DATA NOTE

Ultra-deep, long-read nanopore sequencing of mock microbial community standards

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Abstract

Background: Long sequencing reads are information-rich: aiding de novo assembly and reference mapping, and consequently have great potential for the study of microbial communities. However, the best approaches for analysis of long-read metagenomic data are unknown. Additionally, rigorous evaluation of bioinformatics tools is hindered by a lack of long-read data from validated samples with known composition. Findings: We sequenced 2 commercially available mock communities containing 10 microbial species (ZymoBIOMICS Microbial Community Standards) with Oxford Nanopore GridION and PromethION. Both communities and the 10 individual species isolates were also sequenced with Illumina technology. We generated 14 and 16 gigabase pairs from 2 GridION flowcells and 150 and 153 gigabase pairs from 2 PromethION flowcells for the evenly distributed and log-distributed communities, respectively. Read length N50 ranged between 5.3 and 5.4 kilobase pairs over the 4 sequencing runs. Basecalls and corresponding signal data are made available (4.2 TB in total). Alignment to Illumina-sequenced isolates demonstrated the expected microbial species at anticipated abundances, with the limit of detection for the lowest abundance species below 50 cells (GridION). De novo assembly of metagenomes recovered long contiguous sequences without the need for pre-processing techniques such as binning.

Conclusions: We present ultra-deep, long-read nanopore datasets from a well-defined mock community. These datasets will be useful for those developing bioinformatics methods for long-read metagenomics and for the validation and comparison of current laboratory and software pipelines.

Keywords: bioinformatics; metagenomics; mock community; nanopore; single-molecule sequencing; real-time sequencing; benchmark; Illumina; de novo assembly

Data Description

Whole-genome sequencing of microbial communities (metagenomics) has revolutionized our view of microbial evolution and diversity, with numerous potential applications for microbial ecology, clinical microbiology, and industrial biotechnology [1,2]. Typically, metagenomic studies use high-throughput sequencing platforms (e.g., Illumina) [3], which generate very high yield, but of limited read length (100–300 base pairs [bp]).

In contrast, single-molecule sequencing platforms such as the Oxford Nanopore MinION, GridION, and PromethION are able to sequence very long fragments of DNA (>10 kilobase pairs [kb]), with >2 megabase pairs [Mb] reported [4,5], and with recent improvements to the platform making metagenomic studies using nanopore more viable, such studies are increasing in frequency [6–9]. Long reads help with alignment-based assignment of taxonomy and function owing to their increased information content [10,11]. Additionally, long reads permit bridging of
repetitive sequences (within and between genomes), aiding genome completeness in de novo assembly [12]. However, these advantages are constrained by a high error rate (∼10%), necessitating the use of specific long-read alignment and assembly methods, which either are not specifically designed for metagenomics or have not been extensively tested on real data [13].

Mock community standards are useful for the development of genomics methods [14] and for the validation of existing laboratory, software, and bioinformatics approaches. For example, validating the accuracy of a taxonomic identification pipeline is important because the consequences of erroneous taxonomic identification from a metagenomic analysis may be severe, e.g., in public health microbiology [15,16] or incorrect diagnoses in clinical microbiology diagnostics. Mock community standards can also be used as positive controls during laboratory work, e.g., to validate that DNA extraction methods will yield the expected representation of a sampled community [14].

Here, we present 4 nanopore sequencing datasets of 2 microbial community standards, providing a state-of-the-art benchmark to accelerate the development of methods for analysing long-read metagenomics data.

**Background information**

The ZymoBIOMICS Microbial Community Standards (CS and CSII) are each composed of 10 microbial species: 8 bacteria and 2 yeasts (Table 1). The organisms in CS (hereafter referred to as "Even") are distributed equally (12%), with the exception of the 2 yeasts, which are each present at 2%. Cell counts from organisms in the CSII ("Log") community are distributed on a log scale, ranging from 89.1% (Listeria monocytogenes) down to 0.000089% (Staphylococcus aureus).

**Methods**

**DNA extraction**

DNA was extracted from 75 μl ZymoBIOMICS Microbial Community Standard (Zymo Research Corporation, Irvine, CA, USA). Product D6300, Lot ZRC190633 and 375 μl ZymoBIOMICS Microbial Community Standard II (Product D6310, Lot ZRC190842) using the ZymoBIOMICS DNA Miniprep extraction kit according to the manufacturer’s instructions, with the following modifications to increase fragment length and maintain the expected representation of the Gram-negative species that are already lysed in the DNA/RNA Shield storage solution. The standard was centrifuged at 8,000×g for 5 min before removing the supernatant and retaining. The cell pellet was resuspended in 750 μl lysis buffer and added to the ZR BashingBead lysis tube (Zymo Research Corporation). Bead-beating was performed on a FastPrep-24 (MP Biomedicals, Solon, OH, USA) instrument for 2 cycles of 40 seconds at 6.0 ms⁻¹, with 5 min sitting on ice between cycles. The bead tubes were centrifuged at 10,000×g for 1 min and 450 μl of supernatant was transferred to a Zymo Spin III-F filter before being centrifuged again at 8,000×g for 1 min. 45 μl (Even) and 225 μl (Log) of the supernatant retained earlier was combined with 450 μl filtrate before adding 1485 μl (Even) or 2025 μl (Log) Binding Buffer and mixing before loading onto the column. Methods are available online via protocols.io [18].

