Assignment - QAA

Angela Crabtree

9/1/2021

```
library(ggplot2)
library(tidyr)
```

Part 1 – Read quality score distributions

Samples

- $2_2B_control$
- Undetermined

Running FastQC

```
/usr/bin/time -v fastqc \
    -o . \
    /projects/bgmp/shared/2017_sequencing/demultiplexed/2_2B_control_S2_L008_R1_001.fastq.gz

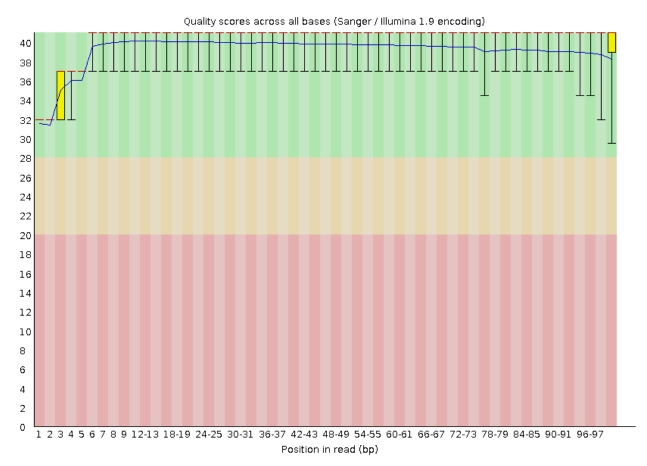
/usr/bin/time -v fastqc \
    -o . \
    /projects/bgmp/shared/2017_sequencing/demultiplexed/2_2B_control_S2_L008_R2_001.fastq.gz

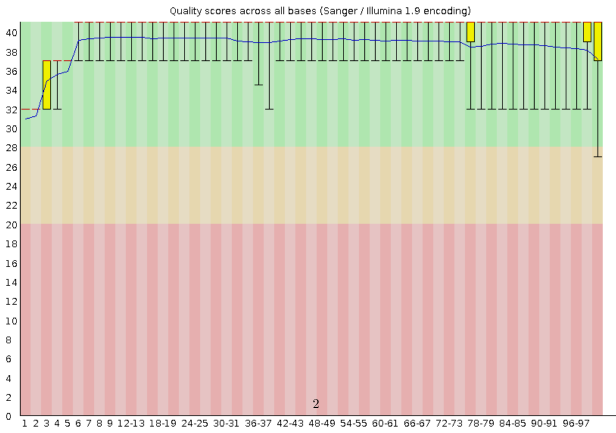
/usr/bin/time -v fastqc \
    -o . \
    /projects/bgmp/shared/2017_sequencing/demultiplexed/Undetermined_S0_L008_R1_001.fastq.gz

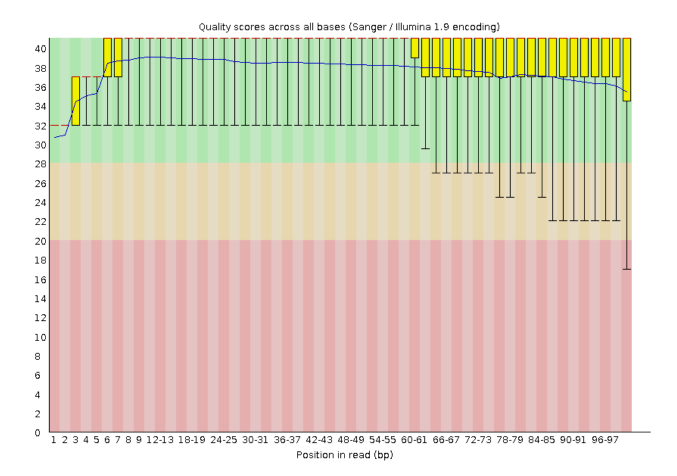
/usr/bin/time -v fastqc \
    -o . \
    /projects/bgmp/shared/2017_sequencing/demultiplexed/Undetermined_S0_L008_R1_001.fastq.gz

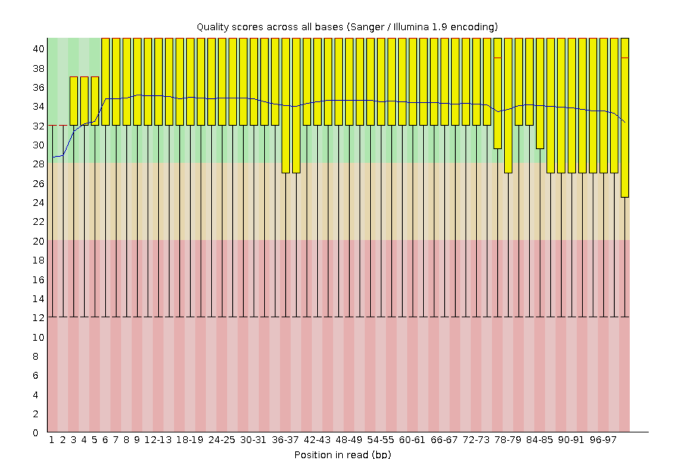
/usr/bin/time -v fastqc \
    -o . \
    /projects/bgmp/shared/2017_sequencing/demultiplexed/Undetermined_S0_L008_R2_001.fastq.gz
```

