Single-Cell-Genomics-Facilitated Read Binning of Candidate Phylum EM19 Genomes from Geothermal Spring Metagenomes

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The vast majority of microbial life remains uncatalogued due to the inability to cultivate these organisms in the laboratory. This “microbial dark matter” represents a substantial portion of the tree of life and of the populations that contribute to chemical cycling in many ecosystems. In this work, we leveraged an existing single-cell genomic data set representing the candidate bacterial phylum “Calescamantes” (EM19) to calibrate machine learning algorithms and define metagenomic bins directly from pyrosequencing reads derived from Great Boiling Spring in the U.S. Great Basin. Compared to other assembly-based methods, taxonomic binning with a read-based machine learning approach yielded final assemblies with the highest predicted genome completeness of any method tested. Read-first binning subsequently was used to extract Calescamantes bins from all metagenomes with abundant Calescamantes populations, including metagenomes from Octopus Spring and Bison Pool in Yellowstone National Park and Gongxiaoshe Spring in Yunnan Province, China. Metabolic reconstruction suggests that Calescamantes are heterotrophic, facultative anaerobes, which can utilize oxidized nitrogen sources as terminal electron acceptors for respiration in the absence of oxygen and use proteins as their primary carbon source. Despite their phylogenetic divergence, the geographically separate Calescamantes populations were highly similar in their predicted metabolic capabilities and core gene content, respiring O2, or oxidized nitrogen species for energy conservation in distant but chemically similar hot springs.

The vast majority of the diversity of microbial life on Earth remains undiscovered; the core metabolisms of yet-uncultivated species, interguild interactions within natural and managed ecosystems, and the contributions of microbial populations to the geochemistry of the environment remain poorly understood (1, 2). There are currently over 60 bacterial and archaeal phylum-level groups that have been observed through the use of 16S rRNA gene sequencing and phylogenetics, with over half containing no cultivated representatives (3). This so-called microbial dark matter comprises a substantial proportion of the tree of life and of microbial communities that likely play significant roles in biogeochemical cycles in a variety of environments (4–9).

Metagenomic analyses of low-diversity microbial communities have yielded robust, near-complete genomic assemblies representative of the abundant populations, expanding the knowledge of the metabolic potential of predominant organisms in these ecosystems (10–14). Nucleotide word frequencies calculated from metagenomic contiguous assembled sequences (contigs) have been used to separate population-specific clusters, or “bins,” from the community DNA pool (15–17), which has greatly advanced our understanding of the genomic diversity in natural environments and how populations differ between chemically distinct environments and along environmental gradients (18, 19). Even using modern sequencing techniques, however, it can be challenging to confidently separate and assemble genomes of community members using metagenomic data alone due to problems such as low population abundance, high community diversity, or similarities in nucleotide word frequencies between phylogenetically distant taxa.

High-throughput, semiautomated isolation and sequencing of single cells representing candidate phyla has opened the door for systematic analysis of environmental DNA that can be unambiguously assigned to these uncultured organisms (4, 20). Great Boiling Spring (GBS), located in the U.S. Great Basin, harbors abundant populations of several candidate phyla, and the low biological diversity in high-temperature sediments facilitates access to their genomes (4, 21–24). Previously, 10 separate single cells representing a deeply branching lineage in the bacterial domain, candidate phylum EM19, were isolated from GBS sediments, sequenced, and assembled into draft single amplified genomes (SAGs) ranging from 0.3 to 1.9 Mbp in length (4). The coassembly of eight of the 10 SAGs yielded a draft genome of 2.24 Mbp that was estimated to be 94% complete based on the proportion of 139 single-copy conserved bacterial markers (SCMs) observed in the assembly (4). Phylogenomic analyses confirmed the independence of EM19 from other bacterial phyla, justifying a
proposal to name the lineage more formally as candidate phylum “Calescamantes” and the genus and species as “Candidatus Calesciibacterium nevadense” [4]. Although core metabolic genes were identified, this work was part of a much larger single-cell genomic survey, and did not describe the metabolic capabilities of “Ca. Calesciibacterium nevadense” or encompass Calescamantes genomes from other locations.

