Analysis procedures for assessing recovery of high quality, complete, closed genomes from Nanopore long read metagenome sequencing

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Abstract

New long read sequencing technologies offer huge potential for effective recovery of complete, closed genomes from complex microbial communities. Using long read (MinION) obtained from an ensemble of activated sludge enrichment bioreactors, we 1) describe new methods for validating long read assembled genomes using their counterpart short read metagenome assembled genomes; 2) assess the influence of different correction procedures on genome quality and predicted gene quality and 3) contribute 21 new closed or complete genomes of community members, including several species known to play key functional roles in wastewater bioprocesses: specifically microbes known to exhibit the polyphosphate– and glycogen–accumulating organism phenotypes (namely Accumulibacter and Dechloromonas, and Micropruina and Defluviicoccus, respectively), and filamentous bacteria (Thiothrix) associated with the formation and stability of activated sludge flocs. Our findings further establish the feasibility of long read metagenome–assembled genome recovery, and demonstrate the utility of parallel sampling of moderately complex enrichments communities for recovery of genomes of key functional species relevant for the study of complex wastewater treatment bioprocesses.

The development of long read sequencing technologies, such as the Oxford Nanopore Technology MinION and Pacific Biosciences SMRT are presenting new opportunities for the effective recovery of complete, closed genomes [1, 2]. While these new approaches have been mostly applied to single species isolates [3, 4], the ability of this new methodology to recover genomes of member taxa from complex microbial communities (microbiome) data is now actively being explored.

After long read sequencing technologies first became available, several studies pioneered the collection of long read data, or combined long and short read data, on complex microbial communities, for example from moderately to highly enriched bioreactor communities [5, 6], co–culture enrichments [7], marine holobionts [8] or from full scale anaerobic digester communities [9], as well as several datasets which provided benchmarking data from long and short read sequencing of mock communities [10, 11, 12]. New long read analysis methods [13, 14] and binning algorithms designed for long read metagenome data [15] have also appeared, anticipating the future expansion of metagenome data generated from these new instruments. More recent studies [16, 17, 18, 19, 20, 21, 22] have collectively established that full length (or near full length genomes) can be recovered from long read sequencing of complex communities, which sets the stage for further development of genome–resolved long read metagenomics.

Here we extend our previous work [16, 22] on recovering metagenome-assembled genomes from long read data obtained from enrichment (continuous culture) reactors inoculated with activated sludge microbial communities. Enrichment reactor communities [23, 24] offer a moderate level of complexity compared to the inoculum communities [25] and so are realistic, yet tractable, systems to use for developing approaches for recovery and vali-
Validation of MAG analysis using long read data. We report results and methodology of long read sequencing from multiple sets of reactor communities. We have obtained short read metagenome data (Illumina) from either the same DNA aliquots as used for long read sequencing, or the same biomass. Specifically we 1) describe new methods for validating long read assembled genomes using their counterpart short read metagenome assembled genomes; 2) assess the influence of different correction procedures on genome quality and predicted gene quality and 3) contribute 21 new closed or complete genomes of community members, including several species known to play key functional roles in wastewater bioprocesses.

Methods

Overview, biomass and data availability

We employed the biomass from a series of enrichment reactor microbial communities, each from activated sludge sourced from wastewater treatment plants located in Singapore. We sampled the following enrichment reactor communities:

(i) A lab-scale sequencing batch reactor, inoculated with activated sludge from a full scale wastewater treatment plant (Public Utilities Board, Singapore), was operated using acetate as the primary carbon source to enrich for polyphosphate accumulating organisms (PAO). The reactor was sampled on day 267 of the operation, with both long read (Nanopore MinION) and short read (Illumina Miseq 301bp PE) sequencing data from the same DNA aliquot. These data have been previously published by us [16] and are available via NCBI BioProject accession PRJNA509764. This data set is referred to below as the PAO1 data.

(ii) A lab-scale sequencing batch reactor, inoculated with activated sludge from a full scale wastewater treatment plant (Public Utilities Board, Singapore), was operated using alternative carbon sources to enrich for polyphosphate accumulating organisms (PAO). The reactor was sampled on April 6, 2018, gDNA extracted and both long read (Nanopore MinION) and short read (Illumina HiSeq2500 251bp PE) data obtained from the same DNA aliquot. These data are available available via NCBI BioProject accession PRJNA611629. This data set is referred to below as the PAO2 data.

(iii) Enrichment targeting putative PAO species, namely members of genera Tetrasphaera and Dechloromonas. Following inoculation with activated sludge from a full scale wastewater treatment plant (Public Utilities Board, Singapore), the reactor was fed with synthetic wastewater containing either glutamate or glucose as the main carbon source, with the feed type switched in a weekly manner, and operated at 31°C. Short read data (Illumina HiSeq2500 251bp PE) had been previously obtained from
sampled biomass on days 272, 279 and 286 of operation, and long read data (Nanopore MinION) obtained from samples taken on days 264 and 293 of operation. These data are available via NCBI BioProject accession PRJNA606905. The long read data obtained from each sampling day is referred to below as the PAO3A and PAO3B data, respectively.

(iv) Enrichment targeting PAO species capable of performing denitrification. A lab–scale sequencing batch reactor was inoculated with activated sludge from a full–scale wastewater treatment plant (PUB, Singapore). The reactors were operated at 35°C using acetate as the primary carbon source fed under anaerobic conditions, but without the addition of allyl–thiourea (ATU) in order to suppress the growth of ammonia oxidizing bacteria, with the aim of targeting polyphosphate–accumulating organisms that could also reduce nitrogen oxides (nitrite and/or nitrate). For this study we obtained long read data (Nanopore MinION) and short read data (Illumina HiSeq2500 251bp PE) from the same DNA aliquot extracted from biomass sampled on day 292 of operation. These data are available via NCBI BioProject accession PRJNA607349. These data are referred to below as the PAO4 data.

Using these data, our main objective was to obtain complete bacterial chromosomes, via assembly of long read data, and use draft genomes obtained from metagenome assembly of corresponding short read data for the purposes of evaluation. This approach takes advantage of the fact of our having obtained data from both sequencing modalities and takes advantage of current understanding of short read metagenome assembly binning and quality assessment procedures [26, 27]. We highlight that it may be possible to recover further genomes from these data by the use of binning procedures adapted to long read data, however here our focus is on analysing contigs directly obtained from the assembly that plausibly represent complete bacterial chromosomes.

