QAA

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Part 1 - Read quality score distributions

Files - referred to as 'Fox' and 'Control' files throughout report

Fox files

- $\bullet \ /projects/bgmp/shared/2017_sequencing/demultiplexed/31_4F_fox_S22_L008_R1_001.fastq.gz$
- $\bullet \ /projects/bgmp/shared/2017_sequencing/demultiplexed/31_4F_fox_S22_L008_R2_001.fastq.gz \\$

Control files

- $\bullet \ / projects/bgmp/shared/2017_sequencing/demultiplexed/23_4A_control_S17_L008_R1_001.fastq.gz$
- $\bullet \ / projects/bgmp/shared/2017_sequencing/demultiplexed/23_4A_control_S17_L008_R2_001.fastq.gz$

1. Fastqc run

```
# load fastqc module
module load fastqc/0.11.5
```

Fastqc graphs

Fox files

```
fastqc /projects/bgmp/shared/2017_sequencing/demultiplexed/31_4F_fox_S22_L008_R1_001.fastq.gz \
    /projects/bgmp/shared/2017_sequencing/demultiplexed/31_4F_fox_S22_L008_R2_001.fastq.gz \
    -o fox \
    -t 8
```

Read 1:

Read 2:

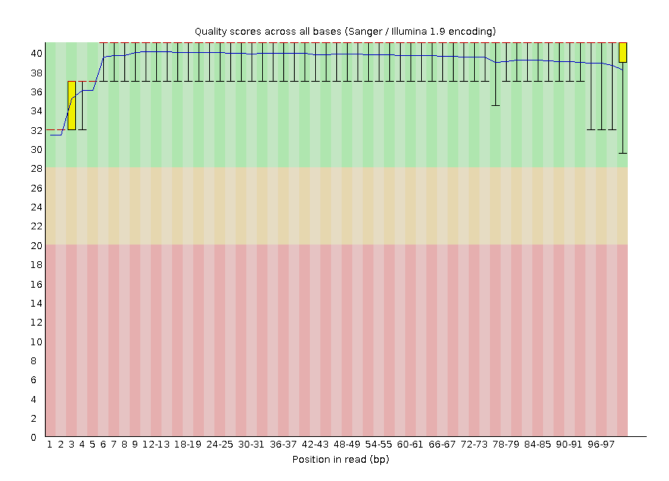


Figure 1: Per-base quality content '31_4F_fox_S22_L008_R1_001'

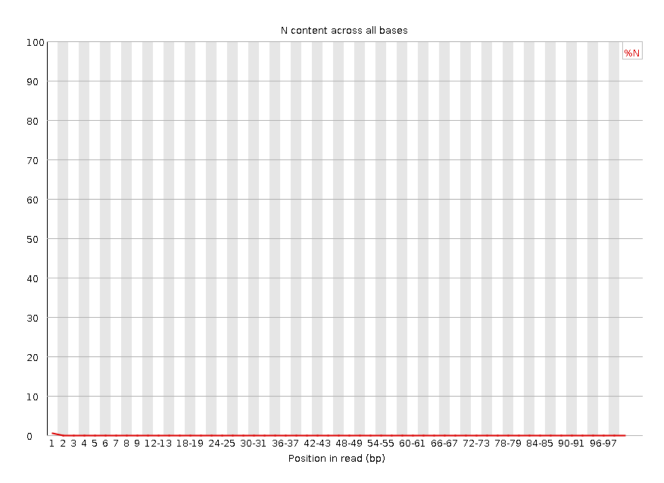


Figure 2: Per-base N content '31_4F_fox_S22_L008_R1_001'

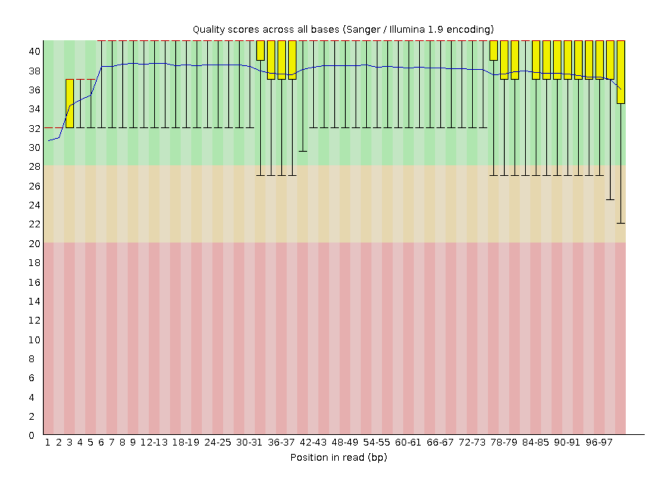


Figure 3: Per-base quality content '31_4F_fox_S22_L008_R2_001'