The goals of the current study were to obtain greater phylogenetic, genetic, and genomic coverage of members of the Calescamantes. Although metagenonomic read binning utilizing read abundances (25) and k-mer frequencies (25, 26) have been utilized previously in whole-community analyses, none have used SAGs as environmentally relevant anchors to improve clustering and genome quality. Here, we assess the effectiveness of a SAG-assisted, read-based binning approach for identifying Calescamantes sequences in metagenomes prior to assembly and to use the resulting genomic data sets to predict the metabolic potential and conserved features of the Calescamantes in detail. Nucleotide word frequencies obtained from SAGs representative of several candidate phyla abundant in GBS and other environmentally relevant genomes were used as anchors to confidently separate Calescamantes sequence reads from environmental metagenomes prior to assembly using a multilayer perceptron (MLP) machine learning approach. Because this approach of read-binning prior to metagenomic assembly has the potential advantage of avoiding chimeric artifacts that can occur during assembly, we compared it with other commonly used assembly-based binning methods on the GBS metagenome [15, 27, 28]. As read-binning outperformed assembly-based technologies on the GBS metagenome, these read-binning methods were also used to confidently predict taxonomic bins from all available metagenomes that contained relatives of “Ca. Calesciibacterium nevadense,” including metagenomes from geothermal springs in Yellowstone National Park (YNP) (Bison Pool and Octopus Spring) and Yunnan Province, China (Gongxiaoshe Spring), expanding both the geographic and phylogenetic coverage of Calescamantes genomic data sets. Comparative analysis of these distinct Calescamantes populations identified a predicted core metabolism that couples the oxidation of Ca

MATERIALS AND METHODS

**SAG data sets, metagenome sampling, and metagenomic DNA extraction.** Ten individual Calescamantes SAGs and a combined SAG assembly obtained from GBS, described previously [4] were retrieved from the IMG system. Previously published [14, 19] metagenomes from Octopus Spring and Bison Spring in YNP were accessed from the IMG system as summarized in Table S1 in the supplemental material. Associated sampling sites, dates, and temperatures are shown in Table S1 in the supplemental material.

Sediment samples for metagenomic analyses were collected from the north edge of the source pool of GBS (site A [80°C] in reference [23]) on 2 December 2008 and from the bottom of Gongxiaoshe Spring (Tengchong, China) on 10 January 2011 as described by Hou et al. [14]. DNA was extracted from both sediments using the FastDNA spin kit for soil (MP Biomedicals, Solon, OH) according to the manufacturer’s protocol.

**DNA sequencing and assembly of whole metagenomes.** Library preparation and sequencing of the GBS metagenome using the 454 FLX platform with Titanium chemistry (Roche, Branford, CT) was performed at the Joint Genome Institute (JGI). GBS metagenomic reads were assembled at the JGI with Newbler 2.4 using a minimum nucleotide identity of 98% and a minimum overlap of 80 bp [29].

For the Gongxiaoshe metagenome, the extracted DNA was sheared to 500 bp using a Covaris ultrasonicator (Covaris Inc., Woburn, MA) according to the manufacturer’s recommendations. The sheared DNA was end repaired, adapter ligated with multiplexing, and purified using the Ovation SP ultralow DR multiplex system (NuGEN Technologies Inc., CA), and the resulting libraries were sequenced (2 by 250) using the Illumina MiSeq platform (Illumina, San Diego, CA). Gongxiaoshe metagenome reads with no ambiguous bases, quality score of >20, and length of >74 bases were assembled using the CLC Genomic Workbench, version 6.0, de Bruijin graph assembler (CLC bio, Boston, MA), taking into account paired reads. After binning, only contigs of at least 500 bases long were included in the final curated assembly.

**Binning and assembly of Calescamantes reads from GBs based on machine learning.** For read-binning prior to assembly, metagenomic reads were binned by nucleotide trimer frequencies using the MLP machine learning package in WEKA, version 3.6, with default parameters [30]. This package uses back-propagation to facilitate the training of nonlinear networks. The MLP initially was trained using nucleotide frequencies of clipped genomic segments (2,000 bp; also see Fig. S1 in the supplemental material) from the GBS Calescamantes SAG coassembly. SAGs and isolate genomes representing other abundant populations in GBS included Calescamantes (JGI_2527291514), “Ferribidibacter” (JGI_0000001-G10), a novel crenarchaeota, “Geoarchaeta” [JGI_2264867219], and the Thermoflexus hugenolitzi strain JAD2 draft genome [JGI_2140918011] [31]. This approach provides multiple points of reference during training of the MLP algorithm. Metagenomic reads were assigned to the GBS Calescamantes population if their MLP confidence score was ≥0.9 (a score of 1 indicates 100% confidence), assessed as the point at which false positives were minimized while maximizing true positives (data not shown). This cutoff was designed to be relatively liberal in order to be inclusive of divergent or novel sequences with the intent to curate postassembly. To augment the MLP-based Calescamantes bin and provide access to tRNA/ rRNA [32] and other genomic regions with anomalous nucleotide frequency, predicted coding regions from the GBS Calescamantes SAG coassembly were queried against an unassembled GBS metagenomic nucleotide database using BLASTN [33], and matches with an E value of ≤1E−15 were selected for further incorporation into the assembly. All reads belonging to the Calescamantes bin were assembled with SPAdes using the “careful” setting and step k-mer values of 77, 99, 111, and 127. Sequence reads that did not assembles were removed (38,340 of 129,215), and assembled contigs were further curated as described below.

