

Description of 41 unrecorded bacterial species in Korea, isolated from freshwater in 2021

Jung-Hun Jo¹, Seung-Bum Kim², Ji-Eun Im¹, Se-Yoon Chun¹ and Wan-Taek Im^{1,*}

¹Department of Biotechnology, Hankyong National University, Anseong 17579, Republic of Korea ²Department of Microbiology, Chungnam National University, Daejeon 34134, Republic of Korea

*Correspondent: wandra@hknu.ac.kr

Here we describe indigenous prokaryotic species in Korea, a total of 41 bacterial strains were isolated from freshwater from the Republic of Korea. From the high 16S rRNA gene sequence similarity (>98.7%) and formation of a robust phylogenetic clade with the closest species, it was determined that each strain belonged to predefined bacterial species. There is no official report that these 41 species belonged to 5 phyla, 10 classes, 18 orders, 23 families, and 29 genera, which were assigned to *Streptomyces, Mycolicibacterium, Smaragdicoccus, Nocardiopsis*, and *Nocardia* of the phylum *Actinobacteria*; *Runella, Flavobacterium, Algoriphagus, Sphingobacterium*, and *Aequorivita* of the phylum *Bacteroidota*; *Paenibacillus, Bacillus, Metabacillus*, and *Fredinandcohnia* of the phylum *Firmicutes*; *Sphingobium, Erythrobacter, Duganella, Methylopila, Novosphingobium, Azospirillum, Simplicispira, Corallococcus, Pseudomonas, Devosia, Pseudorhodoferax, Pseudomonas, Prolinoborus, Pectobacterium*, and *Aquabacterium* of the phylum *Proteobacteria*; *Proshecobacter* of the phylum *Verrucomicrobia*. Gram reaction, colony and cell morphology, basic biochemical characteristics, isolation source, and strain IDs are also described in the species description section.

Keywords: 16S rRNA, bacterial diversity, freshwater, indigenous prokaryotic species in Korea, unrecorded species

© 2024 National Institute of Biological Resources DOI:10.12651/JSR.2024.13.4.353

INTRODUCTION

Fresh water accounts for 2.5% of the water-based environment, and seawater accounts for 97.5%. Fresh water is slightly different from seawater containing 3.5% salt, and includes various types of ecosystems, including rivers and river soils, where water flows depending on the appeal and altitude difference where the flow is stopped. And it exists in the form of rivers, lakes, and groundwater, and accounts for only about 0.6% of the total water system present on earth except for glaciers, but it is an essential resource for human survival (Wetzel, 2001). In a freshwater environment, most dominant prokaryotes are not cultured except for some groups. Microorganisms are more common in downstream water than in upstream water, and there are many low-nutritive microorganisms in upstream water, and many high-nutritive microorganisms multiply in downstream water. In general, most groups of bacteria form biofilm, but fungi and birds are often found. Microorganisms living in poor and eutrophobic lakes are also different. In eutrophobic lakes, photosynthetic bacteria such as Cyanobacteria multiply in summer. So far, bacterial clusters in the freshwater environment have been known as about 20 phyla, and among them, Actinomycetota, Bacteroidota, Pseudomonadota, Cyanobacteriota, and Verrucomicrobiota were dominant (Newton et al., 2011). According to GOLD statistics, environmental bacteria account for about 10% of all genetic projects, and most of environmental ecological research is biased toward digestive organs, oceans, and soil, accounting for 9.2% of fresh water (Mukherjee et al., 2022). Like the marine and land environment, freshwater-derived prokaryotes basically serve as decomposers to convert organic matter into inorganic substances by transferring organic matter to higher nutritional levels through microfed networks, and are well-known for their function in the circulation of greenhouse gases. Therefore, research on the analysis of microbial diversity derived from freshwater, which has a close impact on human life, but relatively insignificant excavation research, can contribute to the role of freshwaterspecific prokaryotes and thus the expansion of domestic microbial species discovery research.

In 2021, we collected environmental samples from freshwater in Korea and isolated many novel and unrecorded bacterial species during a research program supported by NNIBR (Nakdonggang National Institute of Biological Resources) of Korea. The identified bacterial species belongs to the classes *Bacilli*, *Actinomycetia*, *Alphaproteobacteria*, *Betaproteobacteria*, *Deltaproteobacteria*, *Gammaproteobacteria*, *Verrucomicrobiae*, *Shpingobacteriia*, *Cytophagia*, and *Flavobacteriia*. Here we shortly describe 41 unrecorded bacterial species in the 10 classes belonging to 18 orders.

