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Authors

Chen, Lin-Xing Anantharaman, Karthik Shaiber, Alon <u>et al.</u>

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Accurate and complete genomes from metagenomes

Lin-Xing Chen,¹ Karthik Anantharaman,^{1,7} Alon Shaiber,^{2,3} A. Murat Eren,^{3,4} and Jillian F. Banfield^{1,5,6}

¹ Department of Earth and Planetary Sciences, University of California, Berkeley, California 94720, USA; ²Graduate Program in Biophysical Sciences, University of Chicago, Chicago, Illinois 60637, USA; ³Department of Medicine, University of Chicago, Chicago, Illinois 60637, USA; ⁴Bay Paul Center, Marine Biological Laboratory, Woods Hole, Massachusetts 02543, USA; ⁵Department of Environmental Science, Policy, and Management, University of California, Berkeley, California 94720, USA; ⁶Earth and Environmental Sciences, Lawrence Berkeley National Laboratory, University of California, Berkeley, California 94720, USA

Genomes are an integral component of the biological information about an organism; thus, the more complete the genome, the more informative it is. Historically, bacterial and archaeal genomes were reconstructed from pure (monoclonal) cultures, and the first reported sequences were manually curated to completion. However, the bottleneck imposed by the requirement for isolates precluded genomic insights for the vast majority of microbial life. Shotgun sequencing of microbial communities, referred to initially as community genomics and subsequently as genome-resolved metagenomics, can circumvent this limitation by obtaining metagenome-assembled genomes (MAGs); but gaps, local assembly errors, chimeras, and contamination by fragments from other genomes limit the value of these genomes. Here, we discuss genome curation to improve and, in some cases, achieve complete (circularized, no gaps) MAGs (CMAGs). To date, few CMAGs have been generated, although notably some are from very complex systems such as soil and sediment. Through analysis of about 7000 published complete bacterial isolate genomes, we verify the value of cumulative GC skew in combination with other metrics to establish bacterial genome sequence accuracy. The analysis of cumulative GC skew identified potential misassemblies in some reference genomes of isolated bacteria and the repeat sequences that likely gave rise to them. We discuss methods that could be implemented in bioinformatic approaches for curation to ensure that metabolic and evolutionary analyses can be based on very high-quality genomes.

[Supplemental material is available for this article.]

In an opinion paper published relatively early in the microbial genomics era, Fraser et al. (2002) stated "you get what you pay for." The investigators argued the lower scientific value of draft (partial) versus complete genomes, noting for example higher error rates, potential contaminant sequences, loss of information about gene order, lower ability to distinguish additional chromosomes and plasmids, and most importantly, missing genes. Despite the clarity of this view, the field moved toward the generation of draft isolate genomes to optimize the rate of supply of new sequence information and to lower the cost. Genome-resolved metagenomics has almost exclusively settled for uncurated draft genomes, now often referred to as metagenome-assembled genomes (MAGs). A summary of the basic methods for generating MAGs was provided by Sangwan et al. (2016). A more recent review provides an overview of assembly methods and offers some insights into the complexity of genome recovery from metagenomes and a valuable overview of certain types of assembly errors that can occur (Olson et al. 2017).

The first MAGs were published in 2004 (Tyson et al. 2004), and there are now hundreds of thousands of them in public databases. The ever increasing depth of high-throughput sequencing now makes even the most challenging environments with low archaeal, bacterial, and viral biomass, such as insect ovaries

⁷Present address: Department of Bacteriology, University of Wisconsin, Madison, WI 53706, USA Corresponding authors: jbanfield@berkeley.edu,

meren@uchicago.edu

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(Reveillaud et al. 2019), human gut tissue biopsies (Vineis et al. 2016), hospital room surfaces (Brooks et al. 2017), and even human blood (Moustafa et al. 2017) amenable to shotgun metagenomic surveys and recovery of MAGs. Although incomplete, draft MAGs represent a major advance over knowing nothing about the genes and pathways present in an organism and led to the discovery of new metabolisms. For example, the complete oxidation of ammonia to nitrate via nitrite (i.e., comammox) was determined by the detection of necessary genes in a single MAG (Daims et al. 2015; van Kessel et al. 2015). MAGs are often derived from uncultivated organisms that can be quite distantly related to any isolated species, which is a clear advantage of MAGs (e.g., Becraft et al. 2017). For this reason, genome-resolved metagenomics has been critical for more comprehensive descriptions of bacterial and archaeal diversity and the overall topology of the Tree of Life (Hug et al. 2016).

Counter to this view, there is some sentiment that MAGs are not useful because they are composites and thus not representative of their populations (Becraft et al. 2017). However, a genome reconstructed from a clonal microbial culture also does not represent the cloud of biologically important variation that exists in the natural population from where the isolate was derived. Population diversity can be analyzed by comparing all individual sequences (or short reads) to the metagenome-assembled reference genome (Simmons et al. 2008; Delmont et al. 2019). Although some populations are near-clonal, others are very complex strain mixtures, and yet others fall on the continuum between these (Lo et al.

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2007; Chivian et al. 2008; Simmons et al. 2008). As strain divergence leads to assembly fragmentation (expanded on below), high-quality genomes are unlikely to be generated for relatively heterogeneous populations. Assembly of exceptionally long fragments (i.e., >100 kbp) from short-read data (i.e., <500 bp) is only anticipated when within-population diversity is low, as may occur following a recent bloom, selective sweep, or because of recent colonization by a single cell or a small cluster of closely related cells. In such cases, the genomes that assemble well are typically highly representative of the population from which they are derived, and the vast majority of reads report the same base at the same position. For example, in one recently published complete 4.55 Mbp genome (Banfield et al. 2017), the frequency of single-nucleotide variants (SNVs) is ~ 0.12% (Fig. 1), not substantially different from the expected sequencing error rate (0.04%-0.12%) (Schirmer et al. 2016).

Assembly and binning are important steps in metagenomic studies

Assembly of short metagenomic reads into contiguous segments of DNA is a computationally intensive task, and its effectiveness often depends on the complexity of the environment (Sharon and Banfield 2013). However, assembly of contigs/scaffolds offers many advantages over short-read-based analyses. First, they enable the identification of complete open reading frames. Second, assemblies provide larger genomic contexts (e.g., operons). In combination, these considerations improve predictions of metabolic capacities. Further, assembled sequences provide information about gene synteny and better resolve taxonomic profiles (e.g., by providing sets of proteins for taxonomy based on concatenated proteins encoded in the same genome (e.g., Hug et al. 2016; Parks et al. 2018). These improvements can overcome misleading interpretations of short-read data (Ackelsberg et al. 2015; Afshinnekoo et al. 2015). We acknowledge, however, that reads-based analyses may be informative for undersampled populations, although the information carried on each read is limited for short-read data.

The critical step required to establish a genome from a metagenomic assembly is binning. This involves assignment of assembled fragments to a draft genome based on detection on any scaffold of some signal(s) that occur(s) locally within a genome and persists genome-wide. Most commonly used features that can facilitate accurate binning of scaffolds include depth of sequencing measured by read coverage; sequence composition measured, for example, by tetranucleotide composition; and phylogenetic profile measured by the "best taxonomic hits" for each predicted protein on each scaffold. Sometimes, and mostly in data sets from very simple communities or for highly abundant organisms, the process of binning can be as easy as collecting together all fragments that share a single clearly defined feature (Supplemental Fig. S1), such as a discrete set of scaffolds with similar coverage, or unique and well-defined tetranucleotide patterns or GC content. In other cases, a combination of a few well-defined signals, such as GC content, coverage, and phylogenetic profile of scaffolds, are sufficient to clearly define a bin. However, over-reliance on phylogenetic profiles can be misleading, especially if the genome is for an organism that is only distantly related to those in the databases used for profiling. Further, some fragments can have an unexpected phylogenetic profile relative to the rest of the genome because the region has not been encountered previously in genomes of related organisms, possibly because it was acquired by lateral gene transfer. Thus, the most robust bins will draw on a combination of multiple clear signals.

If a study includes a set of samples with related community membership, an important constraint for bin assignment can be provided by the shared patterns of abundance of a fragment across a sample series. The use of series samples (collected at different depths/time points/treatments from the same experiment setup) data for binning was first proposed by Sharon et al. (2013), and

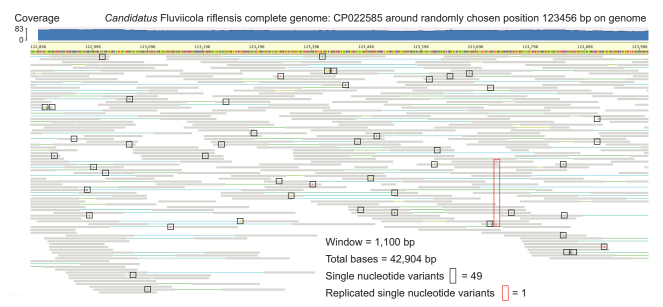


Figure 1. The low frequency of single-nucleotide variants (SNVs) of a recently published CMAG. A randomly chosen region, centered on position 123,456 (1100 bp in length) of the CMAG of *Candidatus* Fluviicola riflensis is shown with mapped reads (Banfield et al. 2017). SNVs that only occur once are indicated by black boxes, and the one replicated SNV is indicated by a red box. Clearly, the consensus sequence is well supported. The mapping of reads to the genome was performed by Bowtie 2 and visualized via Geneious.

this strategy is now a central feature in most automated binning algorithms, including CONCOCT (Alneberg et al. 2014), MaxBin (Wu et al. 2014), ABAWACA (Brown et al. 2015), and MetaBAT (Kang et al. 2015), as well as manual binning and MAG refinement strategies (Wrighton et al. 2012; Shaiber and Eren 2019). Seriesbased binning can exclude contaminant scaffolds from a MAG whose abundance shows a different pattern over time/space/treatment. We have found that no single binning algorithm is the most effective for all sample/environment types or even for all populations within one sample. The recently published method DAS Tool tests a flexible number of different binning methods, evaluates all outcomes, and chooses the best bin for each population (Sieber et al. 2018). A similar strategy has been used in a modular pipeline software called MetaWRAP (Uritskiy et al. 2018).