**Nanopore sequencing library preparation**

Quantification steps were performed using the dsDNA HS assay for Qubit. DNA was size-selected by cleaning up with 0.45× volume of Ampure XP (Beckman Coulter, Brea, CA, USA) and eluted in 100 μl EB (Qiagen, Hilden, Germany). Libraries were prepared from 1,400 ng input DNA using the SQK-LSK109 kit (Oxford Nanopore Technologies, Oxford, UK) in accordance with the manufacturer’s protocol, except incubation times for end repair, dA-tailing, and ligation were increased to 30 min to improve ligation efficiency. The Even and Log libraries were split and used on both the GridION and PromethION flowcells.

**Sequencing**

Sequencing libraries were quantified and 2 aliquots of 50 and 400 ng were prepared for GridION and PromethION sequencing, respectively. The GridION sequencing was performed using FLOMIN106 (rev.C) flowcells, MinKNOW 1.15.1, and standard 48-hour run script with active channel selection enabled. The PromethION sequencing was performed using FLO-PR002 flowcells, MinKNOW 1.14.2, and standard 64-hour run script with active channel selection enabled.

Refuelling was performed approximately every 24 hours (GridION, PromethION) by loading 75 μl (GridION) or 150 μl (PromethION) refuelling mix (sequencing buffer diluted 1:1 with nuclease-free water). In addition, after the standard scripts had completed, the PromethION was restarted several times to utilize the remaining active pores and maximize total yield.

**Table 1. Description of the 10 organisms comprising the ZymoBIOMICS Mock Community Standards**

| Species                        | Type | Estimated size (Mb) | NRRL accession | ATCC accession | Sequence type | Illumina FASTQ | PacBio RSII FASTQ | PacBio Sequel FASTQ |
|--------------------------------|------|---------------------|----------------|----------------|---------------|----------------|--------------------|---------------------|
| Bacillus subtilis              | Gram + | 4.05                | B-354          | 6633           | ST7           | ERR2935851      | SRR7498042         | SRR7415629          |
| Cryptococcus neoformans × Cryptococcus deneiformis | Yeast | 18.90               | Y-2534         | 32045          |               | ERR2935856      |                    |                    |
| Enterococcus faecalis          | Gram + | 2.85                | B-537          | 7080           | ST55          | ERR2935850      | SRR7415622         | SRR7415630          |
| Escherichia coli               | Gram – | 4.88                | B-1109         |                | ST10          | ERR2935852      | SRR7498041         |                    |
| Lactobacillus fermentum        | Gram + | 1.91                | B-1840         | 14931          |               | ERR2935857      |                    |                    |
| Listeria monocytogenes         | Gram + | 2.99                | B-33116        | 19117          | ST449         | ERR2935854      | SRR7415624         | SRR7415635          |
| Pseudomonas aeruginosa         | Gram – | 6.79                | B-3509         | 15442          | ST252         | ERR2935853      | SRR7498043         |                    |
| Saccharomyces cerevisiae        | Yeast | 12.10               | Y-567          | 9763           | ST139         | ERR2935855      | SRR7498048         | SRR7415638          |
| Salmonella enterica            | Gram + | 4.76                | B-4212         |                | ST9           | ERR2935849      | SRR7415627         | SRR7415637          |

Table adapted from ZymoBIOMICSTM Microbial Community Standard II (Log Distribution) Instruction Manual v1.1.2 Table 2 and Appendix A. The S. enterica genome is listed at Agricultural Research Service Culture Collection (NRRL) (B-4212) as Serovar Typhimurium LT2, but our genomic analysis shows it is likely to be Serotype Choleraesuis, indicating possible mis-annotation. ATCC: American Type Culture Collection.
Nanopore basecalling

Reads were basecalled on-instrument using the Guppy v2.2.2 GPU basecaller (Oxford Nanopore Technologies) with the supplied dna-r9.4.1.450bps_flipflop_prom.cfg configuration (PromethION) and dna_r9.4.1.450bps_flipflop.cfg (GridION).

Illumina sequencing

DNA was extracted from pure cultures of each species using the ZymoBIOMICS DNA Miniprep Kit. Library preparation was performed using the Kapa HyperPlus Kit (Kapa Biosystems, Wilmington, MA, USA) with 100 ng DNA as input and TruSeq Y-adapters (Illumina, San Diego, CA, USA). The purified library derived from each sample was quantified with the 4200 TapeStation System (Agilent Technologies, Santa Clara, CA, USA) and pooled together in an equimolar fashion. The multiplexed isolates were sequenced on an Illumina HiSeq 15000 sequencing system using 2×101 bp (paired-end) sequencing, over 4 lanes. Raw reads were demultiplexed using bc12fastq v2.17. Shotgun sequencing of the Even and Log communities was performed with the same protocol, with the exception that the Log community was sequenced individually on 2 flowcell lanes and the Even community was instead sequenced on an Illumina MiSeq using 2×151 bp (paired-end) sequencing.