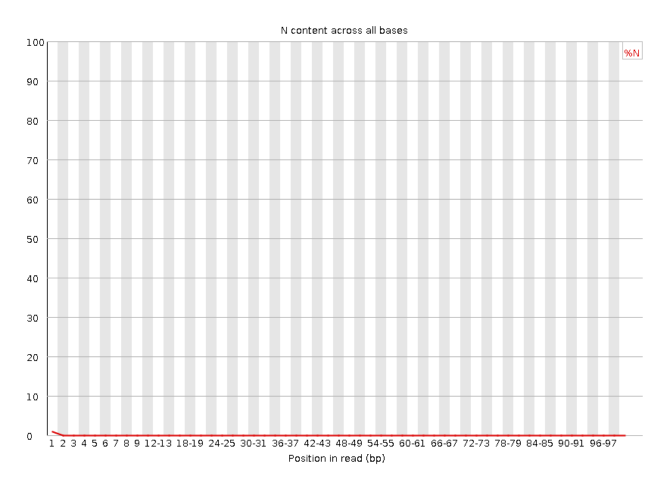


Figure 4: Per-base N content '31_4F_fox_S22_L008_R2_001'

Control files

fastqc /projects/bgmp/shared/2017_sequencing/demultiplexed/23_4A_control_S17_L008_R1_001.fastq.gz \
 /projects/bgmp/shared/2017_sequencing/demultiplexed/23_4A_control_S17_L008_R2_001.fastq.gz \
 -o control \
 -t 8

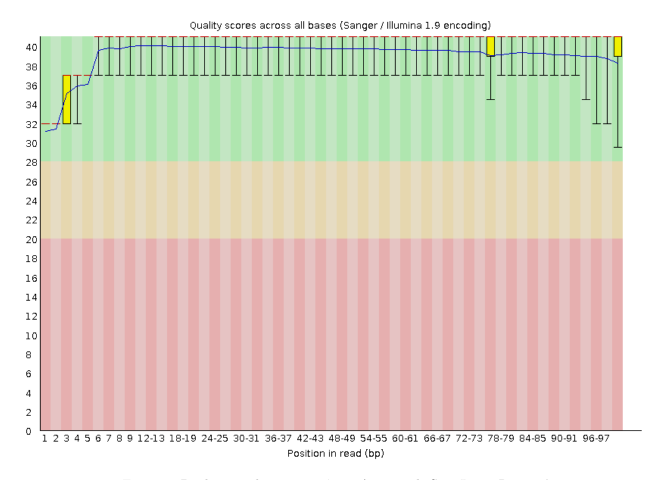


Figure 5: Per-base quality content '23_4A_control_S17_L008_R1_001'

Read 1:

Read 2: The per-base N quality graphs are consistent with the per-base quality graphs in that base 1 has higher N content relative to the other bases. This is in alignment with the lower per-base quality score relative to the other bases.

2. Histogram generator

Fox files

Read 1

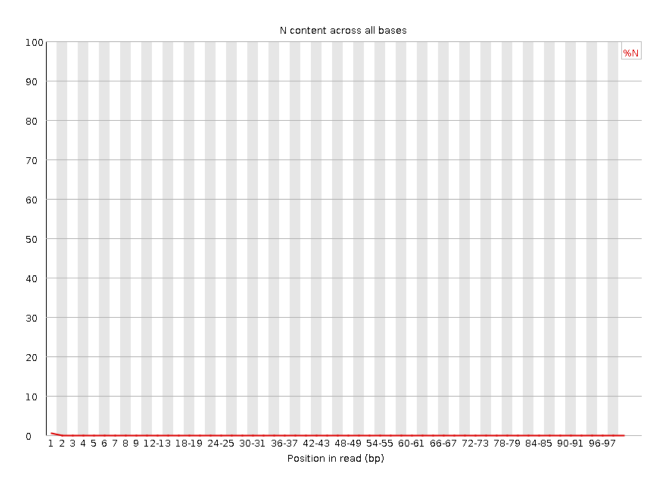


Figure 6: Per-base N content '23_4A_control_S17_L008_R1_001'