A case study: Binning can greatly improve data interpretation

Contigs that do not represent entire chromosomes may not be appropriate proxies for microbial populations without binning, and claims made based on unbinned contigs can lead to erroneous conclusions. For instance, a recent study focusing on human blood used shotgun metagenomic sequencing of circulating cell-free DNA from more than 1000 samples and recovered a large number of contigs with novel bacterial and viral signatures (Kowarsky et al. 2017), suggesting that "hundreds of new bacteria and viruses" were present in human blood, and that this environment contained more microbial diversity than previously thought. Although the investigators performed PCR experiments to independently confirm the existence of some of these signatures in blood samples, they did not attempt to assign assembled contigs to genome bins. Here, we studied contigs from these blood metagenomes with a genome-resolved strategy to investigate the presence of previously unknown bacterial populations.

To explore the origin of bacterial signatures found in the novel set of contigs recovered from cell-free DNA blood metagenomes (Fig. 2A), we first searched for the 139 bacterial single-copy core genes (SCGs) described by Campbell et al. (2013). This analysis identified 76 bacterial SCGs among all contigs; of these, 56 occurred only once, suggesting that a single microbial population may explain a large fraction of the bacterial signal found among novel contigs (Fig. 2B). Of the 56 genes that occurred only once, 18 were ribosomal proteins. Comparison of the amino-acid seguences of these ribosomal proteins to those in the NCBI's nonredundant protein sequence database revealed that the vast majority of them best matched to proteins from genomes that fall within the recently described "Candidate Phyla Radiation" (CPR) (Brown et al. 2015), a group of microbes with rather small genomes, reduced metabolic capacities (Rinke et al. 2013; Brown et al. 2015), and at least in some cases very small cell sizes (Luef et al. 2015), which suggest largely symbiotic lifestyles (He et al. 2015; Nelson and Stegen 2015). Even though ribosomal proteins found in blood metagenomes best matched to CPR genomes, the levels of sequence identity of these matches were very low, and taxonomic affiliations of best hits were divergent within the CPR (Supplemental Table S1), which could simply reflect the novelty of a single population rather than the presence of multiple populations. To investigate the distribution of these proteins, we clustered novel contigs based on their tetranucleotide frequencies (Fig. 2A). We found that most bacterial SCGs occurred in a relatively small group of contigs with similar tetranucleotide composition.

Manual selection of these contigs, and their further refinement using additional "non-novel" contigs that were not included in the original study by Kowarsky et al. (2017), resulted in a single CPR MAG that is 613.5 kbp in size with a completion estimate of 52.5%. Our phylogenomic analysis affiliated this MAG with the superphylum Parcubacteria (previously OD1) of the CPR (Fig. 2C). Regardless of the origins of this population in these metagenomes, our genome-resolved analysis contrasts with the prior interpretation of these data and suggests that Parcubacteria appears to be the only major novel bacterial group whose DNA is present in human blood metagenomes. This finding shows the critical importance of binning-based strategies to justify claims of microbial diversity in metagenomic analyses.

Yet, binning can be an important source of error

A real danger is that conclusions from draft MAGs may be incorrect because of misbinning (the wrong assignment of a genome fragment from one organism to another). It is critical to not rely on MAGs with high levels of contamination because these will likely yield misleading evolutionary and ecological insights (Bowers et al. 2017; Shaiber and Eren 2019). Misbinning is especially likely if scaffolds are short (e.g., <5 kbp), where binning signals can be noisy or unreliable. Thus, for better binning performance, it is helpful to use an assembler that includes a scaffolding step (insertion of Ns in gaps between contigs spanned by paired-end reads), such as IDBA-UD (Peng et al. 2012) or metaSPAdes (Nurk et al. 2017). MAGs can also be screened for short scaffolds with, for example, erroneous rRNA genes, which are often misbinned owing to their anomalous coverage (especially if the scaffolds are short and the genes are present in multicopy). Bins may also be contaminated by phage and plasmid genome fragments with coincidentally similar coverage or GC content, and so forth.

Completeness and contamination are often estimated using the inventory of expected SCGs in a MAG. A set of SCGs is selected based on their presence in all bacterial genomes, or at least all genomes within a taxonomic group (identified based on the phylogeny). In a genome without contamination, they should be present without redundancy. A widely used tool to assay both completeness and contamination is CheckM (Parks et al. 2015), although other methods are in use (Eren et al. 2015; Anantharaman et al. 2016). It has been noted both in the original study and in subsequent studies that CheckM can generate a false sense of bin accuracy, as shown by combining two partial single-cell genome bins (Parks et al. 2015; Becraft et al. 2017). The absence of multiple copies of SCGs does not preclude the presence of fragments from unrelated organisms that will compromise the biological value of the MAGs. Although there are tools for interactive visualization of genome bins in a single sample (Laczny et al. 2015; Raveh-Sadka et al. 2015) or across multiple samples (Eren et al. 2015) that enable manual curation opportunities to identify contamination beyond SCG-based estimates, the scalability of this strategy is limited. For example, recently there have been reports of many thousands, even hundreds of thousands, of draft MAGs from public metagenomic data sets (Parks et al. 2017; Almeida et al. 2019; Nayfach et al. 2019; Pasolli et al. 2019). Such large-scale analyses often rely on simplified procedures, for example, coverage profile of a single sample for binning, use of a single binning algorithm, or completeness/contamination estimates based on SCG inventories. Because these genomes are readily adopted by the scientific community for a wide variety of investigations, errors caused by misbinning will propagate.

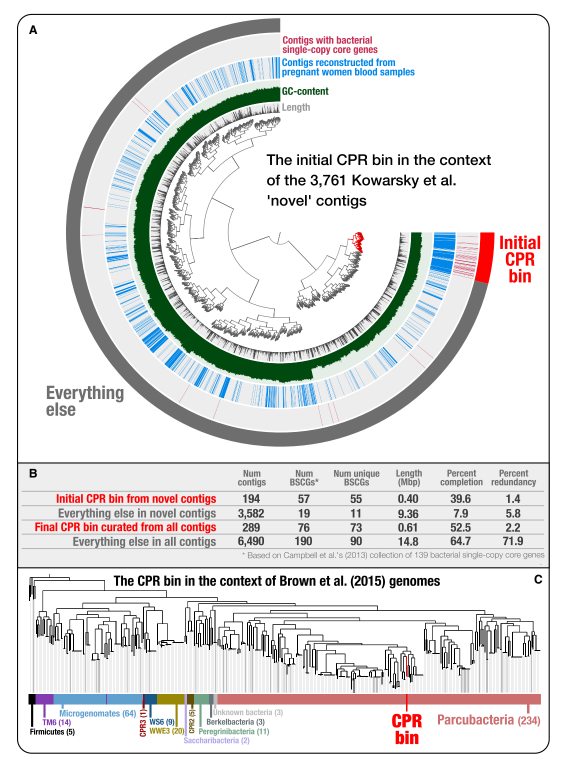


Figure 2. Genome-resolved metagenomics is essential to better investigate microbial diversity. (*A*) The *inner* dendrogram displays the hierarchical clustering of 3761 "novel" Kowarsky et al. contigs based on their tetranucleotide frequency (using Euclidean distance and Ward clustering) with the set of contigs that identify the genome in these data that is a member of the Candidate Phyla Radiation (CPR). Although the two *inner* layers display the length and GC content of each contig, the *outermost* layer marks each contig that contains one or more bacterial single-copy core genes. Finally, the second most *outer* layer marks each contig that contains one or more bacterial single-copy core genes. Finally, the second most of four cohorts of individuals in Kowarsky et al. (2017) (others being heart transplant, lung transplant, and bone marrow transplant patients), most ribosomal proteins we found in the assembly originated from contigs that were assembled from the pregnant women (Supplemental Table S1). The signal in this layer shows that contigs with bacterial single-copy core genes associate very closely with other contigs based on tetranucleotide frequencies, and most of these contigs are assembled from pregnant women blood metagenomes, providing additional confidence that this group of contigs represents a single microbial population genome within the "novel" set of contigs to the final CPR bin we have refined using the entire set of contigs, which included non-novel contigs we obtained from the authors of the original study (M Kowarsky, J Camunas-Soler, M Kertesz, et al., pers. comm.). (*C*) Phylogenetic analyses show the placement of the CPR bin in the context of CPR genomes released by Brown et al. (2015). More details of this case study are available at http://merenlab.org/data/parcubacterium-in-hbcfdna/.

A case study: SCGs can fail to predict the quality of MAGs

In a recent publication, Pasolli et al. (2019) used a single-sample assembly approach combined with automatic binning to generate 345,654 MAGs from the human microbiome, of which 154,723 pass a completion and quality threshold based on SCGs. The investigators suggest that the quality of the MAGs they have reconstructed through this pipeline was comparable to the quality of genomes from bacterial isolates or MAGs that were manually curated (Pasolli et al. 2019). However, reconstructing MAGs from single metagenomes and the heavy reliance on SCGs to estimate their quality can yield misleading results.

