SF-seq

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SF-seq read quality score distributions

In this assignment I worked with the following read files from /projects/bgmp/2017_sequencing/demultiplexed:

```
2_2B_control_S2_L008_R1_001.fastq.gz
2_2B_control_S2_L008_R2_001.fastq.gz
29_4E_fox_S21_L008_R1_001.fastq.gz
29_4E_fox_S21_L008_R2_001.fastq.gz
```

To determine quality score distributions for each read the program FastQC was loaded on talapas along with it's dependencies: ml easybuild intel/2017a FastQC

In a working directory called SF-seq, a new directory was created for the output of FastQC. FastQC was then run using the following options on all four files.

```
$ fastqc --noextract -o fastqc_out -t 4 \
2_2B_control_S2_L008_R1_001.fastq.gz 2_2B_control_S2_L008_R2_001.fastq.gz \
29_4E_fox_S21_L008_R1_001.fastq.gz 29_4E_fox_S21_L008_R2_001.fastq.gz
```

Results

Quality score distributions for the forward and reverse reads as well as per-base N content of both treatments were extracted from the FastQC output and compared to result from my quality score distribution plots.

The quality score distributions for the treatment 29_4E_fox_S21 show high average quality scores (approching 40) for read one with lower quality scores at the begining and slowly tapering towards the end of the read (Figure 1). Read two shows a lower quality score across all the bases, not exceeding 38, with low quality at the start and end of the read (Figures 2). The percent N-content across the bases for reads one and two show the highest percent N-content at the first base position for both reads, and undetectable N-content over the remaining base positions (Figures 3, 4). The detectable N-content at the first base position explains the low quality start in the quality score distributuions because an 'N' at that position will decrease the average quality score at that base position. The plots for the treatment 2_2B_control show a similar pattern where read two shows a lower average quality score than read one, and the low quality score at the begining of the read is reflected in the percent N-content of each read at base position one (Figure 5-8).

I then ran my quality score distribution script qscore_dist2.py on the files using the following slurm script:

```
#!/bin/bash
#SBATCH --partition=fat
                                   ### partition
#SBATCH --job-name=QD
                          ### Job Name
#SBATCH --output=QD.out
                           ### File in which to store job output
#SBATCH --error=QD.err
                           ### File in which to store job error messages
#SBATCH --time=0-24:00:00
                                    ### Wall clock time limit in Days-HH:MM:SS
#SBATCH --nodes=1
                                    ### Node count required for the job
#SBATCH --ntasks-per-node=28
                                    ### Nuber of tasks to be launched per Node
#SBATCH --mail-user=jvancamp@uoregon.edu
                                                ### Notifyme
#SBATCH --mail-type=ALL
                                                ### All of it
```

load the necessary modules

```
# run qscore_dist2.py on all files
time ./qscore_dist2.py -f ~/SF-seq/data/2_2B_control_S2_L008_R1_001.fastq
# run qscore_dist2.py on all files
time ./qscore_dist2.py -f ~/SF-seq/data/2_2B_control_S2_L008_R2_001.fastq
# run qscore_dist2.py on all files
time ./qscore_dist2.py -f ~/SF-seq/data/29_4E_fox_S21_L008_R1_001.fastq
# run qscore_dist2.py on all files
time ./qscore_dist2.py -f ~/SF-seq/data/29_4E_fox_S21_L008_R2_001.fastq
The error file from the script held the time of each run, showing that each run took less than 20 minutes,
while the total runtime was just over one hour.
$ cat QD.err
The following have been reloaded with a version change:
  1) zlib/1.2.8 => zlib/1.2.11
        18m24.967s
real
        18m21.792s
user
sys 0m1.324s
        18m28.796s
real
        18m26.856s
user
sys 0m1.334s
real
        15m24.851s
        15m23.165s
user
sys 0m1.163s
real
        15m16.293s
```

ml zlib/1.2.11 python3/3.6.1

15m14.693s

user

sys 0m1.070s

The quality score distribution plots from my script are shown in Figures 9-12. These distributuions show similar patterns as the results from the FastQC output. Read two for both treatments shows a lower average quality score for more base positions than in read one, and the quality score drop-off toward the beginning and end of the read is similar to the results found in the FastQC output as expected.