MATERIALS AND METHODS

A total of 41 bacterial strains affiliated to the classse *Bacilli*, *Actinomycetia*, *Alphaproteobacteria*, *Betaproteobacteria*, *Gammaproteobacteria*, *Deltaproteobacteria*, *Verrucomicrobiae*, *Shpingobacteriia*, *Cytophagia*, and *Flavobacteriia* were isolated from freshwater samples. Freshwater sample was processed separately, spread onto diverse culture media including R2A, 1/5 R2A, nutrient, 1/5 nutrient, and incubated at 20, 25 and 30°C for 5–10 days (Table 1). The designated strain IDs, sources, culture media, and incubation conditions are summarized in Table 1. All strains were purified as single colonies and stored as 20–25% (w/v) glycerol suspension – 80°C as well as lyophilized ampoules.

The morphology was studied on agar plates for 2–5 days under their optimum temperature. Cell size and shape were examined either by transmission electron microscopy or scanning electron microscopy. Gram staining was performed using a Gram-staining kit of the standard procedures. Using API 20NE (bioMérieux) the biochemical characteristics were performed according to the manufacturer's instructions.

Bacterial DNA extraction, PCR amplification and 16S rRNA gene sequencing were performed using the standard procedures described by Lane (1991). The 16S rRNA gene sequences of the strains assigned to the 41 bacterial strains were compared with the sequences held in GenBank by BLASTN and also analyzed using the EzBioCloud blast (https://www.ezbiocloud.net) (Yoon et al., 2017). For phylogenetic analyses, multiple alignments were performed using the Clustal_X program (Thompson et al., 1997) and gaps were edited in the BioEdit program (Hall, 1999). Evolutionary distances were calculated using the Kimura two-parameter model (Kimura, 1983). The phylogenetic trees were constructed by using the neighbor-joining (Saitou et al., 1987) methods with the MEGA X Program (Kumar et al., 2018) with bootstrap values based on 1,000 replications (Felsenstein, 1985).

Table 1. The taxonomic affiliations of isolated strains belong to the class Alphaproteobacteria.

Class	Order	Family	Genus	Strain ID	NNIBR ID	Most closely related species	Similarity (%)	Isolation source	Medium	Incubation conditions (°C, days)
Bacilli	Bacillales Bacillales Bacillales Bacillales Bacillales	Bacillaceae Bacillaceae Bacillaceae Paenibacillaceae Paenibacillaceae Paenibacillaceae	Bacillus Fredinandcolmia Metabacillus Paenibacillus Paenibacillus	MMS21-DH2R7 MMS21-DH2MA11 MMS21-DH3MA7 AS127 AS12 AS51 K11	NNIBR2021642BA1 NNIBR2021642BA49 NNIBR2021642BA3 NNIBR2021642BA40 NNIBR2021642BA18 NNIBR2021642BA18	Bacillus cytotoxicus Fredinandcolmia onubensis Metabacillus mangrovi Paenibacillus ihuae Paenibacillus massiliensis subsp. massiliensis Paenibacillus ottowii	98.91 99.18 99.35 99.73 99.73	Freshwater Freshwater Freshwater Freshwater Freshwater Freshwater	MA MA MA R2A R2A R2A R2A R2A	30°C, 2d 30°C, 3d 30°C, 2d 25°C, 3d 25°C, 3d 25°C, 3d
Actinomycetia	Corynebacteriales Corynebacteriales Streptosporangiales Corynebacteriales Streptomycetales Streptomycetales Streptomycetales	Mycobacteriaceae Nocardiaceae Nocardiapsaceae Streptomycetaceae Streptomycetaceae Streptomycetaceae Streptomycetaceae	Mycolicibacterium Nocardia Nocardiopsis Smaragdicoccus Streptomyces Streptomyces Streptomyces	CG42 MMS21-JYBA42 K69 K21 WG8 AS22 CG18 MMS21-OR1	NNIBR 2021642B A32 NNIBR 2021642B A35 NNIBR 2021642B A35 NNIBR 2021642B A34 NNIBR 2021642B A36 NNIBR 2021642B A19 NNIBR 2021642B A19 NNIBR 2021642B A9 NNIBR 2021642B A9	Mycolicibacterium aichiense Nocardia africana Nocardiopsis dassonvillei subsp. dassonvillei Smaragdicoccus niigatensis Streptomyces bulli Streptomyces shenzhenensis Streptomyces shenzhenensis	98.74 99.29 99.93 100 98.75 98.75 99.17 99.31	Freshwater Freshwater Freshwater Freshwater Freshwater Freshwater Freshwater Freshwater	R2A R2A R2A R2A R2A R2A R2A R2A R2A R2A	25°C, 3d 30°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d 30°C, 3d