${\bf Quality \ score \ distributions}$









Per-base N content

The per-base N content is consistent with the quality score distributions.

Running in-house script for histogram generation

```
/usr/bin/time -v ./histograms.py \
-f /projects/bgmp/shared/2017_sequencing/demultiplexed/2_2B_control_S2_L008_R1_001.fastq.gz \
-r 101 -o ../img/2_2B_control_R1_hist.png

/usr/bin/time -v ./histograms.py \
-f /projects/bgmp/shared/2017_sequencing/demultiplexed/2_2B_control_S2_L008_R2_001.fastq.gz \
-r 101 -o ../img/2_2B_control_R2_hist.png

/usr/bin/time -v ./histograms.py \
-f /projects/bgmp/shared/2017_sequencing/demultiplexed/Undetermined_S0_L008_R1_001.fastq.gz \
-r 101 -o ../img/Undetermined_R1_hist.png

/usr/bin/time -v ./histograms.py \
-f /projects/bgmp/shared/2017_sequencing/demultiplexed/Undetermined_S0_L008_R2_001.fastq.gz \
-r 101 -o ../img/Undetermined_R2_hist.png
```

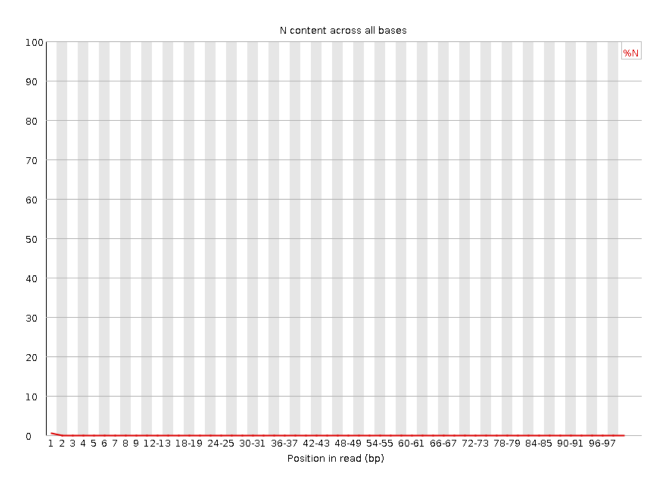


Figure 1: 2_2B_control R1

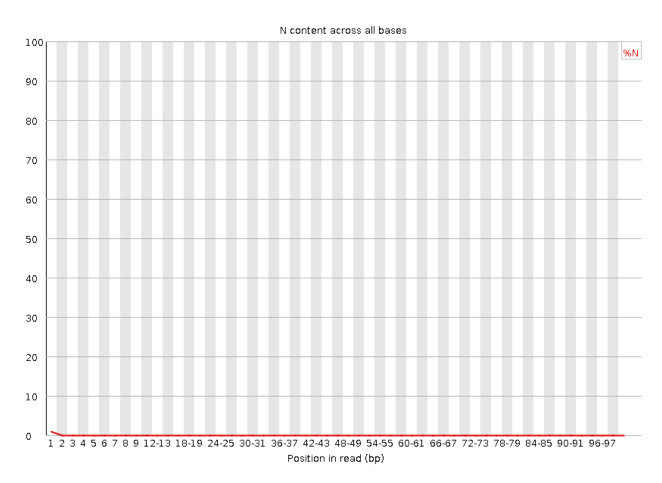


Figure 2: 2_2B_control R2

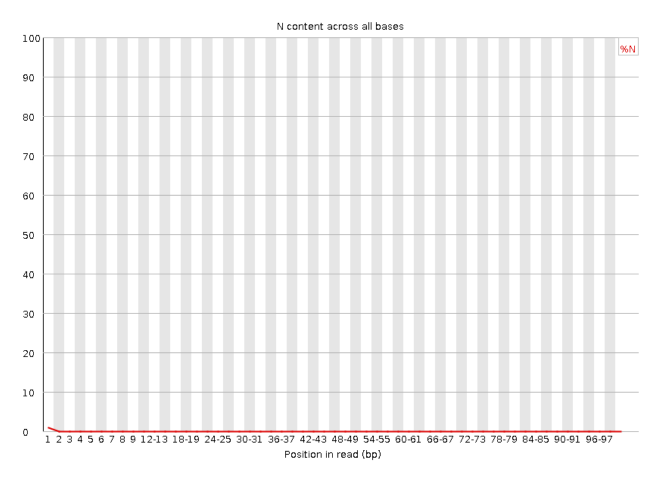


Figure 3: Undetermined R1

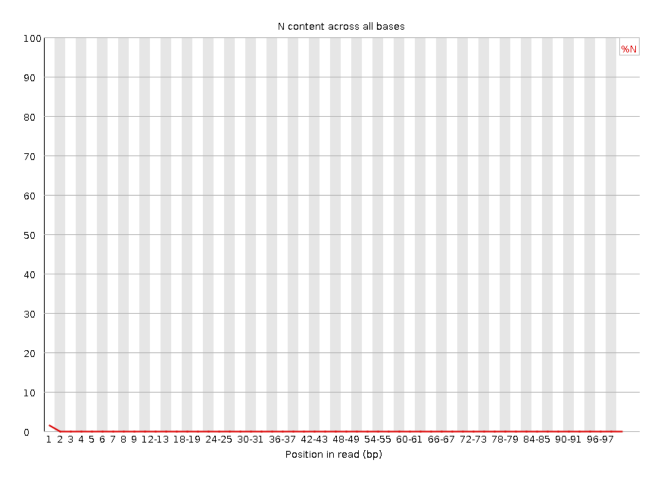


Figure 4: Undetermined R2

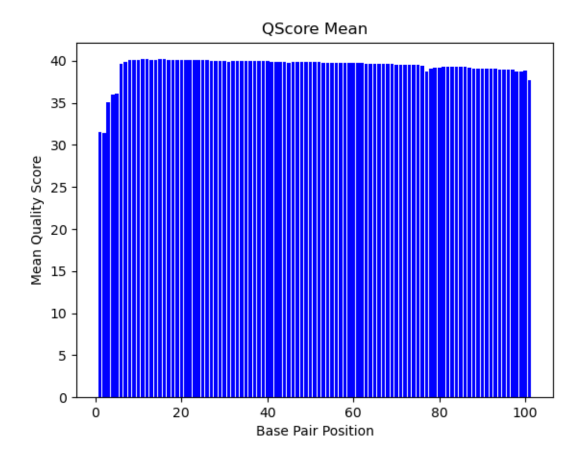


Figure 5: 2_2B_control R1

These quality score distributions generated from the in-house script are very similar to the plots generated by FastQC. The in-house python script ran much slower than FastQC because FastQC is written in java which is a compiled language and is therefore more efficient to run than python (an interpreted language).