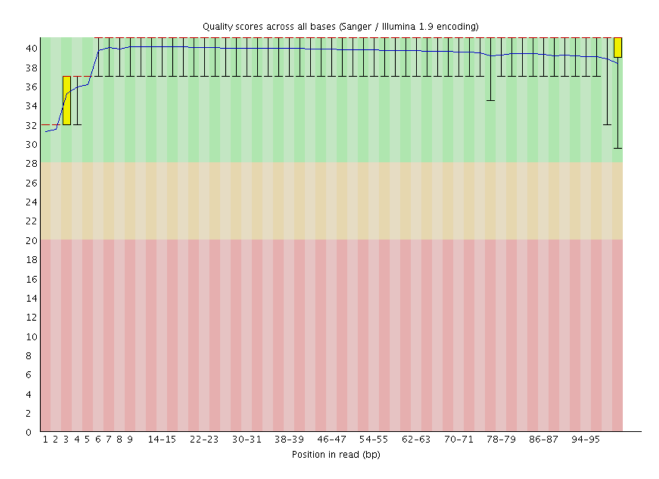


Figure 1: $29_4E_fox_S21$ Read1 QScore distribution

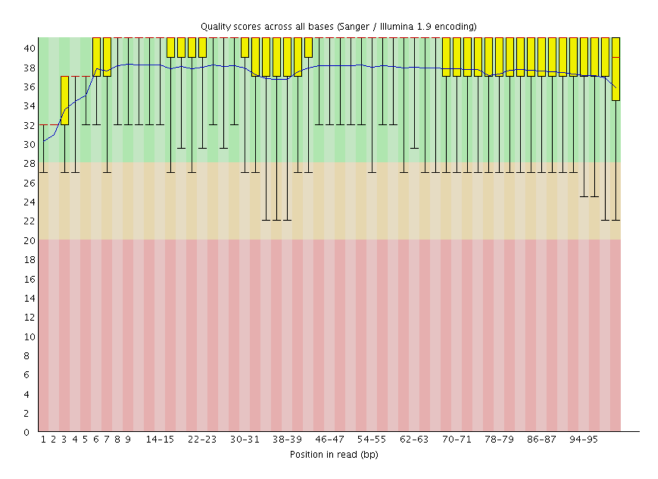


Figure 2: $29_4E_fox_S2$ Read2 QScore distribution

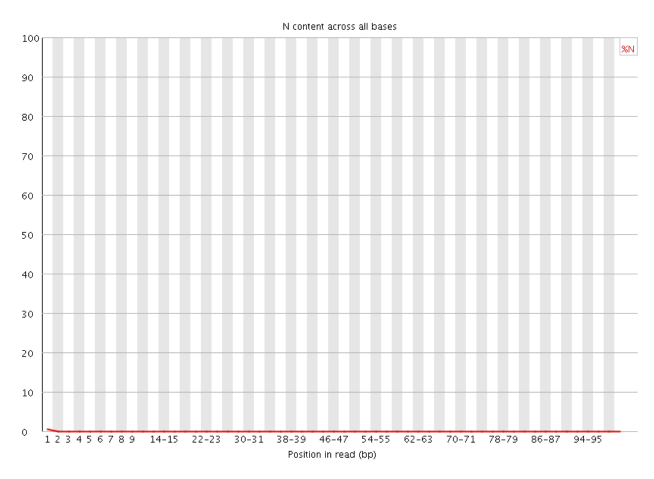


Figure 3: $29_4E_fox_S2$ Read 1 Per-base N-content

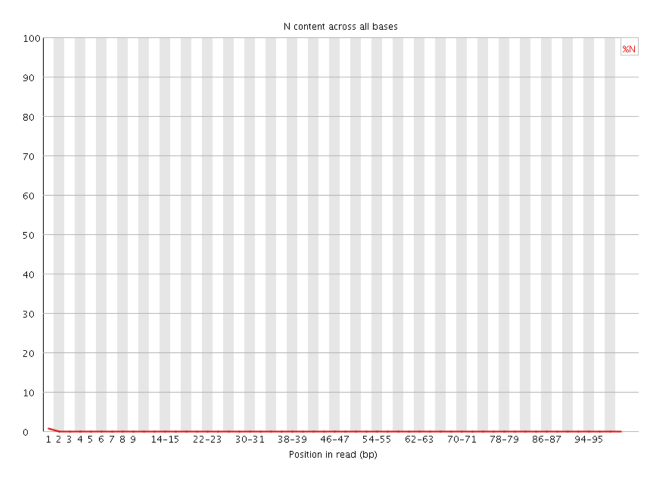


Figure 4: 29_4E_fox_S2 Read 2 Per-base N-content

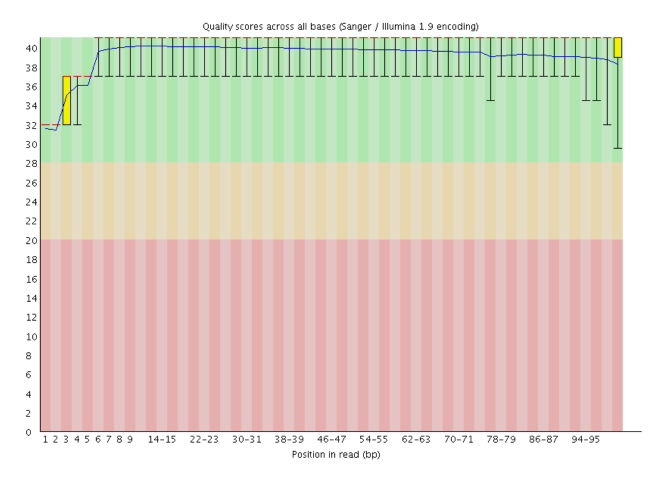