Class	Order	Family	Genus	Strain ID	NNIBR ID	Most closely related species	Similarity (%)	Isolation source	Medium	Incubation conditions (°C, days)
Actinomycetia	Streptomycetales Streptomycetales Streptomycetales	Streptomycetaceae Streptomycetaceae Streptomycetaceae	Streptomyces Streptomyces Streptomyces	MMS21-DH1MA22 MMS21-305 MMS21-MKJW7	NNIBR2021642BA48 NNIBR2021642BA46 NNIBR2021642BA45	Streptomyces speibonae Streptomyces violarus Streptomyces wuyuanensis	98.97 99.59 100	Freshwater Freshwater Freshwater	R2A R2A R2A	30°C, 3d 30°C, 3d 30°C, 3d
Alphaproteobacteria	Rhodospirillales Hyphomicrobiales Sphingomonadales Hyphomicrobiales Sphingomonadales Sphingomonadales	Azospirillaceae Devosiaceae Erythrobacteraceae Methylocystaceae Sphingomonadaceae Sphingomonadaceae Sphingomonadaceae	Azospirillum Devosia Erythrobacter Methylopila Novosphingobium Sphingobium	AS37 AS 105 K14 K14 CG59 WG87 AS 117 K3	NNIBR 2021642B A30 NNIBR 2021642B A38 NNIBR 2021642B A24 NNIBR 2021642B A28 NNIBR 2021642B A39 NNIBR 2021642B A39 NNIBR 2021642B A39 NNIBR 2021642B A31	Azospirilum oryzae Devosia elaeis Erythrobacter tepidarius Methylopila oligotropha Novosphingobium arabidopsis Novosphingobium sediminis Sphingobium suffavum	99.41 99.93 99.64 99.64 99.78 99.78	Freshwater Freshwater Freshwater Freshwater Freshwater Freshwater Freshwater	R2A R2A R2A R2A R2A R2A R2A R2A R2A	25°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d
Betaproteobacteria	Burkholderiales Burkholderiales Neisseriales Burkholderiales Burkholderiales	Comanonadaceae Oxalobacteraceae Neisseriaceae Comanonadaceae Comanonadaceae	Aquabacterium Duganella Prolinoborus Pseudorhodoferax Simplicispira	MMS21-SJ6-8 CG34 MMS21-Sa21 AS150 CG15	NNIBR 202 1642B A47 NNIBR 202 1642B A27 NNIBR 202 1642B A7 NNIBR 202 1642B A42 NNIBR 202 1642B A43 NNIBR 202 1642B A31	Aquabacterium lacunae Duganella rivi Prolinoborus fasciculus Pseudorhodoferax aquiterrae Simplicispira lacusdiani	99.59 99.24 98.83 99.72 99.31	Freshwater Freshwater Freshwater Freshwater Freshwater	MA R2A R2A R2A R2A R2A	30°C, 3d 25°C, 3d 30°C, 3d 25°C, 3d 25°C, 3d
Gammaproteobacteria	Enterobacterales Enterobacterales Enterobacterales Enterobacterales	Pectobacteriaceae Pectobacteriaceae Pectobacteriaceae Pectobacteriaceae	Pectobacterium Pseudomonas Pseudomonas Pseudomonas	MMS21-Er7 AS210 AS199 CG31	NNIBR2021642BA10 NNIBR2021642BA44 NNIBR2021642BA43 NNIBR2021642BA336	Pectobacterium aroidearum Pseudomonas batumici Pseudomonas huaxiensis Pseudomonas neuropathica	99.52 99.04 99.86 98.86	Freshwater Freshwater Freshwater Freshwater	R2A R2A R2A R2A R2A	30°C, 3d 25°C, 3d 25°C, 3d 25°C, 3d
Deltaproteobacteria Verrucomicrobiae	Myxococcales Verrucomicrobiales	Myxococcaceae Verrucomicrobiaceae	Corallococcus Prosthecobacter	CG50 CG69	NNIBR2021642BA33 NNIBR2021642BA37	Corallococcus interemptor Prosthecobacter dejongeii	100 99.39	Freshwater Freshwater	R2A R2A	25°C, 3d 25°C, 3d
Shpingobacteriia Cytophagia	Sphingobacteriales Cytophagales Cytophagales	Sphingobacteriaceae Cyclobacteriaceae Spirosomaceae	Sphingobacterium Algoriphagus Runella	MMS21-DH2R2 AS131 K7	NNIBR2021642BA2 NNIBR2021642BA41 NNIBR2021642BA22	Sphingobacterium canadense Algoriphagus aquaeductus Runella aurantiaca	98.97 98.92 98.74	Freshwater Freshwater Freshwater	R2A R2A R2A	30°C, 2d 25°C, 3d 25°C, 3d
Flavobacteriia	Flavobacteriales Flavobacteriales Flavobacteriales	Flavobacteriaceae Flavobacteriaceae Flavobacteriaceae	Aequorivita Flavobacterium Flavobacterium	MMS21-MC1 MMS21-SJ5-12 WG6	NNIBR 2021642BA11 NNIBR 2021642BA51 NNIBR 2021642BA25	Aequorivita sinensis Flavobacterium difficile Flavobacterium squillarum	9.99 77.79 89.86	Freshwater Freshwater Freshwater	MA R2A R2A	30°C, 2d 30°C, 3d 25°C, 3d

