

Pseudomonas fluorescens SBW25 genome assembly, chromosome: 1

GenBank: OV986001.1

[FASTA Graphics](#)

LOCUS OV986001 2418 bp DNA linear BCT 27-MAR-2022

DEFINITION Pseudomonas fluorescens SBW25 genome assembly, chromosome: 1.

ACCESSION [OV986001](#) REGION: 3777..6194

VERSION OV986001.1

DBLINK BioProject: [PRJEA31229](#)

BioSample: [SAMEA2272316](#)

KEYWORDS .

SOURCE Pseudomonas fluorescens SBW25

ORGANISM [Pseudomonas fluorescens SBW25](#)

Bacteria; Proteobacteria; Gammaproteobacteria; Pseudomonadales;
Pseudomonadaceae; Pseudomonas.

REFERENCE 1

AUTHORS Fortmann-Grote,C.

TITLE Direct Submission

JOURNAL Submitted (23-FEB-2022) Max Planck Institute for Evolutionary
Biology, Microbial Population Biology (Rainey Lab),
August-Thienemann-Strasse 2, 24306 Ploen, Germany

COMMENT On Mar 27, 2022 this sequence version replaced [AM181176.4](#).
De-novo PacBio assembly generated at MPI Evolutionary Biology -
Microbial Population Biology.

FEATURES Location/Qualifiers

source 1..2418
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/db_xref="taxon:[216595](#)"
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mRNA 1..2418
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    similarity:fasta; with=UniProt:GYRB_ECOLI
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ORIGIN

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