Bioinformatics Methods

Illumina draft assembly

For the purposes of estimating sequencing coverage and contiguity, we constructed a draft assembly from our available Illumina sequencing data. Illumina reads for each of the 10 isolates were assembled using SPAdes v3.12.0 [19] with paired-end reads as input, using parameters -m 512 -t 12. Scaffolds from SPAdes <500 bp length or with <10× coverage were removed. The remaining scaffolds were combined into a single mock community draft assembly for downstream analysis. Multilocus sequence typing (MLST) of the scaffolds was conducted with mlst [20].

Pacific Biosciences draft assembly

A recently released orthogonal data set from McIntyre et al. includes individual Pacific BioSciences (PacBio) sequencing of 8 of the 10 organisms that compose the 2 Zymo communities [17]. Assemblies for the 8 isolates that passed quality control (excluding L. fermentum and C. neoformans) were generated with HGAPv2 [21]. Assemblies have been made available by the authors and were downloaded from [22] (Git commit db4e9fd4) for the purposes of assessing metagenomic assembly accuracy for the 7 bacterial species where complete genomes were available.

Sequencing coverage estimation

Nanopore reads were aligned to the illumina draft assembly using minimap2 [23] v2.14-r883 with parameters -ax map-ont -t 12 and converted to a sorted BAM file using samtools [24]. To reduce erroneous mappings, alignment BAM files were filtered using the script bamstats.py according to the following criteria: reference mapping length ≥500 bp, map quality (MAPQ) > 0, there are no supplementary alignments for this read, and read is not a secondary alignment. Per-species coverage summary statistics were generated using the summariseStats.R R script.

Nanopore read accuracy

Read accuracy was determined by calculating BLAST-like identities from the filtered alignments (as per [25]), calculated as (L − NM)/L using the minimap2 number of mismatches (NM) SAM tag and the sum of match, insertion, and deletion CIGAR operations (L).

Metagenomic assembly and contiguity estimation

Metagenomic assemblies were constructed with wtdbg2 v2.2 [26] from the nanopore sequencing of the communities. wtdbg2 was compiled from source via Git commit 904f2b3. For GridION, all nanopore reads were used. For PromethION, a 25% subsample was selected with seqtk [27]. Assemblies were conducted under a variety of parameter values for homopolymer-compressed k-mer size (-p), minimum graph edge weight support (-e), and read length threshold (-l). Global parameters for all runs (-1 -K10000 --node-max 6000) were used to turn off k-mer subsampling (to remove assembly stochasticity) and increase the coverage thresholds applied to k-mers and constructed nodes.

Assembled contigs were assigned to taxa with kraken2 [28] (--use-names -t12) using a database containing all of the archaeal, bacterial, fungal, protozoal, and viral sequences from RefSeq, and UniVec_Core (database download links are in our repository). The kraken2 output was parsed with extracken.py and plotted with contiguity.R to visually assess contiguity. Following assignment, contigs can be extracted into separate FASTA with extract_contigs_with_kraken.py.

Assembly polishing

After inspection of the contiguity.R plot, 8 high-contiguity assemblies were selected for polishing. Polishing consisted of 2 iterations of racon [29], followed by medaka [30] and 2 iterations of pilon [31]. racon v1.3.2 was used to polish contigs with the nanopore reads. medaka v0.5.0 was used to polish the racon polished contigs, with the nanopore reads specifying the r941 flip model. The PromethION assemblies were polished using the same seqtk-derived 25% subset from which the assemblies were constructed. pilon v1.23 was used to polish the medaka polished contigs, with the CS (Even) community Illumina reads.

Estimation of genome completeness

To estimate accuracy of the polished assemblies, contigs were first assigned to taxa and extracted into separate FASTA using kraken2 as previously described. For the 7 bacteria for which corresponding PacBio draft assemblies were available, sequence identity dotplots were generated using a modified version of minidot [32], which uses minimap2 (-x asm10 -no-long-join – dual=yes -p) to align the polished contigs binned by kraken2, to the corresponding PacBio draft genome. Genome completeness was estimated with CheckM v1.0.13 [33] using the taxonomy_wf subcommand, after each phase of the polishing pipeline. CheckM was executed separately for each kraken2 bin that had a corresponding PacBio reference, specifying the appropriate species for the bin to taxonomy_wf. We report the CheckM “Completeness” score, which estimates completeness by identifying colocated marker gene sets on the assembled contigs as a proportion of the total collection of marker gene sets expected for a specific taxa.
Table 2. Summary of the 4 nanopore sequencing experiments