Table 1. Continued.



Fig. 1. Transmission and scanning electron micrographs of cells of the strains isolated in this study. Strains: 1, AS51; 2, AS52; 3, CG18; 4, K3; 5, K7; 6, K11; 7, K14; 8, WG6; 9, WG8; 10, CG34; 11, CG59; 12, WG87; 13, AS37; 14, CG15; 15, CG42; 16, CG50; 17, K21; 18, K69; 19, CG31; 20, CG69; 21, AS105; 22, AS117; 23, AS127; 24, AS131; 25, AS150; 26, AS199; 27, AS210; 28, MMS21-SJ5-12; 29, MMS21-305; 30, MMS21-MKJW7; 31, MMS21-DH1MA22; 32, MMS21-Sa21; 33, MMS21-DH2R7; 34, MMS21-DH2R2; 35, MMS21-DH3MA7; 36, MMS21-DH2MA11; 37, MMS21-MC1; 38, MMS21-Er7; 39, MMS21-SJ6-8; 40, MMS21-OR1; 41, MMS21-JYBA42.

RESULTS AND DISCUSSION

The 41 strains were distributed in 10 classes of the Bacilli, Actinomycetia, Alphaproteobacteria, Betaproteobacteria, Gammaproteobacteria, Deltaproteobacteria, Verrucomicrobiae, Shpingobacteriia, Cytophagia, and Flavobacteriia; 6 strains for the class Bacilli, 11 strains for the Actinomycetia, 7 strains for the Alphaproteobacteria, 5 strains for the Betaproteobacteria, 4 strains for the Gammaproteobacteria, 1 strain for the Deltaproteobacteria, 1 strain for the Verrucomicrobiae, 1 strain for the Shpingobacteriia, 2 strains for the Cytophagia and 3 strains for the Flavobacteriia (Table 1). These strains were Gram-staining-negative or positive, chemoheterotrophic, rod, short-rod and coccoid shaped showing in Fig. 1. The strains in the phylum *Actinobacteria* (Fig. 2) were found to belong to 1 class 5 separate genera: *Streptomyces* (7 species), *Mycolicibacterium* (1 species), *Smaragdicoccus* (1 species), *Nocardiopsis* (1 species), *Nocardia* (1 species).

Fig. 3 shows phylogenetic assignment of 17 strains of the phylum *Proteobacteria* belong to 4 classes 14 separate genera: *Aquabacterium* (1 species), *Azospirillum* (2 species), *Corallococcus* (1 species), *Devosia* (1 species), *Duganella* (1 species), *Erythrobacter* (1 species), *Methylopila* (1 species), *Novosphingobium* (2 species), *Pectobacterium* (1 species), *Prolinoborus* (1 species), *Pseudomonas* (3 species), *Pseudorhodoferax* (1 species), *Simplicispira* (1 species) and *Sphingobium* (1 species).

