

**Brevibacillus laterosporus spore coat-canoe shaped parasporal body complex protein A (cpbA)
gene, complete cds**

GenBank: KY124461.1

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LOCUS KY124461 780 bp DNA linear BCT 14-MAR-2017
DEFINITION Brevibacillus laterosporus spore coat-canoe shaped parasporal
body
complex protein A (cpbA) gene, complete cds.
ACCESSION KY124461
VERSION KY124461.1
KEYWORDS .
SOURCE Brevibacillus laterosporus
ORGANISM [Brevibacillus laterosporus](#)
Bacteria; Firmicutes; Bacilli; Bacillales; Paenibacillaceae;
Brevibacillus.
REFERENCE 1 (bases 1 to 780)
AUTHORS Marche,M.G., Mura,M.E., Falchi,G. and Ruiu,L.
TITLE Spore surface proteins of Brevibacillus laterosporus are involved
in insect pathogenesis
JOURNAL Sci Rep 7, 43805 (2017)
PUBMED [28256631](#)
REMARK Publication Status: Online-Only
REFERENCE 2 (bases 1 to 780)
AUTHORS Ruiu,L., Mura,M.E. and Marche,M.G.
TITLE Direct Submission
JOURNAL Submitted (11-NOV-2016) Dipartimento di Agraria, Universita degli
Studi di Sassari, Via E. de Nicola, Sassari 07100, Italy
COMMENT ##Assembly-Data-START##
Sequencing Technology :: Sanger dideoxy sequencing
##Assembly-Data-END##
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