| Signal accession | FASTQ accession | Sequencer   | Standard (lot) | Time (h) | Reads (M) | N50 (kb) | Quality (median Q) | Yield (Gb) | Q>7 (Gb) |
|------------------|-----------------|-------------|----------------|----------|-----------|-----------|-------------------|------------|----------|
| ERR2887847       | ERR3152364      | GridION     | Zymo CS Even ZRC190633 | 48       | 3.49      | 5.3       | 10.3              | 14.38      | 12.39    |
| ERR2887850       | ERR3152366      | GridION     | Zymo CSII Log ZRC190842 | 48       | 3.67      | 5.4       | 9.8               | 16.51      | 13.97    |
| ERR2887848       | ERR3152365      | PromethION  | Zymo CS Even ZRC190633 | 64       | 35.7      | 5.4       | 10.5              | 150.88     | 130.32   |
| ERR2887849       | ERR3152365      | PromethION  | Zymo CS Even ZRC190633 | 64       | 34.5      | 5.4       | 10.7              | 153.31     | 133.68   |
| ERR2887851       | ERR3152367      | PromethION  | Zymo CSII Log ZRC190842 | 64       | 35.7      | 5.4       | 10.5              | 150.88     | 130.32   |
| ERR2887852       | ERR3152367      | PromethION  | Zymo CSII Log ZRC190842 | 64       | 34.5      | 5.4       | 10.7              | 153.31     | 133.68   |

PromethION runs were restarted following the standard 64-hour protocol. The table reflects total yield across both the standard run and subsequent restarts.

Figure 1 Summary plots for the 4 generated data sets: (a) collector’s curve showing sequencing yield over time for each of the 4 sequencing runs, (b) density plot showing sequence accuracy (BLAST-like identities), (c) density plot showing sequencing speed over time by sequencing experiment.

Table 3. Summary statistics for Illumina sequencing data

| Dataset | Pairs (M) | Yield (Gb) | phred ≥ 30 (%) | Accession |
|---------|-----------|------------|----------------|-----------|
| Isolates | 13.53 ± 5.23 | 2.73 ± 1.06 | 87.72 ± 5.43 | See Table 1 |
| CS (Even) | 8.8 | 2.65 | 95.12 | ERR2984773 |
| CSII (Log) | 47.8 | 9.66 | 95.71 | ERR2935805 |

Illumina sequencing was performed on an Illumina HiSeq 1500, with the exception of the Even community, which was sequenced on an Illumina MiSeq.

Results

Nanopore sequencing metrics

We generated a total of 335.1 gigabase pairs (Gb) of sequence from the 4 nanopore sequencing runs (Table 2, Fig. 1a). PromethION flowcells generated ~10 times more sequencing data than the comparative GridION runs and showed equivalent read length N50 and read accuracy (Fig. 1b). We observed a difference in sequencing speed between the PromethION (mean speed, 419 and 437 bps for Even and Log, respectively) and the GridION (mean speed, 352 and 372 bp for Even and Log) (Fig. 1c).

Illumina sequencing metrics

Illumina datasets for the 10 individually sequenced isolates averaged 13.53 million pairs of reads (ranging between 7.1 and 23.2 million), with proportions of reads with a mean phred score ≥ 30 ranging between 75.51% and 93.09% (Table 3). Illumina sequencing generated 8.8 million pairs of reads (2×151 bp, MiSeq) and 47.8 million pairs of reads (2×101 bp, HiSeq) for the Even and Log community, respectively (Table 3).

Nanopore mapping statistics

We identify the presence of all 10 microbial species in the community, for both Even and Log samples, in expected proportions (Fig. 2). For the Even community, the GridION results provide sufficient depth (i.e., ≥30× coverage) to potentially assemble all 8 of the bacteria. The coverage of the yeast genomes was lower (10× and 17×), potentially sufficient for assembly scaffolding. On the PromethION all genomes had >100× mean coverage (Tables 4 and 5).

For the log-distributed community, 3 taxa have sufficient coverage for assembly on GridION, compared with 4 on PromethION. On PromethION, a further 2 genomes (S. enterica and E. coli) have sufficient coverage for assembly scaffolding. We were able to detect S. aureus, the lowest abundance organism on both platforms, with 19 reads from PromethION (from 400-cell input) and 4 reads from GridION (from 50-cell input).

Nanopore metagenomic assemblies

We assessed the contiguity of our nanopore metagenomic assemblies for each run with different assembly parameters.

For the Even community, genomes of the expected size were present for each of the bacterial species, contained in small numbers of large contigs (Fig. 3). However, the 2 yeasts are highly fragmented, consistent with their low read depth.
Table 4. Read alignment statistics for Even samples, showing absolute measurements and proportion of sequencing yield and the estimated genome coverage obtained for each organism in the mock community