We examined one of the Pasolli et al. (2019) MAGs, "HMP_2012_SRS023938_bin.39" (hereafter referred to as Pasolli MAG), which resolves to the candidate phylum Saccharibacteria (formerly known as TM7), a poorly understood branch of the Tree of Life that contains members that are common in the human oral cavity (Bor et al. 2019). This MAG, 897,719 bp in length with 57 contigs (N50: 34,012 bp) (Supplemental Table S2), was recovered by Pasolli et al. (2019) from a supragingival plaque sample (experiment accession: SRR060355; sample accession: SRS023938) collected and sequenced by the Human Microbiome Project (HMP) (Turnbaugh et al. 2007). Anvi'o estimated the Pasolli MAG to include 84% of bacterial SCGs with very low redundancy (2.8%), in comparison, CheckM reported 63.39% completeness and 0.85% contamination (Supplemental Table S2).

The HMP data set included two additional plaque metagenomes from the same individual, providing an opportunity to investigate the distribution patterns of contigs binned together in

this MAG across multiple samples from the same person through metagenomic read recruitment. Organizing contigs based on their sequence composition and differential coverage patterns across three samples revealed two distinct clusters (Fig. 3), the smaller one of which contained 11 contigs that added up to a total length of 53.5 kbp (Fig. 3, outer circle, orange). Although the average mean coverage of contigs in these clusters were relatively comparable in the metagenome from which the MAG was reconstructed (24.6× vs. 31.1×), the average coverages differed in the other two plaque metagenomes (99.4× vs. 20.7× in SRS013723 and 9.7× vs. 33.56× in SRS078738), which suggest that the emergence of these two clusters was due to the improved signal for differential coverage with the inclusion of additional samples (Fig. 3). A BLAST search on the NCBI's nonredundant database matched genes found in 10 of 11 contigs in the smaller cluster to genomes of Veillonella (belonging to Firmicutes) (Supplemental Table S3), a genus that is common to the human oral cavity (Mark Welch et al. 2014) and includes members that are present in multiple oral sites (Eren et al. 2014). Genes in the remaining contig in the smaller cluster lacked a strong match (contig 0000000028) (Supplemental Table S3), yet best matched to genes in Selenomonas genomes instead of Saccharibacteria, suggesting that the smaller cluster represented contamination. Because these contaminating contigs did not include any SCGs, their inclusion did not influence SCG-based completeness and contamination estimates. Thus, they remained invisible to the quality assessment. Although the contamination in this case will unlikely influence the placement of this particular MAG in the Tree of Life owing to the lack of SCGs in it, the contamination does change the functional makeup of the MAG: Our annotation of 54 genes in the 11 contaminating contigs using the

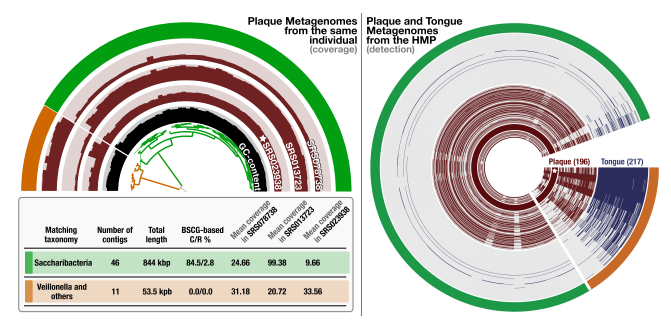


Figure 3. Contamination in MAG without extra copies of SCGs. In the *left* panel, the half-circle displays the mean coverage of each contig in Pasolli MAG across three plaque metagenomes that belong to the same individual, for which the "star" symbol denotes the sample from which the original MAG was reconstructed. The dendrogram in the *center* represents the hierarchical clustering of the 57 contigs based on their sequence composition and differential mean coverage across the three metagenomes, and the *innermost* circle displays the GC content for each contig. The *outermost* circle marks two clusters: one with 46 contigs (green) and another one with 11 contigs (orange). The table *underneath* this display summarizes various statistics about these two clusters, including the best matching taxonomy, total length, completion and redundancy (C/R) estimations based on SCGs, and the average mean coverage of each cluster across metagenomes. In the *right* panel, the distribution of the same contigs and clusters are shown across 196 plaque (brown) and 217 tongue (blue) metagenomes generated by the Human Microbiome Project (HMP). Each concentric circle in this display represents a single metagenome, and data points display the detection of the contigs in Pasolli MAG.

NCBI's Clusters of Orthologous Groups (COGs) revealed 30 functions that were absent in the MAG after the removal of the contamination (Supplemental Table S4). In addition to misleading functional profiles, contamination issues often influence ecological insights. Our read recruitment analysis to characterize the distribution of the Pasolli MAG contigs across all 196 plaque and 217 tongue metagenomes from 131 HMP individuals showed that although this Saccharibacteria population appears to be restricted to plaque samples, contigs that contaminated this MAG recruited reads also from the tongue samples (Fig. 3; Supplemental Table S5).

We did not investigate the quality of the full set of 154,723 MAGs described by Pasolli et al. (2019) or the genomes reported in other studies that relied on similar automated strategies (Almeida et al. 2019; Nayfach et al. 2019). Nevertheless, this example shows that SCGs alone cannot predict the lack of contamination in a given MAG or characterize the extent of contamination in genomic collections (for another example, see Supplemental Fig. S2). Overall, it is essential for our community to note that computational analyses that rely heavily on SCGs to assess the quality of MAGs can promote erroneous insights.

Genome curation: moving toward complete genomes

The opportunity to recover huge numbers of new genomes from metagenomic data sets motivates the development of new tools to more comprehensively curate draft MAGs, ideally to completion. Although the term "complete" should be reserved for genome sequences with (usually) circular chromosomes reported in single scaffolds, in contemporary genome-resolved metagenomics studies the term is commonly used to describe bacterial and archaeal genomes that have all the expected SCG markers used to evaluate completeness. This use of the term "complete" does not exclude genomes that are extremely fragmented, which can suffer from contamination issues, as we show above. Here, we use the term "complete" explicitly to describe multiple properties of a genome: (1) circular (assuming the chromosome is circular) and single chromosomal sequence, with (2) essentially perfect read coverage support throughout (i.e., the majority [e.g., >50%] of bases in mapped reads at any position matches to the consensus base), and (3) no gaps. To avoid any confusion, we will use the term "CMAGs" to describe complete MAGs that meet the three criteria.

The first genome of an uncultivated bacterium to our knowledge, appeared in 2000 and was for an insect symbiont (Shigenobu et al. 2000), but the DNA of only one microorganism was sampled so binning was not required. The first CMAGs appeared in 2008, but this was for a bacterium that comprised >99.9% of the sample (Chivian et al. 2008). Another genome published in the same year was for a candidate phylum bacterium in an anaerobic digester and was reconstructed by sequencing of a fosmid library (Pelletier et al. 2008). In addition, a genome for a member of Elusimicrobia was reported from a Termite gut in 2008 (Hongoh et al. 2008). It was not until 2012 and 2013 that a series of CMAGs from multispecies natural communities began to appear (Iverson et al. 2012; Castelle et al. 2013; Di Rienzi et al. 2013; Kantor et al. 2013). In most cases, these genomes were very close to complete upon de novo assembly, although some effort was required to finish them. Near complete de novo assembly is a very rare outcome, given that most genomes are assembled using short paired-end reads (e.g., 150 bp with a few hundred base pair insert size). However, given that many samples generate hundreds of draft genomes, very highquality de novo assembly of a genome is not uncommon overall.

Nevertheless, the curation of even very well-assembled MAGs is very rarely undertaken, perhaps owing to the involvement of typically manual and generally not well-understood steps. Here, we describe the methods that can be used for genome curation and provide examples to illustrate potential caveats along with their likely solutions. Our hope is that the following sections will motivate the development of new tools to enable routine curation of genomes from metagenomes.

A limited number of published complete metagenome-assembled genomes

To the best of our knowledge, as of September 10, 2019, 59 bacterial and three archaeal CMAGs from microbial community data sets are publicly available (Table 1). Of these, four CMAGs were finished using Pacific Biosciences (PacBio) reads. The published CMAGs are primarily for members of the Candidate Phyla Radiation (CPR; 36 genomes) and DPANN (two genomes), which have unusually small genomes (average genome size of 1.0 Mbp) (Table 1). Other reported CMAGs include those for Proteobacteria (eight genomes), Saganbacteria (WOR-1; 4), Bacteroidetes (two), *Candidatus* Bipolaricaulota (two), Firmicutes (two), and one from each of Dependentiae (TM6; also small genomes), Elusimicrobia, Melainabacteria, Micrarchaeota, Nitrospirae, Zixibacteria, and *Candidatus* Cloacimonetes (Table 1).

CMAGs are not limited to bacteria and archaea. Because all of the extracted DNA is sequenced, genomes are also reconstructed for phage and plasmids. In fact, the tool VirSorter (Roux et al. 2015) predicts circularized sequences suitable for verification and curation to remove gaps and local assembly errors. Two recent studies reported unusually large complete phage genomes. In the first case, 15 complete megaphage genomes, each >540 kbp in length, were reconstructed and curated from human and animal microbiomes (Devoto et al. 2019). In the second case, 35 complete genomes >200 kbp derived from phage, including the largest phage genomes yet reported (Al-Shayeb et al. 2020). The distinction of these sequences from prophage and the accurate size determinations could not be made without circularized genomes, and the complete, accurate inventory of genes would be precluded with only draft genomes.

Genome curation: filling scaffolding gaps and removal of local assembly errors

Genome curation requires the identification and correction of local assembly errors and removal of gaps at scaffolding points. However, the exclusion of these steps in current genome-resolved metagenomics studies propagate errors such as incomplete or incorrect protein-coding gene sequences in public databases.

