dir/27 4C mbnl S19 L008 R2 001.trimmed.paired.fastg.gz \
$out_dir/27_4C_mbnl_S19_L008_R2_001.trimmed.unpaired.fastq.gz \
LEADING:3 TRAILING:3 SLIDINGWINDOW:5:15 MINLEN:35
/usr/bin/time -v trimmomatic PE -phred33 \
$in_dir/28_4D_mbnl_S20_L008_R1_001.adaptertrimmed.fastq.gz \
$in dir/28 4D mbnl S20 L008 R2 001.adaptertrimmed.fastq.gz \
$out dir/28 4D mbnl S20 L008 R1 001.trimmed.paired.fastq.gz \
$out dir/28 4D mbnl S20 L008 R1 001.trimmed.unpaired.fastq.gz \
\ sout_dir/28_4D_mbnl_S20_L008_R2_001.trimmed.paired.fastq.gz \
$out_dir/28_4D_mbnl_S20_L008_R2_001.trimmed.unpaired.fastq.gz \
LEADING:3 TRAILING:3 SLIDINGWINDOW:5:15 MINLEN:35
exit
```

Trimmed read length distribution plots were generated by running FASTQC on the trimmed FASTQs and plotting the data in R. Only paired reads were used for this and all subsequent steps, meaning that the unpaired files output by Trimmomatic were not used.

Script submitted to Talapas to run FASTQC:

### #!/bin/bash

```
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00

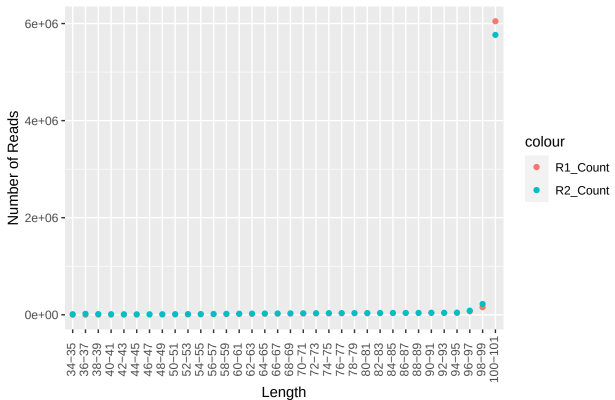
module load fastqc/0.11.5

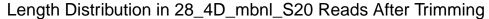
input_dir=/projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/trimmed
output_dir=/projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/trimmed_fastqc_output/
/usr/bin/time -v fastqc \
$input_dir/27_4C_mbnl_S19_L008_R1_001.trimmed.paired.fastq.gz \
$input_dir/27_4C_mbnl_S19_L008_R2_001.trimmed.paired.fastq.gz \
$input_dir/28_4D_mbnl_S20_L008_R1_001.trimmed.paired.fastq.gz \
$input_dir/28_4D_mbnl_S20_L008_R1_001.trimmed.paired.fastq.gz \
$input_dir/28_4D_mbnl_S20_L008_R2_001.trimmed.paired.fastq.gz \
$input_dir/28_4D_mbnl_S20_L008_R1_001.trimmed.paired.fastq.gz \
$input_dir
```

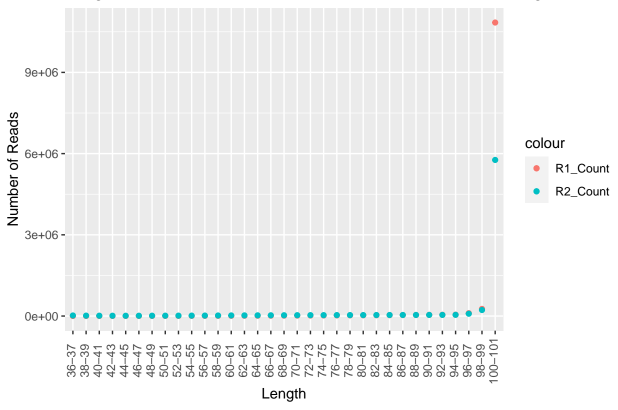
## -o \$output\_dir

exit

# Length Distribution in 27\_4C\_mbnl\_S19 Reads After Trimming







From these plots, we can see that in both libraries read 2 is generally trimmed more often than read 1. We expect the adapter trimming rates to be about the same, since it is dependent on insert length and we expect read 1 and read 2 of the same read pair to have the same insert length. This is shown in the adapter-trimming rates reported by cutadapt, shown above. Therefore, the discrepancy between read 1 and read 2 trimming rates is due to a difference in quality trimming rates. This makes sense because read 2 tends to be of lower quality than read 1, due to the extra time sitting on the flow cell and enduring harsh washes during read 1. This is also reflected in the per-base quality scores, which we already saw to be lower in read 2. The discrepancy between read 1 and read 2 trimming rates is especially pronounced in the second library, 28 4D mbnl S20.

## Part 3: Alignment and strand-specificity

A Mus musculus reference genome was generated from a FASTA file and a GTF file, both downloaded from Ensembl release 104. Both reference genome generation and alignment of the RNA-seq data to the reference database were performed using STAR version 2.7.9a. STAR was run by submitting scripts to Talapas' queuing system.

Genome generation script:

## #!/bin/bash

#SBATCH --account=bgmp #SBATCH --partition=bgmp #SBATCH --cpus-per-task=8 #SBATCH --nodes=1 #SBATCH --time=1-0:00:00

conda activate QAA

