Draft Genome Sequences from a Novel Clade of *Bacillus cereus Sensu Lato* Strains, Isolated from the International Space Station


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ABSTRACT

The draft genome sequences of six *Bacillus* strains, isolated from the International Space Station and belonging to the *Bacillus anthracis*-*B. cereus*-*B. thuringiensis* group, are presented here. These strains were isolated from the Japanese Experiment Module (one strain), U.S. Harmony Node 2 (three strains), and Russian Segment Zvezda Module (two strains).

Among the six *Bacillus cereus sensu lato* strains reported here, three U.S. Harmony Node 2 isolates (ISSFR-3F, ISSFR-9F, and ISSFT-23F) and one Japanese Experiment Module isolate (JEM-2) were sequenced and assembled with both Illumina MiSeq and PacBio RSII sequence data. The remaining assemblies, including two Russian isolates, were sequenced and assembled with only MiSeq data. The MiSeq runs yielded on average 24 to 54 million 300-bp reads (from 1,402× to 3,093× average coverage), while PacBio yielded 4,000 to 116,000 reads (from 7× to 202× average coverage) (Table 1). Due to the extremely high coverage (>1,000×), Illumina MiSeq reads were randomly down-sampled to 100× using an estimated genome size of 5.3 Mb, resulting in an average of 1.2 to 1.5 million paired-end reads per isolate. Next, the down-sampled reads were assembled using iMetAMOS (1) with IDBA_UD and SPAdes (2). IDBA_UD was selected as the best assembly for all six isolates. Low confidence bases within the selected IDBA_UD (3) assemblies were masked out by mapping all reads to the assembled contigs and detecting conflicting variants with FreeBayes (4). The PacBio reads were assembled following the methods described by Berlin et al. (5) with Celera Assembler version 8.3rc1 and polished with Quiver (6). A second round of polishing was performed post-Quiver using the available MiSeq data as input to Pilon (7).

The six International Space Station (ISS) isolates were aligned (NUCmer (8), Parsnp (9)) against members of the *B. cereus sensu lato* group genomes (10). The PacBio assemblies were used for all isolates with sufficient read coverage, and Illumina assemblies were used for the remaining isolates. Based on genome size estimates (5.2 to 5.3 Mb), NUCmer pairwise alignments (>99.9% average pairwise nucleotide identity) and maximum-likelihood phylogenetic placement, all six isolates were found to exhibit a very high degree of similarity.

Received 31 May 2017 Accepted 5 June 2017 Published 10 August 2017


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Finally, due to their high genomic similarity to the \textit{B. anthracis} type strain (\textit{H}1\textsubscript{11022}: 98\% average nucleotide identity), all six genomes were examined for evidence of pathogenicity. However, none of the commonly known \textit{B. anthracis} signature elements were identified. Specifically, all six ISS isolates (i) contain the \textit{plcR} (11) ancestral “C” allele, which has been used in large-scale phylogenetic analyses to distinguish \textit{B. anthracis} strains from the rest of the \textit{B. cereus} group; (ii) lack significant hits to \textit{pXO1} and \textit{pXO2} plasmids; and (iii) are phylogenetically placed outside of the \textit{B. anthracis} clade. Results were consistent with a comparative genomic analysis performed using the Lawrence Livermore National Laboratory Microbial Threat Characterization Pipeline. Altogether, the collective genomic evidence supports the conclusion that the six ISS isolates represent a novel \textit{Bacillus} sp. located within the \textit{B. cereus sensu lato} group.

\textbf{Accession number(s).} The complete genome sequences were deposited in NCBI under the accession numbers listed in Table 1 and can be accessed from the NASA GeneLab system (GLDS-64; https://genelab-data.ndc.nasa.gov/genelab/accession/GLDS-64).

\textbf{ACKNOWLEDGMENTS}

Part of the research described in this publication was carried out at the Jet Propulsion Laboratory (JPL), California Institute of Technology, under a contract with NASA. The contributions of S.R., R.K.P., T.E.B., S.K., A.P., J.K., T.J.T., M.J.R., and N.H.B. were funded under contract no. HSHQDC-07-C-00020 awarded by the Department of Homeland Security (DHS) Science and Technology Directorate (S&T) for the management and operation of the National Biodefense Analysis and Countermeasures Center (NBACC), a Federally Funded Research and Development Center. This research was also funded by 2012 Space Biology (NNH12ZTT001N) grant no. 19-12829-26 under Task Order NNN13D111T awarded to K.V., which also funded the postdoctoral fellowship for A.C.S., and by JPL subcontract 1506453 to G.E.F. The research carried out at Lawrence Livermore National Laboratory was funded by 2014 Space Biology (NNH14ZTT002N) grant no. NNX15AJ29G awarded to C.J., which also funded J.A. and C.T. The contribution of S.P.V.T. was supported by the European Space Agency (MAP Project no. AO-LS-99-MAP-LSS-018 “biofilms”) and SRON (MG-064/MG-068).

We thank Kazuyuki Tasaki, Director of the JEM Utilization Center, and Julie Robinson, Chief Scientist of the ISS (U.S.) for arranging Japanese Aerospace Exploration Agency (JAXA) approval to use information on strains collected during the routine operation and maintenance of the JEM-Kibo Module. We also thank Hermie J. M. Harmsen and Hendrik I. J. Roest for providing the Russian isolates and logistical support, as well as Jonathan Hnath, Kathy Fronda, Gregory Horn, and Henry Lupari from NBACC for technical assistance.

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