Automatic tools like GapFiller (Nadalin et al. 2012) may be useful for the filling of "N" gaps at scaffold joins (read pairs should span the gap if the scaffolding was done correctly). Our primary approach to gap filling makes use of unplaced pairs for reads adjacent to the gaps. When reads are mapped to genome fragments that compose a bin, a file of unplaced paired reads is generated for each fragment. By mapping these unplaced paired reads to the corresponding fragment, it is usually possible to incrementally close the gap (so long as there is sufficient depth of coverage). After the first round of mapping of unplaced paired reads, the consensus sequence must be extended into the gap before remaining unplaced paired reads are remapped. The newly introduced paired

Archaes; DPANN; Dispherotries Condidute Forterrea multitransposorum Archaes; DPANN; Micrarchaeota RMAN 2 Archaes; DPANN; Micrarchaeota RMAN 2 Bactenia; CPR, Bail 4 Battenium RFOXYA1; FULL 37.12 Bail 4 Condidatus Roizmanbacteria bacterium RFOXYA1; PULL 38.14 Condidatus Roizmanbacteria bacterium RFOXYD1; PULL 38.14 Condidatus Roizmanbacteria bacterium RFOXYD1; PULL 38.12 Bacteria; CPR; Peregrinibacteria BC Condidatus Peribacter BC 28.12 Bacteria; CPR; Collierbacteria BC 28.12 28.12 28.12 28.12 Bacteria; CPR; Collierbacteria BC 29.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.12 28.11 <t< th=""><th></th><th>Size (bp)</th><th>accession</th><th>GC_skew</th><th>Reference</th></t<>		Size (bp)	accession	GC_skew	Reference
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	F	1,216,967	MGBN01000001	As expected	Anantharaman et al. 2016
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		878,109	CP006914	Uncertain/uneven replichores	Kantor et al. 2013
		585,024	MEZB01000001	As expected As expected	Anantharaman et al. 2016 Anantharaman et al. 2016
		699,981 699,981	METB01000001 METD01000001	Uncertain/uneven replichores Uncertain/uneven replichores	Anantharaman et al. 2016 Anantharaman et al. 2016
				-	
6		602,646 1,013,781 1,038,683	CP011216 CP005957 CP011211	Uncertain/uneven replichores As expected As expected	Brown et al. 2015 Albertsen et al. 2013 Brown et al. 2015
2		1,450,269 845,464	CP025011 CP006915	As expected As expected	Starr et al. 2018 Kantor et al. 2013
			PHACUUUUUUUU ERS4269226	As expected As expected	Hernsdorf et al. 2017 This study
Bacteria; CPR; Beckwithbacteria Candidatus Beckwithbacteria bacterium		1,177,760 1,049,888	CP011210	Noisy, probably as expected No skew	Kantor et al. 2015 Brown et al. 2015
Bacteria; CPR; Berkelbacteria Berkelbacteria bacterium GW2011_GWE1_39_12 SB 2 sc 141 Backelbacteria circularizad		915,059 768 550	CP011213 EPC4770530	As expected	Brown et al. 2015 This study
Candidatus Pacebacteria	-3_36_13 ^b	853,053 1,133,667	SAMN03319330 ERS4266120	As expected As expected	Brown et al. 2015 This study
Bacteria; CPR; Campbellbacteria Candidatus Campbellbacteria bacterium CW2011 ODI 34 28		752,630	CP011215	As expected	Brown et al. 2015
Bacteria; CPR; Gracilibacteria BD02T64_BD1-5(BD1-5)	-	1,343,103	SUB5359196	As expected	Sieber et al. 2019

Taxonomy	Genome	Size (bp)	GenBank/ENA accession	GC_skew	Reference
Bacteria; CPR; Kaiserbacteria	Candidatus Kaiserbacteria bacterium BIECERI OMO2 12 ETII 45 36	962,580	MFMM01000001	As expected	Anantharaman et al. 2016
Bacteria; CPR; Woesebacteria Bacteria; CPR; Wolfebacteria Bacteria; Bacteroidetes	Candidatus Woesebacteria bacterium GW2011_GWF1_31_35 Candidatus Wolfebacteria bacterium GW2011_GWB1_47_1 Candidatus Fluviicola riflensis	819,458 984,447 4,551,443	CP011214 CP011209 CP022585	As expected As expected As expected	Brown et al. 2015 Brown et al. 2015 Banfield et al. 2017
Bacteria; <i>Candidatus</i> Bipolaricaulota Bacteria: <i>Candidatus</i> Cloacimonetes	bacteroiaetes UKLI 3-3 Candidatus Acetothermia bacterium Ran1 Candidatus Bipolairaudis sp. Ch78 Candidatus Cloacimonas acidaminovorans (via fosmid library)	3,236,529 1,324,338 1,701,655 2,246.820	CP012155 LS483254 CP034928 NC 020449.1	As expected Uncertain/low skew As expected As expected	Driscoll et al. 2017 Hao et al. 2018 Kadnikov et al. 2019 Pelletier et al. 2008
(WWE1) Bacteria; <i>Candidatus Z</i> ixibacteria Bacteria; Elusimicrobia Bacteria; Firmicutes	Candidate division Zixibacteria bacterium RBG-1 Candidate phylum Termite Group 1 Rs-D17 Candidatus Desulfortudis audaxviator ^c	2,122,767 1,125,857 2,349,476	AUYT01000001 NC_020419 NC_010424	As expected As expected As expected	Castelle et al. 2013 Hongoh et al. 2008 Chivian et al. 2008
Bacteria; Melainabacteria Bacteria; Nitrospirae Bacteria; Proteobacteria	candidatus Melainabacteria bacterium MEL.A1 Nitrospinatio Candidatus Liberibacter asiaticus str. Psy62 ^c Buchnera sp. APS SCNpilot_BF_INOC_Rickettsiales_complete_39_4 Ca. <i>Riegeria sontandreae</i> (contain ambigous bases) Hyphomonadoceae UKL13-1	7,20,200 3,295,117 1,227,204 640,681 988,358 1,342,908 3,501,508	CP017245 NZ_LN885086 CP001677.5 NC_002528 ERS4267492 LR026963 CP012156 PRIMA557483	As expected As expected As expected As expected As expected As expected Low skew / uncertain As expected	Di Rienzi et al. 2013 Di Rienzi et al. 2013 Daims et al. 2015 Duan et al. 2009 Shigenobu et al. 2000 This study jäckle et al. 2019 Driscoll et al. 2017 Chen et al. 2017
Bacteria; Dependentiae (TM6) Bacteria; WOR-1	Betaproteobacterium UKL13.2 Candidatus Pseudomonas sp. strain JKJ-1 Candidatus Sulfuricurvum sp. RIFRC-1 Candidate division TM6 bacterium TM6SC1 Candidate division WOR-1 bacterium	3,387,087 6,408,606 2,358,861 1,088,795 1,668,697	CP012157 PRJNA320198 CP003920 — METT01000001	As expected No genome available As expected As expected As expected	Driscoll et al. 2017 White et al. 2016 Handley et al. 2014 Antipov et al. 2016 Anantharaman et al. 2016
	Candidate division WOR-1 bacterium RIFOXYA2_FULL_52_19 Candidate division WOR-1 bacterium RIFOXYB2_FULL_52_9 Candidate division WOR-1 bacterium RIFOXYC12_FULL_52_18	1,668,697 1,668,697 1,668,697	METY01000001 MEUD01000001 MEUG01000001	As expected As expected As expected	Anantharaman et al. 2016 Anantharaman et al. 2016 Anantharaman et al. 2016

constructed for organisms from 30 different phylum-level groups. Five of the listed genomes were completed in this study. ^bWrongly labeled in NCBI as TM6. Note that many genomes show asymmetric patterns of GC skew, which is attributed to uneven length replichores (also seen in isolate genome analysis). ^cThe CMAG was reconstructed from a sample with only one organism present.

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reads should be placed at an appropriate distance from their existing pairs, given the fragment insert size. Often a few iterations are needed for gap closure. However, if the gap does not close and no further extension can be accomplished using the existing collection of unplaced pairs, the full metagenomic read data set can be mapped to the new version of the scaffold and another round of extension performed until the gap is closed.

If a gap cannot be closed using the unplaced paired reads owing to low coverage, one solution may be to include reads from another sample in which the same population occurs (this may not be appropriate for some investigations), or by performing a deeper sequencing of the same sample. In other cases, the necessary reads are misplaced, either elsewhere on that scaffold or on another scaffold in the bin. This happens because the reads have been "stolen" thus the true location sequence is not available to be mapped to. This often leads to read pileups with anomalously high frequencies of SNVs in a subset of reads. However, anomalously high read depths can also occur owing to mapping of reads from another genome. The misplaced reads can be located based on read names and extracted for gap filling. Other indications of misplacement of reads include read pairs that point outward (rather than toward each other, as expected) or with unusually long paired read distances. One of these reads is misplaced and the other read normally constrains the region to which the pair must be relocated. Relocation of the misplaced read can often lead to filling of scaffolding gaps. In some cases, gap filling cannot be easily achieved despite sufficient read depth. This can occur, for example, because of complex repeats. Sometimes these repeat regions can be resolved by careful read-by-read analysis, often requiring relocation of reads based on the placement of their pairs as well as sequence identity.

Another important curation step is the removal of local assembly errors (Supplemental Fig. S3). We suspect that these errors are particularly prominent in IDBA-UD assemblies, although it is likely that all assemblers occasionally make local assembly errors. Local assembly errors can be identified because the sequence in that region lacks perfect support, by even one read. The region should be opened up and each read within that region separated to the appropriate side of the new gap (so that all reads match the consensus sequence). Unsupported consensus sequence should be replaced by Ns. The new gap can be filled using the procedure for filling scaffolding gaps, as described above.