**Binning and assembly of Calescamantes reads from Gongxiaoshe Spring, Bison Pool, and Octopus Spring.** The above-described procedure was repeated for sequence reads from all other sites using MLP genomic anchors determined by the closest matching genomes in NCBI (those which had BLAST hits with the highest identity to metagenomic reads) and were representative of predominant populations in those environments. In addition to the Calescamantes SAG coassembly, the MLP was trained using Thermofluxus hugenolitzi [JGI_2140918011], Acetothermus autotrophicum [AP011800.1 to AP011803.1], Thermus aquaticus [NZ_ABV00000000.2], and the Fervidibacter SAG [JGI_0000001-G10] for Gongxiaoshe Spring; Thermus aquaticus [NZ_ABV00000000.2], Thermofluxus hugenolitzi [JGI_2140918011], Pyrobaculum islandicum [CP0005004], Thermocrinis ruber [CP000708.1], and the Archaeoaeta SAG [JGI_2264867219] for Octopus Spring [JGI_2264867219]; and Ther- mus aquaticus [NZ_ABV00000000.2], Hydrogenobacter thermophilus [AP011112], Pyrobaculum islandicum [CP000504.1], and Thermofluxus hugenolitzi [JGI_2140918011] for Bison Pool. Calescamantes bins from Gongxiaoshe and Octopus Spring were assembled using SPAdes, as described above. Bison Pool metagenome reads were assembled using Dragon (QuantiTreeX, Germany) using default settings, as SPAdes had difficulty assembling small numbers of Sanger sequences. Accession num-
FIG 1 Flow chart of the progression of data analysis, showing (left to right) sample collection; sequencing of single-cell genomes and reference GenBank genomes (stars) and metagenomes (circles); nucleotide trimer frequency calculation; MLP model training; and read-binning, assembly, and postassembly analyses (annotation, metabolic reconstruction, and estimation of genome completeness). The MLP pipeline (top and right) subsequently was conducted on all geographic sites analyzed in this study.
frequency using the MLP approach is shown in Fig. 1. Calescamantes SAG nucleotide trimer frequencies from GBS were similar to those of an archaeal population (Geoarchaeota) present in GBS at ~5% of the population (average Pearson correlation of ~0.92), complicating the recovery of Calescamantes sequences from the metagenome (4, 21, 49). This coincidental similarity in nucleotide frequency between these distant taxa confounded binning by principal component analysis alone. However, the availability of multiple SAGs from both Calescamantes and Geoarchaeota allowed us to train the MLP algorithm for confident bin assignment of the metagenome reads. The MLP classified 148,637 of 1,203,155 reads in the metagenome, and BLASTN separately identified an additional 14,098 nonredundant reads, for a total of 162,735 GBS metagenomic reads belonging to the Calescamantes population. After quality-control filtering, the SPAdes assembly of the bin containing reads identified by MLP and BLASTN consisted of 319 contigs assembled from 50,774 sequence reads. Reads that were included in this assembly totaled 4.2% of the metagenome, whereas Calescamantes had a relative abundance of 1.21% in a previous 16S rRNA gene analysis of the same DNA sample from which the GBS metagenome was derived (23). It is likely that the abundance based on metagenome reads more closely reflects the abundance of Calescamantes, because differences in ribosomal copy number and amplification bias can skew organisms’ relative abundances (50).