The overall data quality of the 2_2B_control library is great, while the quality of the Undetermined library is worse but probably acceptable for most applications. R2 reads of the Undetermined library were quite poor.

Part 2 – Adaptor trimming comparison

Illumina TruSeq Adapter Sequences found here

Running Cutadapt

cutadapt -a AGATCGGAAGAGCACACGTCTGAACTCCAGTCA -A AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT \
 -o 2_2B_control.1.fastq.gz \

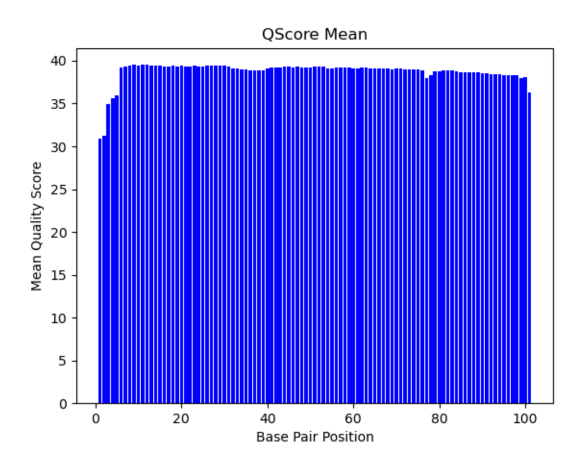


Figure 6: 2_2B_control R2

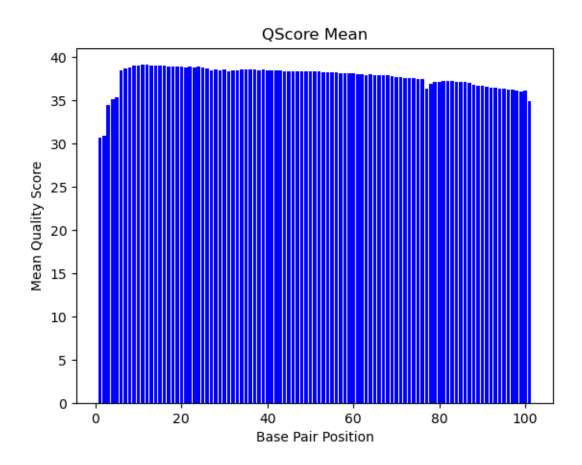


Figure 7: Undetermined R1

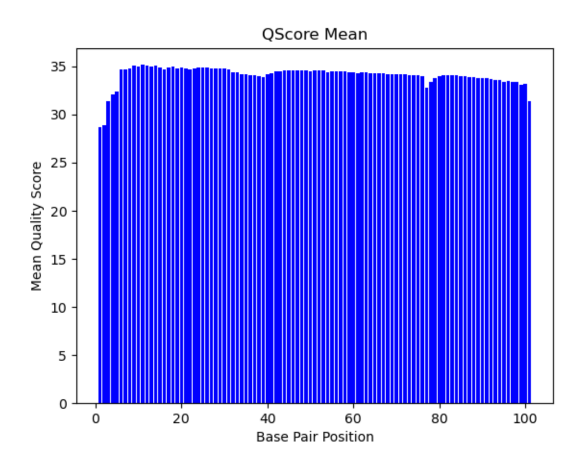


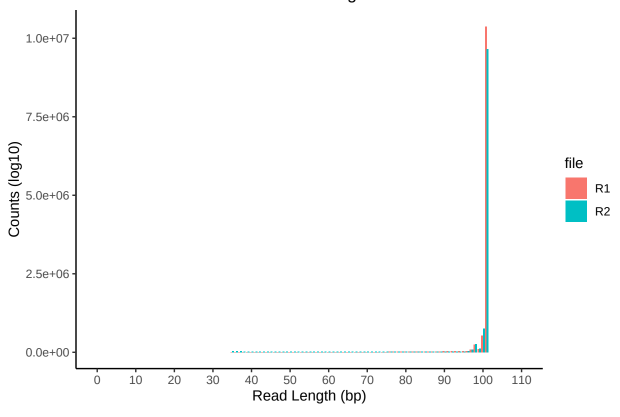
Figure 8: Undetermined R2

```
-p 2_2B_control.2.fastq.gz \
  /projects/bgmp/shared/2017_sequencing/demultiplexed/2_2B_control_S2_L008_R1_001.fastq.gz \
  /projects/bgmp/shared/2017 sequencing/demultiplexed/2 2B control S2 L008 R2 001.fastq.gz
cutadapt -a AGATCGGAAGAGCACACGTCTGAACTCCAGTCA -A AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT \
  -o Undetermined.1.fastq.gz \
  -p Undetermined.2.fastq.gz \
  /projects/bgmp/shared/2017_sequencing/demultiplexed/Undetermined_SO_L008_R1_001.fastq.gz \
  /projects/bgmp/shared/2017_sequencing/demultiplexed/Undetermined_SO_L008_R2_001.fastq.gz
Running Trimmomatic
trimmomatic PE -threads 4 \
  ../cutadapt/2_2B_control.1.fastq.gz \
  ../cutadapt/2_2B_control.2.fastq.gz \
  2_2B_control.1.trim.fastq.gz 2_2B_control.1.untrim.fastq.gz \
  2_2B_control.2.trim.fastq.gz 2_2B_control.2.untrim.fastq.gz \
  LEADING:3 \
  TRAILING:3 \
  SLIDINGWINDOW:5:15 \
  MINLEN:35
trimmomatic PE -threads 4 \
  ../cutadapt/Undetermined.1.fastq.gz \
  ../cutadapt/Undetermined.2.fastq.gz \