A second type of local assembly error is where Ns have been inserted during scaffolding despite overlap between the flanking sequences (Supplemental Fig. S4). We have observed this problem with both IDBA-UD and CLC workbench assemblies. The solution is simply to identify the problem and close the gap, eliminating the Ns and the duplicate sequence.

Another common assembly error involves local repeat regions in which an incorrect number of repeats has been incorporated into the scaffold sequence. This situation may be detected by manual inspection of read mapping profile, as it leads to anomalous read depth over that region. Sometimes the correct number of reads may only be approximated based on the consistency of the coverage within the repeat region and other parts of the scaffold (see example below).

Rarely, in our experience, assemblers create scaffolds that are chimeras of sequences from two different organisms (e.g., Mineeva et al. 2020). These joins typically lack paired read support and/or can be identified by very different coverage values and/or phylogenetic profiles on either side of the join. Another seemingly rare error involves the artificial concatenation of an identical sequence, sometimes of hundreds of base pairs in length, repeated up to (or more than) three times. This has been a problem with some sequences of seemingly large phage deposited in public databases, as discussed by Devoto et al. (2019) and Al-Shayeb et al. (2020). This phenomenon is easily identified by running a repeat finder, a step that should also be included in the curation to completion pipeline (see below).

Using GC skew as a metric for checking genome correctness

GC skew is a form of compositional bias-imbalance of guanosine (G) relative to cytosine (C) on a DNA strand—that is an inherent feature of many microbial genomes, although some are known to display little or no GC skew (e.g., certain Cyanobacteria) (Nakamura 2002). The phenomenon of strand-specific composition was described by Lobry (1996), who observed that the sign of the relative GC skew changes crossing the oriC and terC regions. Thus, the inflection point in genome GC skew at the origin of replication is often close to the *dnaA* gene and typically contains a small repeat array. GC skew is calculated as (G - C/G + C) for a sliding window along the entire length of the genome (suggested window = 1000 bp, slide = 10 bp). The skew is also often summed along the sequence to calculate cumulative GC skew. This was proposed by Grigoriev (1998), who showed that the calculation of the cumulative GC skew over sequential windows is an effective way to visualize the location of the origin and terminus of replication. For complete genomes, the GC skew is often presented starting at the origin of replication, proceeding through the terminus and back to the origin (i.e., as if the chromosome was linear). The pattern of the cumulative GC skew, where the function peaks at the terminus of replication, indicates that the genome undergoes bidirectional replication. The pattern is fairly symmetrical unless the replichores are of uneven lengths. Because the magnitude of the cumulative GC skew varies from genome to genome, the magnitude of the skew could potentially be used as a binning signal.

The explanation for the origin of GC skew is not fully agreed upon. It may arise in large part because of differential mutation rates on the leading and lagging strands of DNA. Enrichment in G over C occurs because of C deamination to thymine $(C \rightarrow T)$, the rates of which can increase at least 100-fold when the DNA is in a single-stranded state. In the process of DNA replication, the leading strand remains single stranded while the paired bases are incorporated by the DNA polymerase into its complementary strand. However, the Okazaki fragments on the lagging strand protect a fraction of the DNA from deamination. Thus, the leading strand becomes enriched in G relative to C compared to the lagging strand. The magnitude of the GC skew can be impacted by the speed of the DNA polymerase processivity (which impacts the length of time that the DNA is single stranded) and the length of the Okazaki fragments. GC skew has been linked to strand coding bias (Rocha et al. 1999). Concentration of genes on the leading strand would afford protection against nonsynonymous mutations (as $C \rightarrow T$ mutations in the wobble position of codons are always synonymous), whereas $G \rightarrow A$ on the lagging strand (following $C \rightarrow T$ on the leading strand) in two cases results in nonsynonymous mutations (AUA for Ile vs. AUG for Met, and UGA for stop codon vs. UGG for Trp). The potential for deamination in the noncoding strand during transcription, another source of GC skew, would also favor genes on the leading strand. GC skew persists

because the leading strand is maintained as such through subsequent replication events.

Given that a well-defined pattern of GC skew is anticipated across many bacterial (and some archaeal) genomes, we wondered whether plots of cumulative GC skew for putative complete genomes can be confidently used to test for genome assembly errors. For this metric to be useful, it would be imperative to establish the extent to which GC skew is indeed a feature of complete bacterial genomes. To our knowledge, the now extensive set of complete isolate genomes has not been leveraged to do this.

We undertook benchmarking of GC skew, and more specifically cumulative GC skew, using all approximately 7000 complete genomes in the RefSeq database. We found that the majority of RefSeq bacterial genomes show the expected pattern of cumulative GC skew. The magnitude of the origin to terminus skew varies substantially, from ± 0.4 excess G relative to C to close to zero (Supplemental Fig. S5). A small subset of the approximately 7000 complete genomes essentially lack GC skew (as reported for some Cyanobacteria, see above) (Supplemental Table S6). Poorly defined (noisy) patterns are often associated with low total cumulative skew. About 15% of genomes have notably asymmetric patterns (i.e., the cumulative skew is substantially larger for one half of the chromosome relative to the other), presumably because the two replichores are of substantially uneven length. Moreover, some bacterial genomes had a GC skew pattern indicating rolling circle replication (Supplemental Table S7). We did not detect a strong correlation between the magnitude of GC skew and bias for genes on the leading strand.

Some complete genomes have quite aberrant skew patterns, with inversions in the cumulative skew within a single replichore or exceedingly uneven predicted replichore lengths. We considered the possibility that a subset of these isolate genomes may contain misassemblies. Such a phenomenon was already shown by Olm et al. (2017) in the case of a Citrobacter koseri isolate genome that was clearly wrongly assembled across rRNA operons (and a PacBio assembly for a closely related strain showed the expected pattern of cumulative GC skew). To test for the possibility that these other complete genomes contained errors, we posited that misassemblies would likely occur at perfect repeats that are longer than the distance spanned by paired reads. Further, we predicted that the pair of repeats flanking the wrongly assembled sequence region would be in reverse complement orientations so that the intervening DNA segment could be flipped at the repeats and that the flipped version would show the expected GC skew pattern. In five of twelve cases that we scrutinized, it was possible to show that reverse complementing the sequence spanned by repeats indeed resulted in genomes with exactly the expected form of cumulative GC skew (Fig. 4; Supplemental Figs. S6, S7). In one case, that is, Flavobacterium johnsoniae UW101 (NC_009441.1), the original assembly notes indicated assembly

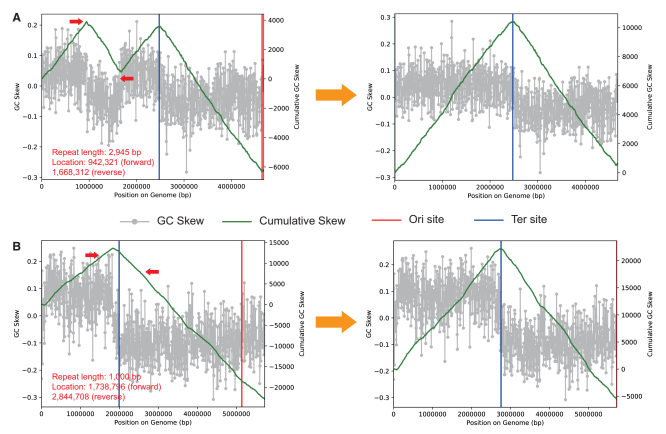


Figure 4. Examples of probable assembly errors in RefSeq bacterial genomes. (*A*) Salmonella enterica subsp. enerica (CP009768.1). (*B*) Desulfitobacterium hafniense Y51 (NC_007907.1). The diagrams show the GC skew (gray) and cumulative GC skew (green line) of the original (*left*) and the modified (*right*) versions of the genomes (all calculated with window size of 1000 bp, and slide size of 10 bp). The location and direction of repeat sequences leading to the abnormal GC skew are indicated by red arrows. After flipping the repeat-bounded sequences, the genomes show the pattern expected for genomes that undergo bidirectional replication (*right*). For more examples, see Supplemental Figures S6 and S7.

uncertainty (although the complete genome was deposited at NCBI).

We acknowledge the possibility that a recent major rearrangement could also give rise to inflexions in GC skew; however, major rearrangements typically have a well-defined placement relative to the origin of replication that is inconsistent with the patterns observed (Eisen et al. 2000). Although we cannot state that these isolate genomes are wrongly assembled, we suggest that it is a distinct possibility. Incorrect assemblies in isolate genomes can be of high significance, given the trust placed in them for evolutionary and metabolic analyses that make use of synteny and gene context. They are also used as references for calculation of growth rates via the PTR method (Korem et al. 2015), and incorrect reference sequences will corrupt such measurements.

It is well known that some archaea replicate their genomes from multiple origins (Barry and Bell 2006). In such cases, the cumulative GC skew pattern is not a useful test of overall genome

accuracy. However, some archaea do show the peaked pattern that is typical of bacteria, thus indicative of bidirectional replication. Overall, we found 18 of 224 RefSeq archaeal genomes tested that show this pattern, and all of them are Euryarchaeota (Supplemental Table S8). In addition, this pattern was reported for a DPANN archaeon (Probst and Banfield 2018).

From high-quality draft sequences to complete genomes from metagenomes

Genome curation to completion is rarely undertaken (Table 1) because there is no single tool available to accomplish it, and there can be confusing complications. The procedure requires the steps described in the previous section as well as extension of scaffolds (or contigs, if no scaffolding step was undertaken) so that they can be joined, ultimately into a single sequence (assuming the genome is a single chromosome). With currently available tools, this is time consuming, sometimes frustrating, and often does not result in a CMAG (usually because of indistinguishable multiple options for scaffold joins typically resulting from repeats such as identical copies of transposons). However, when it can be done, the resulting genome solution should be essentially unique, as we will show below. There is nothing "arbitrary" about the process, except occasionally the choice of which set (usually a pair) of subequal locus variants (e.g., SNVs) will represent the final genome. Even in those cases, depending on the availability of multiple appropriate metagenomes for read recruitment analyses, tools for haplotype deconvolution such as

DESMAN (Quince et al. 2017) may offer quantitative support for such decisions.