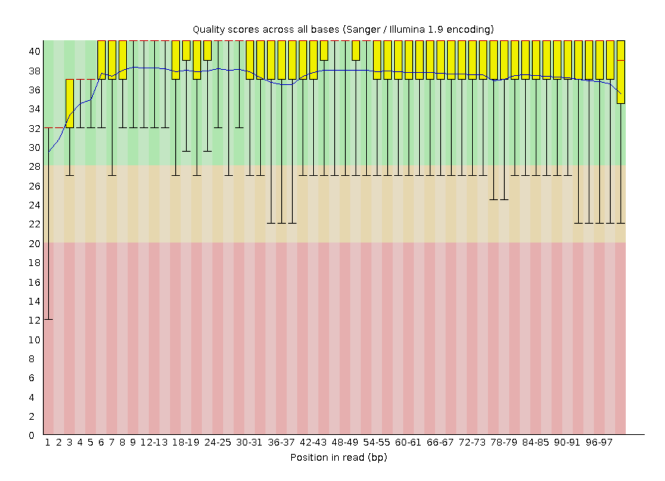


Figure 7: Per-base quality content '23_4A_control_S17_L008_R2_001'

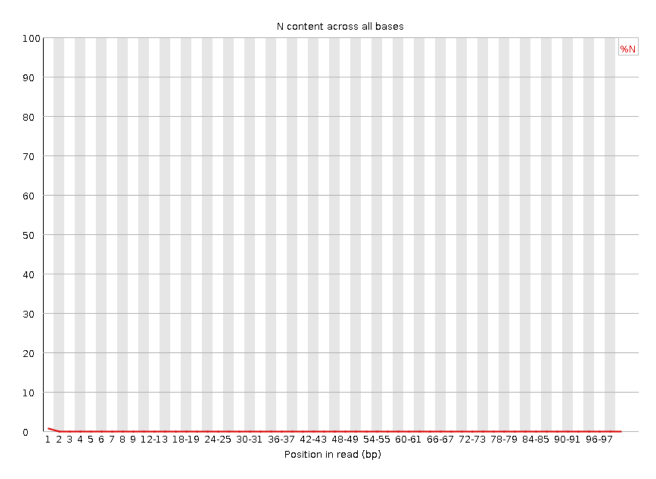


Figure 8: Per-base N content '23_4A_control_S17_L008_R2_001'

```
/projects/bgmp/jadler2/bioinfo/Bi623/QAA/python_scripts/histogram_generator.py \
    -f /projects/bgmp/shared/2017_sequencing/demultiplexed/31_4F_fox_S22_L008_R1_001.fastq.gz \
    -1 101 \
    -n 3788343 \
    -o images/31_4F_fox_S22_L008_R1_001.histogram.png \
    -t "Fox Read 1"
```

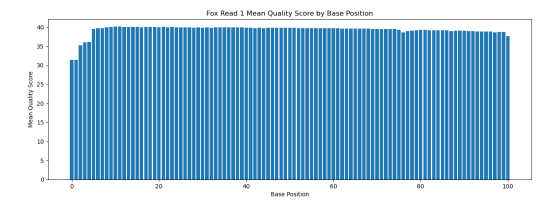


Figure 9: Figure: Distribution of lengths for untrimmed reads '31_4F_fox_S22_L008_R1_001'

Read 2

```
/projects/bgmp/jadler2/bioinfo/Bi623/QAA/python_scripts/histogram_generator.py \
    -f /projects/bgmp/shared/2017_sequencing/demultiplexed/31_4F_fox_S22_L008_R2_001.fastq.gz \
    -l 101 \
    -n 3788343 \
    -o images/31_4F_fox_S22_L008_R2_001.histogram.png \
    -t "Fox Read 2"
```