The Calescamantes MLP assembly from the GBS metagenomic reads binned using SAGs as training models, but without the inclusion of any SAG contigs in the assembly, shared a similar nucleotide trimer composition (average Pearson correlation of >0.98) with the GBS SAG coassembly and had near-identical G+C content (34.1% and 34.2%, respectively). An updated SAG coassembly provided by JGI and the MLP assembly were annotated on the RAST online server (40), identifying 2,205 coding regions in the SAG coassembly, 46% of which were classified as hypothetical proteins, and 2,721 coding regions in the GBS MLP assembly, of which 47% were classified as hypothetical proteins (Table 1 and Fig. 2A). The higher number of gene regions in the MLP assembly compared to that of the SAG coassembly at GBS

### Table 1: Statistics for the Calescamantes SAG coassembly and GBS, Gongxiaoshe Spring, Octopus Spring, and Bison Pool metagenome MLP assemblies

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value for:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>GBS SAG coassembly</td>
</tr>
<tr>
<td>Total no. of reads</td>
<td>NA&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>MLP binned reads (no.)</td>
<td>NA</td>
</tr>
<tr>
<td>Assembly size (Mbp)</td>
<td>2.25</td>
</tr>
<tr>
<td>No. of contigs</td>
<td>138</td>
</tr>
<tr>
<td>Largest contig (bp)</td>
<td>103,150</td>
</tr>
<tr>
<td>N&lt;sub&gt;90&lt;/sub&gt; content</td>
<td>25,634</td>
</tr>
<tr>
<td>G+C content</td>
<td>34.1</td>
</tr>
<tr>
<td>No. of coding sequences (RAST)</td>
<td>2,205</td>
</tr>
<tr>
<td>Assigned KO no. (KAAS)</td>
<td>851</td>
</tr>
<tr>
<td>No. (%) of hypothetical genes (RAST)</td>
<td>945 (46)</td>
</tr>
<tr>
<td>tRNAs (RAST)</td>
<td>49</td>
</tr>
<tr>
<td>tRNAs (KAAS)</td>
<td>49</td>
</tr>
<tr>
<td>23S</td>
<td>1</td>
</tr>
<tr>
<td>16S</td>
<td>1</td>
</tr>
<tr>
<td>5S</td>
<td>1</td>
</tr>
<tr>
<td>SCM duplicates&lt;sup&gt;b&lt;/sup&gt; (%)</td>
<td>5</td>
</tr>
<tr>
<td>Estimated completeness&lt;sup&gt;d&lt;/sup&gt; (%)</td>
<td>94</td>
</tr>
</tbody>
</table>

<sup>a</sup> Gongxiaoshe and Octopus Spring metagenomes contained paired-end sequences.

<sup>b</sup> Conserved gene copy number and genome completeness were estimated with MetaGeneMark and Prodigal using single-conserved markers (SCMs) described by Rinke et al. (4).

<sup>c</sup> If the values differed between programs, both values are given.

<sup>d</sup> NA, not applicable.
most likely is due to the presence of approximately 200 more contigs in the MLP assembly, which typically contained gene fragments on their discontinuous ends, supported by the fact that 64% of these additional truncated genes are duplicated elsewhere in the MLP assembly. The GBS MLP assembly contained 367 predicted genes that were not present in the SAG coassembly, although it is important to note that the MLP (and any metagenome) assembly is an assemblage of population-level diversity in the environmental sample. In contrast, there were 292 predicted coding regions in the SAG coassembly that were not detected in the MLP assembly. The GBS MLP assembly contained several predicted coding regions that closed some apparent metabolic gaps in the annotated SAG coassembly (namely, cases where some, but not all, genes in a pathway were present), including genes encoding proteins involved in fatty acid biosynthesis (fabZ), NADH-quinone oxidoreductase subunit F (nuoF), and DNA gyrase subunit A (gyrA). Likewise, the SAG coassembly identified metabolic gaps in the GBS MLP assembly: protein biosynthesis gene 3-dehydroquinate dehydratase I (aroE), flagellar basal body rod gene (floB), and glycol-tRNA synthetase (glyQ). Both the MLP and SAG assemblies contained ribosomal proteins that were not detected in the other assembly (see Table S5 in the supplemental material).