In our experience, the most important first step in the path toward recovery of a CMAG is to start from a well-defined bin that appears to comprise the vast majority of the genome of interest (Fig. 5, step 1). As above, this is usually determined based on genome completeness evaluation (Fig. 5, step 2) and/or a very strong set of binning signals (e.g., Supplemental Fig. S1). It should be noted that some genomes (e.g., CPR bacteria) may naturally lack certain SCGs that are otherwise considered universal in other bacteria (Brown et al. 2015), and may require a modified list of universal SCGs such as those proposed for CPR genomes for more accurate evaluations of completion (Anantharaman et al. 2016). Importantly, the targeted bin should be polished to remove contamination scaffolds, as noted above (Fig. 5, step 3).

Given currently available tools, it is probably wise to choose a bin with no more than 10 pieces (Fig. 5, step 4), although a MAG

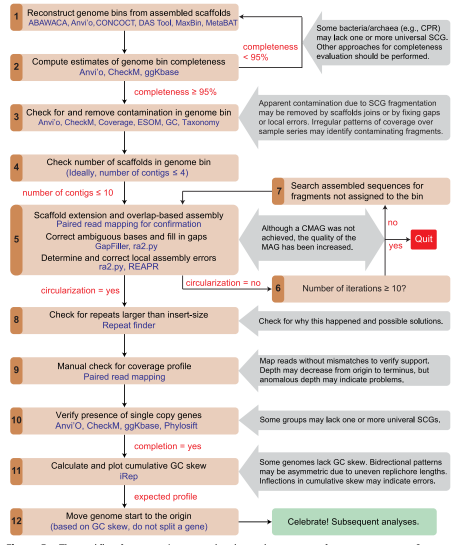


Figure 5. The workflow for generating curated and complete genomes from metagenomes. Steps are shown in black, and the tools or information used in blue. Notes for procedures are shown in gray boxes. The detailed procedures for scaffold extension and gap closing are available in the Supplemental Methods and also online (https://ggkbase-help.berkeley.edu/genome_curation/scaffold-extension-and-gap-closing/).

with larger number of scaffolds can be curated to completion if necessary (Chen et al. 2019). The best possible case is when the genome is de novo assembled into a single piece. In some cases, the genome is already circularized, based on overlap sequences at the scaffold ends, with paired-end reads that span the scaffold ends. Although rare, this does occur, mostly for small genomes (e.g., Saccharibacteria) (Albertsen et al. 2013; Starr et al. 2018). In other cases, a modest amount of end extension may be required for circularization (see below). The single scaffold should be checked for complete coverage and support of the consensus. Gaps or local assembly errors must be dealt with before the genome is classified as curated and complete (some additional checks are described below).

Some assemblers (e.g., IDBA-UD, metaSPAdes) retain sequences that are nonunique at scaffold ends. Assembly termination presumably happens because assembly algorithms are designed to stop at points of uncertainty rather than risk making incorrect joins (Supplemental Fig. S8A). Incidentally, because different assemblers can yield different results, there can be value in comparing the results for the same data assembled using different tools and/or parameters (see examples below). Also, in some cases, assembly of scaffolds representing the same organism (or a closely related organism) from a related sample, could help scaffold extension and/or linkage (Supplemental Fig. S8B). Potential scaffold joins can be made by identifying perfect overlaps at the ends of scaffolds of a MAG ("overlap-based assembly") (Fig. 5, step 5). Often, the length of perfect overlap of scaffolds assembled using IDBA-UD and metaSPAdes is n and n-1, respectively, where n is the largest k-mer size used in de novo assembly. Although the assembler chose not to make these joins (possibly owing to confusion involving even a single read), seemingly unique joins involving scaffolds in a bin can be made tentatively during curation. Ultimately, nonunique joins can be eliminated or resolved at the end of the curation process. It is important to note that nonuniqueness of a join may not be evident in an initial scaffold set owing to failure to include a relevant scaffold in the bin or lack of de novo assembly of relevant regions. Thus, it is important to test for repeated regions that cannot be spanned by paired reads at the end of curation (either in the potentially complete genome or curated scaffold set if completion is not achieved). Failure to identify perfect repeats can also lead to problems in isolate genomes, as we show above.

Scaffolds within a bin that do not overlap at the start of curation may be joined after one or more rounds of scaffold extension (Supplemental Fig. S9). This process of extending, joining, and remapping may continue until all fragments compose a single circularized sequence. It should be noted that read-by-read scaffold extension is very time consuming. If an extended scaffold cannot be joined to another scaffold after a few rounds of extension it may be worth testing for an additional scaffold (possibly small, thus easily missed by binning) by searching the full metagenome for overlaps (Fig. 5, steps 6 and 7). Sometimes, the failure of scaffold extension is caused by missing paired reads, which may be found at the end of another fragment. If they are pointing out but the sequences cannot be joined based on end overlap, a scaffolding gap can be inserted in the joined sequence (reverse complementing one of the scaffolds may be necessary). Closure of the new scaffolding gap uses the approach described above.

During the attempt to obtain a circularized sequence, it is important to note that if the genome has a single pair of duplicated sequences that are larger than can be spanned by paired reads, a reasonable solution can be found if the genome bin is curated into just two pieces. In this case, the only solutions are either resolution into two chromosomes or generation of a single genome (Supplemental Fig. S10). This observation underlines the importance of curation from a high-quality bin, because curation from a full metagenome would leave open the existence of other scaffolds that also bear that repeat.

Phage genomes present an additional challenge given the complexity and variability in their genome structures and replication mechanisms (Lo Piano et al. 2011). For example, phages generally have linear genomes although most circularize during replication. Circularization is an important criterion for genome completion unless the case can be made that the genome is linear. Genome linearity can be established based on reads that terminate in a defined region, with all paired reads pointing inward from the termini (Supplemental Fig. S11). When phage genomes have terminal repeated sequences, the assembled genome will appear circular. If the phage was sampled during replication, the read coverage will be consistent over the entire genome after trimming one end to remove repeated sequence. If a genome with terminal repeats was packaged when it was sampled, it will show doubled coverage at one end after trimming of the repeated sequence.

Once the genome is circularized, it is important to check for repeats larger than spanned by paired-end reads (as noted above) (Fig. 5, step 8). Assuming a seemingly CMAG is achieved, several steps to further verify the accuracy of the assembly path may be warranted (Fig. 5, step 9). First, reads may be mapped to the sequence allowing no mismatch to confirm no coverage gap attributable to base miscall and to verify that no region has abnormal coverage. Tools that provide interactive visualization and inspection of coverage patterns, such as anvi'o, Geneious (Kearse et al. 2012), or Integrative Genomics Viewer (Robinson et al. 2011), may be used for this task. Second, we advocate verification of paired read placements over the entire assembly to check for problem areas that may have been missed in automated procedures. Abnormally low coverage may result from a subpopulation variant, whereas higher than expected coverage could indicate the existence of a block of sequence that was pinched out from the genome at a repeated region. Systematic decline of coverage from origin to terminus of replication is expected if genome replication was ongoing at the time of sampling (see below). Third, the presence of expected genes (e.g., universal SCGs) should be verified. The genome can be classified using phylogenetic analyses (e.g., based on 16S rRNA gene or concatenated ribosomal proteins sequences) (Fig. 5, step 10). After the completion of MAG, the start of the genome should be moved to the noncoding region near the origin of replication (Fig. 5, steps 11 and 12). See below for details regarding how GC skew can be used to locate the origin.

An important consideration in genome curation to completion is knowing when to give up. In some cases, failure to circularize after a few rounds of curation may be an indication that the effort could be better invested in other activities. If alternative assembly paths that cannot be distinguished by the unique placement of paired reads are identified, failure may be on the horizon. However, as noted above (Supplemental Fig. S10), it can be appropriate to continue curation as a final unique solution may be possible even in the presence of a repeat that cannot be spanned by paired reads.

Case studies illustrating the curation of draft MAGs

Here, we illustrate how a draft MAG can be curated to completion or into better quality status, with step-by-step procedures detailed in the Supplemental Methods. The genomes in cases one and two are not published (for details, see Methods), and case three includes a published genome.

Case one: curation of a CPR genome to completion

ALT_04162018_0_2um_scaffold_13, length of 1,128,909 bp, was the only scaffold in the binned MAG (i.e., bin.56) from MetaBAT (Supplemental Methods). CheckM reported 70.1% completeness without contamination, and preliminary analyses based on 16S rRNA and rpS3 genes identified it as a Parcubacteria genome. This genome was likely near complete based on the detection of all CPR universal SCGs, although we did not identify overlap at the ends of the scaffold that would circularize it. This scaffold could be circularized after a single round of scaffold end extension, with read pairs placed at the ends of the scaffold. In fact, we found two very small assembled sequences that were variants of each other, and both could be used for circularization. The nonuniqueness of this region terminated the original assembly. We chose the dominant variant to represent the population genome. No repeat sequence longer than the sequencing insert size was detected. A total of 13 local assembly errors were reported by ra2.py. All these errors were manually fixed and validated, including a complicated error in the sequence of a protein-coding gene that contains multiple repeat regions. The complete genome has a length of 1,133,667 bp, and encodes 1147 protein-coding genes, 47 tRNA, and a copy each of 5S/16S/23S rRNA genes.