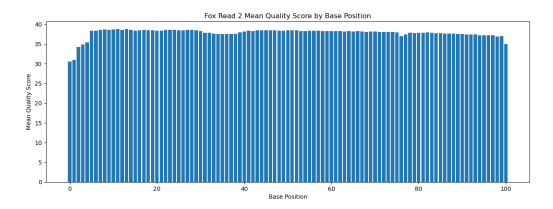


Figure 10: Figure: Distribution of lengths for untrimmed reads '31_4F_fox_S22_L008_R2_001'

Control files

Read 1

```
/projects/bgmp/jadler2/bioinfo/Bi623/QAA/python_scripts/histogram_generator.py \
-f /projects/bgmp/shared/2017_sequencing/demultiplexed/23_4A_control_S17_L008_R1_001.fastq.gz \
-l 101 \
-n 44303262 \
-o images/23_4A_control_S17_L008_R1_001.histogram.png \
-t "Control Read 1"
```

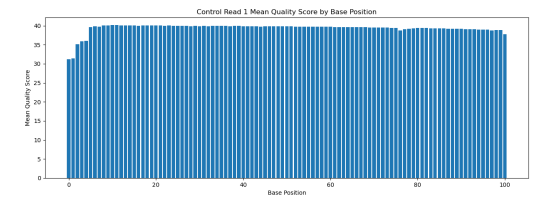


Figure 11: Figure: Distribution of lengths for untrimmed reads '23_4A_control_S17_L008_R1_001'

Read 2

```
/projects/bgmp/jadler2/bioinfo/Bi623/QAA/python_scripts/histogram_generator.py \
-f /projects/bgmp/shared/2017_sequencing/demultiplexed/23_4A_control_S17_L008_R2_001.fastq.gz \
-l 101 \
-n 44303262 \
-o images/23_4A_control_S17_L008_R2_001.histogram.png \
-t "Control Read 2"
```



Figure 12: Figure: Distribution of lengths for untrimmed reads '23_4A_control_S17_L008_R2_001'

Yes, the output and runtimes do differ. The fastqc graphs include box and whiskers to demonstrate the range of values found at each base. The fastqc charts also include green (28-41), yellow (20-28), and red (0-20) areas that correspond to quality level. This is helpful in providing the viewer with a quick reference of quality level at each base.

Fox files

The histogram generator runs for the fox files each took about 3 minutes 45 seconds and only produced the one histogram. The fastqc runs can be run with multiple CPUs using the -t flag. Running the two files with -t 8 results in total runtime of 29 seconds and produces substantially more information relative to our histogram generator script.

Control files

The histogram generator runs for the control files each took about 40 minutes. Running fastqc on the two controls files with 8 CPUs result sin total runtime of 3 minutes 45 seconds and produces substantially more information relative to our histogram generator script.

3. Comment on the overall data quality of the two libraries

Fox files The quality of each of the read files is high. Neither was flagged for poor quality. The mean score is in the green for each of the read files. The 31_4F_fox_S22_L008_R2_001.fastq.gz file does have some whiskers that drop down into the 'yellow' zone (less than 28 mean quality score), but the mean and majority quality scores hover between 34-41.

Control files The quality for each of the read files is high, though R2 does have quite a few whiskers extending into the yellow range and one whisker that extends into the red range. The mean quality score across all bases are all in the green, though the 1st base mean quality is around 30, the lowest mean quality score of any of the four files we processed.

Part 2 - Adaptor trimming comparison

4. New conda env, install new packages

```
# create new conda env
conda create --name QAA python=3.9
conda activate QAA
# install new packages
conda install cutadapt
conda install Trimmomatic
```

Version of cutadapt and Trimmomatic

```
cutadapt = 3.4Trimmomatic = 0.39
```

5. Adapter trim with cutadapt

Fox files

```
cutadapt -a AGATCGGAAGAGCACACGTCTGAACTCCAGTCA \
   -A AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT \
```

```
-o /projects/bgmp/jadler2/bioinfo/Bi623/QAA/trimmed_31_4F_fox_S22_L008_R1_001.fastq.gz \ -p /projects/bgmp/jadler2/bioinfo/Bi623/QAA/trimmed_31_4F_fox_S22_L008_R2_001.fastq.gz \ /projects/bgmp/shared/2017_sequencing/demultiplexed/31_4F_fox_S22_L008_R1_001.fastq.gz \ /projects/bgmp/shared/2017_sequencing/demultiplexed/31_4F_fox_S22_L008_R2_001.fastq.gz
```

