Atabeyarchaeia classis nov.

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Abstract

Atabeyarchaeia and Freyarchaeia represent two distinct lineages within the Asgard superphylum, expanding our understanding of archaeal diversity and metabolic capabilities. Genomic analyses reveal that Atabeyarchaeia are likely non-methanogenic acetogenic chemoheterotrophs with potential for lithoautotrophic growth using H2 as an electron donor. They possess pathways for degrading amino acids and other carbon compounds, including the ability to metabolize C1, C3, and C5 compounds, which is uncommon among Asgard archaea. Freyarchaeia, originally described as a separate lineage, shares some metabolic features with Atabeyarchaeia but exhibits distinct characteristics. Both groups are positioned within the Asgard clade, with phylogenetic analyses using multiple marker gene sets (including 47 archaeal clusters of orthologous genes, 15 ribosomal protein genes, and the 16S rRNA gene) consistently placing Atabeyarchaeia as a sister group to Freyarchaeia. Despite sharing only 40-45% average amino acid identity with other Asgard genomes, both lineages are classified within the Asgardarchaeota phylum, adhering to the Genome Taxonomy Database (GTDB) standards. The proposed classification establishes Atabeyarchaeum as the type genus for the family Atabeyarchaeaceae, order Atabeyarchaeales, and class Atabeyarchaeia, while maintaining Freyarchaeia as a distinct lineage within the same phylum. This research highlights the metabolic diversity within the Asgard superphylum and provides insights into the potential roles of these archaea in carbon cycling and their interactions with other microbial communities in various ecosystems.

Class Freyarchaeia

Etymology

[Fre.y.ar.chae'i.a] **N.L. neut. n.** *Freyarchaeum*, referring to the type genus Freyarchaeum; *-ia*, ending to denote a class; **N.L. neut. pl. n.** *Freyarchaeia*, the Freyarchaeum class

Nomenclatural type

Genus Freyarchaeum

Description

The type species for this taxonomic group were initially reconstructed by manual curation of Illumina short-read assemblies and then validated using both Nanopore and PacBio long reads. Phylogenetic analyses of this fully curated genome using several sets of marker genes placed the complete genomes in a monophyletic group within the Asgard clade as a sister group to Atabeyarchaeia. We performed phylogenetic analyses using concatenated marker sets of 47 archaeal clusters of orthologous genes (arCOGs) and 15 ribosomal protein (RP15) gene cluster, as well as the 16S rRNA gene. The new genomes share only <45% AAI when compared to other Asgard genomes, consistent with their assignment to a new phylum. Although our analyses provide evidence for distinction at the phylum level, we chose to adhere to the Genome Taxonomy Database (GTDB) for standardized microbial genome nomenclature. We establish a robust classification and propose Freyarchaeum as the type genus for the family Freyarchaeaceae, the order Freyarchaeales, the class Freyarchaeia with the Asgardarchaeota phylum. Based on protein content and compositional features, we infer that Freyarchaeia is a non-methanogenic acetogenic chemoheterotroph, with potential to grow lithoautotrophically by using H2 as an electron donor. This lineage was originally reconstructed and named from Guaymas Basin, located in the Gulf of California, México, and from Jinze Hot Spring (Yunnan, China). Subsequently, another group used the original data to recover similar genomes and referred to them as Jordarchaeia. Here, we retain the original nomenclature.

Classification

Archaea » Asgardarchaeota » Freyarchaeia

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48235

Class Atabeyarchaeia

Etymology

[A.ta.be.y.ar.chae'i.a] **N.L. neut. n.** *Atabeyarchaeum*, referring to the type genus Atabeyarchaeum; *-ia*, ending to denote a class; **N.L. neut. pl. n.** *Atabeyarchaeia*, the Atabeyarchaeum class

Nomenclatural type

Genus Atabeyarchaeum

Description

Based on protein content and compositional features, we infer that Atabeyarchaeia is a non-methanogenic acetogenic chemoheterotroph, with potential to grow lithoautotrophically by using H2 as an electron donor. Phylogenetic analyses using several sets of marker genes placed two complete genomes in a monophyletic group within the Asgard clade as a sister group to Freyarchaeia. We performed phylogenetic analyses using concatenated marker sets of 47 archaeal clusters of orthologous genes (arCOGs) and 15 ribosomal protein (RP15) gene cluster, as well as the 16S rRNA gene. The new genomes share only 40–45% AAI when compared to other Asgard genomes, consistent with their assignment to a new phylum. Although our analyses provide evidence for distinction at the phylum level, we chose to adhere to the Genome Taxonomy Database (GTDB) for standardized microbial genome nomenclature. We establish a robust classification and propose Atabeyarchaeum as the type genus for the family Atabeyarchaeaceae, the order Atabeyarchaeales, the class Atabeyarchaeia with the Asgardarchaeota phylum.