  Undetermined.1.trim.fastq.gz Undetermined.1.untrim.fastq.gz \
  Undetermined.2.trim.fastq.gz Undetermined.2.untrim.fastq.gz \
  LEADING:3 \
  TRAILING:3 \
  SLIDINGWINDOW:5:15 \
  MINLEN:35
Outputting Distribution Data for R
zcat 2_2B_control.1.trim.fastq.gz | awk 'NR % 4 == 2 { print length() }' | sort -n | uniq -c >
  2_2B_control_R1_distr.txt
zcat 2_2B_control.2.trim.fastq.gz | awk 'NR % 4 == 2 { print length() }' | sort -n | uniq -c >
  2 2B control R2 distr.txt
zcat Undetermined.1.trim.fastq.gz | awk 'NR % 4 == 2 { print length() }' | sort -n | uniq -c >
  Undetermined_R1_distr.txt
zcat Undetermined.2.trim.fastq.gz | awk 'NR % 4 == 2 { print length() }' | sort -n | uniq -c >
  Undetermined_R2_distr.txt
r1 = read.table("data/Undetermined_R1_distr.txt", header = F, sep="")
r2 = read.table("data/Undetermined_R2_distr.txt", header = F, sep="")
r1$file = "R1"
r2$file = "R2"
r1r2 = rbind(r1, r2)
colnames(r1r2)=c("rcount", "rlength", "file")
```

head(r1r2)

```
##
     rcount rlength file
## 1
       2617
                 35
                      R1
## 2
       2799
                 36
                      R1
## 3
       2979
                 37
                      R1
## 4
       3086
                 38
                      R1
## 5
       3253
                 39
                      R1
## 6
       3223
                 40
ggplot(r1r2, aes(x=rlength, y=rcount, fill=file)) +
 geom_bar(stat="identity", position=position_dodge()) +
 theme_classic() +
  scale_x_continuous(breaks = seq(0, 110, by=10), limits=c(0,110)) +
 labs(title="Undetermined Reads After Trimming",
       x="Read Length (bp)", y="Counts (log10)", color="Read File")
```