Control files

cutadapt -a AGATCGGAAGAGCACACGTCTGAACTCCAGTCA \

- -A AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTG \
- -o /projects/bgmp/jadler2/bioinfo/Bi623/QAA/trimmed 23 4A control S17 L008 R1 001.fastq.gz \
- -p /projects/bgmp/jadler2/bioinfo/Bi623/QAA/trimmed_23_4A_control_S17_L008_R2_001.fastq.gz \ /projects/bgmp/shared/2017_sequencing/demultiplexed/23_4A_control_S17_L008_R1_001.fastq.gz \ /projects/bgmp/shared/2017_sequencing/demultiplexed/23_4A_control_S17_L008_R2_001.fastq.gz

Confirm adapter sequences were removed

```
zcat trimmed_31_4F_fox_S22_L008_R1_001.fastq.gz | grep "AGATCGGAAGAGCACACGTCTGAACTCCAGTCA"

zcat trimmed_31_4F_fox_S22_L008_R2_001.fastq.gz | grep "AGATCGGAAGAGCGTCGTGTAGGGAAGAGTGT"

zcat trimmed_23_4A_control_S17_L008_R1_001.fastq.gz | grep "AGATCGGAAGAGCACACGTCTGAACTCCAGTCA"

zcat trimmed_23_4A_control_S17_L008_R2_001.fastq.gz | grep "AGATCGGAAGAGCGTCGTGTAGGGAAGAGTGT"
```

Results of adapter check:

All returns to the above commands returned nothing, confirming that adapter sequences are not present in the adapter-trimmed files.

Reasoning for choosing commands for adapter check:

I chose to use these commands because I want to confirm that the adapter sequence was removed from the reads and the trimmed reads output to a new file.

What proportion of reads were trimmed?

Fox files Total read pairs processed: 3,788,343

Read 1 with adapter: 456,168 (12.0%) Read 2 with adapter: 482,503 (12.7%)

Control files Total read pairs processed: 44,303,262

Read 1 with adapter: 1,359,563 (3.1%) Read 2 with adapter: 1,657,295 (3.7%)

6. Trimmomatic runs

Fox files

```
java -jar /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trimmomatic-0.39/trimmomatic-0.39.jar PE \
   -threads 8 \
   /projects/bgmp/jadler2/bioinfo/Bi623/QAA/adapter_trimmed/trimmed_31_4F_fox_S22_L008_R1_001.fastq.gz \
   /projects/bgmp/jadler2/bioinfo/Bi623/QAA/adapter_trimmed/trimmed_31_4F_fox_S22_L008_R2_001.fastq.gz \
   -baseout /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_31_4F_fox_S22_L008_001.f
   LEADING:3 \
   TRAILING:3 \
   SLIDINGWINDOW:5:15 \
   MINLEN:35
```

Control files

```
java -jar /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trimmomatic-0.39/trimmomatic-0.39.jar PE \
   -threads 8 \
   /projects/bgmp/jadler2/bioinfo/Bi623/QAA/adapter_trimmed/trimmed_23_4A_control_S17_L008_R1_001.fastq.
   /projects/bgmp/jadler2/bioinfo/Bi623/QAA/adapter_tirmmed/trimmed_23_4A_control_S17_L008_R2_001.fastq.
   -baseout /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_23_4A_control_S17_L008_R
   LEADING:3 \
   TRAILING:3 \
   SLIDINGWINDOW:5:15 \
   MINLEN:35
```

7. Distribution of length of trimmed sequences in each read for each group

Fox reads

```
./python_scripts/dual_histogram.py \
-i1 /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_31_4F_fox_S22_L008_001_1P.fas
-n1 3597908 \
-i2 /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_31_4F_fox_S22_L008_001_2P.fas
-n2 3597908 \
-o images/trimmed_lengths_fox.histogram.png \
-t "Fox Read 1, Read 2 Distribution of Trimmed Lengths"
```

