



Draft Genome Sequence of *Bacillus coagulans* GBI-30, 6086, a Widely Used Spore-Forming Probiotic Strain

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Bacillus coagulans GBI-30, 6086 is a safe strain, already available on the market, and characterized by certified beneficial effects. The draft genome sequence presented here constitutes the first pillar toward the identification of the molecular mechanisms responsible for its positive features and safety.

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Bacillus coagulans strain GBI-30, 6086 is a safe, spore-forming strain, as testified by the GRAS (Generally Recognized As Safe) status received in 2012 from the United States FDA; the strain is authorized for human consumption and already available in a wide selection of functional foods or as dietary supplement (1). This strain is characterized by certified beneficial effects in gastrointestinal disorders, such as irritable bowel syndrome, intestinal gas, and colitis (2, 3, 4, 5); in rheumatoid arthritis (6); and in common viral infections of the respiratory tract (7). As a spore-former, B. coagulans GBI-30, 6086 can be incorporated into foods, where it can survive the mild heat-treatments used for sterilization andwithstand the harsh conditions of the gastrointestinal tract, i.e., the low pH of the gastric barrier (8).

Here we report the draft genome sequence of *B. coagulans* GBI-30, 6086 in order to unveil the genetic basis of its safety and probiosis. To the best of our knowledge, this is the first published fully assembled genome of a commercial *B. coagulans* probiotic strain.

The whole-genome sequencing was performed using the Illumina GAIIx platform at CRA-Genomics Research Centre (Piacenza, Italy) with a paired-end library; the reads were *de novo* assembled using the CLC Genomic Workbench version 7.0. The genome sequence was annotated by the NCBI Prokaryotic Genomes Annotation Pipeline.

A total of 14,500,000 paired-end reads of 110-bp length on average (genome coverage of 840 \times) were assembled into 224 contigs (N_{50} length of 44,706 bp), with the largest assembled contig of 125,999-bp length. The draft genome consists of 3,458,655 bp with GC % content of 46.38.

A total of 3,373 genes were predicted, of which 3,197 are coding sequences (CDS), 18 are rRNAs, and 82 are tRNAs; 79 were identified to be pseudogenes, and 1 was identified as ncRNA. The genome also contains 3 CRISPR arrays, which could be involved as a defense mechanism toward foreign genetic elements (9).

The strain is predicted to encode for about 500 proteins in-

volved in central carbohydrate metabolism, including glycolysis, pentose phosphate, and xylose utilization pathways (9, 10). As expected, about 80 genes regarding dormancy and sporulation were annotated, in particular, spore DNA protection, sporulation mechanisms, spore core dehydration, and spore germination. Furthermore, the genome contains determinants involved in the adhesion (i.e., fibronectin- and mucus-binding proteins) and active metabolism in the host (as a biotin biosynthesis pathway).

The complete genome of *B. coagulans* GBI-30, 6086 obtained in the present study will contribute to a wider and deeper insight into the safety features of this strain, and the comparative genomic analysis with other *Bacillus* strains genomes (11, 12, 13) might shed new light on the molecular mechanisms at the basis of its probiotic and beneficial properties.

Nucleotide sequence accession numbers. This whole-genome shotgun project has been deposited in DDBJ/EMBL/GenBank under the accession number JPSK00000000. The version described in this paper is the first version, JPSK01000000.

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