```
/usr/bin/time -v STAR \
--runThreadN 8 \
--runMode genomeGenerate \
--genomeDir Mus_musculus.GRCm39.dna.ens104.STAR_2.7.9a \
--genomeFastaFiles /projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/mus/Mus_musculus.GRCm39.dna.primary
--sjdbGTFfile /projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/mus/Mus_musculus.GRCm39.104.gtf
exit
Alignment script:
#!/bin/bash
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=8
#SBATCH --nodes=1
#SBATCH --time=1-0:00:00
conda activate QAA
in_dir=/projects/bgmp/sgrindst/bioinformatics/Bi623/QAA/trimmed
/usr/bin/time -v STAR --runThreadN 8 --runMode alignReads \
--outFilterMultimapNmax 3 \
--outSAMunmapped Within KeepPairs \
--alignIntronMax 1000000 --alignMatesGapMax 1000000 \
--readFilesCommand zcat \
--readFilesIn $in_dir/27_4C_mbnl_S19_L008_R1_001.trimmed.paired.fastq.gz $in_dir/27_4C_mbnl_S19_L008_R2
--genomeDir Mus_musculus.GRCm39.dna.ens104.STAR_2.7.9a \
--outFileNamePrefix 27_4C_mbnl_S19_L008_
/usr/bin/time -v STAR --runThreadN 8 --runMode alignReads \
--outFilterMultimapNmax 3 \
--outSAMunmapped Within KeepPairs \
--alignIntronMax 1000000 --alignMatesGapMax 1000000 \
--readFilesCommand zcat \
--readFilesIn $in_dir/28_4D_mbnl_S20_L008_R1_001.trimmed.paired.fastq.gz $in_dir/28_4D_mbnl_S20_L008_R2
--genomeDir Mus_musculus.GRCm39.dna.ens104.STAR_2.7.9a \
--outFileNamePrefix 28_4D_mbnl_S20_L008_
exit
From STAR's output SAM files, counts of mapped and unmapped reads were obtained by running the
following Python script:
#!/usr/bin/env python
import argparse
import re
def get_args():
```

```
parser = argparse.ArgumentParser(description='Takes SAM as input and returns count of reads mapping
    parser.add_argument('-f', '--file', help='input SAM file')
    return parser.parse_args()
args = get_args()
mapped_sum = 0 #counts number of unique aligned reads
unmapped_sum = 0 #counts number of unique unaligned reads
with open(args.file, 'r') as fh:
    for line in fh:
        line=line.strip()
        if line[0] == '@': #ignore header section
        flag = int(re.sub(r'[^t]+\t([^t]+).*',r'\1',line)) #extract bitwise flag
        if((flag & 4) != 4 and (flag & 256 != 256)):
            mapped_sum += 1
        if((flag & 4) == 4 \text{ and } (flag & 256 != 256)):
            unmapped_sum += 1
print('Number of mapped reads: ', mapped_sum)
print('Number of unmapped reads: ', unmapped_sum)
The above Python script was run on Talapas with this wrapper script:
#!/bin/bash
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00
#determine the number of reads mapping and not mapping to reference
/usr/bin/time -v /projects/bgmp/sgrindst/bioinformatics/Bi621/PS/ps8-sally-grindstaff/sam_mapping_count
-f aligned/27_4C_mbnl_S19_L008_Aligned.out.sam > aligned/27_4C_mbnl_S19_L008_mapped_counts.txt
/usr/bin/time -v /projects/bgmp/sgrindst/bioinformatics/Bi621/PS/ps8-sally-grindstaff/sam_mapping_count
-f aligned/28_4D_mbnl_S20_L008_Aligned.out.sam > aligned/28_4D_mbnl_S20_L008_mapped_counts.txt
For library 27_4C_mbnl_S19,
Number of mapped reads: 13320042
Number of unmapped reads: 433870
For library 28_4D_mbnl_S20,
Number of mapped reads: 22657659
Number of unmapped reads: 793147
Before running HTSeq-Count, SAMTools version 1.13 was used to convert SAMs to BAMs, sort by name,
and convert back to SAMs (as recommended by HTSeq-Count) with the following script:
#!/bin/bash
#SBATCH --account=bgmp
```

