Register list for 16 new names including Andeanibacterium gen. nov.

Submitted by Díaz-García, Laura

Genus Andeanibacterium

Etymology

[An.de.a.ni.bac.te'ri.um] **N.L. masc. adj.** *andeanus*, pertaining to the Andes; **N.L. neut. n.** *bacterium*, small rod; **N.L. neut. n.** *Andeanibacterium*, Andean rod

Nomenclatural type

Species Andeanibacterium colombiense^{Ts}

Description

A genus established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction, and phylogenomic analyses and is classified as a member of the *Sphingomonadaceae* family. Genomic metrics include ANI (79.88), AAI (61.59) and dDDH d4 (20.1). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 54 CAZymes, 15 of them involved in lignocellulose degradation. In addition, putative genes for PET hydrolisis were found within the MAG.

Classification

Bacteria » Pseudomonadota » Alphaproteobacteria » Sphingomonadales » Sphingomonadaceae » Andeanibacterium

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24097

Species Andeanibacterium colombiense^{Ts}

Etymology

[co.lom.bi.en'se.] N.L. neut. adj. colombiense, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119316 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction, and phylogenomic analyses and is classified as a member of the *Sphingomonadaceae* family. The type species of the genus is *Andeanibacterium colombiense*. Genomic metrics include ANI (79.88), AAI (61.59) and dDDH d4 (20.1). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 54 CAZymes, 15 of them involved in lignocellulose degradation. In addition, putative genes for PET hydrolisis were found within the MAG.

Classification

Bacteria » Pseudomonadota » Alphaproteobacteria » Sphingomonadales » Sphingomonadaceae » Andeanibacterium » Andeanibacterium colombiense^{Ts}

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:31306

Species *Pedobacter colombiensis*

Etymology

[co.lom.bi.en'sis] N.L. masc. adj. colombiensis, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119313 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_3858. Genomic metrics include ANI (81.7), AAI (81.02) and dDDH d4 (29.6). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 254 CAZymes, 63 of them involved in lignocellulose degradation. In addition, putative genes for PET and PUR hydrolisis were found within the MAG.

Classification

Bacteria » Bacteroidota » Sphingobacteriia » Sphingobacteriales » Sphingobacteriaceae » Pedobacter » Pedobacter colombiensis

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24104

Species Brevundimonas colombiensis

Etymology

[co.lom.bi.en'sis] N.L. fem. adj. colombiensis, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119326 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_833. Genomic metrics include ANI (87.3), AAI (82.62) and dDDH d4 (29.7). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 73 CAZymes, 13 of them involved in lignocellulose degradation. In addition, putative genes for protocatechuate and phtalate catabolism were found within the MAG.

Classification

Bacteria » Pseudomonadota » Alphaproteobacteria » Caulobacterales » Caulobacteraceae » Brevundimonas » Brevundimonas colombiensis

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24108

Species Pseudomonas colombiensis

Etymology

[co.lom.bi.en'sis] N.L. fem. adj. colombiensis, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119322 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_425. Genomic metrics include ANI (86.68), AAI (86.03) and dDDH d4 (29.3). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 103 CAZymes, 19 of them involved in lignocellulose degradation. In addition, putative genes for PUR hydrolisis were found within the MAG.

Classification

Bacteria » Pseudomonadota » Gammaproteobacteria » Moraxellales » Pseudomonadaceae » Pseudomonas » Pseudomonas colombiensis

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24106

Species Sphingomonas colombiensis

Etymology

[co.lom.bi.en'sis] N.L. fem. adj. colombiensis, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119315 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_27. Genomic metrics include ANI (85.46), AAI (66.97) and dDDH d4 (20.8). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 59 CAZymes, 10 of them involved in lignocellulose degradation. In addition, putative genes for PET hydrolisis were found within the MAG.