Classification

Archaea » Asgardarchaeota » Atabeyarchaeia

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:33330

Order Freyarchaeales

Etymology

[Fre.y.ar.chae.a'les] **N.L. neut. n.** *Freyarchaeum*, referring to the type genus Freyarchaeum; *-ales*, ending to denote an order; **N.L. fem. pl. n.** *Freyarchaeales*, the Freyarchaeum order

Nomenclatural type

Genus Freyarchaeum

Description

The type species for this taxonomic group were initially reconstructed by manual curation of Illumina short-read assemblies and then validated using both Nanopore and PacBio long reads. Phylogenetic analyses of this fully curated genome using several sets of marker genes placed the complete genomes in a monophyletic group within the Asgard clade as a sister group to Atabeyarchaeia. We performed phylogenetic analyses using concatenated marker sets of 47 archaeal clusters of orthologous genes (arCOGs) and 15 ribosomal protein (RP15) gene cluster, as well as the 16S rRNA gene. The new genomes share only <45% AAI when compared to other Asgard genomes, consistent with their assignment to a new phylum. Although our analyses provide evidence for distinction at the phylum level, we chose to adhere to the Genome Taxonomy Database (GTDB) for standardized microbial genome nomenclature. We establish a robust classification and propose Freyarchaeum as the type genus for the family Freyarchaeaceae, the order Freyarchaeales, the class Freyarchaeia with the Asgardarchaeota phylum. Based on protein content and compositional features, we infer that Freyarchaeia is a non-methanogenic acetogenic chemoheterotroph, with potential to grow lithoautotrophically by using H2 as an electron donor. This lineage was originally reconstructed and named from Guaymas Basin, located in the Gulf of California, México, and from Jinze Hot Spring (Yunnan, China). Subsequently, another group used the original data to recover similar genomes and referred to them as Jordarchaeia. Here, we retain the original nomenclature.

Classification

Archaea » Asgardarchaeota » Freyarchaeia » Freyarchaeales

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48236

Order Atabeyarchaeales

Etymology

[A.ta.be.y.ar.chae.a'les] **N.L. neut. n.** *Atabeyarchaeum*, referring to the type genus Atabeyarchaeum; *-ales*, ending to denote an order; **N.L. fem. pl. n.** *Atabeyarchaeales*, the Atabeyarchaeum order

Nomenclatural type

Genus Atabeyarchaeum

Description

Based on protein content and compositional features, we infer that Atabeyarchaeia is a non-methanogenic acetogenic chemoheterotroph, with potential to grow lithoautotrophically by using H2 as an electron donor. Phylogenetic analyses using several sets of marker genes placed two complete genomes in a monophyletic group within the Asgard clade as a sister group to Freyarchaeia. We performed phylogenetic analyses using concatenated marker sets of 47 archaeal clusters of orthologous genes (arCOGs) and 15 ribosomal protein (RP15) gene cluster, as well as the 16S rRNA gene. The new genomes share only 40–45% AAI when compared to other Asgard genomes, consistent with their assignment to a new phylum. Although our analyses provide evidence for distinction at the phylum level, we chose to adhere to the Genome Taxonomy Database (GTDB) for standardized microbial genome nomenclature. We establish a robust classification and propose Atabeyarchaeum as the type genus for the family Atabeyarchaeaceae, the order Atabeyarchaeales, the class Atabeyarchaeia with the Asgardarchaeota phylum.

Classification

Archaea » Asgardarchaeota » Atabeyarchaeia » Atabeyarchaeales

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48234

Family Freyarchaeaceae

Etymology

[Fre.y.ar.chae.a'ce.ae] **N.L. neut. n.** *Freyarchaeum*, referring to the type genus Freyarchaeum; *-aceae*, ending to denote a family; **N.L. fem. pl. n.** *Freyarchaeaceae*, the Freyarchaeum family

