Sample Quality Results

1 Basic Statistics

Sample: 20-D15-41762-MC-Nextera-Myeloid-Val1-Repeat_S20_L001

File type: Conventional base calls
Encoding: Sanger / Illumina 1.9

	Before trimming	After trimming
Total Sequences	842667	468362
Sequences flagged as poor quality	0	0
Sequence length	151	50-150
%GC	43	41

2 FastQC

Figure 1: Per base sequence quality

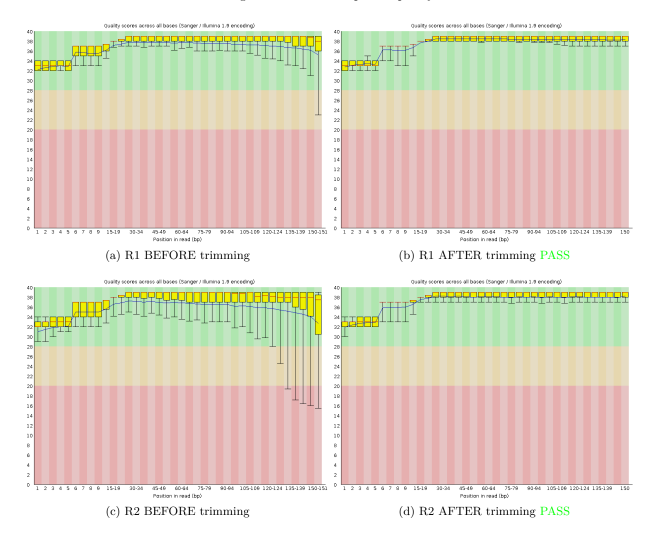
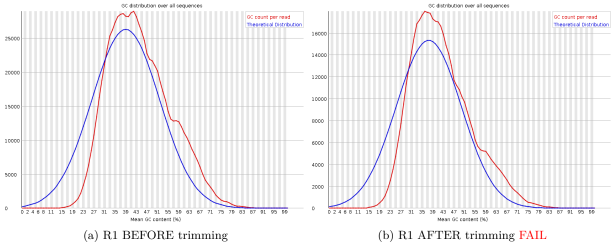
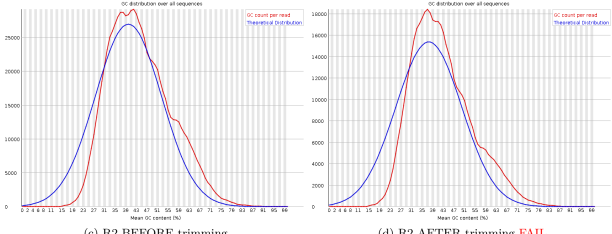


Figure 2: Per sequence GC content





(c) R2 BEFORE trimming

(d) R2 AFTER trimming FAIL

Figure 3: Sequence Length Distribution

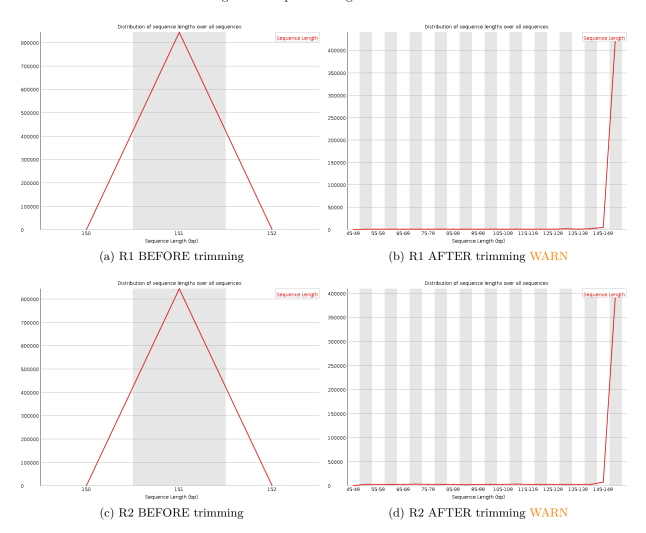
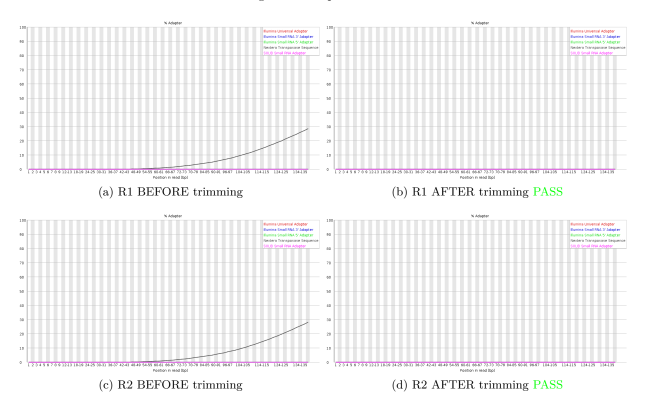
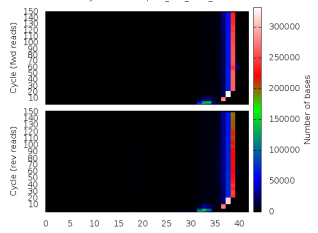


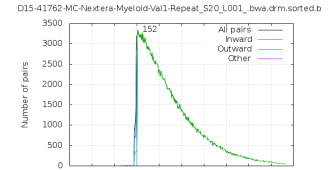
Figure 4: Adapter Content



3 BamStats

 $1762\text{-MC-Nextera-Myeloid-Val1-Repeat_S20_L001_.bwa.drm.sorted.bam.st} \epsilon$





100 150 200 250 300 350 400 450 500

Insert Size

(b) Fragment size

0

(a) Base quality per cycle

