Draft Genome Sequence of *Salipaludibacillus keqinensis* ACCC 60430T, an Aerobic Halophilic Bacterium Isolated from a Salt Lake

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**ABSTRACT** The haloalkaliphilic bacterium *Salipaludibacillus keqinensis* ACCC 60430T, which grows optimally at 8.0% (wt/vol) Na+/H11001 and pH 9.0, was isolated from Keqin Lake in Qiqihaer, China. The draft genome includes 4,006 predicted genes and 3,784 coding sequences (CDSs). Genomic analysis showed that various genes may explain the mechanism of salt and alkali resistance.

*Salipaludibacillus keqinensis* ACCC 60430T is a strictly aerobic Gram-positive bacterium isolated using serial dilutions from Keqin Lake in Heilongjiang Province, China, and characterized and described as *Salipaludibacillus keqinensis* by Wang et al. (1). Growth is observed at 4 to 40°C (optimum, 25°C), at pH 8.0 to 10.0 (optimum, 9.0), and with up to 16.0% (wt/vol) NaCl (optimum, 8.0%) (1). To understand the resistance mechanism for survival under saline-alkaline conditions, genome sequencing of *S. keqinensis* ACCC 60430T was performed using an Illumina HiSeq 4000 sequencer.

Cells of *S. keqinensis* ACCC 60430T grown on LB medium at pH 9.0 and 25°C with 8% (wt/vol) NaCl for 48 h were collected (1), and genomic DNA was extracted using a microbial DNA isolation kit (Aidlab Biotech, Beijing, China) according to the manufacturer’s instructions. The genome sequencing library was constructed using the TruSeq DNA Nano library prep kit for Illumina. Sequencing was performed by the Illumina HiSeq 4000 sequencer with a paired-end read length of 2 × 150 bp at approximately 240× coverage. The reads were quality inspected with the default program parameters using FastQC v0.11.8 and Quake (2). Genomic sequence contigs were de novo assembled using the MicrobeTrakr Plus v0.9.1 (Zeta Biosciences, Shanghai, China). BWA-MEM v0.7.12 (3) was used post-assembly sequence correction according to the manufacturer’s instructions. Automatic annotations were performed using the NCBI Prokaryotic Genome Annotation Pipeline (PGAP) (4) (https://www.ncbi.nlm.nih.gov/genome/annotation_prok/), and checked by the GenBank curation team for the standard requirements. A total of 4,805,090 reads were assembled into 15 contigs, with a total length of 4,150,426 bp, G+C contents of 39.6%, and an N50 value of 1,053,560 bp. Among the 4,006 genes predicted, 3,874 were potential protein-coding genes (coding sequences [CDSs]). We also identified 92 RNAs, including 12 rRNAs (8 5S RNAs, 3 16S RNAs, and 1 23S RNA), 75 tRNAs, and 5 noncoding RNAs (ncRNAs).

Genome sequence analysis of *S. keqinensis* ACCC 60430T revealed its life characteristics of high salt and alkali resistance. The genome includes 1 gene cluster (*betL betP betS betT betT1 betT2*) responsible for glycine betaine synthesis from choline and 6 ectT genes responsible for ectoine biosynthesis from aspartate semialdehyde. Eight genes encode glycine betaine-binding protein OpuAC, and 9 genes encode glycine betaine/proline betaine-binding periplasmic protein. All of them are very effective osmotic pressure regulators (5, 6). Furthermore, the presences of 5 genes (3 TrkA type and 2 TrkH type) are responsible for K⁺ uptake systems, implying that *S. keqinensis* ACCC
60430T can use K+ as an osmolyte to obtain isosmotic cytoplasm by a salt-in strategy when coping with a rapid osmotic shock (7). Alkaliphilic S. keqinensis ACC 60430T has 6 genes encoding Na+/H+ antiporters (8), 1 asp23 gene encoding alkaline shock protein 23, and 2 genes encoding Ca2+/H+ antiporter (9), which may enable S. keqinensis ACC 60430T survival in alkaline environments (10). These predicted genes likely play a role in the salt and alkaline tolerance of S. keqinensis ACC 60430T.

Data availability. The draft genome assembly of S. keqinensis ACC 60430T has been deposited at DDBJ/ENA/GenBank under the accession number PDOD00000000. Raw sequencing reads have been submitted to the Sequence Read Archive (SRA accession number SRR8759055).

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