Nomenclatural type

Genus Freyarchaeum

Description

The type species for this taxonomic group were initially reconstructed by manual curation of Illumina short-read assemblies and then validated using both Nanopore and PacBio long reads. Phylogenetic analyses of this fully curated genome using several sets of marker genes placed the complete genomes in a monophyletic group within the Asgard clade as a sister group to Atabeyarchaeia. We performed phylogenetic analyses using concatenated marker sets of 47 archaeal clusters of orthologous genes (arCOGs) and 15 ribosomal protein (RP15) gene cluster, as well as the 16S rRNA gene. The new genomes share only <45% AAI when compared to other Asgard genomes, consistent with their assignment to a new phylum. Although our analyses provide evidence for distinction at the phylum level, we chose to adhere to the Genome Taxonomy Database (GTDB) for standardized microbial genome nomenclature. We establish a robust classification and propose Freyarchaeum as the type genus for the family Freyarchaeaceae, the order Freyarchaeales, the class Freyarchaeia with the Asgardarchaeota phylum. Based on protein content and compositional features, we infer that Freyarchaeia is a non-methanogenic acetogenic chemoheterotroph, with potential to grow lithoautotrophically by using H2 as an electron donor. This lineage was originally reconstructed and named from Guaymas Basin, located in the Gulf of California, México, and from Jinze Hot Spring (Yunnan, China). Subsequently, another group used the original data to recover similar genomes and referred to them as Jordarchaeia. Here, we retain the original nomenclature.

Classification

Archaea » Asgardarchaeota » Freyarchaeia » Freyarchaeales » Freyarchaeaceae

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48237

Family Atabeyarchaeaceae

Etymology

[A.ta.be.y.ar.chae.a'ce.ae] **N.L. neut. n.** *Atabeyarchaeum*, referring to the type genus Atabeyarchaeum; *-aceae*, ending to denote a family; **N.L. fem. pl. n.** *Atabeyarchaeaceae*, the Atabeyarchaeum family

Nomenclatural type

Genus Atabeyarchaeum

Description

Based on protein content and compositional features, we infer that Atabeyarchaeia is a non-methanogenic acetogenic chemoheterotroph, with potential to grow lithoautotrophically by using H2 as an electron donor. Phylogenetic analyses using several sets of marker genes placed two complete genomes in a monophyletic group within the Asgard clade as a sister group to Freyarchaeia. We performed phylogenetic analyses using concatenated marker sets of 47 archaeal clusters of orthologous genes (arCOGs) and 15 ribosomal protein (RP15) gene cluster, as well as the 16S rRNA gene. The new genomes share only 40–45% AAI when compared to other Asgard genomes, consistent with their assignment to a new phylum. Although our analyses provide evidence for distinction at the phylum level, we chose to adhere to the Genome Taxonomy Database (GTDB) for standardized microbial genome nomenclature. We establish a robust classification and propose Atabeyarchaeum as the type genus for the family Atabeyarchaeaceae, the order Atabeyarchaeales, the class Atabeyarchaeia with the Asgardarchaeota phylum.

Classification

Archaea » Asgardarchaeota » Atabeyarchaeia » Atabeyarchaeales » Atabeyarchaeaceae

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48233

Genus Freyarchaeum

Etymology

[Fre.y.ar.chae'um] N.L. fem. n. *Freya*, the Norse Goddess most commonly associated with love and fertility; N.L. neut. n. *archaeum*, ancient, archaeon; N.L. neut. n. *Freyarchaeum*, an archaeal genus named for Freya, the Norse Goddess most commonly associated with love and fertility

Nomenclatural type

Species Freyarchaeum deiterrae^{Ts}

Description

The type species for this taxonomic group were initially reconstructed by manual curation of Illumina short-read assemblies and then validated using both Nanopore and PacBio long reads. Phylogenetic analyses of this fully curated genome using several sets of marker genes placed the complete genomes in a monophyletic group within the Asgard clade as a sister group to Atabeyarchaeia. We performed phylogenetic analyses using concatenated marker sets of 47 archaeal clusters of orthologous genes (arCOGs) and 15 ribosomal protein (RP15) gene cluster, as well as the 16S rRNA gene. The new genomes share only <45% AAI when compared to other Asgard genomes, consistent with their assignment to a new phylum. Although our analyses provide evidence for distinction at the phylum level, we chose to adhere to the Genome Taxonomy Database (GTDB) for standardized microbial genome nomenclature. We establish a robust classification and propose Freyarchaeum as the type genus for the family Freyarchaeaceae, the order Freyarchaeales, the class Freyarchaeia with the Asgardarchaeota phylum. Based on protein content and compositional features, we infer that Freyarchaeia is a non-methanogenic acetogenic chemoheterotroph, with potential to grow lithoautotrophically by using H2 as an electron donor. This lineage was originally reconstructed and named from Guaymas Basin, located in the Gulf of California, México, and from Jinze Hot Spring (Yunnan, China). Subsequently, another group used the original data to recover similar genomes and referred to them as Jordarchaeia. Here, we retain the original nomenclature.