**DNA extraction**

Genomic DNA in the case of the samples from PAO1, PAO3A, PAO3B and PAO4 was extracted from the sampled biomass as described previously by us [16], briefly, we used the FastDNA™ SPIN Kit for Soil (MP Biomedicals), using 2× bead beating with a FastPrep homogenizer (MP Biomedicals). Extracted gDNA from the PAO1, PAO3A and PAO3B samples was then size–selected on a BluePippin DNA size selection device (SageScience) using a BLF–7510 cassette with high pass filtering with a 8 kbp cut–off. The gDNA from the PAO4 sample was size–selected using Circulomics Short Read Eliminator XS kit (Circulomics Inc). Size–selected DNA was then taken for Nanopore library construction (see below).

From the biomass from PAO2 sampling, high molecular weight (HMW) DNA was extracted using a modified xanthogenate–SDS protocol [28]. Briefly, 2 mL of biomass from lab-scale sequencing batch reactor was harvested by centrifugation, the pellet was
resuspended in 0.6 mL of DNA/RNA shield (Zymo Research) and added to 5.4 mL of pre-heated (65°C) XSP buffer (1:1 volumes of XS buffer and phenol). The tubes were incubated at 65°C for 15 min, vortexed for 10–15 sec, placed on ice for 15 min and centrifuged at 14000 rpm for 5 min. The aqueous phase was transferred to a fresh tube and extracted with equal volume of phenol:chloroform:isoamyl alcohol (25:24:1) followed by extraction with chloroform:isoamyl alcohol (24:1). The aqueous phase after chloroform:isoamyl alcohol (24:1) extraction was ethanol precipitated and resuspended in TE buffer. The extracted DNA was further treated with RNase A (Promega) then extraction with phenol, followed by phenol:chloroform:isoamyl alcohol (25:24:1), and ethanol precipitation. Purified DNA was taken to library construction for Nanopore sequencing.

**Short read sequencing**

Genomic DNA Library preparation was performed using a modified version of the Illumina TruSeq DNA Sample Preparation protocol. We then performed a MiSeq sequencing run with a read length of 301 bp (paired-end) or a HiSeq2500 sequencing run with a read length of 251 bp (paired-end) as specified above.

**Long read sequencing**

Nanopore sequencing was performed on a MinION Mk1B instrument (Oxford Nanopore Technologies) using a SpotON FLO MIN106 flowcells and R9.4 chemistry. Data acquisition was performed using MinKNOW software, without live basecalling, running on a HP ProDesk 600G2 computer (64–bit, 16 GB RAM, 2 Tb SSD HD; Windows 10). The runs were continued until active pores in flowcells were depleted. For PAO1, PAO3A and PAO3B extractions, the sequencing library was constructed from approximately 4–4.5 µg of size–selected genomic DNA using SQK–LSK108 Ligation Sequencing Kit and approximately 900 ng of the library was loaded onto each flowcells. For PAO2 data set, sequencing libraries were constructed from HMW DNA using two different sequencing kits from ONT. The first kit was the Rapid Sequencing kit SQK–RAD004, for which the library was constructed from 400 ng of HMW DNA and the entire library loaded onto the flow cell. The second kit was the Ligation Sequencing kit SQK–LSK 108, for which 1.0 µg of genomic DNA was used for library construction, and 400 ng was loaded onto the flow cell. For PAO4 data set, the sequencing library was constructed from 1.2–1.3 µg of size selected DNA using SQK–LSK109 Ligation Sequencing Kit (Oxford Nanopore Technologies). The library was diluted to allow 250 ng of the library to be loaded on the flowcell.

**Analysis of long read sequence data**

Basecalling was performed with guppy (CPU version 3.2.1, 3.2.2 or 3.3.0 for Linux 64–bit machines; see Table S1). Adaptor trimming was performed using Porechop (version 0.2.2)
with default settings except \texttt{-v 3 -t 20}. We assembled long read data using \textit{Canu} (version 1.8 or 1.9, default settings except \texttt{corOutCoverage=10000, corHmapSensitivity=high, corMinCoverage=0, redMemory=32, oeaMemory=32 and batMemory=200 useGrid=false}) [30]. \textit{Unicycler} (version 0.4.7 or version 0.4.8 with default settings except \texttt{-t 20 --keep 3}) [31] and \textit{Flye} (version 2.4 with default settings except \texttt{-t 20 --plasmids --debug --meta}) [32]. Contigs generated from long read data are hereafter denoted as long read assembled contigs (LRAC). The number of reads used in each assembly was estimated by mapping long read to LRAC sequence with \textit{minimap2} (version 2.17) [33] and using \textit{samtools-1.6} to calculate the number of aligned reads [34]. We used \textit{DIAMOND} (version 0.9.24) [35] to perform alignment of LRAC sequences (with default settings except \texttt{-f 100 -p 40 --log --long-reads -c1 -b12}) against the NBCI–NR database (February, 2019) [36]. From the MEGAN Community Edition suite (version 6.17.0) [37] we used \textit{daa-meganizer} (run with default settings except \texttt{--longReads, --lcaAlgorithm longReads, --lcaCoveragePercent 51, --readAssignmentMode alignedBases} and the following settings for mapping files: \texttt{--acc2taxa prot_acc2tax-Nov2018X1.abin, --acc2eggnog acc2eggnog-Oct2016X.abin, --acc2interpro2go acc2interpro-June2018X.abin, --acc2vfdb acc2vfdb-Feb2019.map}) to format the .daa output file for use in the MEGAN GUI (version 6.17.0). Within MEGAN, LRAC sequences were exported with the ‘Export Frame-Shift Corrected Reads’ option to obtain frameshift corrected sequence. LRAC sequence that was at least 1Mb in length were categorised as potential whole chromosome sequence and from thereon described as LR–chr sequence. We processed LR–chr sequences with \textit{CheckM} (version 1.0.11) [38] and \textit{Prokka} (version 1.13) [39] to assess genome quality. LR–chr sequences that demonstrated \textit{CheckM-SCG} completeness 90% and contamination < 5% were classified as putative genomes. The entire set of putative genomes were dereplicated using the \textit{dRep} (version 2.2.3) workflow [40] with the following settings: \texttt{-p 44 -comp 90 -con 5 -str 100 --genomeInfo}. We performed taxonomic annotation on recovered genome sequence using \textit{GTDB-Tk} (version v0.3.2, running default parameters except \texttt{--cpus 40 -x fasta}) [41]. Coverage profiles were generated from both long read and short read data, by mapping each of these to LR–chr sequences using \textit{minimap2} (version 2.17) using the following flags \texttt{-ax map-ont} for long read data, and \texttt{-ax sr -a -t 20} for short read data. Sorted .bam files were subsequently processed using \textit{bedtools genomeCoverageBed} (version 2.26.0) with the following flags \texttt{-d}. We extracted 16S–SSU rRNA genes as identified with \textit{Prokka} and annotated them against SILVA database (SURREf_NR99_132_SILVA_13_12_17_opt.arb) [42] using \textit{sina-1.6.0-linux} [43] running default settings except \texttt{--t -v --log-file --meta-fmt csv} and with \texttt{--lca-fields} set for all five databases, namely \texttt{tax_slv, tax_embl, tax_ltp, tax_gg} and \texttt{tax_rdp}.