Classification

Bacteria » Pseudomonadota » Alphaproteobacteria » Sphingomonadales » Sphingomonadaceae » Sphingomonas » Sphingomonas colombiensis

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24102

Species Kaistia colombiensis

Etymology

[co.lom.bi.en'sis] N.L. fem. adj. colombiensis, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119318 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_2176. Genomic metrics include ANI (86.41), AAI (77.29) and dDDH d4 (28.1). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 95 CAZymes, 18 of them involved in lignocellulose degradation. In addition, putative genes for PET and PUR hydrolisis were found within the MAG.

Classification

Bacteria » Pseudomonadota » Alphaproteobacteria » Hyphomicrobiales » Kaistiaceae » Kaistia » Kaistia colombiensis

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24101

Species Microbacterium colombiense

Etymology

[co.lom.bi.en'se.] N.L. neut. adj. colombiense, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119308 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_1911. Genomic metrics include ANI (85.9), AAI (81.1) and dDDH d4 (28.3). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 82 CAZymes, 30 of them involved in lignocellulose degradation. In addition, putative genes for PET and PUR hydrolisis were found within the MAG.

Classification

Bacteria » Actinomycetota » Actinobacteria » Micrococcales » Microbacteriaceae » Microbacterium » Microbacterium colombiense

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24100

Species Chryseobacterium colombiense

Etymology

[co.lom.bi.en'se.] N.L. neut. adj. colombiense, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119310 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_1654. Genomic metrics include ANI (88.96), AAI (95) and dDDH d4 (36.2). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 139 CAZymes, 41 of them involved in lignocellulose degradation. In addition, putative genes for PUR hydrolisis were found within the MAG.

Classification

Bacteria » Bacteroidota » Flavobacteriia » Flavobacteriales » Weeksellaceae » Chryseobacterium » Chryseobacterium colombiense

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24098

Species Cohnella colombiensis

Etymology

[co.lom.bi.en'sis] N.L. fem. adj. colombiensis, belonging to Colombia

Nomenclatural type

INSDC Nucleotide: CP119317 Ts

Description

The species is established on the basis of MiGA taxonomic novelty analysis, the taxonmic placement using maximum likelihood trees with 120 bacterial marker genes (bac120) and and the type material is the genome MAG_2441. Genomic metrics include ANI (67.06), AAI (67.21) and dDDH d4 (30.4). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 149 CAZymes, 44 of them involved in lignocellulose degradation. In addition, putative genes for PET and PUR hydrolisis were found within the MAG.

Classification

Bacteria » Bacillota » Bacilli » Caryophanales » Paenibacillaceae » Cohnella » Cohnella colombiensis

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24096

Species Pseudobacter hemicellulosilyticus

Etymology

[he.mi.cel.lu.lo.si.ly'ti.cus] **N.L. neut. n.** hemicellulosum, hemicellulose; **N.L. masc. adj.** lyticus, able to dissolve; **N.L. masc. adj.** hemicellulosilyticus, hemicellulose dissolving

Nomenclatural type

INSDC Nucleotide: CP119311 Ts

Description

The species is established on the basis of MiGA taxonomic novelty analysis, the taxonomic placement using maximum likelihood trees with 120 bacterial marker genes (bac120) and and the type material is the genome MAG_7. Genomic metrics include ANI (69.21), AAI (65.96) and dDDH d4 (22.1). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 237 CAZymes, 99 of them involved in lignocellulose degradation. In addition, putative genes for PET and PUR hydrolisis were found within the MAG.

Classification

Bacteria » Bacteroidota » Chitinophagia » Chitinophagales » Chitinophagaceae » Pseudobacter » Pseudobacter hemicellulosilyticus

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24095

Species Sphingomonas phytovorans

Etymology

[phy.to.vo'rans.] **Gr. neut. n.** *phyton*, plant; **L. pres. part.** *vorans*, devouring, destroying; **N.L. fem. part. adj.** *phytovorans*, plant-devouring

Nomenclatural type

INSDC Nucleotide: CP119314 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_3606. Genomic metrics include ANI (80.99), AAI (80.97) and dDDH d4 (22.2). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 95 CAZymes, 29 of them involved in lignocellulose degradation. In addition, putative genes for PET and PUR hydrolisis were found within the MAG.