Undetermined Reads After Trimming

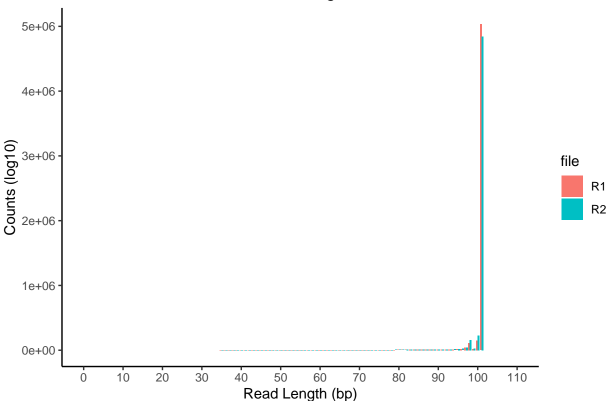


```
r1 = read.table("data/2_2B_control_R1_distr.txt", header = F, sep="")
r2 = read.table("data/2_2B_control_R2_distr.txt", header = F, sep="")
r1$file = "R1"
r2$file = "R2"
r1r2 = rbind(r1, r2)
colnames(r1r2)=c("rcount","rlength","file")
head(r1r2)
```

```
## rcount rlength file
## 1 723 35 R1
```

```
## 2
         826
                    36
                         R1
## 3
         890
                    37
                         R1
## 4
         942
                    38
                         R1
        1010
## 5
                    39
                         R1
## 6
        1148
                    40
                         R1
```

2_2B_control Reads After Trimming



R2 reads should be of lower quality than R1 reads because they are sequenced after many cycles have already been completed on the sequencer, and after there has been another round of bridge amplification. This is reflected in the distribution of read lengths after trimming, above.

Part 3 – Alignment and strand-specificity

SLURM: STAR alignment & HTSeq

```
#!/bin/bash
#SBATCH --partition=bgmp ### Partition (like a queue in PBS)
#SBATCH --job-name=qaa ### Job Name
```

```
#SBATCH --output=star-%j.log
                                  ### File in which to store job output
#SBATCH --error=star-%j.err
                                  ### File in which to store job error messages
#SBATCH --time=0-20:00:00
                                  ### Wall clock time limit in Days-HH:MM:SS
#SBATCH --nodes=1
                                  ### Number of nodes needed for the job
#SBATCH --cpus-per-task=8
                                  ### Number of CPUs to be used per task
#SBATCH --account=bgmp
                                  ### Account used for job submission
#SBATCH --mail-user=acrabtre@uoregon.edu
                                          ### email for job submission notifications
#SBATCH --mail-type=ALL
                                  ### specifies types of notification emails to send
## load conda environment
conda activate gaa
## assign variables
d=/projects/bgmp/acrabtre/bioinfo/Bi623/QAA/star
f_read=/projects/bgmp/acrabtre/bioinfo/Bi623/QAA/trim/Undetermined.1.trim.fastq.gz
r_read=/projects/bgmp/acrabtre/bioinfo/Bi623/QAA/trim/Undetermined.2.trim.fastq.gz
prfx="Undetermined_Mmus_"
samfile=$d/${prfx}Aligned.out.sam
cd $d
## run STAR to generate genome indexes
/usr/bin/time -v STAR \
   --runThreadN 8 \
   --runMode genomeGenerate \
   --genomeDir $d/mmus \
   --genomeFastaFiles $d/mmus/Mus_musculus.GRCm39.dna.primary_assembly.fa \
   --sjdbGTFfile $d/mmus/Mus_musculus.GRCm39.104.gtf
## run STAR to assemble reads to reference genome
/usr/bin/time -v STAR --runThreadN 8 --runMode alignReads \
   --outFilterMultimapNmax 3 \
   --outSAMunmapped Within KeepPairs \
   --alignIntronMax 1000000 --alignMatesGapMax 1000000 \
   --readFilesCommand zcat \
   --readFilesIn $f_read $r_read \
   --genomeDir $d/mmus \
   --outFileNamePrefix $d/$prfx
## run python script to get counts of mapped and unmapped reads
python /projects/bgmp/acrabtre/bioinfo/Bi623/QAA/scripts/inspector sam.py $samfile >
 $d/${prfx}map_counts.txt
## run HTseq to count reads that map to features
htseq-count --stranded=no $samfile $d/mmus/Mus_musculus.GRCm39.104.gtf > $d/${prfx}hts.genecounts
htseq-count --stranded=yes $samfile $d/mmus/Mus_musculus.GRCm39.104.gtf > $d/${prfx}hts_str.genecounts
## assign variables
d=/projects/bgmp/acrabtre/bioinfo/Bi623/QAA/star
f_read=/projects/bgmp/acrabtre/bioinfo/Bi623/QAA/trim/2_2B_control.1.trim.fastq.gz
```