**Table 2** SAG coassembly statistics compared to those of different binning methods on *Calescamantes* populations from GBS

<table>
<thead>
<tr>
<th>Binning method</th>
<th>N&lt;sub&gt;cont&lt;/sub&gt;</th>
<th>Maximum (bp)</th>
<th>Minimum (bp)</th>
<th>Avg. contig size (bp)</th>
<th>Genome size (Mb)</th>
<th>No. of contigs</th>
<th>%cov&lt;sup&gt;a&lt;/sup&gt;</th>
<th>No. of SCMs at ≥1 copy</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAG</td>
<td>40,523</td>
<td>204,003</td>
<td>320</td>
<td>16,262</td>
<td>2.24</td>
<td>138</td>
<td>95</td>
<td>93</td>
</tr>
<tr>
<td>MLP plus BLAST&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>11,262</td>
<td>49,870</td>
<td>513</td>
<td>6,937</td>
<td>2.21</td>
<td>319</td>
<td>90</td>
<td>89</td>
</tr>
<tr>
<td>MLP contig bin</td>
<td>11,223</td>
<td>44,899</td>
<td>129</td>
<td>5,375</td>
<td>2.33</td>
<td>435</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td>MLP&lt;sup&gt;c&lt;/sup&gt;</td>
<td>5,510</td>
<td>30,007</td>
<td>135</td>
<td>3,843</td>
<td>2.03</td>
<td>528</td>
<td>83</td>
<td>83</td>
</tr>
<tr>
<td>ESOM plus BLAST&lt;sup&gt;d&lt;/sup&gt;</td>
<td>4,968</td>
<td>22,905</td>
<td>100</td>
<td>1,867</td>
<td>2.19</td>
<td>1,177</td>
<td>84</td>
<td>83</td>
</tr>
<tr>
<td>MaxBin plus BLAST&lt;sup&gt;d&lt;/sup&gt;</td>
<td>4,057</td>
<td>22,905</td>
<td>103</td>
<td>2,509</td>
<td>2.13</td>
<td>805</td>
<td>84</td>
<td>83</td>
</tr>
<tr>
<td>Metawatt</td>
<td>4,023</td>
<td>22,905</td>
<td>900</td>
<td>3,076</td>
<td>2.04</td>
<td>138</td>
<td>81</td>
<td>81</td>
</tr>
<tr>
<td>BLASTN</td>
<td>4,109</td>
<td>14,181</td>
<td>501</td>
<td>2,832</td>
<td>1.89</td>
<td>668</td>
<td>79</td>
<td>77</td>
</tr>
</tbody>
</table>

<sup>a</sup>Percent total genome coverage (%cov) was estimated with MetaGeneMark (M) and Prodigal (P) using single conserved markers (SCMs) as described by Rinke et al. (4).

<sup>b</sup>BLAST indicates the inclusion of nonredundant SAG-identified BLAST reads in the assembly.

<sup>c</sup>MLP-only assembly without including nonredundant SAG-identified BLAST reads in the assembly.

<sup>d</sup>MLP-only assembly without including nonredundant SAG-identified BLAST reads in the assembly.
SCMs, indicating that this assembly was composed of two distinct but closely related *Calescamantes* populations (Table 1). A comparison of 16S rRNA gene sequences recovered by BLASTN from the Octopus metagenome to the 16S rRNA gene in the corresponding assembled genome exhibited a bimodal pattern of sequence identity (see Fig. S3 in the supplemental material), lending further evidence to the concept of the presence of two very closely related *Calescamantes* populations in Octopus Spring. To obtain a higher-fidelity assembly, contigs from the *Calescamantes* populations in Octopus Spring were separated by top BLASTN hits to the predicted gene-coding regions identified in the GBS SAG. The resulting Octopus Spring assembly consisted of 799 contigs, a largest contig of 26,269 bp, and was estimated to be 87% complete. RAST identified a total of 2,040 predicted coding regions, 40% of which were classified as hypothetical proteins, and 363 genes were predicted gene-coding regions identified in the GBS SAG. The Octopus metagenomic assemblies all code for complete or near-complete pathways for purine and pyrimidine nucleotide metabolism, fatty acid biosynthesis and degradation, peptidoglycan and cell membrane biosynthesis, and crucial genes involved in the cell cycle (e.g., *ftsZ*). Genes predicted to encode ABC transporters in-include components of the NADH dehydrogenase complex I (*nuo*), succinate/fumarate dehydrogenase complex II (*sdh*), and the cytochrome *c* oxidase complex IV (*cox*). Whereas bc complex III proteins were not annotated, an alter-native complex III (ACHI) that performs the same function, despite being evolutionarily distinct, was identified (57). Genes encoding an A-type oxygen reductase and all but an epsilon subunit of a putative F-type ATP synthase were also annotated. The Gongxiaoshe assembly contained nitrate reductase (*nara*) and nitrous oxide reductase (*nosZ*) genes, but nitrite reductase (*nirS* or *nirK*) was not identified. North American assemblies contained putative oxygen limitation-sensing genes (*fisL*), as well as nitrite reductase (*nirS*), nitric oxide reductase (*eNOR* family), and nitrous oxide reductase (*nosZ*) genes. The *nirS* gene was predicted to encode an enzyme using a heme-heme binding site (58). *nosZ* and the *eNOR* gene both contained copper-copper (CUA/CUB) binding sites (59, 60). The putative *eNOR* gene was not closely related to anything in NCBI GenBank, with a closest BLAST hit to *Hydrogenobacter thermophilus* TK-6 (E value of 1E−22). North American assemblies did not contain *naraG*. The gene encoding nitric oxide reductase (*nora*), catalyzing the reduction of nitric oxide to nitrous oxide, was not detected in any of the SAG or metagenomic assemblies, although the *eNOR* gene, an energy-conserving alternate nitric oxide reductase, was identified in both the GBS and Gongxiaoshe assemblies (61).