Figure 5: 2_2B_control_S2 Read1 QScore distribution

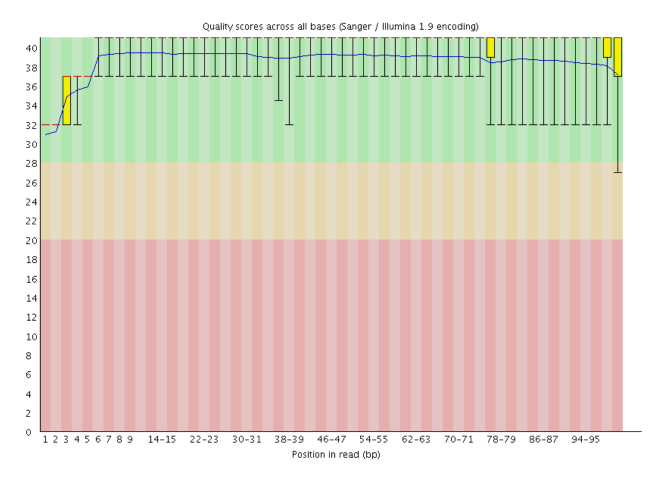


Figure 6: 2_2B_control_S2 Read2 QScore distribution

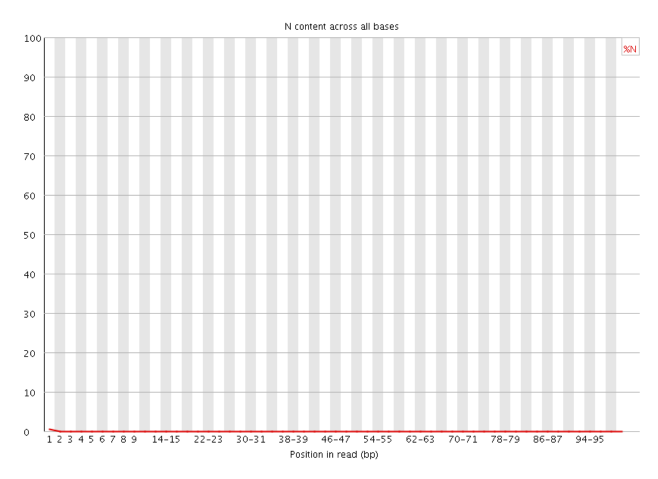


Figure 7: 2_2B_control_S2 Per-base N-content

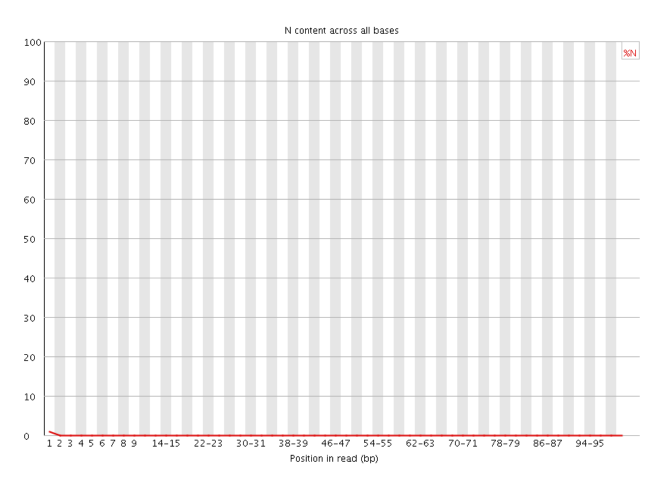


Figure 8: 2_2B_control_S2 Read2 Per-base N-content

/home/jvancamp/SF-seq/data/29_4E_fox_S21_L008_R1_001.fastq Mean quality score for each Base Position