Control reads

```
./python_scripts/dual_histogram.py \
-i1 /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_23_4A_control_S17_L008_R1_001
-n1 42056563 \
-i2 /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_23_4A_control_S17_L008_R1_001
-n2 42056563 \
-o images/trimmed_lengths_control.histogram.png \
-t "Control Read 1, Read 2 Distribution of Trimmed Lengths"
```

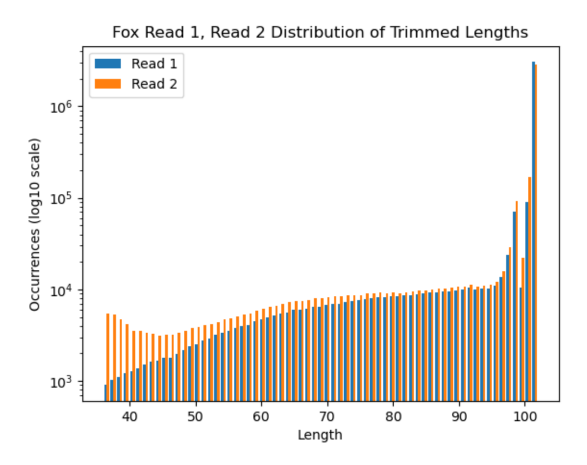


Figure 13: **Figure**: Distribution of lengths of '31_4F_fox_S22_L008_R1_001' and '31_4F_fox_S22_L008_R2_001' reads following adapter and quality trimming.

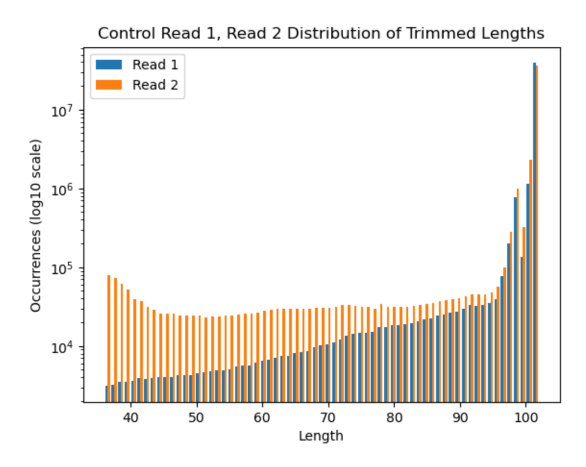


Figure 14: **Figure**: Distribution of lengths of '23_4A_control_S17_L008_R1_001' and '23_4A_control_S17_L008_R2_001' reads following adapter and quality trimming.

R1 or R2 adapter-trimmed at different rates?

I would expect R2 to be adapter-trimmed at a higher rate. Because this read sits on the sequencer for a longer period of time, the strand is exposed to a greater amount of harsh chemicals and there are more opportunities for errors to be made, including erroneous adapter binding.

Part 3 - Alignment and strand-specificity

8. Install new packages

```
conda install star -c bioconda
conda install numpy
conda install pysam
conda install matplotlib
pip install HTSeq
```

9. STAR mouse database creation and mapping

Grab files from ensembl and construct database

```
# wget files
wget http://ftp.ensembl.org/pub/release-104/fasta/mus_musculus/dna/Mus_musculus.GRCm39.dna.primary_asser
wget http://ftp.ensembl.org/pub/release-104/gtf/mus_musculus/Mus_musculus.GRCm39.104.gtf.gz

# generate mouse db
STAR --runThreadN 8
    --runMode genomeGenerate
    --genomeDir /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.dna.ens.104.STAR_2.7.9a
    --genomeFastaFiles /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.dna.primary_assembly.:
    --sjdbGTFfile /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.104.gtf
```

Align quality trimmed reads

Fox reads

```
STAR --runThreadN 8 \
--runMode alignReads \
--outFilterMultimapNmax 3 \
--outSAMunmapped Within KeepPairs \
--alignIntronMax 1000000 \
--alignMatesGapMax 1000000 \
--readFilesCommand zcat \
--readFilesIn /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_31_4F_fox_S22_L008_v
/projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_31_4F_fox_S22_L008_v
--genomeDir /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.dna.ens.104.STAR_2.7.9a \
--outFileNamePrefix ./align/fox/
```