The SAG coassembly and the GBS, Gongxiaoshe, and Octopus Spring metagenomic assemblies all code for complete or near-complete pathways for pyruvate and pyrimidine nucleotide metabolism, fatty acid biosynthesis and degradation, peptidoglycan and cell membrane biosynthesis, and crucial genes involved in the cell cycle (e.g., *ftsZ*). Genes predicted to encode ABC transporters involved in lipopolysaccharide and lipoprotein transport across the cytoplasmic membrane used for outer membrane and lipopolysaccharide (LPS) assembly were present (e.g., *rfb*, *lmbB* and *lol*), as were other outer membrane proteins typically associated with Gram-negative (diderm) organisms (see Table S4 in the supplemental material) (47). Genes involved in DNA replication, nucleotide excision and mismatch repair, and homologous recombination were also present (e.g., *DNA polIII*, *uvrA*, *mutA*, and *recA*). With the exception of assemblies from Bison Pool, putative genes encoding all tRNA synthetases and at least one tRNA for each amino acid were identified in all assemblies.

**Calescamantes** contained near-complete pathways for amino acid biosynthesis and peptide ABC transporters (e.g., *psl* and *liv*)
FIG 3 Maximum-likelihood phylogeny based on partial 16S rRNA genes (953 bp) representing all sequenced members within the Calescamantes phylum (≥85% identity), as well as members of other well-known bacterial phyla. Hot spring locations from which Calescamantes sequences were obtained are shown on the right. Black dots indicate bootstrap values of ≥90, and white dots indicate values of ≥50. The SAG coassembly and MLP assembly 16S rRNA gene sequences are shown together on the same branch, as they were 100% identical. 16S rRNA gene sequences from Cole et al. (23), which branch as a close sister to the SAG/MLP 16S rRNA gene (99.2 to 100% identical), were not included in the phylogeny due to their short length (200 to 400 bp). The scale bar indicates 0.05 substitutions per site. Sequences originating from MLP assemblies (MLP), SAGs, and/or amplified 16S rRNA gene clone libraries (Clone) are indicated in parentheses.
and the putative capability to import long branched-chain amino acids and oligopeptides (e.g., opp). These transport proteins could be the primary mechanism for carbon uptake in these organisms. Besides being a probable electron donor for respiration, peptides and amino acids could also be an important resource for anabolic reactions, as no key enzymes for autotrophic metabolism were identified (e.g., RubisCO, ATP citrate lyase, pyruvate:ferredoxin oxidoreductase, acetyl-coenzyme A synthase, or carbon monoxide dehydrogenase). Calescamantes also contained a putative D-xylose transporter (xyl) but no other known xylose utilization enzymes, suggesting that any xylose utilization would have to be through novel means. Additional membrane transport proteins putatively identified in Calescamantes assemblies include those specific to heme (ccm), potassium (kdp), copper (cus), phosphate (pst), and alkanesulfonate (ssu) uptake, and all assemblies had complete assimilatory sulfate reduction pathways (cys), indicating the ability to transport and anabolize trace elements and cofactors necessary for enzymatic activity.

