## Illumina vs. AVITI

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We performed two sequencing experiments on a library prepared from a pool of samples. In one experiment, Illumina sequencing was used; in the other, AVITI. To compare results, we restrict our analysis to *E. coli* samples from a single donor (Baby 2, B002).

### 1 Number of reads

We used fastp to trim, filter, and tally reads with pseudocode:

```
$ fastp {input FASTQs} \
--cut_front \
--cut_tail \
--trim_poly_x \
--cut_mean_quality 30 \
--qualified_quality_phred 30 \
--unqualified_percent_limit 10 \
--length_required 50
```

Results were collected using MultiQC and parsed using custom code.

### Results

• AVITI has  $\sim 10^9$  reads, even after filtering (figure 1)

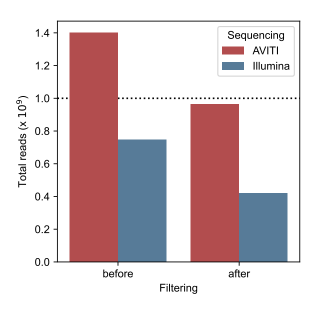


Figure 1: Total number of reads for AVITI or Illumina, before and after filtering with fastp.

### 2 Variant quality scores

We used bcftools to generate pile-ups and bcftools stats to extract variant quality scores with pseudocode:

Results were collected using MultiQC and parsed using custom code.

#### Results

• AVITI has slightly higher, though comparable, variant quality scores compared with Illumina (figure 2)

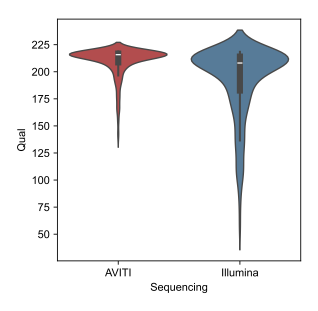


Figure 2: Variant quality scores for AVITI or Illumina.

### 3 Sequence typing

We used srst2 to perform sequence typing for E. coli (MLST database name Escherichia coli#1) with pseudocode:

```
$ srst2 --input_pe {trimmed reads} --mlst_* '{Escherichia_coli#1}'
```

#### Results

- $\frac{250}{256} \approx 98\%$  agreement between AVITI and Illumina (table 1)
- AVITI ( $\diamond$ ) has higher seq. depth at core genes used for sequence typing than Illumina ( $\bullet$ ) (figure 3)
- (ST)73 is the dominant sequence type (figure 3)

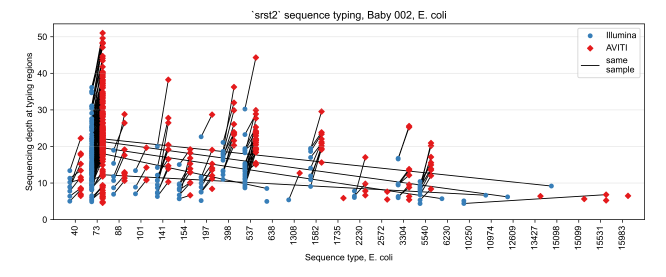


Figure 3: **Sequence typing results** for successful srst2 sequence typing of identical samples (–) prepared using AVITI (•) or Illumina (•).

	Count
AVITI & Illumina (total)	317
AVITI, typed (total)	309
Illumina, typed (total)	264
AVITI & Illumina, typed (total)	256
AVITI & Illumina, typed (agree)	250

Table 1: **Summary of sequence typing results**, tallying the number of samples for different criteria. The top row provides the total number of samples; the bottom row provides the number of samples *successfully* sequence typed for *both* AVITI *and* Illumina *and agree* in the designated sequence type.

## 4 Reconstructing phylogenies of dominant STs

# 5 Appendix

Code to reproduce is available on GitHub at t-silvers/sequencing-comparison-aviti-illumina.