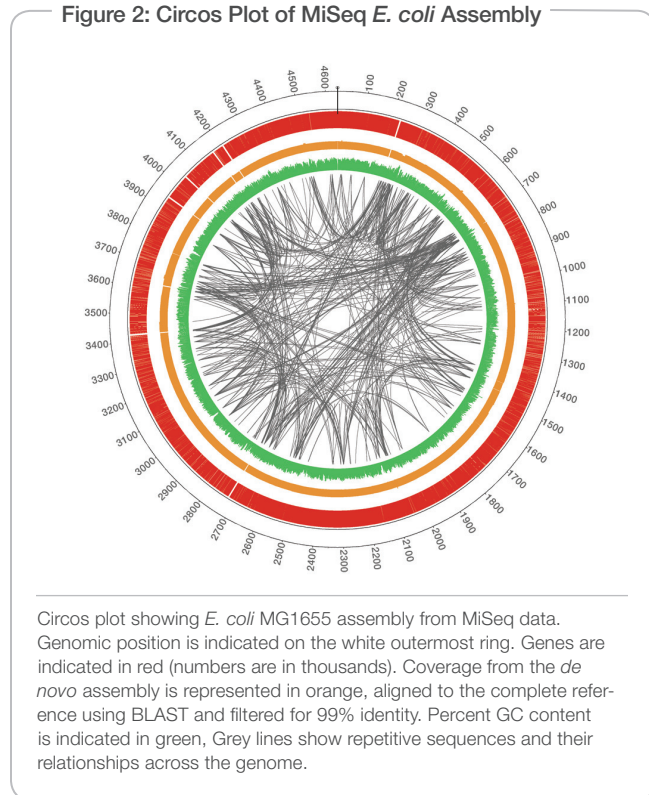


Figure 2: Circos Plot of MiSeq *E. coli* Assembly



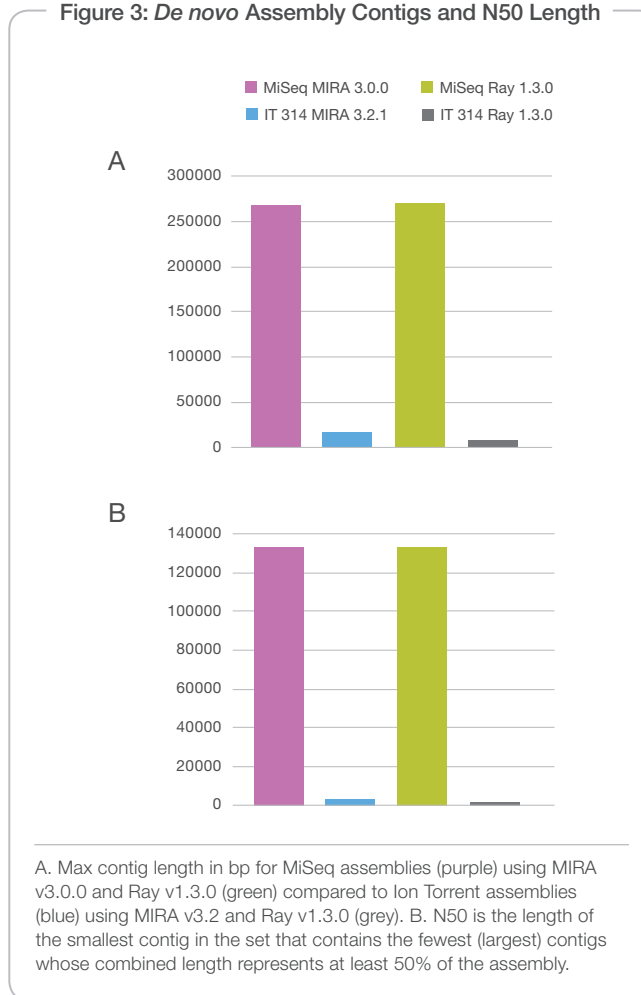
analyzed using CASAVA 1.8a5, and *de novo* assembly was completed using Velvet. For the *de novo* assembly comparison between MiSeq and Ion Torrent, the open access assemblers MIRA¹ and Ray² were used on MiSeq data down-sampled to 50x and compared to the entire data set from Ion Torrent reads³. These open source assembly tools are reported to work well with both Illumina and Ion Torrent data³, and produced results comparable to Velvet for the MiSeq data.

Results and Data Analysis

Data generated from the MiSeq and HiSeq systems showed similar cluster density and numbers of clusters passing filter. *De novo* assembly metrics from the HiSeq and MiSeq reads are very similar (Table 1). Comparison of HiSeq and MiSeq data with the reference sequence illustrates equivalent coverage over a range of GC content (Figure 1). Data from the MiSeq assembly overlaying the *E. coli* reference sequence are shown in a Circos plot (Figure 2), demonstrating excellent coverage over the entire genome.

De novo assembly data from the 2 × 150 bp MiSeq run was compared with Ion Torrent data³. To make an equal comparison, MiSeq data was down-sampled to 50x coverage, comprising 231 Mb, or approximately 1/7th of the data. Both the max contig length and N50 values were vastly superior in the down-sampled MiSeq data compared to the entire Ion Torrent data set (Figures 3A and B).

Figure 3: *De novo* Assembly Contigs and N50 Length



Conclusions

Using the same library preparation from bacterial DNA, sequencing on MiSeq was shown to be very comparable to HiSeq; both platforms yield high-quality data with > 85% bases above Q30 with even GC coverage. *De novo* assembly with these data also produce similar results, with excellent coverage of the reference sequence. Sequencing results generated on the MiSeq System are highly predictive of those delivered by the high-throughput HiSeq 2000 sequencing platform, making MiSeq ideal for piloting larger studies or performing independent experiments requiring speed and accuracy. For *de novo* assembly, the importance of high-quality, paired-end MiSeq reads is readily apparent compared to Ion Torrent. The high quality assembly produced from MiSeq paired-end reads show that better data give a more accurate picture of the genome.

References

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