Case two: curation of a Betaproteobacteria genome without completion

Bin.19 contained seven scaffolds (3.6 Mbp in size) and was evaluated by CheckM to be 98.42% complete with 0.12% contamination (Supplemental Methods). Analyses of the 16S rRNA gene sequence indicated it was a Betaproteobacteria (92% similarity to that of Sulfuricella denitrificans skB26). After the first round of scaffold extension and assembly, only two scaffolds could be combined (i.e., scaffolds 21 and 25). We searched for the pieces that could be used to link the scaffolds together using the newly extended parts of the scaffolds via BLASTN against the whole scaffold set. This approach retrieved four short (584-1191 bp in length) and one longer piece (15,678 bp in length) that encodes several bacterial universal SCGs including rpS7, rpS12, rpL7/L12, rpL10, rpL1, and rpL11 (which were absent from bin.19), and whose two ends both encode elongation factor Tu (EF-Tu). Two of the four short pieces could be perfectly joined in two possible places to the extended scaffold set of Bin.19. Based on comparison with the Sulfuricella denitrificans skB26 genome, we hypothesized the linkage patterns for these fragments and then considered the two choices for how the resulting two large genome fragments could be arrayed. The linkage choices were supported based on the overall pattern of GC skew (see above; Supplemental Methods). Technically, however, the bin remains as two contigs with two internal joins unsupported by unique paired read placement. Based on the GC skew of the pair of contigs linked by Ns, the genome is near complete. After correcting the local assembly errors, the genome has a total length of 3.72 Mbp, encodes 3544 protein-coding genes, 41 tRNA and one copy of each of the 5S/16S/23S rRNA genes, and is clearly of higher quality than the original bin owing to scaffold extension, new scaffold inclusion, and correction of local assembly errors.

Case three: curation of a published incomplete genome to completion

Here, we completed a published curated (for local assembly errors) but incomplete genome belonging to the order Rickettsiales (Kantor et al. 2017). This genome was assembled de novo into a single circularizable 988 kbp scaffold, with two closely spaced gaps (Supplemental Methods). Closing of these gaps required relocation of unplaced paired reads.

In addition to the aforementioned case studies, we curated three additional bacterial genomes to completion as part of our methods refinement. These genomes are listed in Table 1.

Other approaches, future opportunities and challenges

Single-cell genomics

Microbial single-cell sequencing is a family of strategies that typically uses microfluidics and whole-genome amplification to physically isolate individual cells and sequence their genomes without cultivation (Stepanauskas 2012). The resulting single-amplified genomes (SAGs) can offer critical insights into microbial lifestyles (Swan et al. 2011) and shed light on intra-population structures of complex microbial consortia (Kashtan et al. 2014) or naturally occurring host-virus interactions (Labonté et al. 2015), where short-read and assembly-based strategies may not be effective. However, state-of-the-art single-cell sequencing strategies typically generate highly fragmented and incomplete genomes because of the need for random amplification arising from small quantities of DNA present in a single cell (Kalisky and Quake 2011). In some cases, sequences from other organisms may contaminate individual wells (Rinke et al. 2013), in other cases combining sequences from different cells into single draft genomes based on sequence identity thresholds of phylogenetic markers (i.e. >97% 16S rRNA identity) (Rinke et al. 2013), may result in hybrid genomes. In fact these hybrids are potentially from different species, given that many consider 97.9% 16S rRNA sequence divergence as a proxy for the species boundary (Newton et al. 2007; Garcia et al. 2018). Probst et al. (2018) indicate that although the cells are often chosen for single-cell sequencing based on their amplified 16S rRNA genes, the sequences recovered do not always match the amplified genes. Some of these problems may be ameliorated with additional steps of binning and refinement, and similar to MAGs, SAGs can also be curated to completion as shown by at least one study that used long (Sanger) reads in conjunction with shortread assemblies (Woyke et al. 2010). Given the fast pace of improvements in microfluidics technologies as well as whole-genome amplification and sequencing chemistry (Woyke et al. 2017), we anticipate that single-cell genomics will continue to gain popularity, and its joint use with other genome-resolved metagenomics strategies will become increasingly frequent.

Complete genomes from long-reads

Among the published CMAGs, four were obtained by assembly of PacBio reads, including three proteobacterial and one Bacteroidetes genome (White et al. 2016; Driscoll et al. 2017). Especially Oxford Nanopore Technologies offers affordable, easyto-operate, and portable sequencers for long-read sequencing. Although improving, errors from nanopore sequencing can exceed state-of-the-art short-read sequencing (Laver et al. 2015); however, new approaches for long-read correction (Rang et al. 2018; Arumugam et al. 2019), hybrid assembly (Wick et al. 2017), and mock community standards (Nicholls et al. 2019) are emerging. Short-read-based assembly strategies often report fragmented contigs caused by repeat elements that exceed short-read lengths, which is an issue long-read sequencing overcomes, improving the quality of genomes from metagenomics (Arumugam et al. 2019). We anticipate that the combination of short reads and long-reads sequencing will be an increasingly common strategy for recovery of highly curated and complete genomes from microbial community samples.

Chromosome conformation capture method

The chromosome conformation capture (i.e., 3C) is a method that enables the determination of physical contacts between different regions of a chromosome and between the different chromosomes of a cell (Dekker et al. 2002). The initial applications of this strategy focused on eukaryotic genomes and revealed, for example, the folding principles (Lieberman-Aiden et al. 2009) and the chromatin looping (Rao et al. 2014) of the human genome. The 3C approach has recently been developed into multiple derivative proximity ligation methods, such as Hi-C (Lieberman-Aiden et al. 2009) and meta3C (Marbouty et al. 2014), and applied to individual microbial populations (Le et al. 2013) as well as complex assemblages of environmental microbes (Marbouty et al. 2014). As these approaches offer physical linkage between DNA fragments that are proximal to each other, they can improve metagenomic binning (Baudry et al. 2019; DeMaere and Darling 2019). Although promising, the additional complexity of library preparations and additional cost because of the need for separate metagenomic libraries (Liu and Darling 2015) prevent their routine application to metagenomic studies. In addition, distinct populations that are in close proximity in the input sample and repeat sequences may yield misleading contact signals and result in chimeric assemblies (Marbouty and Koszul 2015). Nevertheless, the application of proximity ligation strategies to naturally occurring complex microbial consortia can provide important insights (Bickhart et al. 2019; Stalder et al. 2019).

Eukaryotes and even macroorganisms

The assembly of draft eukaryotic genomes from shotgun metagenomes is possible, despite the large genome sizes of most eukaryotes. However, eukaryotic MAGs can be readily contaminated by fragments of genomes from coexisting bacteria and archaea (Boothby et al. 2015; Arakawa 2016), so careful evaluation is needed to avoid misleading conclusions (Delmont and Eren 2016). We have found that phylogenetic profiling of contigs based on best matches in reference databases can be an effective way to identify contaminating bacterial and archaeal sequences.

An important step for recovery of reasonable quality eukaryotic genomes from metagenomes is to separate assembled eukaryotic from prokaryotic genome fragments before binning. Then, eukaryote-specific gene predictions can be established and gene annotations used to estimate genome completeness. The *k*-merbased classifier, EukRep, was developed to accomplish this separation (West et al. 2018). Although eukaryote genome recovery from metagenomes is increasingly reported (Quandt et al. 2015; Mosier et al. 2016; Olm et al. 2019), to our knowledge, none have been extensively curated or completed. A major limitation on the quality of MAGs relates to genome fragmentation. Fragmentation is doubly problematic because small fragments are hard to bin accurately, and gaps result in incomplete gene inventories. Fragmentation can arise as a result of the presence of duplicated sequences (e.g., transposases, rRNA operons), but the most pronounced problems usually are the result of coexisting closely related strains that confuse de Bruijn graph-based assemblers (Olson et al. 2017). For example, although Prochlorococcus and SAR11 are among the most abundant bacteria in ocean habitats, the co-occurrence of closely related strains (Giovannoni 2017) leads to very fragmented MAGs and poor representation in the final data sets (Delmont et al. 2018; Tully et al. 2018). Of the three commonly used metagenomic assemblers, IDBA-UD, MEGAHIT, and metaSPAdes (Greenwald et al. 2017), metaSPAdes was best designed to handle microvariations between fragments from related strains to generate longer composite sequences (Olson et al. 2017). However, care should be taken when undertaking detailed analyses (e.g., biochemical testing) of open reading frames generated in this way because they may be chimeric.

Practically, another approach that can sometimes address the problem of assembly fragmentation caused by strain variety is collections of sequences from related samples (e.g., along a geochemical gradient) to identify communities in which there is much reduced complexity of related strains. For example, opportunities can arise because of the recent proliferation of one strain over the background of numerous closely related strains following changes in environmental conditions. In other words, if a genome cannot be recovered from one sample, look for it in related samples. We anticipate that this approach will be most effective for genome recovery from soil environments, where strain diversity can be extreme and environmental heterogeneity provides access to different strain mixtures.

Conclusions

Genomes derived from metagenomes have advanced our understanding of microbial diversity (Anantharaman et al. 2016; Hug et al. 2016; Parks et al. 2017) and metabolism (e.g., van Kessel et al. 2015; Anantharaman et al. 2018). These genomes are readily adopted by the scientific community for a wide variety of investigations, and errors will propagate. In fact, the proposal of a new nomenclature for large swaths of the Tree of Life based largely on MAGs (Parks et al. 2018) brings a potential crisis into focus. We conclude that it is imperative that complete, curated genomes are recovered for all major lineages (including those that lack any isolated representative). The increased span of phylogenetic coverage by complete genomes will provide a valuable reference set against which newly recovered genomes can be confidently compared and augment what has been achieved by the isolate-based Genome Encyclopedia of Bacteria and Archaea program (Wu et al. 2009). New complete sequences from previously genomically undescribed lineages will also improve understanding of how protein families and functions are distributed, facilitate more powerful analyses of evolutionary processes such as lateral gene transfer, and enable more accurate phylogenetic representations of life's diversity. Finally, we advocate for the development of methods to routinely curate assemblies and draft genomes (if not to completion) at scale to ensure the accuracy of evolutionary and ecosystem insights.