Fig. 4 shows phylogenetic assignment of 13 strains of



Fig. 2. Neighbor-joining phylogenetic tree, based on 16S rRNA gene sequences, showing the relationship between the strains in the phylum *Actinobacteria*. Bootstrap values (>60%) are shown in the neighbor-joining method. Bar, 0.02 substitutions per nucleotide position.



Fig. 3. Neighbor-joining phylogenetic tree, based on 16S rRNA gene sequences, showing the relationship between the strains in the phylum *Proterobacteria*. Bootstrap values are greater than 60% are shown the neighbor-joining tree method. Bar, 0.02 substitutions per nucleotide position.



Fig. 4. Neighbor-joining phylogenetic tree, based on 16S rRNA gene sequences, showing the relationship between the strains in the phylum *Firmicutes*, *Bacteroidota* and *Verrucomicrobia*. Bootstrap values (>60%) are shown in the neighbor-joining tree. Bar, 0.05 substitutions per nucleotide position.

the phyla Firmicutes, Bacteroidota and Verrucomicrobia belong to 5 classes 10 separate genera: Aequorivita (1 species), Algoriphagus (1 species), Bacillus (1 species), Flavobacterium (2 species), Fredinandcohnia (1 species), Metabacillus (1 species), Paenibacillus (3 species), Pros*thecobacter* (1 species), *Runella* (1 species) and *Sphingobacterium* (1 species).

Here we report 41 unrecorded bacterial species in Korea belonging to 10 classes of 5 phyla.

Description of *Paenibacillus massiliensis* subsp. *massiliensis* AS51

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are circular and white color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction, esculin hydrolysis, β -galactosidase, but negative for indole production, glucose fermentation, arginine dihydrolase, urease, gelatinase. Utilizes D-glucose, L-arabinose, D-mannose and D-mannitol. Does not utilize *N*-acetyl-glucosamine, D-maltose, potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Strain AS51 (=NNIBR2021642BA18) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Streptomyces bullii AS52

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are filamentous and white color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction, esculin hydrolysis, β -galactosidase, but negative for indole production, glucose fermentation, arginine dihydrolase, urease, gelatinase. Utilizes D-glucose, L-arabinose, *N*-acetyl-glucosamine, D-maltose, potassium gluconate and malic acid. Does not utilize D-mannose, D-mannitol, capric acid, adipic acid, trisodium citrate and phenylacetic acid. Strain AS52 (=NNIBR2021642BA19) has been isolated from a freshwater algae sample, Anseongsi, Korea.

Description of Streptomyces cellostaticus CG18

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are filamentous and white \rightarrow gray color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction, urease, esculin hydrolysis, β -galactosidase, but negative for indole production, glucose fermentation, arginine dihydrolase, gelatinase. Utilizes D-glucose, L-arabinose, D-mannose, N-acetyl-glucosamine, D-maltose, potassium gluconate, adipic acid and malic acid. Does not utilize D-mannitol, capric acid, trisodium citrate and phenylacetic acid. Strain CG18 (=NNIBR2021 642BA20) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Sphingobium sufflavum K3

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and yellow color after 3 days on R2A at 25°C. In API 20NE, negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis, gelatinase and β -galactosidase. Utilizes D-glucose and malic acid. Does not utilize L-arabinose, D-mannose, D-mannitol, *N*-acetylglucosamine, D-maltose, potassium gluconate, capric acid, adipic acid, trisodium citrate and phenylacetic acid. Strain K3 (=NNIBR2021642BA21) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Sphingobium sufflavum K7

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and pink color after 3 days on R2A at 25°C. In API 20NE, positive for esculin hydrolysis, β -galactosidase, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, gelatinase. Utilizes D-glucose, D-mannose, *N*-acetyl-glucosamine and D-maltose. Does not utilize L-arabinose, D-mannitol, potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Strain K7 (=NNIBR2021642BA22) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Paenibacillus ottowii K11