r_read=/projects/bgmp/acrabtre/bioinfo/Bi623/QAA/trim/2_2B_control.2.trim.fastq.gz

prfx="2_2B_control_Mmus_"

```
samfile=$d/${prfx}Aligned.out.sam
## run STAR to assemble reads to reference genome
/usr/bin/time -v STAR --runThreadN 8 --runMode alignReads \
    --outFilterMultimapNmax 3 \
    --outSAMunmapped Within KeepPairs \
    --alignIntronMax 1000000 --alignMatesGapMax 1000000 \
    --readFilesCommand zcat \
    --readFilesIn $f read $r read \
    --genomeDir $d/mmus \
    --outFileNamePrefix $d/$prfx
## run python script to get counts of mapped and unmapped reads
python /projects/bgmp/acrabtre/bioinfo/Bi623/QAA/scripts/inspector_sam.py $samfile >
  $d/${prfx}map_counts.txt
## run HTseq to count reads that map to features
htseq-count --stranded=no $samfile $d/mmus/Mus musculus.GRCm39.104.gtf > $d/${prfx}hts.genecounts
htseq-count --stranded=yes $samfile $d/mmus/Mus_musculus.GRCm39.104.gtf > $d/${prfx}hts_str.genecounts
Read counts for each genecounts file (use to determine if library was stranded):
$ cat 2_2B_control_hts_counts.txt | grep -v "^__" | awk '{sum+=$2} END {print sum}'
4702902
$ cat 2_2B_control_Mmus_hts_counts_str.txt | grep -v "^__" | awk '{sum+=$2} END {print sum}'
216262
$ cat Undetermined_Mmus_hts_counts.txt | grep -v "^__" | awk '{sum+=$2} END {print sum}'
$ cat Undetermined_Mmus_hts_counts_str.txt | grep -v "^__" | awk '{sum+=$2} END {print sum}'
299593
Total number of reads respectively for each sample file:
$ cat 2_2B_control_hts.genecounts | awk '{sum+=$2} END {print sum}'
5652541
$ cat Undetermined_Mmus_hts_counts.txt | awk '{sum+=$2} END {print sum}'
12160073
htseq-count -stranded=yes
  • 2 2B control: htseq mapped ~3.8% of total reads to reference genome
  • Undetermined: htseq mapped \sim 2.5\% of total reads to reference genome
```

 $htseq\text{-}count \ -stranded = no$

- 2 2B control: htseq mapped ~83% of total reads to reference genome
- Undetermined: htseq mapped $\sim 52\%$ of total reads to reference genome

I propose this library is stranded because the number of mapped reads (in gene counts files) after running HTSeq are extremely low. If the kit was non-stranded, the counts should be about half of the stranded=no counts. For example, in the "Undetermined" sample, only 2.5% of the reads mapped to the reference genome stranded=yes, whereas 52% of the reads were mapped when stranded=no. If we ran the stranded=reverse option, we would expect to find the missing $\sim 50\%$ because only the reverse complement of the reads are able to the reference genome.

2 2B control

mapped reads: 11078823 (98%)umapped reads: 226259 (2%)

Undetermined

mapped reads: 15584518 (64%)umapped reads: 8735628 (36%)

Additionally, the number of mapped and unmapped reads were calculated from the SAM file for both libraries (above). The proportion of mapped read counts from htseq-count non-stranded option more closely refected the proportion of mapped read counts calculated directly from the SAM file.