\textbf{Analysis of short read sequence data}

The raw FASTQ files were processed using \textit{cutadapt} (version 1.14) [44] with the following arguments: \texttt{--overlap 10 -m 30 -q 20,20 --quality-base 33}. We performed
metagenome assemblies from short read data using SPAdes (version 3.12.0-Linux or 3.14.0-
Linux, with default settings except -k 21,33,55,77,99,127 --meta) [45] either as single
sample assemblies, in the case of short read data from PAO1, PAO2 and PAO4, or as
co-assembly of all short read in the case of the PAO3A and PAO3B samples. The con-
tigs generated from short read data are hereafter denoted as short read assembled con-
tigs (SRAC). We identified putative member genomes using MetaBAT2 [46], after filtering
for contigs at least 2000 bp in length. We identified 16S genes within contigs using the
--search16 module of USEARCH (version 10.0.240, 64 bit) [47], and annotated them using
the SILVA::SINA webserver (using default parameters) [42, 43]. For each identified bin we
performed genome quality estimation using CheckM (version 1.0.11). We performed tax-
ononomic annotation on recovered member genomes using GTDB-Tk (version 0.3.2, running
default parameters except --cpus 40 -x fasta).

Comparative analysis of long and short read assemblies

We used BLASTN (version 2.7.1+) [48] to examine the degree of sequence alignment between
LRAC and SRAC sequences. We treated the LRAC as the subject sequences and the SRAC
as the query sequences, using default BLASTN parameters (except -outfmt 6). From the
BLASTN tabular output, we retained the highest bit–score from each unique combination of
query and subject pair. In order to identify the short reads bin(s) that are cognate to a given
LR–chr sequence, we then computed the concordance statistic (κ), as previously described
by us in [16], for all combinations of short read contigs (categorised by bin membership) and
LRAC sequence, that were present in the BLASTN output. We then compute the following
component statistics:

(1) \( \hat{\rho}_{id} \): The mean of the percent identity (PID), calculated across alignments, and
quantified as a proportion. \( \hat{\rho}_{id} \) is defined on the interval \([0,1]\)

(2) \( \hat{\alpha}_{2ql} \): The mean of the quotient of the alignment length to the query length, cal-
culated across alignments, and quantified as a proportion. \( \hat{\alpha}_{2ql} \) is always \( \geq 0 \) and
while values \( > 1 \) can be observed, in practice the maximum observed value be ap-
proximately 1.

(3) \( p_{srac} \): the quotient of the number of short read contigs in the bin that produce
alignments and the total number of short read contigs in the bin. \( p_{srac} \) is defined on
interval \([0,1]\)

(4) \( p_{aln} \): the proportion of the long read contig that is covered by an alignment. \( p_{aln} \) is
defined on the interval \([0,1]\).

Collectively these statistics contain information on how well a set of short read contigs will
tile a LR–chr sequence, namely, completeness of coverage (captured by \( p_{srac} \) and \( p_{aln} \)), as
well as quality of the alignments (captured by \( \hat{\rho}_{id} \) and \( \hat{\alpha}_{2ql} \)). We can hypothesise that if
the majority of the contigs in a short read MAG completely covered a LR–chr sequence
with high quality alignments, we would predict all four of these statistics would hold values
be close to unity. A simple extension of this prediction is to calculate the mean of the four
statistics, which we denote as the concordance statistic, \( \kappa = (\hat{\text{pid}} + \hat{\text{al}} + \hat{\text{aln}} + \hat{\text{srac}})/4 \),
which provides a single number to screen large numbers of pairwise combinations of short
read and long read derived MAG in an efficient way. The concordance statistic (\( \kappa \)) was
computed using all alignments, as well as after filtering for near–full length alignments
(defined as \( \text{alq} \geq 0.95 \)). We provide an R package \texttt{srac2lrac} to compute \( \kappa \) (along
with all component statistics) following calculation of \texttt{BLASTN}–like alignment statistics and
definition of short read bins.

Analysis of effects of frame-shift correction on coding sequence

The frameshift correction procedures employed from the \texttt{MEGAN-LR} package [13, 16] have
been crucial in permitting the use of genome quality and annotation workflows, namely
\texttt{CheckM} and \texttt{Prokka}, and we further evaluated the extent to which these procedures im-
proved accuracy of the coding gene sequence. To do so we analysed the distribution of
ratio of predicted gene length to the length of the nearest orthologous gene, as suggested
by Watson and colleagues [49], before and after the application of frameshift correction
procedures, as well as against two other sequence correction algorithms, namely \texttt{Medaka}
(version 0.11.5) [50] and \texttt{Racon} (version 1.4.3) [51]. In the first instance, we employ a single
round of correction and in the case of \texttt{Racon} do not use short read data for correction (in
order to maintain independence of each data type, as in the case of the concordance statist-
ic calculations). \texttt{MEGAN-LR} frameshift correction procedure uses the results of alignments
made against RefSeq NR, we did not compare against this same database to avoid positively
biasing the performance, rather we used predicted genes from the cognate short read assem-
bies as a database of genes to use as subject sequences. Specifically, we took the protein
coding sequence of each ORF in each of four versions of the genome generated above, and
performed homology search of each sequence against the short read assembly ORF database
using \texttt{DIAMOND} (version 0.9.24, running in blastp mode with default parameters except \(-f \)
6 \texttt{qseqid qlen slen ssseqid sallseqid -p 40 -v --log --max-target-seqs 1 \)). We
then calculated the quotient of the length of the query sequence to the length of subject
sequence holding the maximum bit–score (best hit). We ran \texttt{CheckM} on each of the four
versions of each putative genome, as described above. We also examined the common prac-
tice of applying multiple rounds of correction, with within and across, different correction
software, by performing both the above analyses on genomes corrected with four sequential
applications of \texttt{Racon} followed by one application of \texttt{Medaka} (denoted as ‘multiple’ from
hereon).
Procedures for refining draft genomes