```
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=8
#SBATCH --nodes=1
#SBATCH --time=1-0:00:00
#sort sam files by name in order to run HTSeq-count later
conda activate QAA
/usr/bin/time -v samtools view -u aligned/27_4C_mbnl_S19_L008_Aligned.out.sam | \
samtools sort -n | \
samtools view -h -o aligned/27_4C_mbnl_S19_L008_Aligned.sorted.sam
/usr/bin/time -v samtools view -u aligned/28_4D_mbnl_S20_L008_Aligned.out.sam | \
samtools sort -n | \
samtools view -h -o aligned/28_4D_mbnl_S20_L008_Aligned.sorted.sam
exit
Then HTSeq-Count version 0.13.5 was run twice per library (once with the stranded parameter set to no
and once with the stranded parameter set to yes) with this script:
#!/bin/bash
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=8
#SBATCH --nodes=1
#SBATCH --time=1-0:00:00
conda activate QAA
/usr/bin/time -v htseq-count -s yes \
aligned/27_4C_mbnl_S19_L008_Aligned.sorted.sam mus/Mus_musculus.GRCm39.104.gtf \
> aligned/27_4C_mbnl_S19_L008.stranded.genecount
/usr/bin/time -v htseq-count -s no \
aligned/27_4C_mbnl_S19_L008_Aligned.sorted.sam mus/Mus_musculus.GRCm39.104.gtf \
> aligned/27_4C_mbnl_S19_L008.unstranded.genecount
/usr/bin/time -v htseq-count -s yes \
aligned/28_4D_mbnl_S20_L008_Aligned.sorted.sam mus/Mus_musculus.GRCm39.104.gtf \
> aligned/28_4D_mbnl_S20_L008.stranded.genecount
/usr/bin/time -v htseq-count -s no \
aligned/28_4D_mbnl_S20_L008_Aligned.sorted.sam mus/Mus_musculus.GRCm39.104.gtf \
> aligned/28_4D_mbnl_S20_L008.unstranded.genecount
exit
HTSeq-Count results:
## # A tibble: 55,420 x 2
```

```
'4'
##
      ENSMUSG0000000001
##
                          <dbl>
      <chr>>
    1 ENSMUSG00000000003
##
                              0
    2 ENSMUSG00000000028
                              6
##
    3 ENSMUSG00000000031
                              0
    4 ENSMUSG0000000037
                              0
##
    5 ENSMUSG00000000049
                             16
##
    6 ENSMUSG00000000056
                              3
##
    7 ENSMUSG00000000058
                              2
                              3
##
    8 ENSMUSG00000000078
    9 ENSMUSG0000000085
                              7
   10 ENSMUSG00000000088
                              4
   # ... with 55,410 more rows
   # A tibble: 55,420 x 2
##
      ENSMUSG0000000001 '1372'
##
      <chr>
                           <dbl>
##
    1 ENSMUSG0000000003
                               0
                             652
##
    2 ENSMUSG00000000028
##
    3 ENSMUSG00000000031
                               0
    4 ENSMUSG00000000037
                               0
    5 ENSMUSG0000000049
##
                               0
    6 ENSMUSG0000000056
                             164
##
    7 ENSMUSG00000000058
                             547
    8 ENSMUSG00000000078
                            1065
##
    9 ENSMUSG0000000085
                             341
## 10 ENSMUSG00000000088
                             314
## # ... with 55,410 more rows
## # A tibble: 55,420 x 2
                            '5'
##
      ENSMUSG0000000001
##
      <chr>
                          <dbl>
    1 ENSMUSG00000000003
##
                              0
    2 ENSMUSG00000000028
                              4
##
##
    3 ENSMUSG0000000031
                              0
    4 ENSMUSG0000000037
                              0
##
    5 ENSMUSG0000000049
                             21
                              4
##
    6 ENSMUSG0000000056
##
    7 ENSMUSG00000000058
                              1
##
    8 ENSMUSG00000000078
                              2
    9 ENSMUSG00000000085
                              4
  10 ENSMUSG00000000088
                              4
## # ... with 55,410 more rows
  # A tibble: 55,420 x 2
      ENSMUSG0000000001 '2901'
##
                           <dbl>
##
      <chr>
##
    1 ENSMUSG00000000003
                               0
##
    2 ENSMUSG00000000028
                             988
##
    3 ENSMUSG0000000031
                               0
                               0
##
    4 ENSMUSG0000000037
##
    5 ENSMUSG00000000049
                               0
    6 ENSMUSG00000000056
                             305
```