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are circular and white color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction, esculin hydrolysis, gelatinase, but negative for indole production, glucose fermentation, arginine dihydrolase, urease, β -galactosidase. Utilizes D-glucose, D-mannose, D-mannitol and D-maltose. Does not utilize L-arabinose, *N*-acetyl-glucosamine, potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Strain K11 (=NNIBR2021642BA23) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Erythrobacter tepidarius K14

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and orange color after 3 days on R2A at 25°C. In API 20NE, negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis, gelatinase and β -galactosidase. Utilizes D-glucose, D-mannose and phenylacetic acid. Does not utilize L-arabinose, D-mannitol, *N*-acetyl-glucosamine, D-maltose, potassium gluconate, capric acid, adipic acid, malic acid and trisodium citrate. Strain K14 (=NNIBR2021642BA24) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Flavobacterium squillarum WG6

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and yellow color after 3 days on R2A at 25°C. In API 20NE, positive for gelatinase, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis, β -galactosidase. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, *N*-acetyl-glucosamine, D-maltose, potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Strain WG6 (= NNIBR2021642BA25) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Streptomyces bryophytorum WG8

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are filamentous and ivory color after 3 days on R2A at 25°C. In API 20NE, negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis, gelatinase and β -galactosidase. Utilizes D-glucose, D-mannose, N-acetylglucosamine, potassium gluconate and adipic acid. Does not utilize L-arabinose, D-mannitol, D-maltose, capric acid, malic acid, trisodium citrate and phenylacetic acid. Strain WG8 (=NNIBR2021642BA26) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Duganella rivi CG34

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and white color after 3 days on R2A at 25°C. In API 20NE, positive for urease, esculin hydrolysis, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, gelatinase and β -galactosidase. Utilizes D-glucose and L-arabinose. Does not utilize D-mannose, D-mannitol, *N*-acetyl-glucosamine, D-maltose, potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Strain CG34 (= NNIBR2021642BA27) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Methylopila oligotropha CG59

Cells are Gram-staining-negative, non-flagellated and oval-shaped. Colonies are circular and white color after 3 days on R2A at 25°C. In API 20NE, negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis, gelatinase and β -galactosidase. Utilizes potassium gluconate and malic acid. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, *N*-acetyl-glucosamine, D-maltose, capric acid, adipic acid, trisodium citrate and phenylacetic acid. Strain CG59 (=NNIBR2021642BA28) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Novosphingobium arabidopsis WG87

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and yellow color after 3 days on R2A at 25°C. In API 20NE, positive for urease, β -galactosidase, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, esculin hydrolysis, gelatinase. Utilizes D-glucose, L-arabinose, D-mannose and potassium gluconate. Does not utilize D-mannitol, *N*-acetyl-glucosamine, D-maltose, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Strain WG87 (=NNIBR2021642BA29) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Azospirillum oryzae AS37

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and white color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction, arginine dihydrolase, urease, esculin hydrolysis, β -galactosidase, but negative for indole production, glucose fermentation, gelatinase. Utilizes D-glucose, L-arabinose, D-mannitol, *N*-acetyl-glucosamine, potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Does not utilize D-mannose and D-maltose. Strain AS37 (=NNIBR2021642BA30) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Simplicispira lacusdiani CG15

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and white color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction, esculin hydrolysis, but negative for indole production, glucose fermentation, arginine dihydrolase, urease, gelatinase, β -galactosidase. Utilizes D-glucose, D-mannose, N-acetyl-glucosamine, D-maltose, adipic acid and malic acid. Does not utilize L-arabinose, D-mannitol, potassium gluconate, capric acid, trisodium citrate and phenylacetic acid. Strain CG15 (=NNIBR2021642 BA31) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Mycolicibacterium aichiense CG42

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are circular and yellow color after 3 days on R2A at 25°C. In API 20NE, negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis, gelatinase and β -galactosidase. Utilizes D-glucose, L-arabinose, D-mannose, D-mannitol, potassium gluconate, adipic acid, malic acid and trisodium citrate. Does not utilize *N*-acetylglucosamine, D-maltose, capric acid and phenylacetic acid. Strain CG42 (=NNIBR2021642BA32) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Corallococcus interemptor CG50

Cells are Gram-staining-negative, non-flagellated and coccoid-shaped. Colonies are swarming and dark orange color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction, indole production, esculin hydrolysis, gelatinase, but negative for glucose fermentation, arginine dihydrolase, urease, β -galactosidase. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, *N*-ace-tyl-glucosamine, D-maltose, potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenyl-acetic acid. Strain CG50 (=NNIBR2021642BA33) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Smaragdicoccus niigatensis K21