The Calescamantes populations also appear capable of chemotactic motility, as they code for near-complete bacterial chemotaxis and flagellar assembly systems. Putative genes were identified for methyl-accepting chemotaxis proteins (mcp) and two-component sensor kinases (che), flagellum motor switch genes (fli), ATP motor proteins (mot), and chemotaxis motor proteins (mot). In addition, multiple flagellar structural assembly genes were identified (fli, flh, and fliG), indicating that these organisms have the capability to both sense and respond to a changing chemical environment, although fliC propeller filament genes were absent from all annotated genome assemblies. Annotated genes are listed in Tables S5A to E in the supplemental material.

Gene content differences between geographically separated populations. Many of the coding regions in the Gongxiaoshe and GBS MLP assemblies were hypothetical (53 and 47%, respectively), and most genes unique to the Calescamantes populations in each location (610 and 663, respectively) were also hypothetical (92 and 94%, respectively). However, the Gongxiaoshe metagenome assembly did contain unique genes with annotated functions that may be ecologically important. Although the nosZ and eNOR genes were present in the Gongxiaoshe assembly, nirS was not identified, suggesting that these populations lack the ability to reduce nitrite to nitric oxide. However, narG (large subunit) was annotated, suggesting that the Gongxiaoshe populations have the capability to reduce nitrate to nitrite, supported by the presence of site-specific genes that code for the transport of molybdenum (modA and modB), which is used as a cofactor in nitrate reductase (NarG) (62, 63), although narH (small β subunit) was not identified. Additionally, formate and short-chain fatty acid transporters were identified (foc and ato) as well as an acetate kinase (ack), indicating that the Calescamantes populations in Gongxiaoshe have more capabilities for carbon assimilation and metabolism.

The North American assemblies (GBS, Bison Pool, and Octopus Spring) all contained nirS as well as nosZ, which clustered phylogenetically with sequences from members of the Aquificae (see Fig. S6 in the supplemental material). The GBS assembly also contained the eNOR gene, which reduces nitric oxide to nitrous...
oxide, completing the denitrification pathway from nitrite to dinitrogen (61). In addition to nirS, GBS and Octopus Spring assemblies also contained uniquely annotated genes for glutamate symport (glt), carbon starvation (csf and dps), heat shock chaperone proteins (groEL), protein metabolism (panD), alcohol dehydrogenase (adh), potassium uptake (kup), and ammonia transport (ant), which could give insight into important environmentally specific metabolisms; the spring at Gongxiaoshe, for instance, is cooler in temperature and ammonia was not detected (14). Additional trehalose metabolism genes (treT and otsA) were annotated exclusively at GBS, which may enable the use of trehalose as an alternative carbon source for these organisms; however, no trehalose-specific transport proteins were annotated. GBS assemblies also exclusively contained unique CRISPR regions (cas 1 and 3), indicating more viral activity at this spring. The Octopus Spring assembly exclusively contained the cell structure gene mreB, suggesting these Calescamantes are rod-shaped.

**DISCUSSION**

*Calescamantes* 16S rRNA gene and protein-coding sequences are representative of a yet-uncultivated bacterial phylum that is very poorly represented in sequence databases (4). To date, *Calescamantes* 16S rRNA gene sequences have been identified and recovered from only a small number of chemically similar, circumneutral to alkaline hot springs in North America and Tengchong, China (14, 18, 29, 51, 53).

As DNA sequencing technology advances at an extremely rapid pace, it has become possible to assemble nearly complete genomes from uncultured candidate phylum populations in environmental metagenomes. However, even with these advances, small population sizes and assembly artifacts can make analysis difficult for rare community members. Single-cell genomics allows for confident anchoring and binning of metagenomic reads by facilitating the identification and analysis of genomic data from low-abundance populations in diverse environmental metagenomes.

Compared to commonly used postassembly binning methods, the MLP read-binning approach used in this study yielded a slightly more complete genomic assembly from metagenome sequences in each case tested without increasing SCM redundancy or decreasing the N50 value, although SAG-assisted postassembly binning with MLP yielded the most comparable results to read-first binning with MLP. The coordinated analysis of metagenomic assemblies and SAG coassemblies has allowed us to expand upon the limitations of each technique, fill apparent gaps in metabolic pathways, and confidently predict and contrast the metabolic potential of the *Calescamantes* populations in several geothermal environments, even when the genomes used for MLP training were genetically divergent from the populations in those environments, as was the case at Gongxiaoshe. While 16S rRNA gene sequencing, single-cell genome sequencing, and community metagenomics are informative on their own, the intersection of these sequencing approaches is greater than the sum of their parts. As sequencing continues to expand at exponential rates, it has become increasingly important to adopt a multifaceted approach in order to improve read-binning, assembly, and annotation for the ultimate goal of accurately reconstructing metabolic potential for yet-uncultivated microbial populations from the natural environment. The integrative MLP read-based binning approach represents an additional tool for organism-based, cultivation-independent genomic analysis and can potentially enhance the understanding of uncultivated microbes beyond what can be obtained by any individual approach.