```
## 7 ENSMUSG00000000058 698
## 8 ENSMUSG00000000078 1387
## 9 ENSMUSG0000000085 408
## 10 ENSMUSG00000000088 400
## # ... with 55,410 more rows
```

For each HTSeq-Count run, the number of reads mapping to features and the total number of reads were analyzed with the following bash script:

```
#!/bin/bash
#SBATCH --account=bgmp
#SBATCH --partition=bgmp
#SBATCH --cpus-per-task=1
#SBATCH --time=1-0:00:00
#count the number of reads mapping to a feature and the total reads (using htseq-count output) in order
#determine whether the data is stranded or unstranded
#commands are taken from my ICA4 GitHub markdown file from Bi621
echo '27_4C_mbnl_S19_L008.stranded.genecount:' > aligned/htseq_count_stats.txt
echo 'reads mapping to feature:' >> aligned/htseq_count_stats.txt
grep -v '^_' aligned/27_4C_mbnl_S19_L008.stranded.genecount | awk '{sum+=$2}END{print sum}' >> aligned/
echo 'total reads:' >> aligned/htseq_count_stats.txt
awk '{sum+=$2}END{print sum}' aligned/27_4C_mbnl_S19_L008.stranded.genecount >> aligned/htseq_count_sta
echo '' >> aligned/htseq_count_stats.txt
echo '27_4C_mbnl_S19_L008.unstranded.genecount:' >> aligned/htseq_count_stats.txt
echo 'reads mapping to feature:' >> aligned/htseq_count_stats.txt
grep -v '^_' aligned/27_4C_mbnl_S19_L008.unstranded.genecount | awk '{sum+=$2}END{print sum}' >> aligne
echo 'total reads:' >> aligned/htseq_count_stats.txt
awk '{sum+=$2}END{print sum}' aligned/27 4C mbnl S19 L008.unstranded.genecount >> aligned/htseq count s
echo '' >> aligned/htseq_count_stats.txt
echo '28_4D_mbnl_S20_L008.stranded.genecount:' >> aligned/htseq_count_stats.txt
echo 'reads mapping to feature:' >> aligned/htseq_count_stats.txt
grep -v '^_' aligned/28_4D_mbnl_S20_L008.stranded.genecount | awk '{sum+=$2}END{print sum}' >> aligned/
echo 'total reads:' >> aligned/htseq_count_stats.txt
awk '{sum+=$2}END{print sum}' aligned/28_4D_mbnl_S20_L008.stranded.genecount >> aligned/htseq_count_sta
echo '' >> aligned/htseq_count_stats.txt
echo '28_4D_mbnl_S20_L008.unstranded.genecount:' >> aligned/htseq_count_stats.txt
echo 'reads mapping to feature:' >> aligned/htseq_count_stats.txt
grep -v '^_' aligned/28_4D_mbnl_S20_L008.unstranded.genecount | awk '{sum+=$2}END{print sum}' >> aligne
echo 'total reads:' >> aligned/htseq_count_stats.txt
awk '{sum+=$2}END{print sum}' aligned/28_4D_mbnl_S20_L008.unstranded.genecount >> aligned/htseq_count_s
exit
```

27\_4C\_mbnl\_S19\_L008.stranded.genecount: reads mapping to feature: 271747

total reads: 6876956

27\_4C\_mbnl\_S19\_L008.unstranded.genecount: reads mapping to feature: 5545724
total reads: 6876956

28\_4D\_mbnl\_S20\_L008.stranded.genecount: reads mapping to feature: 424716
total reads: 11725403

28\_4D\_mbnl\_S20\_L008.unstranded.genecount: reads mapping to feature: 9414809 total reads: 11725403

Converting these numbers to percentages, we get:  $27\_4C\_mbnl\_S19\_L008$ , stranded – 3.9515594% of reads mapping to features

 $27\_4C\_mbnl\_S19\_L008$ , unstranded -80.6421329% of reads mapping to features

 $28\_4D\_mbnl\_S20\_L008$ , stranded -3.6221868% of reads mapping to features

28 4D mbnl S20 L008, unstranded – 80.2941187% of reads mapping to features

From these data, I propose that the libraries were prepared using a stranded method. I believe this to be true because if the libraries were unstranded, then I would expect about half as many reads to map to features with the stranded parameter set to 'yes' than with the stranded parameter set to 'no'. This is because in an unstranded library reads 1 and 2 can both correspond to either the template or coding strand of the genome, so I would not expect a bias. However, in this case I see an order of magnitude difference (3.9515594% vs. 80.6421329% for the first library and 3.6221868% vs. 80.2941187% for the second library) when I set the stranded parameter to yes vs. no in HTSeq-Count. This indicates that most of the mapping features are coming from one strand over the other, which leads me to believe that the libraries were prepared in a strand-specific manner.