Cells are Gram-staining-positive, non-flagellated and coccoid-shaped. Colonies are circular and dark green color after 3 days on R2A at 25°C. In API 20NE, positive for urease, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, esculin hydrolysis, gelatinase, β -galactosidase. Utilize *N*-acetyl-glucosamine and malic acid. Does not utilize D-glucose, L-arabinose, D-mannitol, D-maltose, potassium gluconate, capric acid, adipic acid, trisodium citrate and phenylacetic acid. Strain K21 (=NNIBR2021642 BA34) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of *Nocardiopsis dassonvillei* subsp. *dassonvillei* K69

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are filamentous and white color after 3 days on R2A at 25°C. In API 20NE, positive for esculin hydrolysis, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, gelatinase, β -galactosidase. Utilize *N*-acetylglucosamine. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, D-maltose, potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Strain K69 (=NNIBR2021642BA35) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Pseudomonas neuropathica CG31

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are irregular raised undulate and cream color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction, arginine dihydrolase and esculin hydrolysis, but negative for indole production, glucose fermentation, urease, gelatinase, β -galactosidase. Utilize capric acid and malic acid. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, *N*-acetyl-glucosamine, D-maltose, potassium gluconate, adipic acid, trisodium citrate and phenylacetic acid. Strain CG31 (=NNIBR2021642BA36) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Prosthecobacter dejongeii CG69

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and cream color after 3 days on R2A at 25°C. In API 20NE, positive for urease, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, esculin hydrolysis, gelatinase, β -galactosidase. Utilize D-glucose, D-mannose, *N*-acetyl-glucosamine, D-maltose, malic acid and trisodium citrate. Does not utilize L-arabinose, D-mannitol, potassium gluconate, capric acid, adipic acid and phenylacetic acid. Strain CG69 (=NNIBR2021642BA37) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Devosia elaeis AS105

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and cream color after 3 days on R2A at 25°C. In API 20NE, positive for esculin hydrolysis, weak positive for β -galactosidase, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, gelatinase. Utilize D-glucose, L-arabinose, D-mannose, D-mannitol, *N*-acetylglucosamine and D-maltose. Does not utilize potassium gluconate, capric acid, adipic acid, malic acid, trisodium citrate and phenylacetic acid. Strain AS105 (=NNIBR 2021642BA38) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Novosphingobium sediminis AS117

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and yellow color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction and β -galactosidase, but negative for indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis and gelatinase. Utilize D-glucose, L-arabinose, D-mannose, D-mannitol, D-maltose, potassium gluconate and adipic acid. Does not utilize *N*-acetylglucosamine, capric acid, malic acid, trisodium citrate and phenylacetic acid. Strain AS117 (=NNIBR2021642BA 39) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Paenibacillus ihuae AS127

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are circular and yellow color after 3 days on R2A at 25°C. In API 20NE, positive for esculin hydrolysis, gelatinase β -galactosidase, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease. Utilize D-glucose, L-arabinose, D-mannose, D-mannitol, D-maltose, potassium gluconate and malic acid. Does not utilize *N*-acetyl-glucosamine, capric acid, adipic acid, trisodium citrate and phenylacetic acid. Strain AS127 (=NNIBR2021642BA40) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Algoriphagus aquaeductus AS131

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and red color after 3 days on R2A at 25°C. In API 20NE, positive for gelatinase and β -galactosidase, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis. Utilize D-glucose, D-mannose, *N*-acetyl-glucosamine, D-maltose and phenylacetic acid. Does not utilize L-arabinose, D-mannitol, potassium gluconate, capric acid, adipic acid, malic acid and trisodium citrate. Strain AS131 (= NNIBR2021642BA41) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Pseudorhodoferax aquiterrae AS150

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and white color after 3 days on R2A at 25°C. In API 20NE, positive for nitrate reduction and urease, but negative for indole production, glucose fermentation, arginine dihydrolase, esculin hydrolysis, gelatinase, β -galactosidase. Utilize L-arabinose, D-mannitol, D-maltose, potassium gluconate, adipic acid and malic acid. Does not utilize D-glucose, D-mannose, N-acetyl-glucosamine, capric acid, trisodium citrate and phenylacetic acid. Strain AS150 (=NNIBR2021642BA 42) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Pseudomonas huaxiensis AS199