Classification

Archaea » Asgardarchaeota » Freyarchaeia » Freyarchaeales » Freyarchaeacea » Freyarchaeum

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48239

Genus Atabeyarchaeum

Etymology

[A.ta.be.y.ar.chae'um] **N.L. fem. n.** *Atabey*, the Mother Goddess in Taíno mythology; **N.L. neut. n.** *archaeum*, ancient, an archaeon from Gr. adj. archaeos; **N.L. neut. n.** *Atabeyarchaeum*, named for Atabey, the Mother Goddess in Taîno Puerto Rican mythology

Nomenclatural type

Species Atabeyarchaeum deiterrae^{Ts}

Description

ased on protein content and compositional features, we infer that Atabeyarchaeia is a non-methanogenic acetogenic chemoheterotroph, with potential to grow lithoautotrophically by using H2 as an electron donor. Phylogenetic analyses using several sets of marker genes placed two complete genomes in a monophyletic group within the Asgard clade as a sister group to Freyarchaeia. We performed phylogenetic analyses using concatenated marker sets of 47 archaeal clusters of orthologous genes (arCOGs) and 15 ribosomal protein (RP15) gene cluster, as well as the 16S rRNA gene. The new genomes share only 40–45% AAI when compared to other Asgard genomes, consistent with their assignment to a new phylum. Although our analyses provide evidence for distinction at the phylum level, we chose to adhere to the Genome Taxonomy Database (GTDB) for standardized microbial genome nomenclature. We establish a robust classification and propose Atabeyarchaeum as the type genus for the family Atabeyarchaeaceae, the order Atabeyarchaeales, the class Atabeyarchaeia with the Asgardarchaeota phylum.

Classification

Archaea » Asgardarchaeota » Atabeyarchaeia » Atabeyarchaeales » Atabeyarchaeacea » Atabeyarchaeum

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48232

Species Atabeyarchaeum deiterrae^{Ts}

Etymology

[de.i.ter'rae] L. fem. n. dea, deity; L. fem. n. terra, of the earth; N.L. gen. n. deiterrae, of an Earth diety

Nomenclatural type

NCBI Assembly: GCA 037310415.1 Ts

Description

Atabeyarchaeum deiteterre has a genome size of about 2.81 Mbp with an average GC% of 49.6%, recovered from California soil. Atabeyarchaeum 16S shares less than 75% similarity with previously described Asgardarchaeota classes. A. deiteterre is an acetogenic chemoheterotroph like other Asgardarchaeota, including Freyarchaeia. Estimated completeness at 97.2%, contamination at 1.87%, and the presence of 16 S rRNA genes (1529 bp) and 40 tRNAs.

Classification

Archaea » Asgardarchaeota » Atabeyarchaeia » Atabeyarchaeales » Atabeyarchaeaceae » Atabeyarchaeum » Atabeyarchaeum deiterrae^{Ts}

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48231

Species Freyarchaeum deiterrae^{Ts}

Etymology

[de.i.ter'rae] L. fem. n. dea, deity; L. fem. n. terra, of the earth; N.L. gen. n. deiterrae, of an Earth diety

Nomenclatural type

NCBI Assembly: GCA 037305845.1 Ts

Description

Freyarchaeum deiteterre has a genome size of about 3.58 Mbp with an average GC% of 40.7%, recovered from California soil. Freyarchaeum 16S shares less than 45% similarity with previously described Asgardarchaeota classes. F. deiteterre is an acetogenic chemoheterotroph like other Asgardarchaeota, including other soil-associated Atabeyarchaeia. Estimated completeness at 98.13%, contamination at 4.21%, and the presence of 16 S rRNA genes (1264 bp) and 38 tRNAs.

Classification

Archaea » Asgardarchaeota » Freyarchaeia » Freyarchaeales » Freyarchaeaceae » Freyarchaeum » Freyarchaeum deiterrae^{Ts}

References

Effective publication: Valentin-Alvarado et al., 2024 [1]

Registry URL

https://seqco.de/i:48238

References

1. Valentin-Alvarado et al. (2024). Asgard archaea modulate potential methanogenesis substrates in wetland soil. *Nature Communications*. DOI:10.1038/s41467-024-49872-z

Register List Certificate of Validation

On behalf of the *Committee on the Systematics of Prokaryotes Described from Sequence Data* (SeqCode Committee), we hereby certify that the Register List **seqco.de/r:igzul8qd** submitted by **Valentin-Alvarado, Luis** and including 10 new names has been successfully validated.

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