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7-19-2018

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Aria Underriner Seacoast School of Technology

Tyler Silverwood Seacoast School of Technology

Carolyn Kelley Seacoast School of Technology

Kyle S. MacLea *University of New Hampshire, Manchester*, kyle.maclea@unh.edu

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Recommended Citation

Underriner A, Silverwood T, Kelley C, MacLea KS. 2018. Genome sequence of the halophilic bacterium Kangiella spongicola ATCC BAA-2076T. Microbiol Resour Announc 7:e00847-18. https://doi.org/10.1128/MRA.00847-18.

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Genome Sequence of the Halophilic Bacterium Kangiella spongicola ATCC BAA-2076^T

Aria Underriner, a Tyler Silverwood, a Carolyn Kelley, a to Kyle S. MacLeab, c, d

- ^aBiomedical Science and Technology Program, Seacoast School of Technology, Exeter, New Hampshire, USA
- ^bBiology Program, University of New Hampshire, Manchester, New Hampshire, USA
- ^cBiotechnology Program, University of New Hampshire, Manchester, New Hampshire, USA
- ^dDepartment of Life Sciences, University of New Hampshire, Manchester, New Hampshire, USA

ABSTRACT The Gram-negative genus Kangiella contains a number of halophilic species that display high levels of iso-branched fatty acids. Kangiella spongicola was isolated from a marine sponge, Chondrilla nucula, from the Florida Keys in the United States. A genome assembly of 2,825,399 bp with a 44.31% G+C content was generated from strain A79^T (=ATCC BAA-2076^T).

acteria within the family Alcanivoraceae and class Gammaproteobacteria include the genus Kangiella. Kangiella is notable for being a newly described genus of halophilic bacteria with an unusually high production of iso-branched fatty acids (1, 2). Kangiella spongicola ATCC BAA-2076^T is a member of this genus recently discovered from the marine sponge Chondrilla nucula, collected from the Florida Keys in the United States (2). This organism is a Gram-negative nonmotile rod-shaped bacterium that is able to survive in environments with up to 15% NaCl and a wide range of growth temperatures and pH values. The genome sequence generated for K. spongicola may be a useful data point in the understanding of sponge microbiota in marine environments, especially in combination with the five other members of the genus (1, 3, 4) that have been sequenced thus far (5, 6).

Kangiella spongicola ATCC BAA-2076^T was bought in lyophilized form from ATCC (Manassas, VA, USA). It was rehydrated in marine broth 2216 (BD, Franklin Lakes, NJ, USA) and incubated in a shaking incubator at 30°C for 24 h. Inoculated marine broth was spread on marine agar 2216 (BD), incubated at the same temperature, and, from the streak plate, a single colony was selected from which to grow a larger liquid culture for production of genomic DNA (gDNA) using a DNA minikit (Qiagen, Valencia, CA, USA). Pure gDNA was tagged with sequence adapters concurrent with fragmentation using the Nextera library prep kit (Illumina, San Diego, CA, USA). Tagged fragments were then sequenced on an Illumina HiSeq 2500 (Hubbard Center for Genome Studies, Durham, NH, USA) instrument. The generated 250-bp paired-end read sequences were bioinformatically trimmed before assembly using Trimmomatic (7). Paired and trimmed reads were assembled into a draft genome using the default settings of SPAdes version 3.11.1 (8). Small contigs and contaminants were removed, and the 140 remaining contigs were analyzed with the NCBI Prokaryotic Genome Annotation Pipeline (PGAP) process for gene prediction and annotation (9) and found to have a total sequence length of 2,825,399 bp, representing an average coverage of 326×. Its closest relatives, K. koreensis and K. aquimarina, are of similar size at 2.85 Mbp and 2.68 Mbp, respectively. Of the 140 contigs, the bulk of the sequence data were found in four contigs with lengths of 1,220,304 bp, 675,233 bp, 545,949 bp, and 83,637 bp. Of the remaining 136 contigs, all were less than 6,001 bp in length. The N_{50} value was 675,233 bp with an L_{50} value of 2, as determined by QUAST (10). The G+C content of 44.31% was consistent

Received 12 June 2018 Accepted 26 June 2018 Published 19 July 2018

Citation Underriner A, Silverwood T, Kelley C, MacLea KS. 2018. Genome sequence of the halophilic bacterium Kangiella spongicola ATCC BAA-2076^T. Microbiol Resour Announc 7:e00847-18. https://doi.org/10.1128/MRA

Editor Jason Stajich, University of California,

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Address correspondence to Kyle S. MacLea. Kyle.MacLea@UNH.edu.

A.U. and T.S. contributed equally to this work.

with the initial report of 44.9% (2). A total of 2,544 genes, 2,460 coding DNA sequences (CDS), 36 pseudogenes, 5 rRNAs, and 38 tRNAs were annotated using NCBI PGAP.

Data availability. This *Kangiella spongicola* ATCC BAA-2076^T whole-genome shot-gun sequence (WGS) project has been deposited in DDBJ/ENA/GenBank under accession number QICH00000000. The version described in this paper is the first version, QICH01000000.

ACKNOWLEDGMENTS

Sequencing and bioinformatics analysis were undertaken at the Hubbard Center for Genome Studies at UNH, supported by NH-INBRE, with the kind assistance of Kelley Thomas, Toni Westbrook, and Stephen Simpson. K.S.M. thanks lab associate Oliver Harmon for the inspiration to sequence a bacterium from a marine sponge. This work was a project of the Microbiology Education through the Genome Annotation-New Hampshire (MEGA-NH) program.

The Department of Life Sciences at UNH Manchester provided funds for purchasing the bacterial strain. Sequencing costs were supported by New Hampshire-INBRE through an Institutional Development Award (IDeA), P20GM103506, from the National Institute of General Medical Sciences of the NIH. The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

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