| Species                  | Expected proportion | GridION: Yield (Gb) | Measured proportion | GridION: Alignment N50 (kb) | Coverage (×) | PromethION: Yield (Gb) | Measured proportion | PromethION: Alignment N50 (kb) | Coverage (×) |
|--------------------------|---------------------|---------------------|---------------------|-----------------------------|--------------|------------------------|---------------------|-------------------------------|--------------|
| Bacillus subtilis        | 12                  | 2.12                | 19.32               | 4.30                        | 524.51       | 21.55                  | 19.02               | 4.40                          | 5,326.44     |
| Listeria monocyto genes  | 12                  | 1.60                | 14.56               | 4.47                        | 534.26       | 16.23                  | 14.33               | 4.58                          | 5,424.46     |
| Enterococcus faecalis    | 12                  | 1.34                | 12.24               | 4.45                        | 472.47       | 13.67                  | 12.07               | 4.57                          | 4,805.60     |
| Staphylococcus aureus    | 12                  | 1.24                | 11.28               | 4.47                        | 453.84       | 12.59                  | 11.11               | 4.59                          | 4,611.61     |
| Salmonella enterica      | 12                  | 1.10                | 9.99                | 8.55                        | 230.51       | 11.69                  | 10.32               | 8.95                          | 2,456.19     |
| Escherichia coli         | 12                  | 1.09                | 9.93                | 8.31                        | 223.59       | 11.62                  | 10.26               | 8.71                          | 2,382.59     |
| Pseudomonas aeruginosa   | 12                  | 1.07                | 9.70                | 8.98                        | 156.85       | 11.45                  | 10.11               | 9.38                          | 1,686.34     |
| Lactobacillus fermentum  | 12                  | 1.02                | 9.28                | 8.62                        | 534.73       | 10.34                  | 9.13                | 7.33                          | 5,425.69     |
| Saccharomyces cerevisiae | 2                   | 0.21                | 1.92                | 4.09                        | 17.46        | 2.12                   | 1.87                | 4.18                          | 175.23       |
| Cryptococcus neoformans  | 2                   | 0.20                | 1.78                | 4.45                        | 10.37        | 2.00                   | 1.77                | 4.54                          | 105.82       |

Table 5. Read alignment statistics for Log samples, describing sequencing yield and estimated genome coverage obtained for each organism in the mock community

| Species                  | GridION: Yield (Gb) | Alignment N50 (kb) | Coverage (×) | PromethION: Yield (Gb) | Alignment N50 (kb) | Coverage (×) |
|--------------------------|---------------------|--------------------|--------------|------------------------|--------------------|--------------|
| Listeria monocytogenes   | 12.10               | 4.95               | 4,043.90     | 110.09                 | 4.97               | 36,796.21    |
| Pseudomonas aeruginosa   | 1.10                | 9.38               | 161.45       | 9.99                   | 9.33               | 1,471.41     |
| Bacillus subtilis        | 0.16                | 5.03               | 38.67        | 1.44                   | 5.04               | 356.00       |
| Saccharomyces cerevisiae | 0.08                | 4.78               | 6.93         | 0.75                   | 4.75               | 62.33        |
| Salmonella enterica      | 0.01                | 9.20               | 2.20         | 0.10                   | 9.17               | 20.04        |
| Escherichia coli         | 0.01                | 8.65               | 2.14         | 0.09                   | 9.17               | 19.24        |
| Lactobacillus fermentum  | 4E−4                | 3.40               | 0.210        | 7E−4                   | 3.37               | 2.03         |
| Enterococcus faecalis    | 2E−4                | 7.62               | 0.055        | 1E−3                   | 6.05               | 0.34         |
| Cryptococcus neoformans  | 6E−5                | 4.41               | 0.003        | 5E−5                   | 3.58               | 0.020        |
| Staphylococcus aureus    | 1E−5                | 7.12               | 0.005        |                        |                    |              |

Note that expected and measured proportions are illustrated in Fig. 2.

Figure 2 Proportion of sequenced bases assigned by minimap2 to each of the 10 organisms that were sequenced (x-axis), against the proportion of yield expected given the known composition (y-axis) of the Zymo CSII (Log) standard.
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Figure 3 Bar plots demonstrating total length and contiguity of genomic assemblies obtained with wtdbg2 from each of the long-read nanopore data sets. For each organism in the community (coloured columns), contigs longer than 10 kb are horizontally stacked along the x-axis. Each row represents a run of wtdbg2, with the parameters for edge support, read length threshold, and homopolymer-compressed k-mer size labelled on the left. Assemblies are grouped by the data set on which they were run (row facets). Additionally, assemblies may be compared to the estimated true genome size, the available McIntyre et al. [17] PacBio assemblies, and per-isolate Illumina SPAdes assembly. Estimated genomes sizes are the same as those found in Table 1; however, to display approximate chromosomes, the 2 yeasts were replaced by their corresponding canonical National Center for Biotechnology Information references for visualization purposes only. The C. neoformans strain used by the Zymo standards is a diploid genetic cross, which may explain the larger assemblies, compared to the represented estimated haploid size.

Discussion

There are several noteworthy aspects of this dataset: we generated >300 Gb of sequence data from the Oxford Nanopore PromethION and 30 Gb from the Oxford Nanopore GridION, on a well-characterized mock community sample, and we have made basecalls and electrical signal data for each of the 4 runs presented here available: a combined dataset size of >4 terabytes. The availability of the raw signal permits future basecalling of the data (an area under rapid development), as well as signal-level polishing and the detection of methylated bases [34].

Individual sequencing libraries were split between the GridION and PromethION, permitting direct comparisons of the instruments to be made. We observed high concordance between the datasets from each platform. We note that the sequencing speed of the PromethION is faster than the GridION, which we attribute to different running temperatures on these instruments (39°C vs 34°C, respectively).

Confident detection of S. aureus was demonstrated for the GridION run to <50 cells using the Log community. The PromethION generated ~5 times more S. aureus reads than the GridION; however, we loaded 8 times as much library, making it seem less sensitive. It may be possible to reduce the input to PromethION flowcells, but we have not attempted this.

