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<th>Citation</th>
<th>Te, Shu Harn, Boon Fei Tan, Janelle R. Thompson, and Karina Yew-Hoong Gin. “Draft Genome Sequences of Two Benthic Cyanobacteria, Oscillatoriales USR 001 and Nostoc sp. MBR 210, Isolated from Tropical Freshwater Lakes.” Genome Announcements 4, no. 5 (October 13, 2016): e01115–16.</th>
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</thead>
<tbody>
<tr>
<td>As Published</td>
<td><a href="http://dx.doi.org/10.1128/genomeA.01115-16">http://dx.doi.org/10.1128/genomeA.01115-16</a></td>
</tr>
<tr>
<td>Publisher</td>
<td>American Society for Microbiology</td>
</tr>
<tr>
<td>Version</td>
<td>Final published version</td>
</tr>
<tr>
<td>Accessed</td>
<td>Mon Nov 06 12:56:45 EST 2017</td>
</tr>
<tr>
<td>Citable Link</td>
<td><a href="http://hdl.handle.net/1721.1/108243">http://hdl.handle.net/1721.1/108243</a></td>
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Draft Genome Sequences of Two Benthic Cyanobacteria, Oscillatoriales USR 001 and Nostoc sp. MBR 210, Isolated from Tropical Freshwater Lakes

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Genomes of two filamentous benthic cyanobacteria were obtained from cocultures obtained from two freshwater lakes. The cultures were obtained by first growing cyanobacterial trichome on solid medium, followed by subculturing in freshwater media. Subsequent shotgun sequencing, de novo assembly, and genomic binning yielded almost complete genomes of Oscillatoriales USR 001 and Nostoc sp. MBR 210.

Benthic cyanobacteria inhabit the bottom of a diverse range of bodies of water, including lakes, wetlands, estuaries, and oceans, forming benthic mats in these environments (1). Under favorable conditions, they can proliferate rapidly and synthesize undesirable secondary metabolites, including toxins and odors (2). However, they are less studied compared to planktonic cyanobacteria even though they are able to cause similar ecological and water quality impacts on affected waters (3). Two filamentous benthic cyanobacteria isolated from tropical freshwater lakes in Singapore were identified as Oscillatoriales USR 001 and Nostoc sp. MBR 210, based on morphological traits (4). Here, we present additional genomic information about these isolates, which is important for functional annotation, pathways analysis, comparative genomics and for better understanding of their roles in bloom formation.

The two filamentous cyanobacteria were acquired through an agar culturing method (5). Briefly, lake sediment samples were streaked across agar plates enriched with McBride Listeria agar (MLA) medium (6). The agar plates were incubated (25°C, light intensity: 25 μmol/m²s) until green filaments appeared; then individual filaments were aseptically cut, transferred, and cultivated in sterile MLA media for 2 weeks. The genomic DNA extraction, Illumina HiSeq 2000 sequencing, and read quality controls were conducted following a method described previously (7). Subsequently, the two metagenomes were de novo assembled separately into scaffolds using CLC Genomics Workbench version 8. Contigs belonging to Cyanobacteria in each metagenome were separated from those of heterotrophic bacteria using MetaBAT (8), following which genome completeness and sequence contaminants were determined using CheckM (9), and the lack of a sequence contaminant was confirmed using a BLAST-based approach (10). The two genomes were annotated using RAST (11) and NCBI PGAP (http://www.ncbi.nlm.nih.gov/genome/annotation_prok).

Genomes of the two cyanobacteria have GC contents of 41% and completeness of 99%, assessed using checkM by comparing 579 to 583 reference marker genes in 79 to 82 lineage-specific reference genomes. The draft genome for Oscillatoriales USR 001 comprises 5.9 Mbp contained in 96 scaffolds, while Nostoc sp. MBR 210 has a genome size of 6.9 Mbp contained in 36 scaffolds. The 16S rRNA of Nostoc sp. MBR 210 (1,482 bp) is 98% identical to that of Nostoc piscinale CENA21 (CP012036.1), whereas the 16S rRNA of Oscillatoriales USR 001 (1,492 bp) is 98% identical to three members of the family Oscillatoriales: Kamptonema animale (EF654087.1), Phormidium animale CCAP (HF678514.1), and Oscillatoria lutea (KM019965.1). Further comparison between the genome of Oscillatoriales USR 001 with all reference genomes of the three genera currently available in the NCBI and JGI IMG databases (12) revealed a two-way average nucleotide identity of <90%, precluding the classification of USR 001 to the genus level. As members of these two organisms are known to be potential toxin (microcystin and anatoxin) and odor (geosmin and 2-methylisoborneol) producers, we used antiSMASH version 3.0.5 (13) and BLASTp to search for potential gene- or gene cluster-encoding secondary metabolites, to which no toxin and off-flavor-producing gene (as above) was identified in both genomes. Both genomes carry genes for photosynthesis and CO₂ and nitrogen fixation (e.g., carboxysome and nitrogenase, respectively) and a limited number of genes encoding sulfur utilization (e.g., betagalactosidase in USR 001; alpha-mannosidase for mannose utilization in MBR 210), suggesting their potential roles as photoheterotrophic nitrogen fixers.

Accession number(s). These whole-genome shotgun projects have been deposited in DDBJ/ENA/GenBank under the accession numbers MBRE00000000 (Oscillatoriales USR 001) and MBRD00000000 (Nostoc sp. MBR 210). The versions described in this paper are the first versions, MBRE01000000 and MBRD01000000.
ACKNOWLEDGMENTS
We thank Boo Chek Yin for providing cultures and the Public Utilities Board of Singapore (PUB) for their collaboration in this project. This research grant is supported by the Singapore National Research Foundation under its Environmental and Water Technologies Strategies Research Programme and administered by PUB. B.F.T. and J.R.T. were supported by the National Research Foundation of Singapore through the Singapore MIT Alliance for Research and Technology’s (SMART) Center for Environmental Sensing and Modeling (CENSAM) research program.

FUNDING INFORMATION
This work, including the efforts of Shu Harn Te and Karina Yew-Hoong Gin, was funded by National Research Foundation Singapore (NRF) (1102-IRIS-14-02). This work, including the efforts of Boon Fei Tan and Janelle R. Thompson, was funded by Singapore-MIT Alliance for Research and Technology Centre (SMART).

REFERENCES