In LR–chr sequence we screened regions of potential misassembly by identifying genomic intervals of at least 10bp in length, where long read coverage was either abnormally high or abnormally low, defined as >1.5 of the median coverage and <0.5 of the median coverage, respectively. We then examined alignments of both long and short read data to the genomes using the Integrated Genome Viewer (IGV version 2.4.14) [52] to identify low coverage regions that showed evidence of misconnection between reads, or weakly supported connection, or in the case of high coverage regions, to disambiguate types of read connections likely to arise from non-cognate sources. We generated VCF files for short read alignments using BCFtools (version 1.9 run with flags -mv) [53] to identify likely single nucleotide variants and presence of insertion/deletion variants, and subsequently used the aligned short read contig sequences to remove false nucleotide calls. We then align the entire genome against itself using BLASTN to check the integrity of the corrected genome sequence. For completeness, we have provided raw LR–chr sequence, frame-shift–corrected sequence and, for the subset of genomes subjected to further refinement, the fully completed versions.

Data availability

Raw sequence data is available at NCBI Short Read Archive (SRA) via BioProject accession identifiers listed above. The R code for performing the concordance statistic analysis are available at https://github.com/rbhwilliams/srac2lrac including test data and scripts taken from the PAO2 data.

A Zenodo submission (https://doi.org/10.5281/zenodo.3695987) contains key secondary data, including: 1) LRAC sequence from each dataset; 2) whole genome sequence from the 21 genomes listed in Table 1 for each of the five correction procedures (FASTA sequence, Prokka and CheckM results); 2) short read assembled sequence and binning results; 3) concordance statistic data and results; 4) short and long read per–base coverage data for the 21 genomes and 5) two manually corrected genomes of Candidatus Accumulibacter (also being submitted to NCBI) along with detailed notes explaining the procedures that were applied.

Results

Long read sequencing depth improved from the beginning of the study period, reflecting rapid improvement in experimental protocols and flow cell technology (Table S1), with the total amount of sequence generated ranging from around 1Gbp/run to just under 12Gbp/run (Table S1). These data were assembled using each of the three workflows as described above. The Canu assembly workflow generated a greater number of LR–chr (n=90) on these data than did either Unicycler (n=44) or Flye (n=60) (Table S2).
As Canu generated a substantially larger number of LR–chr sequences, we subsequently focused attention on the results obtained with this workflow (see Table S3 for comparative summary of LR–chr sequences from each workflow).

We next applied the truncated MIMAG criteria for estimating high quality MAG status (SCG–estimated completeness > 90% and contamination < 5%) and observed a total of 23 LR–chr generated from Canu that could be considered plausible candidates for being whole chromosomal sequence, from here on referred to as putative genomes for convenience. A further 13 LR–chr sequences from Canu were classifiable as medium quality (SCG–estimated completeness ≥ 50% and contamination < 10%, including one that was LR–chr was circular). We de–replicated the entire set of 23 putative genomes using the dRep workflow with a relatedness threshold of ANImf > 99 (Figure S1), obtaining a reduced set of 21 putative genomes (Table 1). The two redundant genomes were obtained from the PAO3A and PAO3B datasets, consistent with the fact that they are the same community sampled at different times.

We then studied each of these 21 dereplicated putative genomes in more detail to establish whether they were, or were not, likely to represent whole chromosomes. Using annotations from the Prokka workflow, all 21 putative genomes met the complete MIMAG criteria for being classified as high quality metagenome assembled genomes, including a minimum number of tRNA encoding genes, and the presence of each of the genes encoding 5S, 16S and 23S SSU–rRNA genes detected in each genome. Estimated SCG completeness was on average 95.87% (range 93.47–99.04%) and mean contamination was 0.37% (range 0.00–1.09%). Nine of the 21 sequences where classified as circular by Canu (Table 1). Coverage profiles generated using both long and short read data within a given community showed uniform coverage, with no substantive gaps observed (see Panel C of Figure 1 and Supplementary Figures 2–30). The proportion of long reads utilised to produce the putative genomes varied with dataset (Table S4) but from conservative estimation (based on subsetting alignments of all long reads against all LRAC sequence), on average 32.6% of reads across all 5 data sets (range: 23.3–55.4). At the individual genome level as few as 1% of reads in a dataset could generate a complete genome (PAO1–tig00000003; see Table S4).

To gain further insight into the quality and completeness of detected genomes, we used the concordance statistic (κ), previously developed by us [16] to identify metagenome–assembled genomes obtained from short read sequence data that were cognate to a long read assembled genome (summary data of each short read assembly is provided in Table S5). The κ–statistic is computed for all combinations of short read MAG and LR–chr sequences. A observed value of κ close to unity will imply that the LR–chr sequence is tiled by the contigs from the short read MAG, and the latter can be considered a likely candidate for being the cognate genome. For 20 of the 21 genomes in Table 1 the maximum observed κ values were high (mean: 0.95 range: 0.83–1.00) (Table S6). If we only considered near full length alignments (al2ql > 0.95), this reduced by around 0.5 units (mean 0.89, median 0.91, range: 0.80–0.97). In Figure 1 we provide a comprehensive visualisation of the
concordance statistic analysis for the case of the PAO3A–tig00018026 genome against its cognate short read MAG (bin 114). Related plots for all 21 genomes are available in the Supplementary Figures 2–30). We observed a genome recovered from PAO4 (PAO4–tig00000079), annotated at species level to *Exiguobacterium profundum*, which held a $\kappa$ value of 0.3 and from which there appeared to be no corresponding complete short read MAG (Supplementary Figure 29).