Early results of metagenomic assembly show promise for reconstruction of whole microbial genomes from mixed samples without a binning step. We focused on the developing wtdbg2 software because the established minimap2 and miniasm method resulted in excessively large intermediate files (tens of terabases per analysis) that were impractical to store and analyse.

For the Even community, using wtdbg2 with varying parameter choices, we were able to assemble 4 of the bacteria into single contigs. However, no single parameter set was found to be optimum for both total genome size and contig length. Increasing -e improved contiguity for the Even community; however, this resulted in the loss of yeasts from the assembly. Increasing the read length threshold (-L) improved contiguity for all sample and platform combinations, at the cost of genome size. Increasing the homopolymer-compressed k-mer size (-p) from the default of 21 to 23 also seemed to improve contiguity.

We found that wtdbg2 expects a maximum of 200× sample coverage and discards sequence k-mers and de Bruijn graph nodes with >200× support. Although these limits can be lifted by specifying higher -K and -node-max, we still observe more fragmented assemblies on the PromethION data (especially for the 100% PromethION data [not shown]), potentially indicating a need to further tune the algorithm to account for the large differences in coverage between genomes. It should be noted that wtdbg2 is still under active development, making it difficult to make concrete recommendations for parameters.
Table 6. Sequence identity dotplots and CheckM genome completeness scores for each of the 7 bacterial species for which there was a corresponding PacBio assembly from McIntyre et al. [17]

|                   | GridION | PromethION |
|-------------------|---------|------------|
|                   |          | Assembly   |          |          |          | Assembly   |          |          |          |
|                   |          | -p 21 -e 3| L50/L95   |          |          | L50/L95   |          |          |          |
|                   |          |          | Base      | +Racon 2| +Medaka  | +Racon 2| +Medaka  | +Pilon 2 | +Pilon 2 |
|                   |          |          |           |         |          |           |         |          |          |
|                   |          |          |           |         |          |           |         |          |          |
|                   | 2 / 5    | 1 / 1    | 5 / 17    | 1 / 3   | 1 / 6   | 6 / 21    | 1 / 1    |          |          |
|                   | 74.27    | 76.10    | 70.14    | 76.11   | 78.64   | 66.78    | 78.48    |          |          |
|                   | 86.65    | 88.08    | 84.21    | 83.41   | 92.96   | 82.74    | 90.27    |          |          |
|                   | 97.45    | 99.07    | 94.46    | 97.50   | 97.33   | 95.10    | 98.35    |          |          |
|                   | 98.42    | 99.66    | 95.46    | 98.57   | 99.77   | 96.98    | 98.88    |          |          |
|                   |          |          |          |         |         |          |          |          |
|                   | 2 / 4    | 1 / 1    | 2 / -     | 1 / 4   | 1 / 1   |          |          |          |          |
|                   | 73.42    | 75.81    | 67.44    | 74.44   | 81.08   | 70.54    | 78.20    |          |          |
|                   | 84.81    | 87.30    | 79.83    | 84.57   | 92.53   | 85.83    | 88.64    |          |          |
|                   | 96.60    | 98.80    | 89.41    | 98.06   | 98.04   | 97.46    | 98.23    |          |          |
|                   | 97.83    | 99.66    | 90.36    | 99.15   | 99.82   | 98.67    | 98.88    |          |          |
|                   |          |          |          |         |         |          |          |          |
|                   | 1 / 1    | 1 / 2    | 3 / 8     | 1 / -   | 1 / 7   | 1 / 1    | 1 / 2    |          |          |
|                   | 74.29    | 76.07    | 72.21    | 57.20   | 79.64   | 68.99    | 78.28    |          |          |
|                   | 85.36    | 86.11    | 84.53    | 62.94   | 92.28   | 84.46    | 90.46    |          |          |
|                   | 97.14    | 99.11    | 95.95    | 71.21   | 98.06   | 96.57    | 98.95    |          |          |
|                   | 98.27    | 99.66    | 97.17    | 72.24   | 99.53   | 98.34    | 98.78    |          |          |
|                   |          |          |          |         |         |          |          |          |
|                   | 2 / 3    | 1 / 1    | 2 / -     | 1 / 3   | 1 / 2   | 1 / 2    | 1 / 3    |          |          |
|                   | 73.71    | 77.52    | 73.73    | 75.42   | 82.06   | 60.74    | 79.24    |          |          |
|                   | 86.42    | 88.31    | 84.69    | 85.14   | 92.81   | 71.29    | 87.81    |          |          |
|                   | 97.16    | 98.83    | 94.26    | 96.86   | 97.86   | 82.42    | 98.45    |          |          |
|                   | 98.44    | 99.66    | 98.13    | 98.69   | 99.83   | 83.58    | 98.86    |          |          |

Four wtdbg2 assembly conditions are represented, varying the homopolymer-compressed k-mer parameter “p” and the graph minimum edge weight threshold “e.” The read length threshold “L” was fixed at 5,000 bp. The left and right halves of the table correspond to the same assembly condition for the GridION and 25% PromethION sequencing data, respectively. The L50/L95 refers to the number of assembled contigs required to span ≥50% and ≥95% of the estimated genome size (see Table 1). A minus sign indicates that the set of assembled contigs assigned to a taxon were not of sufficient total length to cover 95% of the estimated size. CheckM genome completeness scores are expressed as a percentage and were calculated per organism at the end of each polishing phase.

bs: B. subtilis; ef: E. faecalis; ec: E. coli; lm: L. monocytogenes; pa: P. aeruginosa; se: S. enterica; sa: S. aureus.

