Species Pluralincolimonas frigidipaludosa^{Ts}

Etymology

[fri.gi.di.pa.lu.do'sa] L. masc. adj. frigidus, cold; L. fem. adj. paludosa, boggy or marshy; N.L. fem. adj. frigidipaludosa, of a cold marsh or bog

Nomenclatural type

NCBI Assembly: GCA_003141835.1 Ts

Description

This species is the type for the genus *Pluralincolimonas*. Estimate genome size for the species is 1.6 Mb, with a %GC of 51.75 %. This species encode an ADP/ATP translocase, and respiratory complexes I and V. All components investigated for the production of a Type-4a pilus is encoded by genomes of this species, as well as a cytoplasmic Group A3 [FeFe] hydrogenase. Coupled with the Group A3 [FeFe] hydrogenase, a reversible acetyl-CoA synthetase may indicate acetogenesis in this species, through the production of acetate from acetyl-CoA by the fermentation of glucose and nucleotides. Through the use of respiratory complex I, oxidized NAD+ is provided and a proton gradient across the membrane is maintained, while the respiratory F-type ATPase (complex V) produces ATP. Genome data for this species originate from a permafrost sample from Stordalen Mire, Sweden, where a pH of 4.8 was reported. The nomenclatural type for the species is the genome GCA 003141835.1.

Classification

Bacteria » Omnitrophota » "Gorgyraia" » Pluralincolimonadales » Pluralincolimonadaceae » Pluralincolimonas » Pluralincolimonas frigidipaludosa^{Ts}

References

Effective publication: Seymour et al., 2023 [1]

Registry URL

https://seqco.de/i:23728

References

 Seymour et al. (2023). Hyperactive nanobacteria with host-dependent traits pervade Omnitrophota. Nature Microbiology. <u>DOI:10.1038/s41564-022-01319-1</u>