On average, for a given LR–chr sequence, $\kappa$–statistics were generated from around two thirds of available short read MAGs, but in most cases the magnitude of the $\kappa$–statistic itself was low. Of the four component statistics, $\hat{\text{pid}}$ and $\hat{p_{srac}}$ showed consistently higher values in the bulk of associations than either $\text{al}^2$ or $p_{aln}$, with the latter two measures provided greater visual discrimination between the short read MAG holding the maximum $\kappa$ value and the bulk distribution of (lower) $\kappa$ scores. As expected, cognate short read MAGs were generally drawn from among the most abundant members of a given reactor community. Contigs from short read bins with related taxonomy usually scored highly on one or more component scores (data not shown), but in combination, only one short read MAG generated a high value $\kappa$ score with component statistics that supported it being the cognate. In several cases, we observe two short MAG that tile two adjacent fragments of a single LR–chr sequence, which we determined to be due to underlying genome being split by MetaBat2 into two or more component sub–MAG (bin–splitting; see Supplementary Figures 2–3, 5–6, 10–11, 20–22).

Identification of probable mis-assembles among LR–chr sequences

Among the complete set of LR–chr we identified several examples of LR-chr that are clearly mis-assemblies. In the PAO3A data, we observed one contig (tig00000001; assembled by Canu) that appeared to be comprised of two separate complete genomes joined together (see Supplementary Figures 31–34 for further dissection). In this case, the proximal two thirds of the LR–chr arises from one genome, while distal third from another, as evidenced by different GC proportions and divergent short bin associations, respectively. In the case of the PAO4 data we observed several LR–chr that were classified by CheckM to have completeness over 90% but which demonstrated substantial degrees of contamination (namely tig00017984, tig00017990 and tig00017987 from Canu), most likely as the results of reads from closely related strains being combined.

Effect of sequence error correction on coding sequence and genome quality

Although the recovered genomes are consistent with being *bone fide* whole bacterial chromosomal sequence, the high error rate present in current nanopore–based sequencing implies these constructs may not meet current expectations of reference genome quality. Examining the length ratio histograms of the predicted genes from long read assembled genomes, against their best hit counterparts from the cognate short read assemblies, we observed
that application of any of the three sequence correction procedures provided some degree of improvement compared to the case of raw sequence, with an increased frequency of the length ratio being located around a value of unity (Figure 2 and Supplementary Figure 35). The performance of Racon was highly variable but always less effective than either MEGAN–LR or Medaka. MEGAN–LR generally provided the best performance, followed by the multiple procedure approach, than Medaka.

We further examined the influence of sequence correction on genome quality statistics, as estimated by CheckM (Table 2). Of the 21 frame–shift corrected genomes classifiable as high quality (Table 1), 3, 13, 7 and 16 of these were also classifiable as high quality when examined in their uncorrected, Medaka–corrected, Racon–corrected and multiple procedure corrected forms (Table 2), with the mean completeness being 76.5% (range: 41.9–93.0), 91.7% (range: 78.0–98.0), 86.8% (range: 66.0–97.0) and 92.0% (range: 77.2–98.0), respectively, compared to a mean of 95.3% (range: 92.6–99.0) in the case of MEGAN–LR. Contamination was never observed to be greater than 3% in any version of the 21 genomes.

Taxonomic analysis of recovered genomes

We inferred taxonomy of the recovered genomes using GTDB–Tk, as provided in Table S7 and summarised below ( additionally we provide a complementary analysis of recovered 16S-SSU rRNA gene sequence annotated against the SILVA database in Table S8, and GTDB–Tk annotations for all short reads bins in Table S9). Of the 21 long read genomes, 5 had sufficiently high degree of similarity to be classified to species level and 10 to genus level, 5 to family level and 1 to class level.

We recovered genomes of four taxa that hold known relevance to wastewater bioprocess, namely two genomes from the PAO species Accumulibacter: the PAO1–tig00000001 genome was closely related to Candidatus Accumulibacter sp. SK–02 [54], and found in our previous analysis of the PAO1 data [16], and the other (PAO2–tig00000001) related to Ca. Accumulibacter sp. BA–94 [54]; Figure 3) and a short read MAG previously recovered by us and denoted as Candidatus Accumulibacter clade IIF Strain SCSELSE–1 [55].

We recovered two genomes for Dechloromonas, generally considered as exhibiting the PAO phenotype [56], and one of genus Micropruina, previously shown to exhibit the glycogen accumulating organism (GAO) phenotype [57, 58]. The PAO1–tig00026549 sequence, annotated to the novel GTDB–derived family 2–12–FULL–67–15 and harbouring a 16S gene annotated to Defluviicoccus (Table S8), represents a novel member of the latter genus, whose members exhibit the GAO phenotype [59]. We also recovered a genome from a member of genus Thiobrix, a filamentous bacterium associated with the maintenance of floccular structure in activated sludge biomass [60, 61].

A set of four genomes recovered here have been previously identified in temperate climate activated sludge, namely 3 members of the CFB group recovered from the PAO1 data, OLB8 [62], OLB11 and OLB12 [62], as well as a genome classified to the Rhodobacteraceae genus UBA1943 [63]. Thiobacillus has been previously identified in activated sludge from
industrial wastewater treatment plants [64, 65]. In the PAO4 community, we recovered a genome close to that of *Exiguobacterium profundum*, originally discovered in deep-sea hypothermal vents [66]. Members of this genus, namely *Exiguobacterium alkaliphilum* and *Exiguobacterium* sp. YS1, have been studied in relation to treatment of high alkaline brewery wastewater and solubilisation of waster activated sludge, respectively [67, 68]. The genome of a member of family *Parachlamydiaceae*, an environmental Chlamydia [69] that was previously recovered by us [16] in the PAO1 data, and is probably a symbiont species of protists that are known to inhabit activated sludge [70]. The genome of *Bre-vundimonas* was closely related to a short read MAG previously obtained from an activated sludge metagenome in Hong Kong [71], and members of this genus have been observed previously in activated sludge systems [72], where they have been associated with quinoline degradation from coking wastewater [73].

A genome from a member of genus *Pseudoxanthomonas* was also recovered. The remaining genomes had no close references, and likely represent novel members of the microbial groups, namely family *Nocardioidaceae* (tig00157979 from PAO3B), from class *Anaerolineae* (tig00018026 from PAO3A), family *Burkholderiaceae* (tig00000024 from PAO3B), and a genome from the novel UBA6002 family (tig00000117 from PAO1).