We found that any form of polishing improves the completeness of assemblies, likely due to the correction of frameshifts caused by indels. Short-read polishing with pilon also improves the assemblies, despite low coverage of the Illumina Even community data, and the results might be expected to improve further with increased coverage.

The availability of this dataset should help with further improvements to long-read assembly techniques.

Other mock microbial samples are available that we did not test here. A notable alternative mock community sample is from the Human Microbiome Project (HMP) and consists of 20 microbial samples (available from BEI Resources). This mock community have been sequenced as part of other studies, although the datasets are much smaller than the ones presented here [9,35]. Bertrand et al. [12] presented a synthetic mock community of their own construction to demonstrate hybrid nanopore-illumina metagenome assemblies.

Re-use potential

The provision of Illumina reads for each isolate permits a ground-truth to be obtained for the individual species contained in the mock community. This will be useful for training new nanopore basecalling and polishing models, long-read aligners, variant callers, and validating taxonomic assignment and assembly software and pipelines.

Availability of source code and requirements

Python and R scripts used to generate the summary information and analyses are open source and freely available via our repository (https://github.com/LomanLab/mockcommunity), under the MIT license. Our pipeline was orchestrated with Snakemake [36]; the workflow is available from our repository.
Ultra-deep, long-read nanopore sequencing of mock microbial community standards

Availability of supporting data and materials

This manuscript, and its supporting data are available under a Creative Commons Attribution 4.0 International license. Unprocessed FASTQ from the Illumina sequencing of the 10 isolates is available at the European Nucleotide Archive, via the identifiers listed in Table 1; identifiers for the Even and Log community illumina sequencing can be found in Table 3. Both the raw signal, and basecalled FASTQ for our nanopore sequencing experiments are available at the European Nucleotide Archive, via the identifiers listed in Table 2. The SPAdes-assembled illumina draft reference, and the collection of nanopore assemblies for each wtdbg2 condition are linked to from our GitHub repository (https://github.com/Loman Lab/mockcommunity), along with the kraken2 database used for taxonomic classification of the assembled contigs. Further updates (such as updated references, or new assemblies) will be made available through our project website https://loman nlab.github.io/mockcommunity/. An archival snapshot of our GitHub repository and associated assembly FASTA files are also available via GigaDB [37].

Abbreviations

ATCC: American Type Culture Collection; bp: base pairs; CS: Community Standards; Gb: gigabase pairs; kb: kilobase pairs; Mb: megabase pairs; MLST: multilocus sequence typing; NRRL: Northern Regional Research Laboratory; ARSCC: Agricultural Research Service Culture Collection; PacBio: Pacific Biosciences.

Competing interests

Cambridge Biosciences provided ZymoBIOMICS products free of charge. SJT is an employee of Zymo Research Corporation. N.J. has received Oxford Nanopore Technologies (ONT) reagents free of charge to support his research programme. N.J. and J.Q. have received travel expenses to speak at ONT events. N.L. has received an honorarium to speak at an ONT company meeting.

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Authors’ contributions

Conceptualization: N.L.; Methodology: N.L., J.Q., S.N., S.T.; Software: S.N., N.L.; Validation: S.N., N.L.; Formal analysis: S.N., N.L.; Investigation: N.L., J.Q., S.N.; Resources: N.L., S.T.; Data curation: S.N., N.L., S.T.; Writing—original draft preparation: S.N.; Writing—review and editing: S.N., N.L., J.Q., S.T.; Visualization: S.N., N.L.; Supervision: N.L.; Project administration: N.L.; Funding acquisition: N.L., S.T.

