**Genome Sequence of the Edible Cyanobacterium *Arthrospira* sp. PCC 8005**

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Received 2 February 2010/Accepted 24 February 2010

We determined the genome sequence of *Arthrospira* sp. PCC 8005, a cyanobacterial strain of great interest to the European Space Agency for its nutritive value and oxygenic properties in the Micro-Ecological Life Support System Alternative (MELiSSA) biological life support system for long-term manned missions into space.

*Arthrospira* is a genus of nonheterocystous filamentous cyanobacteria that typically reside in alkaline lakes (9). Although they clearly form a separate taxonomic unit (4), various *Arthrospira* species were originally included in the genus *Spirulina*, and some are still being commercialized under that name. *Arthrospira* species have a high protein content, are rich in essential fatty acids, and produce a variety of minerals, vitamins, and nutritional pigments (3). Consequently, they have a long history of human consumption and are used worldwide as feedstock in agriculture and aquaculture (5). Besides their nutritional value, *Arthrospira* species have clinical relevance as an antioxidant (7) and are used in biohydrogen technology (1). The PCC 8005 strain was selected by the European Space Agency (ESA) as an oxygen producer and as a nutritional end product of the Micro-Ecological Life Support System Alternative (MELiSSA) life support system (6).

Whole-genome shotgun sequencing of strain PCC 8005 was performed using 454 pyrosequencing technology (amounting to 400,000 reads) and Sanger sequencing (up to 96,000 longer reads), leading to a final assembly of 119 contigs into 16 scaffolds representing 6,279,260 bases with an average G+C content of 44.7%. These scaffolds were processed by the MaGe annotation platform (8) and predicted 5,856 protein-coding sequences (CDSs) and 176 genes encoding RNA.

The genome of *Arthrospira* sp. PCC 8005 displays the highest overall synteny with the genomes of *Arthrospira maxima* CS-328, with 4,618 proteins as bidirectional best hits (BDBHs), and *Lyngbya* PCC8106, with 3,083 BDBHs. The MaGe annotation system assigned 63% of the CDSs to one or more functional COGs (clusters of orthologous groups) and reported 1,704 conserved hypothetical and 884 hypothetical proteins. Functional annotation by MaGe showed that many genes belonging to the same pathway are dispersed over the genome.

The PCC 8005 genome is highly repetitive in nature, with more than 300 kb present as tandem sequences, and contains four clustered, regularly interspaced short palindromic repeats (CRISPRs), which may provide a cellular defense against phages and plasmids (2). The genome also contains at least 140 complete insertion elements (ISs) belonging to various families and eight copies of a putative genomic island.

Several genes encoding desaturase were identified, including those required for the production of β-carotene and the two essential fatty acids linoleic acid (LA-18:2ω6Δ9,12) and γ-linoleic acid (GLA-18:3ω6,9,12). Genome data confirm the inability of PCC 8005 to fix nitrogen, as it lacks essential *nif* genes, and indicate that nitrogen metabolism in PCC 8005 follows classic routes utilizing nitrate, nitrite, urea, and ammonium, with *NtcA* as the global nitrogen regulator. An intact *nifH* operon encoding a nitrite hydratase and its regulator would allow strain PCC 8005 to utilize nitriles (R—C≡N) as the sole source of nitrogen. Hydrogen production in strain PCC 8005 is governed by a soluble bidirectional hydrogenase encoded and activated by intact *fox* and *hyp* loci. No *hyp* genes for a separate uptake hydrogenase were detected.

The PCC 8005 genome sequence and its curated annotation are important assets to better understand the physiology and metabolic potential of *Arthrospira* and will open up new opportunities in the functional genomics of this species.

**Nucleotide sequence accession numbers.** This whole-genome shotgun project was deposited at DDBJ/EMBL/GenBank under accession no. ADDH0000000. The version described here
is the first version, ADDH01000000. MaGe annotation data will become publicly available from the ArthroScope project site within 6 months of the GenBank submission date.

We thank C. Médigue and D. Vallenet of Génoscope (Centre National de Séquençage, Evry, France) for essential advice and training in genome annotation using the MaGe platform.

This work was supported by the European Space Agency ESA/ESTEC under MELGEN-2 contract no. 20397/06/NL/SFe in the frame of the MELiSSA project. A.W. is a research associate of the FRS-FNRS.

This Genome Announcement is dedicated to the lasting memory of our colleague Larissa Hendrickx (deceased 3 March 2008), who initiated the PCC 8005 genome project as part of MELiSSA.

REFERENCES