Further refinement of genomes of *Candidatus Accumulibacter*

We applied manual refinement procedures, as described in Materials and Methods, to the two recovered genomes of *Candidatus Accumulibacter*, namely PAO1–tig00000001 and PAO2–tig00000001, in order to obtain submission quality finished genomes. Detailed notes on refinement and manually curation procedures are provided in the Zenodo submission.

Discussion

In this paper we explore how long read metagenome data, generated by a Nanopore MinION sequencer, can enhance the recovery of member genomes of microbial communities. Building on our previous analyses [13, 16, 22], we obtain further data from activated sludge enrichment bioreactor communities, and obtain 21 non-redundant complete genomes, of which nine are closed (circular) and six are from species with key functional relevance to wastewater bioprocesses. Additionally, we present further details of methodology for assessing whether genomes obtained from short read assemblies recapitulate those obtained from assembled long read data (the concordance statistic, briefly introduced by us in [16]), and examine aspects of genome quality not previously covered, including the quality of gene level coding sequence and the sequence rising from the mis-assembly and related artefacts. These new results highlight that by using long read sequencing in microbial communities of moderate complexity, it is clearly feasible to capture sequence constructions that are close to the requirements of high quality, closed genomes, for the most abundance community members, without the use of contig binning procedures. However careful evaluation
of such genomes still appears mandatory to assess quality and the presence of artefactual constructs.

The widespread use of metagenome-assembled genomes (MAG) methodology on short read metagenome data has provided a tremendous number of new draft genomes from diverse microbiomes and microbial communities (for example [20, 54] among others). However substantial limitations of these approaches have become evident, including problems related to the use of multi-sample co-assemblies [20, 74, 75], the challenges of resolving genomes to strain level [76], difficulties related to extracting MAGs from communities of high ecological complexity [77, 78], and the limitations of automated binning procedures, requiring careful evaluation of recovered genomes [79]. In response to these challenges, recent efforts have combined short read with emerging complementary techniques such as HiC metagenomics [80, 81, 82], synthetic long reads [83, 84], or long read sequencing, and collectively these results suggest substantial improvement can be made in the quality and completeness of metagenome assembled genomes using multiple types of sequence data. In the present study, we make use of DNA extractions that are co-assayed with both long and short sequencing, or DNA extractions from sampling events close enough together in time, that we can discount the influence of the ecogenomic differences as major influence on any observed differences between the types of sequence data.

Our analysis proceeds on the basis that neither short read nor long read data can be assumed to provide an accurate reference genome, and so we seek to understand and characterize the degree of agreement between assembled sequence generated from each data source. Although error prone MinION sequence can be corrected using higher quality short read sequences [41, 42], we have deliberately kept the two sources of data separate so as to not introduce any positive bias in the calculation of the concordance statistics. The concordance statistic was developed to provide a straightforward screening procedure for identifying short read MAG that are cognate to assembled genomes from long read data, by capturing information from alignment statistics. The concordance statistic also may have broader utility, for example we can observe several instances of 'split' bins from the short read assembly that are cognate to a given long read assembled genome, and cases where assembled long read sequence is demonstrably artefactual. We highlight that the concordance statistics capture more information than are contained in dot-plots (which require the imposition of arbitrary decision thresholds on alignment statistics), albeit at the cost of increased complexity. We provide R code for computing the concordance statistics from alignment results, and example workflows for visualisation.

While these are clearly vast improvements on the working models of genomes available from short read MAG analysis, several problems are still present that require attention and/or explicit correction. Firstly, the high error rate implicit in MinION sequence (not less than 5% sequencing error [85]) requires correction procedures to be applied either pre- or post-assembly. In the present case, we are relying on the frameshift correction algorithm implemented in MEGAN-LR [13, 16], which appears to perform slightly better than the next best correction procedure ('multiple'). As previously discussed [16], this correction
procedure permit the application of existing genome quality workflows (CheckM in the present case), and the resulting sequence can be considered to be at least a high quality assembly under currently accepted criteria (MIMAG as defined in [27]). However further analysis of the corrected gene content suggests there remains a substantial proportion of genes that remain inadequately corrected when compared against genes predicted from the cognate short read assemblies. Because the MEGAN-LR correction is dependent on aligned sequence from database comparisons, a combination of false positives alignments and a lack of closely related reference genomes could result in inappropriate or inadequate correction of the query sequences in our analysis, and additionally mis-assembly of genes in the short read assembly (subject sequences) could also be a factor in patterning these findings. A second factor relates to the inclusion of artefactual sequence (mis-assembly), which we identify and remove using examination of read alignment and coverage profiles, in line with recent calls for the continuing need for careful evaluation of the output of automated genome recovery procedures [79]. Collectively these results indicate that long read sequencing technology that harbours high error rates should be considered complementary to short read sequencing for the foreseeable future, with self-evident implications for experimental design choices.

We have deliberately focused on long read assembled contigs that form single contiguous sequences that are consistent with being whole bacterial chromosomes, which plays to the full strengths of long read sequencing. The remaining, much larger set of contigs, that do not meet our criteria for being considered putative genomes, will be in part comprised of genome fragments that could be recovered into draft genomes using binning methods. Although the amount of long read metagenome data collected from microbial communities of high to very high complexity is only just emerging [21], recent work on combining short and long read data from human fecal microbiomes [19] suggests that binning procedures will have to be developed, or adapted from short read methods, for the full potential of these new hybrid data to be realised.

In the present study, we are able to draw strength from the fact that the communities under study are of moderate complexity, and, in ecological terms are of low evenness, compared to the source inoculum, namely activated sludge residing in full scale wastewater treatment plants [25]. This suggests that one way to approach a systematic genome-resolved dissection of such complex communities would be to simply sample a diverse array of such enrichment communities, rather than rely on more deeper, expensive sequencing of a limited number of highly complex source communities. While such an approach may miss some relevant species (due to the biases of enrichment protocols), it would permit the recovery of many near-finished genomes from key species of direct functional relevance to wastewater bioprocess engineering, as obtained here.
List of Supplementary Materials

List of Supplementary Tables (.xlsx format)

- Supplementary Table 1: Summary of long read data.
- Supplementary Table 2: Summary statistics for long read assemblies from three assembly workflows.
- Supplementary Table 3: Comparison of number and CheckM–derived genome quality statistics of LR–chr sequences generated by three assembly workflows.
- Supplementary Table 4: Estimation of long read read count used for assembly of recovered genomes.
- Supplementary Table 5: Summary statistics for short read assemblies.
- Supplementary Table 6: Summary of concordance analysis for recovered genomes.
- Supplementary Table 7: Taxonomic annotation of genomes using GTDB–Tk.
- Supplementary Table 8: Taxonomic annotation of genomes from 16S sequence.
- Supplementary Table 9: Taxonomic annotation of short read MAG using GTDB–Tk.