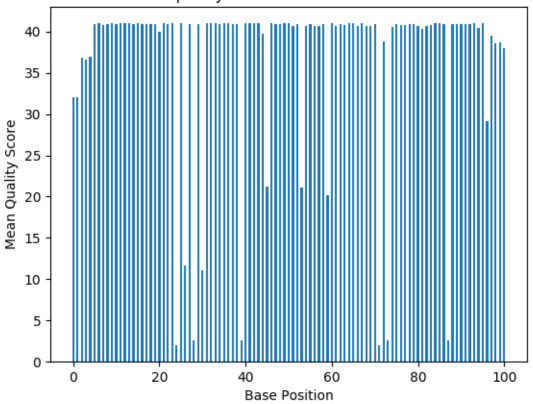


Figure 9: 29_4E_fox_S21 Read1 QScore distribution (My Script)

/home/jvancamp/SF-seq/data/29_4E_fox_S21_L008_R2_001.fastq Mean quality score for each Base Position

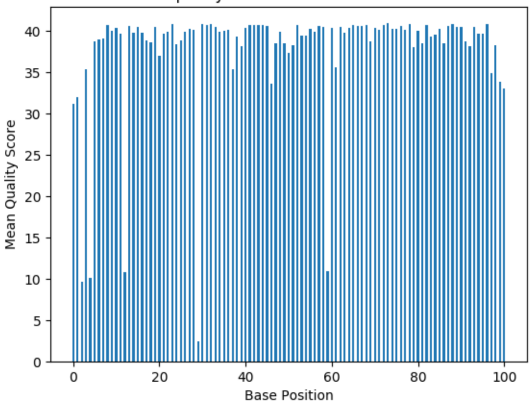


Figure 10: $29_4E_fox_S21$ Read2 QScore distribution (My Script)

/home/jvancamp/SF-seq/data/2_2B_control_S2_L008_R1_001.fastq Mean quality score for each Base Position

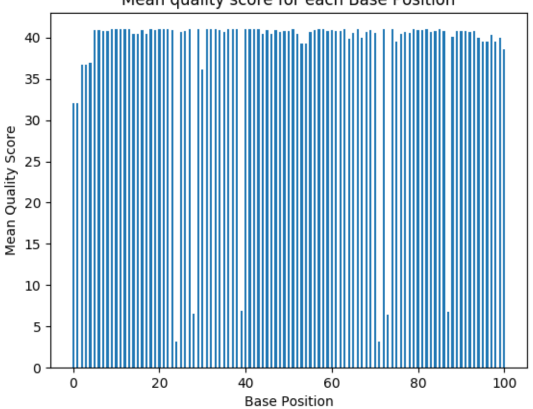


Figure 11: 2_2B_control_S2 Read1 QScore distribution (My Script)

/home/jvancamp/SF-seq/data/2_2B_control_S2_L008_R2_001.fastq Mean quality score for each Base Position

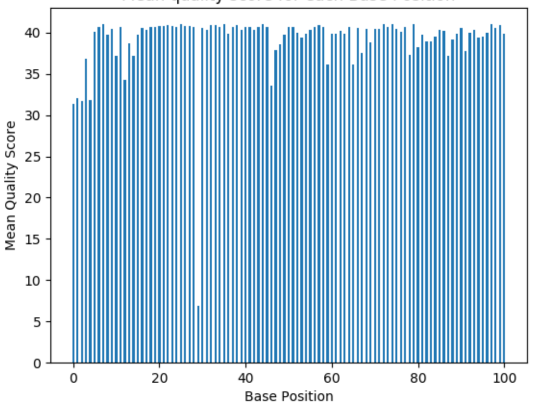


Figure 12: 2_2B_control_S2 Read2 QScore distribution (My Script)