Classification

Bacteria » Pseudomonadota » Alphaproteobacteria » Sphingomonadales » Sphingomonadaceae » Sphingomonas » Sphingomonas phytovorans

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24103

Species Brevundimonas phytovorans

Etymology

[phy.to.vo'rans.] **Gr. neut. n.** *phyton*, plant; **L. pres. part.** *vorans*, devouring, destroying; **N.L. fem. part. adj.** *phytovorans*, plant-devouring

Nomenclatural type

INSDC Nucleotide: CP119309 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_1865. Genomic metrics include ANI (91.28), AAI (87.92) and dDDH d4 (40.3). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 39 CAZymes, 6 of them involved in lignocellulose degradation. In addition, putative genes for protocatechuate and phtalate catabolism were found within the MAG.

Classification

Bacteria » Pseudomonadota » Alphaproteobacteria » Caulobacterales » Caulobacteraceae » Brevundimonas » Brevundimonas phytovorans

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24099

Species Pseudomonas phytovorans

Etymology

[phy.to.vo'rans.] **Gr. neut. n.** *phyton*, plant; **L. pres. part.** *vorans*, devouring, destroying; **N.L. fem. part. adj.** *phytovorans*, plant-devouring

Nomenclatural type

INSDC Nucleotide: CP119325 Ts

Description

A genus established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_876. Genomic metrics include ANI (92.18), AAI (95) and dDDH d4 (47). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 87 CAZymes, 16 of them involved in lignocellulose degradation. In addition, putative genes for PUR hydrolysis were found within the MAG.

Classification

Bacteria » Pseudomonadota » Gammaproteobacteria » Moraxellales » Pseudomonadaceae » Pseudomonas » Pseudomonas phytovorans

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24109

Species Devosia phytovorans

Etymology

[phy.to.vo'rans.] **Gr. neut. n.** *phyton*, plant; **L. pres. part.** *vorans*, devouring, destroying; **N.L. fem. part. adj.** *phytovorans*, plant-devouring

Nomenclatural type

INSDC Nucleotide: CP119312 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_4196. Genomic metrics include ANI (84.84), AAI (74.37) and dDDH d4 (23.5). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 90 CAZymes, 17 of them involved in lignocellulose degradation. In addition, putative genes for PET and PUR hydrolisis were found within the MAG.

Classification

Bacteria » Pseudomonadota » Alphaproteobacteria » Hyphomicrobiales » Devosiaceae » Devosia » Devosia phytovorans

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24105

Species Microbacterium phytovorans

Etymology

[phy.to.vo'rans.] **Gr. neut. n.** *phyton*, plant; **L. pres. part.** *vorans*, devouring, destroying; **N.L. neut. part. adj.** *phytovorans*, plant-devouring

Nomenclatural type

INSDC Nucleotide: CP119321 Ts

Description

A species established on the basis of MiGA taxonomic novelty analyses, AAI, dDDH, 16S rRNA gene phylogenetic reconstruction and the type material is the genome MAG_4610. Genomic metrics include ANI (82.28), AAI (73.07) and dDDH d4 (21.5). The MAG was reconstructed from lignocellulolytic bacterial consortium and encoded for 74 CAZymes, 31 of them involved in lignocellulose degradation. In addition, putative genes for PET and PUR hydrolisis were found within the MAG.

Classification

Bacteria » Actinomycetota » Actinobacteria » Micrococcales » Microbacteriaceae » Microbacterium » Microbacterium phytovorans

References

Proposed: Díaz-García et al., 2024

Registry URL

https://seqco.de/i:24107

References

1. Díaz-García et al. (2024). Andean soil-derived lignocellulolytic bacterial consortium as a source of novel taxa and putative plastic-active enzymes. *Systematic and Applied Microbiology*. DOI:10.1016/j.syapm.2023.126485

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:-kiq_w89** submitted by **Díaz-García**, **Laura** and including 16 new names has been successfully validated.

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