Supplementary Figures

- Supplementary Figure 1: Tree generated from MASH distances between 23 LR–chr classifiable as putative genomes and used to undertake genome dereplication.
- Supplementary Figures 2–30: Summary of concordance statistic analysis for all 21 genomes listed in Table 1.
- Supplementary Figures 31–34: Summary of concordance statistic analysis for an artefactual LR–chr sequence (PAO3A–tig000000001).
- Supplementary Figures 35: Figure 2 presented with a logarithmic scale on the vertical axis.

Author contributions

The study was designed by R.B.H.W and I.B. R.E.Z.M, S.R, G.L.Q, Y.Y.L and S.W setup and operated enrichment reactors, and obtained samples with I.B. I.B and F.L designed long read sequencing experiments and I.B performed DNA extractions and performed long read sequencing. D.I.D–M obtained short read sequencing data. K.A, M.A.S.H, D.H.H.,
X.H.L and R.B.H.W designed analyses, performed data analysis and/or wrote analysis code. All authors contributed to data interpretation. R.B.H.W wrote the manuscript with specific contributions from all other authors. K.A and I.B made equal contributions to this work.

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Table 1: Summary statistics for 21 putative genomes recovered in this study

| Genome identifier | Length (bp) | #CDS | #rRNA | #tRNA | Completeness | Contamination | Taxonomic annotation |
|-------------------|-------------|------|-------|-------|--------------|---------------|-----------------------|
| PAO1–tig00000001  | 5,190,177   | 5,116| 2     | 53    | 95.28        | 1.11          | s_Accumulibacter sp000584975 |
| PAO1–tig00000003  | 4,268,816   | 5,123| 1     | 36    | 92.82        | 0.25          | g_OLB11               |
| PAO1–tig00000117  | 2,656,706   | 3,153| 1     | 38    | 97.53        | 0.15          | f_UBA6002             |
| PAO1–tig00026549  | 4,352,448   | 4,225| 1     | 46    | 94.64        | 0.50          | f_2-12-FULL-67-15     |
| PAO1–tig00026557  | 3,138,394   | 3,619| 2     | 44    | 94.19        | 1.58          | f_Parachlamydiaceae   |
| PAO1–tig00026560  | 4,262,704   | 4,373| 2     | 37    | 93.99        | 0.55          | g_ELB16-189           |
| PAO1–tig00198536  | 3,913,768   | 3,521| 2     | 38    | 92.57        | 1.98          | s_OLB8 sp001567405   |
| PAO2–tig00000001  | 5,027,886   | 4,558| 2     | 46    | 93.85        | 0.00          | g_Accumulibacter      |
| PAO2–tig00000013  | 3,452,123   | 3,382| 2     | 46    | 94.49        | 0.18          | g_Dehchloromonas      |
| PAO3A–tig00000003 | 3,666,458   | 3,454| 1     | 53    | 94.56        | 0.28          | g_Micropruna          |
| PAO3A–tig00000024 | 2,740,818   | 2,753| 1     | 44    | 96.92        | 0.00          | s_Brevundimonas sp002426005 |
| PAO3A–tig00000209 | 3,302,829   | 3,190| 1     | 47    | 95.38        | 0.00          | f_Nocardioidaceae     |
| PAO3A–tig00018026 | 4,685,957   | 4,742| 1     | 48    | 93.47        | 1.09          | c_Aeroliraeae         |
| PAO3A–tig00139797 | 3,548,924   | 3,508| 2     | 47    | 99.04        | 0.48          | s_Thiobacillus sp00189930 |
| PAO3B–tig00000024 | 3,282,734   | 2,937| 2     | 51    | 98.10        | 0.23          | f_Burkholderiaceae    |
| PAO3B–tig00000027 | 3,375,962   | 3,179| 1     | 53    | 93.16        | 0.42          | g_Rhodoblastus        |

a As predicted using Prokka (see Methods). The rRNA count denotes presence of three genes encoding 5S, 16S and 23S sequences.
b Genome quality estimates from CheckM (see Methods).
c Taxonomic assignments from GTDB–Tk (see Methods).
d Sequence classified as circular by Canu.
Table 2: Influence of sequence procedures on CheckM–derived genome quality statistics