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References

1. Handelsman J. Metagenomics: application of genomics to uncultured microorganisms. Microb Mol Biol Rev 2004;68(4):669–85.
2. Hug LA, Baker BJ, Anantharaman K, et al. A new view of the tree of life. Nat Microbiol 2016;1:16048.
3. Quince C, Walker AW, Simpson JT, et al. Shotgun metagenomics, from sampling to analysis. Nat Biotechnol 2017;35(9):833–44.
4. Jain M, Koren S, Miga KH, et al. Nanopore sequencing and assembly of a human genome with ultra-long reads. Nat Biotechnol 2018;36:338.
5. Payne A, Holmes N, Rakyan V, et al. BulkVis: a graphical viewer for Oxford nanopore bulk FAST5 files. Bioinformatics 2018, doi:10.1093/bioinformatics/bty841.
6. Sanderson ND, Street TL, Foster D, et al. Real-time analysis of nanopore-based metagenomic sequencing from infected orthopaedic devices. BMC Genomics 2018;19(1):714.
7. Charalampous T, Richardson H, Kay GL, et al. Rapid diagnosis of lower respiratory infection using nanopore-based clinical metagenomics. bioRxiv 2018, doi:10.1101/387548.
8. Somerville V, Lutz S, Schmid M, et al. Long read-based de novo assembly of low complex metagenome samples results in finished genomes and reveals insights into strain diversity and an active phage system. bioRxiv 2018, doi:10.1101/476747.
9. Leggett RM, Alcon-Giner C, Heavena D, et al. Rapid profiling of the preterm infant gut microbiota using nanopore sequencing aids pathogen diagnostics. bioRxiv 2018, doi:10.1101/180406.
10. Huson DH, Auch AF, Qi J, et al. MEGAN analysis of metagenomic data. Genome Res 2007;17(3):377–86.
11. Wommack KE, Bhavsar J, Ravel J. Metagenomics: read length matters. Appl Environ Microbiol 2008;74(5):1453–63.
12. Bertrand D, Shaw J, Narayan M, et al. Nanopore sequencing enables high-resolution analysis of resistance determinants and mobile elements in the human gut microbiome. bioRxiv 2018, doi:10.1101/456905.
13. Szczurek A, Hofmann P, Belmann P, et al. Critical assessment of metagenome interpretation—a benchmark of metagenomics software. Nat Methods 2017;14(11):1063.
14. Mason CE, Afshinnekooh E, Tighe S, et al. International standards for genomes, transcriptomes, and metagenomes. J Biomol Tech 2017;28(1):8–18.
15. Ackelsberg J, Rakeman J, Hughes S, et al. Lack of evidence for plague or anthrax on the New York City subway. Cell Syst 2015;1(1):4–5.
16. McIntyre AB, Ounit R, Afshinnekoo E, et al. Comprehensive benchmarking and ensemble approaches for metagenomic classifiers. Genome Biol 2017;18(1):182.
17. McIntyre AB, Alexander N, Grigorev K, et al. Single-molecule sequencing detection of N 6-methyladenine in microbial reference materials. Nat Commun 2019;10(1):579.
18. Quick JC, Nicholls SM, Loman NJ, et al. Ultra-deep, long-read nanopore sequencing of mock microbial community standards. protocols.io 2019. dx.doi.org/10.17504/protocols.io.x9tfr6n.
19. Bankerich A, Nurrk S, Antipov D, et al. SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. J Comput Biol 2012;19(5):455–77.
20. mstl. https://github.com/tseemann/mlst, Accessed 22 Feb 2019.
21. Chin CS, Alexander DH, Marks P, et al. Nonhybrid, finished microbial genome assemblies from long-read SMRT sequencing data. Nat Methods 2013;10(6):563.
22. mCaller_analysis_scripts/assemblies. https://github.com/al-mcintyre/mCaller_analysis_scripts/tree/master/assemblies, Accessed 22 Feb 2019.
23. Li H. Minimap2: pairwise alignment for nucleotide sequences. Bioinformatics 2018; 1: 7.
24. Li H, Handsaker B, Wysoker A, et al. The sequence alignment/map format and SAMtools. Bioinformatics 2009;25(16):2078–9.
25. On the definition of sequence identity. http://lh3.github.io/2018/11/25/on-the-definition-of-sequence-identity, Accessed 22 Feb 2019.
26. Ruan J, Li H. Fast and accurate long-read assembly with wtdbg2. BioRxiv 2019, doi:10.1101/530972.
27. seqtk: Toolkit for processing sequences in FASTA/Q formats. https://github.com/lh3/seqtk, Accessed 22 Feb 2019.
28. Wood DE, Salzberg SL. Kraken: ultrafast metagenomic sequence classification using exact alignments. Genome Biol 2014;15(3):R46.
29. Vaser R, Sović I, Nagarajan N, et al. Fast and accurate de novo assembly from long uncorrected reads. Genome Res 2017;27(5):737–46.
30. medaka: Sequence correction provided by ONT Research. https://github.com/nanoporetech/medaka, Accessed 22 Feb 2019.
31. Walker BJ, Abeel T, Shea T, et al. Pilon: an integrated tool for comprehensive microbial variant detection and genome assembly improvement. PloS One 2014; 9(11):e112963.
32. minidot: Fast and pretty dotplots for whole genomes assemblies using minimap2 and R/ggplot2. https://github.com/SamStudio8/minidot, Accessed 22 Feb 2019.
33. Parks DH, Imelfort M, Skennerton CT, et al. CheckM: assessing the quality of microbial genomes recovered from isolates, single cells, and metagenomes. Genome Res 2015;25(7):1043–55.
34. Simpson JT, Workman RE, Zuzarte PC, et al. Detecting DNA cytosine methylation using nanopore sequencing. Nat Methods 2017;14(4):407–10.
35. Huson DH, Albrecht B, Baçi C, et al. MEGAN-LR: new algorithms allow accurate binning and easy interactive exploration of metagenomic long reads and contigs. Bioil Direct 2018;13(1):6.
36. Köster J, Rahmann S. Snakemake—a scalable bioinformatics workflow engine. Bioinformatics 2012;28(19):2520–2.
37. Nicholls SM, Quick JC, Tang S, et al. Supporting data for “Ultra-deep, long-read nanopore sequencing of mock microbial community standards”. GigaScience Database 2019. http://dx.doi.org/10.5524/100580.