Control reads

```
STAR --runThreadN 8 \
--runMode alignReads \
--outFilterMultimapNmax 3 \
--outSAMunmapped Within KeepPairs \
--alignIntronMax 1000000 \
--alignMatesGapMax 1000000 \
--readFilesCommand zcat \
--readFilesIn /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_23_4A_control_S17_L
/projects/bgmp/jadler2/bioinfo/Bi623/QAA/Trim_Reads/quality_trimmed_23_4A_control_S17_L008_R1_001_2P.
--genomeDir //projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.dna.ens.104.STAR_2.7.9a \
--outFileNamePrefix ./align/control/
```

10. Mapped vs. unmapped reads

Fox files

```
# determine mapped vs unmapped
/projects/bgmp/jadler2/bioinfo/Bi623/QAA/python_scripts/sam_filter.py \
   -f /projects/bgmp/jadler2/bioinfo/Bi623/QAA/align/fox/Aligned.out.sam \
   -o /projects/bgmp/jadler2/bioinfo/Bi623/QAA/mapping_summaries/map_v_unmap_fox.txt"
```

Summary of sam file: /projects/bgmp/jadler2/bioinfo/Bi623/QAA/align/fox/Aligned.out.sam

mapped reads: 6969878 unmapped reads: 225938

Control files

```
/projects/bgmp/jadler2/bioinfo/Bi623/QAA/python_scripts/sam_filter.py \
-f /projects/bgmp/jadler2/bioinfo/Bi623/QAA/align/control/Aligned.out.sam \
-o /projects/bgmp/jadler2/bioinfo/Bi623/QAA/mapping_summaries/map_v_unmap_control.txt
```

Summary of sam file: /projects/bgmp/jadler2/bioinfo/Bi623/QAA/align/control/Aligned.out.sam

mapped reads: 79473045 unmapped reads: 4640081

11. Count reads that map to features using htseq-count

Stranded

Fox files

```
htseq-count -q \
  -f sam \
  -s yes \
  /projects/bgmp/jadler2/bioinfo/Bi623/QAA/align/fox/Aligned.out.sam \
  /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.104.gtf
```

Control files

```
htseq-count -q \
  -f sam \
  -s yes \
  /projects/bgmp/jadler2/bioinfo/Bi623/QAA/align/control/Aligned.out.sam \
  /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.104.gtf
```

Unstranded

Fox files

```
htseq-count -q \
  -f sam \
  -s no \
  /projects/bgmp/jadler2/bioinfo/Bi623/QAA/align/fox/Aligned.out.sam \
  /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.104.gtf
```

Control files

```
htseq-count -q \
  -f sam \
  -s no \
  /projects/bgmp/jadler2/bioinfo/Bi623/QAA/align/control/Aligned.out.sam \
  /projects/bgmp/jadler2/bioinfo/Bi623/QAA/Mus_musculus.GRCm39.104.gtf
```

12. Demonstrate whether data are strand-specific RNA-seq libraries or not

Determine number of mapped reads in each file:

```
for f in $(ls); do echo $f; grep -v '^_' $f | awk '{sum+=$2} END {print sum}'; done
control_stranded.genecount
1516759

control_unstranded.genecount
31747007

fox_stranded.genecount
180499

fox_unstranded.genecount
2922418
```

Determine total reads in each file:

```
for f in $(1s); do echo $f; cat $f | awk '{sum+=$2} END {print sum}'; done
control_stranded.genecount
42056563
```

control_unstranded.genecount
42056563

fox_stranded.genecount
3597908

fox_unstranded.genecount
3597908

Determine percentage of reads mapped:

control_stranded.genecount
1516759 / 42056563 = 3.6%

control_unstranded.genecount
31747007 / 42056563 = 75.0%

fox_stranded.genecount
180499 / 3597908 = 5.0%

fox_unstranded.genecount
2922418 / 3597908 = 81.0%

Based on the information above, I propose the data are from unstranded library preps because only 3.6% and 5.0% of the stranded control and fox reads, respectively, map to the mouse genome, whereas, 75.0% and 81.0% of the unstranded control and fox reads, respectively, map to the mouse genome.