| Genomes | Completeness* | Contamination* |
|---------|---------------|----------------|
|         | Uncorrected   | MEGAN  | Medaka\(^b\) | Racon\(^c\) | Multiple\(^d\) | Uncorrected | MEGAN | Medaka\(^b\) | Racon\(^c\) | Multiple\(^d\) |
| PAO1–tig00000001 | 84.28 | 95.28 | 93.91 | 87.88 | 94.71 | 0.66 | 1.11 | 1.59 | 1.62 | 1.11 |
| PAO1–tig00000003 | 64.34 | 92.82 | 84.84 | 78.21 | 87.74 | 1.56 | 0.25 | 0.90 | 0.74 | 0.74 |
| PAO1–tig000000117 | 82.46 | 97.53 | 94.82 | 90.99 | 94.82 | 0.15 | 0.15 | 0.15 | 0.15 | 0.15 |
| PAO1–tig00026549 | 86.57 | 94.64 | 97.13 | 91.45 | 96.73 | 0.50 | 0.50 | 0.50 | 0.50 | 0.50 |
| PAO1–tig00026557 | 75.56 | 94.19 | 95.24 | 86.16 | 95.58 | 2.70 | 1.58 | 1.58 | 1.58 | 1.58 |
| PAO1–tig00026560 | 71.51 | 93.99 | 94.24 | 86.70 | 94.52 | 1.59 | 0.55 | 0.55 | 0.55 | 0.55 |
| PAO1–tig00198536 | 74.83 | 92.57 | 94.64 | 88.68 | 94.39 | 2.74 | 1.98 | 2.48 | 2.23 | 2.48 |
| PAO2–tig00000001 | 69.00 | 93.85 | 89.33 | 83.64 | 90.88 | 0.03 | 0.00 | 0.03 | 0.03 | 0.03 |
| PAO2–tig00000013 | 45.13 | 94.49 | 89.16 | 86.32 | 91.52 | 0.00 | 0.18 | 0.59 | 0.38 | 0.12 |
| PAO3A–tig00000003 | 70.75 | 94.56 | 93.78 | 83.48 | 93.52 | 0.10 | 0.28 | 0.50 | 0.10 | 0.50 |
| PAO3A–tig00000024 | 70.79 | 94.69 | 89.46 | 84.66 | 89.86 | 0.81 | 0.00 | 0.32 | 0.32 | 0.32 |
| PAO3A–tig00000209 | 87.00 | 95.38 | 96.80 | 94.60 | 97.32 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| PAO3A–tig0018026 | 41.99 | 93.47 | 81.67 | 66.02 | 77.17 | 0.28 | 1.09 | 1.09 | 1.27 | 1.37 |
| PAO3A–tig00139797 | 91.56 | 99.04 | 97.05 | 97.02 | 97.05 | 0.48 | 0.48 | 0.55 | 0.48 | 0.48 |
| PAO3B–tig00000024 | 89.02 | 98.10 | 93.63 | 90.53 | 91.71 | 0.23 | 0.23 | 0.23 | 0.00 | 0.39 |
| PAO3B–tig00000027 | 73.78 | 93.16 | 85.11 | 81.80 | 87.97 | 0.42 | 0.42 | 0.42 | 0.16 | 0.73 |
| PAO4–tig00000001 | 93.04 | 98.64 | 97.97 | 96.58 | 97.95 | 1.90 | 2.07 | 2.07 | 2.06 | 2.06 |
| PAO4–tig00000030 | 76.77 | 97.10 | 91.63 | 85.14 | 91.87 | 1.30 | 2.30 | 1.52 | 1.63 | 1.11 |
| PAO4–tig00000046 | 91.85 | 96.30 | 97.64 | 94.90 | 97.40 | 0.47 | 0.47 | 0.47 | 0.50 | 0.47 |
| PAO4–tig00000079 | 80.26 | 94.79 | 89.66 | 89.85 | 90.87 | 0.00 | 0.00 | 0.66 | 0.72 | 0.66 |
| PAO4–tig00000228 | 70.80 | 94.81 | 77.98 | 77.13 | 78.18 | 0.02 | 0.00 | 0.08 | 0.00 | 0.00 |

Genomes shown in bold show CheckM completeness >90% and contamination < 5%.

\(^a\) Genome completeness and contamination estimates obtained from CheckM (see Materials and Methods).

\(^b\) Medaka: Single round of correction applied to each uncorrected genome sequence.

\(^c\) Racon: Single round of correction applied to each uncorrected genome sequence, run with default parameters for all datasets.

\(^d\) Multiple: four sequential applications of Racon (run with default parameters using long read data) and then one application of Medaka (run with default parameters; the following models were used \(-m r941\_min\_high\_g303\) for PAO1, PAO2, PAO3A and PAO3B and \(-m r941\_min\_high\_g330\) for PAO4)
Figure 1: Summary of concordance statistic analysis for an LR–chr (tig00018026) from the PAO3A reactor community (annotated to class *Anaerolineae*) showing close relationship to a short read metagenome assembled genome from the same reactor community (bin 114). (A): Distribution of $\kappa$-scores for tig00018026 against 242 bins recovered from the corresponding short read assembly. Bin 114 has the highest $\kappa$ at 0.97; (B): coverage–GC plot for the short read assembly, with bin 114 highlighted (closed black circles and dark grey convex hull; other bins highlighted by light grey convex hulls); (C): short read (SR, black crosses) and long read (LR, grey crosses) coverage profiles across tig00018026. (D–F): BLASTN statistics for alignments of short read contigs (bin 114) against tig00018026. Horizontal segments show alignment position on LR–chr and height of segment is value of corresponding statistic ($y$-axis) namely percent identity (PID) ($D$), the ratio of alignment length to query length ($al2q_l$) ($D$) and $\log_{10}$ bitscore ($E$). (F): GC content as a function of position on tig00018026 (grey closed circles, computing in adjacent windows of length 46700 bp) and for aligned short read contigs (black closed circles); (G–K): distribution of four component statistics of $\kappa$ (see Methods), with the position of the top scoring short read bin highlighted. (G): proportion of short read contigs in bin aligned to LR–chr ($q_{\text{LR}}$); (H): mean percent identity ($giid$); (I): mean ratio of alignment length to query length $al2q_l$ and (K): proportion of the long read contig that is covered by an alignment ($p_{\text{aln}}$).
Figure 2: Density estimates for the length ratio statistics, computed from the length of predicted genes in long read assemblies (query) and length of their best hit counterparts in cognate short assemblies (subjects), and categorised by type of sequence correction employed (from left to right, raw assembled sequence [uncorrected], frame-shift correction using MEGAN–LR, sequence correction using Medaka, sequence correction using Racon and application of the multiple procedure approach. Results from individual recovered genomes are highlighted by colour, and x-axis truncated at 2.5 units. A version with a log-scale on the vertical axis is provided in Supplementary Figure 35.
Figure 3: Dendrogram generated from MASH distances between draft genomes of Candidatus Accumulibacter, including two genomes recovered in the present study. Genomes from genera Thauera, Azoarcus and Dechloromonas were used as an outgroup. Underscore separated number in brackets refers to dRep secondary cluster assignments (two genomes are in the same secondary cluster if their ANImf ≥ 99). Note the structure of the tree recapitulates previously defined clade associations (Clade IF: BA94, SK11, SK12; Clade IC: BA91, SK02, SK01; Clade I: BA92 and BA93. With UW1 being a singleton for Clade IA). Genome references as follows, from top of tree: SCELSE1 (GCA_005524045.1); BA91 (GCA_000585095.1); UBA2327 (GCA_002345025.1); SK11 (GCA_000584995.1); UBA8770 (GCA_003487685.1); UBA11070 (GCA_003535635.1); UBA9001 (GCA_003542235.1); UBA2315 (GCA_002345285.1); SK02 (GCA_000585015.1); HKU2 (GCA_000584975.1); CANDO2 (GCA_009467855.1); CAMEJO (GCA_003332265.1); UBA2783 (GCA_002352265.1); UBA6658 (GCA_003487685.1); BA93 (GCA_000585075.1); CANDO1 (GCA_009467885.1); Ca. Accumulibacter aalborgensis (GCA_900089955.1); UBA704 (GCA_002304785.1)