*Calescamantes* populations in GBS, Yellowstone, and Gongxiaoshe are predicted to be heterotrophic, motile bacteria that have the capability to respire aerobically and incorporate carbon into biomass through protein transport and metabolism (Fig. 4; also see Fig. S4 and S5 in the supplemental material). Additionally, these *Calescamantes* populations are putative facultative anaerobes, capable of reducing oxidized nitrogen sources in the absence or limitation of oxygen. The sediment environment in GBS is likely to be either hypoxic or anoxic, and the bulk water in GBS has a relatively low oxygen tension (25 to 50 μM), with nitrate and nitrite concentrations of 1.8 to 16 and 0.79 to 10.2 μM, respectively (22). Previous analysis at this spring has shown that chemolithothrophic ammonia oxidation and denitrification are active processes in GBS sediments (64). Although chemolithothrophic nitrite oxidation has not been measured directly in GBS, enrichment cultures for nitrite-oxidizing bacteria (NOB) in GBS and other geothermal springs failed to show evidence of nitrite oxidation above ~65°C (65), and evidence of NOB above these temperatures is absent from 16S rRNA gene censuses (23, 24, 64) and metagenomes (data not shown). These data are consistent with a model in which ammonia from the source pool is oxidized to nitrite by ammonia-oxidizing archaea in the genus “*Nitrosocaldus,*” followed by anaerobic respiration of nitrite to nitrous oxide or dinitrogen (22, 64). In addition to the predicted capacity for nitrite reduction to dinitrogen by *Calescamantes*, *Thermus thermophilus* isolates from GBS are also capable of reduction of nitrate to nitrous oxide but lack the ability to reduce nitrous oxide to dinitrogen (22). As such, *Thermus thermophilus* and *Calescamantes* could be important consumers of the nitrite produced by *Nitrosocaldus*, where the combined activities of these organisms may result in complete denitrification to dinitrogen. The colocalization of the ACII complex gene with the nirS gene provides further evidence of the potential ability of *Calescamantes* to reduce oxidized nitrogen sources. As it is unlikely that the ACIII complex conserves as much energy as the bc complex, the eNOR gene likely conserves more energy in the reduction of nitric oxide to nitrous oxide, which could make up for this loss (57, 61).

Octopus Spring (66) and Gongxiaoshe (14) also harbored abundant populations of *Thermus* and *Nitrosocaldus*, making up 10 to 20% and 5 to 10% of the microbial community, respectively, in the two springs (13, 14) and indicating similar cometabolic roles in these communities. The oxygen tension in all studied springs was between 10 and 50 μM at or near the source. Although ammonia and nitrate were below detectable limits at Gongxiaoshe, nitrite was present (~26 μM). Ammonium and nitrate were present in Octopus Spring (2.3 and 6 μM, respectively) (13), although nitrite was not directly measured. Low ammonium and nitrate levels at Bison Pool (not detected and 0.4 μM, respectively) may contribute to the low abundance of *Calescamantes* in this spring.

The phylogenetic position of *Calescamantes* in the 16S rRNA and concatenated protein trees reaffirms its identity as a candidate phylum in the *Bacteria*, and it is most closely related to the *Aquificae* and *Thermodesulfobacteria* (67, 68). Based on the recent recommendation to allow the expansion of *Candidatus* status to nearly complete genomes conforming to genomic species delineations based on ANI values and other criteria (5), we propose to
expand the candidate taxonomy within the phylum *Calescamantes* (4) to include the lineage inhabiting Gongxiaoshe Spring. **Taxonomy.** We propose the taxonomic epithet “*Candidatus Calescimonas tengchongensis*.” The taxonomic description is “*Calescimonas*” (Cal.es.ci.mo’nas. L. v. calesco, to become warm, grow hot; N.L. n. monas a unit; N.L. n. Calescimonas; a rod-shaped bacterium from an extremely hot environment), “tengchongensis” ( teng.chong.en’sis. N.L. fem. adj. tengchongen- sis of or pertaining to Tengchong County, Yunnan province, China).

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