Methods

Preparation of MAGs as examples for genome curation

This study includes two MAGs that were not previously published as examples for genome curation. These genomes were assembled from samples collected in a mine tailings impoundment (Manitoba, Canada). The raw reads of metagenomic sequencing were filtered to remove Illumina adapters, PhiX and other Illumina trace contaminants with BBTools (https://sourceforge .net/projects/bbmap/), and low-quality bases and reads using Sickle (version 1.33; https://github.com/najoshi/sickle). The high-quality reads were assembled using both IDBA-UD (Peng et al. 2012) and metaSPades (Nurk et al. 2017). For a given sample, the quality trimmed reads were mapped to the assembled scaffolds using Bowtie 2 with default parameters (Langmead and Salzberg 2012). The coverage of each scaffold was calculated as the total number of bases mapped to it divided by its length. The proteincoding genes were predicted from the scaffolds using Prodigal (Hyatt et al. 2010) and searched against KEGG, UniRef100, and UniProt for annotation. The 16S rRNA gene was predicted using a HMM model, as previously described (Brown et al. 2015). The tRNAs were predicted using tRNAscan-SE 2.0 (Lowe and Chan 2016). For each sample, scaffolds with a minimum length of 2.5 kbp were assigned to preliminary draft genome bins using MetaBAT with default parameters (Kang et al. 2015), with both tetranucleotide frequencies (TNF) and coverage profile of scaffolds (from multiple samples) considered. The scaffolds from the obtained bins and the unbinned scaffolds with a minimum length of 1 kbp were uploaded to ggKbase (http://ggkbase.berkeley.edu/). The genome bins were evaluated based on the consistency of GC content, coverage, and taxonomic information, and scaffolds were identified as contaminants were removed.

GC skew evaluation of RefSeq genomes

We analyzed all the NCBI RefSeq genomes downloaded on May 10, 2017, for GC skew. Both skew and cumulative skew were calculated, and patterns were displayed using the publicly available program gc_skew.py (https://github.com/christophertbrown/iRep) (Brown et al. 2016).

Refinement of the CPR genome from blood

For initial characterization of the CPR bin, we used the contigs made publicly available as the "Dataset S6" in the original study (Kowarsky et al. 2017). These contigs represent what remained after the removal of contigs with matches to sequences in any existing public databases (Kowarsky et al. 2017); we will refer to these contigs as "novel contigs." In our study we also had access to the remaining contigs, and we will refer to this data set as "all contigs."

For binning and refinement of the CPR genome, and metagenomic read recruitment analyses, we used anvi'o v5.5 to generate a contigs database from the novel contigs using the program "anvigen-contigs-database," which recovered the tetranucleotide frequencies for each contig; we used Prodigal v2.6.3 (Hyatt et al. 2010) with default settings to identify open reading frames; and HMMER v3.2.1 (Eddy 2011) was used to identify matching genes in our contigs to bacterial single-copy core genes by (Campbell et al. 2013). To visualize all novel contigs, we used the program "anvi-interactive," which computed a hierarchical clustering dendrogram for contigs using Euclidean distance and Ward linkage based on their tetranucleotide frequency (TNF), and displayed additional data layers of contig cohort origin; HMM hits we supplied to the program as a TAB-delimited additional data file. We manually selected a branch of contigs that created a coherent cluster based on the TNF data and the occurrence of bacterial singlecopy core genes. Although this procedure allowed us to identify an initial genome bin with modest completion, its comprehensiveness and purity was questionable because our binning effort (1) used only the novel contigs from Kowarsky et al. (2017), which were a subset of all contigs assembled, and (2) only used tetranucleotide signatures to identify the genome bin, which can introduce contamination as the sequence signatures of short fragments of DNA can be noisy. To address these issues, we first acquired the remaining 3002 contigs that were not included in the original study (Kowarsky et al. 2017) and that might be derived from the same blood-associated CPR population. Then, we used all blood metagenomes for a read recruitment analysis. This analysis allowed us to identify contigs from the non-novel contig collection that match to the distribution patterns of the initial CPR bin. Because the coverage of this population was extremely low, we used a special clustering configuration for anvi'o to use "differential detection" rather than "differential coverage" (see the reproducible workflow for details). This analysis resulted in contigs with similar detection patterns across all metagenomes. We summarized this final collection of contigs using "anvi-summarize," which gave access to the FASTA file for the bin. Anvi'o automated workflows (http://merenlab.org/2018/07/09/anvio-snakemakeworkflows/) that use Snakemake (Köster and Rahmann 2012) performed all read recruitment analyses with Bowtie 2 (Langmead and Salzberg 2012). We profiled all mapping results using anvi'o following the analysis steps outlined in Eren et al. (2015).

To put our CPR bin into the phylogenetic context of the other available CPR genomes, we used the 797 metagenome-assembled CPR genomes (Brown et al. 2015). We used the anvi'o program "anvi-get-sequences-for-hmm-hits" to (1) collect the 21 aminoacid sequences found in the CPR bin (Ribosomal_L10, Ribosomal_L11, Ribosomal_L11_N, Ribosomal_L13, Ribosomal_ L14, Ribosomal L17, Ribosomal L20, Ribosomal L21p, Ribosomal_L27, Ribosomal_L32p, Ribosomal_L5_C, Ribosomal_ Ribosomal_L9_N, Ribosomal_S13, L9_C, Ribosomal_S11, Ribosomal_S16, Ribosomal_S2, Ribosomal_S20p, Ribosomal_S4, Ribosomal_S7, Ribosomal_S9) from all genomes, (2) align them individually, (3) concatenate genes that belong to the same genome, and (4) report them as a FASTA file. Some of the key parameters we used with this program included "--hmm-source Campbell_et_al" to use the single-copy core gene collection defined by Campbell et al. (2013), "--align-with famsa" to use FAMSA (Deorowicz et al. 2016) to align sequences for each ribosomal protein, "--returnbest-hit" to get only the most significant HMM hit if a given ribosomal protein found in multiple copies in a given genome, and "--max-num-genes-missing-from-bin 3" to omit genomes that miss more than three of the 21 genes listed. We used trimAl v1.4.rev22 (Capella-Gutiérrez et al. 2009) to remove positions that were gaps in >50% of the genes in the alignment (-gt 0.50), IQ-TREE v1.5.5 (Nguyen et al. 2015) with the "WAG" general matrix model (Whelan and Goldman 2001) to infer the maximum likelihood tree, and anvi'o was used to visualize the output.

Refinement of the Pasolli MAG

We downloaded the Pasolli MAG ("HMP_2012_SRS023938_ bin_39"; https://opendata.lifebit.ai/table/SGB) and the 481 HMP oral metagenomes from the HMP FTP server (ftp://publicftp.hmpdacc.org/Illumina/). We used anvi'o v6 and the Snakemake-based (Köster and Rahmann 2012) program "anvirun-workflow" to run the anvi'o metagenomics workflow (Eren et al. 2015). Briefly, we generated a contigs database from the Pasolli MAG FASTA file by running "anvi-gen-contigs-database," during which anvi'o calculates tetranucleotide frequencies for each contig, and Prodigal (Hyatt et al. 2010) to identify genes. In order to estimate the completion and redundancy of the Pasolli MAG based on SCGs, we used the program "anvi-run-hmms" with the default HMM profiles, which include 71 bacterial SCGs (HMMs described in anvi'o v6), and annotated genes with functions using "anvi-run-ncbi-cogs," which searches amino-acid sequences using BLASTP v2.7.1+ (Altschul et al. 1990) against the December 2014 release of the COG database (Tatusov et al. 2000). We mapped the paired-end reads from the 481 HMP metagenomes to the Pasolli MAG using Bowtie 2 with default parameters (Langmead and Salzberg 2012) and converted the mapping output to BAM files using SAMtools v1.9 (Li et al. 2009). We used "anvi-profile" to generate profile databases from BAM files, in which coverage and detection statistics for contigs in each metagenome were stored. We used "anvi-merge" to merge the anvi'o profile databases of (1) only the three plaque metagenomes of HMP individual 159268001, which includes the sample from which the Pasolli MAG was constructed (sample accession SRS023938), and (2) all 481 HMP oral metagenomes. In order to manually refine the Pasolli MAG, we ran the anvi'o interactive interface using "anvi-interactive" with the merged anvi'o profile database that included only the three plaque metagenomes of HMP individual 159268001. Refinement was done using hierarchical clustering of the contigs based on sequence composition and differential coverage using Euclidean distance and Ward's method. To estimate the taxonomic assignment, we blasted the protein sequences of genes in the 11 contigs identified as contamination against the NCBI's nonredundant protein sequences database. To visualize the detection values of the contigs of the Pasolli MAG across all 481 HMP oral metagenomes, we used the full merged profile database and the program "anvi-interactive." We used "anvi-summarize" to generate tabular summaries of detection and coverage information of the refined Saccharibacteria bin and the 11 contigs of contamination across the 481 metagenomes.

Data access

All the five complete genomes reconstructed in this study have been submitted to the European Nucleotide Archive (ENA; https://www.ebi.ac.uk/ena) under accession numbers ERS4269018, ERS4269226, ERS4270539, ERS4266120, and ERS4267492.

Competing interest statement

The authors declare no competing interests.

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