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### Draft Genome Sequences of Supercritical CO<sub>2</sub>-Tolerant Bacteria *Bacillus subterraneus* **MITOT1 and** *Bacillus cereus* **MIT0214**

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**We report draft genome sequences of** *Bacillus subterraneus* **MITOT1 and** *Bacillus cereus* **MIT0214 isolated through enrichment** of samples from geologic sequestration sites in pressurized bioreactors containing a supercritical (sc) CO<sub>2</sub> headspace. Their ge**nome sequences expand the phylogenetic range of sequenced bacilli and allow characterization of molecular mechanisms of scCO2 tolerance.**

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**D**uring geologic carbon sequestration (GCS), large quantities of CO<sub>2</sub> are captured, compressed to supercritical (sc) state, and injected underground. Whether microbial activities transform injected  $CO<sub>2</sub>$  is not well understood due to toxic effects of  $s<sub>c</sub>CO<sub>2</sub>$  [\(1](#page-1-0)[–](#page-2-0)[5\)](#page-2-1). Samples from GCS sites at Otway Basin, Australia and Frio-2, Texas, were used as inocula for serial enrichment cultures in bioreactors containing scCO<sub>2</sub>, yielding strains *Bacillus subterraneus* MITOT1 and *Bacillus cereus* MIT0214, respectively [\(6\)](#page-2-2). Tolerance of  $\sec O_2$  was confirmed by growth of spores in pure cultures and was time and inocula density dependent. To investigate mechanisms of growth under  $\mathrm{sCO}_2$ , genomic DNA was sequenced.

MITOT1 was sequenced on the Illumina HiSeq 2000 platform (Beijing Genomics Institute). MIT0214 was sequenced on the Illumina GAIIx platform (MIT Biomicrocenter). Paired-end 100 bp reads were quality trimmed (removing 10 starting and 20 trailing bases) and assembled *de novo* with CLC Genomic Workbench with automatic k-mer sizes of 23 and 21, yielding 185 and 238 contigs of  $>$ 500 bp, respectively. The draft genome of MITOT1 is 3.9 Mbp with  $42.1\%$  G+C content, while the MIT0214 draft genome is 5.6 Mbp with 34.9% G+C content. Annotation using the RAST server [\(7\)](#page-2-3) predicted 4,021 (with 1,235 hypothetical) and 5,640 (with 1,399 hypothetical) coding sequences in MITOT1 and MIT0214.

Phylogenetic analysis of the 16S rRNA gene placed MITOT1 within a clade of bacilli isolated from diverse environments including deep subsurface, soil, manufacturing effluent, and fermented seafood [\(8](#page-2-4)[–](#page-2-5)[12\)](#page-2-6), some of which are capable of anaerobic reduction of Fe(III),  $Mn(IV)$ , Se(VI), and As(V) [\(8,](#page-2-4) [10\)](#page-2-7). The closest relative by BLASTn of the 16S rRNA gene was *B. subterraneus* HWG-A11 (98.6% identity). The nearest genome sequenced strain (98.1% 16S rRNA identity) was *B. boroniphilus* DSM17376, isolated from boron-contaminated soil [\(13\)](#page-2-8) and sharing 83.3% average nucleotide identity (ANI) [\(14\)](#page-2-9) with 2,600 sequence homologs (>60% identity). RAST functional comparison of the MITOT1 and *B. boroniphilus* DSM17376 genomes with closely related bacilli (strain 1NLA3E, *B. infantis* NRRL B-14911,

*B. megaterium* DSM319, and *B. coagulans* 36D1) predicted multiple anaerobic respiratory reductases and terminal cytochrome C oxidases unique to MITOT1 and *B. boroniphilus*, pointing to diverse catabolic potential for this group [\(15,](#page-2-10) [16\)](#page-2-11).

Strain MIT0214 was most similar to *B. cereus* ATCC 14579 by BLASTn of 16S rRNA (99.8% identity), sharing 98.5% ANI and 4,858 sequence homologs (>60% identity). *B. cereus* strains have been isolated from diverse environments, including strain Q1 (92.5% ANI; 4,617 sequence homologs) from an oil reservoir [\(17\)](#page-2-12). Comparisons among genomes of MITOT1, MIT0214, and the closely related sequenced genomes did not reveal clear signatures associated with  $\sec 0<sub>2</sub>$  tolerance, which is unsurprising in light of recent observations that tolerance is widespread among bacilli [\(6\)](#page-2-2). Availability of draft genome sequences for *B. subterraneus* MITOT1 and *B. cereus* MIT0214 from two GCS sites will facilitate future work targeting gene/protein expression to advance mechanistic insights into  $\mathrm{scCO}_2$  tolerance.

**Nucleotide sequence accession numbers.**This whole-genome shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession numbers JXIQ00000000 and JXDH00000000. The versions described in this paper are the first versions.

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