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and red color after 3 days on R2A at 25°C. In API 20NE, negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis, gelatinase and β -galactosidase. Utilize D-glucose, D-mannitol, potassium gluconate, capric acid, malic acid, trisodium citrate and phenylacetic acid. Does not utilize L-arabinose, D-mannose, *N*-acetyl-glucosamine, D-maltose and adipic acid. Strain AS199 (= NNIBR2021642BA43) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Pseudomonas batumici AS210

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular and cream color after 3 days on R2A at 25°C. In API 20NE, negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis, gelatinase and β -galactosidase. Utilize D-glucose, L-arabinose, D-mannose, D-mannitol, potassium gluconate, capric acid, malic acid and trisodium citrate. Does not utilize *N*-acetyl-glucosamine, D-maltose, adipic acid and phenylacetic acid. Strain AS210 (= NNIBR2021642BA44) has been isolated from a freshwater algae sample, Anseong-si, Korea.

Description of Flavobacterium difficile MMS21-SJ5-12

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular, convex, entire and yellow color after 3 days on R2A at 30°C. In API 20NE, positive for esculin hydrolysis and β -galactosidase, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease and gelatinase. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, potassium gluconate, capric acid, malic acid, trisodium citrate, *N*-acetyl-glucosamine, D-maltose, adipic acid and phenylacetic acid. Strain MMS21-SJ5-12 (=NNIBR2021642BA51) has been isolated from a freshwater sample, Okcheon-gun, Korea.

Description of Streptomyces violarus MMS21-305

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are filamentous, raised, undulate and yellow color after 3 days on R2A at 30°C. In API 20NE, positive for nitrate reduction, urease, glucose fermentation, gelatinase and β -galactosidase, but negative for esculin hydrolysis, indole production and arginine dihydrolase. Utilize D-glucose, L-arabinose, D-mannose, D-mannitol, potassium gluconate, malic acid, trisodium citrate, *N*-acetyl-glucosamine, D-maltose and adipic acid. Does not utilize capric acid and phenylacetic acid. Strain MMS21-305 (=NNIBR2021642BA46) has been isolated from a freshwater sample, Gimhae-si, Korea.

Description of *Streptomyces wuyuanensis* MMS21-MKJW7

Cells are Gram-staining-positive and rod-shaped. Colonies are filamentous, raised, undulate and brown color after 3 days on R2A at 30°C. In API 20NE, positive for glucose fermentation, gelatinase and β -galactosidase, but negative for nitrate reduction, indole production, arginine dihydrolase, urease and esculin hydrolysis. Utilize D-glucose, Dmannose, *N*-acetyl-glucosamine, D-maltose, potassium gluconate and malic acid. Does not utilize L-arabinose, adipic acid, D-mannitol, capric acid, trisodium citrate and phenylacetic acid. Strain MMS21-MKJW7 (=NNIBR 2021642BA45) has been isolated from a freshwater sample, Gimhae-si, Korea.

Description of *Streptomyces speibonae* MMS21-DH1MA22

Cells are Gram-staining-positive and rod-shaped. Colonies are circular, convex, entire and gray color after 3 days on R2A at 30°C. In API 20NE, positive for nitrate reduction and glucose fermentation, but negative for indole production, esculin hydrolysis, β -galactosidase, arginine dihydrolase, urease and gelatinase. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, potassium gluconate, capric acid, malic acid, trisodium citrate, *N*-acetyl-glucosamine, D-maltose, adipic acid and phenylacetic acid. Strain MMS21-DH1MA22 (=NNIBR2021642BA 48) has been isolated from a freshwater sample, Gongjusi, Korea.

Description of Prolinoborus fasciculus MMS21-Sa21

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are circular, convex, entire and white color after 3 days on R2A at 30°C. In API 20NE, positive for nitrate reduction and urease, but negative for esculin hydrolysis, β -galactosidase, indole production, glucose fermentation, arginine dihydrolase and gelatinase. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, potassium gluconate, capric acid, malic acid, trisodium citrate, *N*-acetyl-glucosamine, D-maltose, adipic acid and phenylacetic acid. Strain MMS21-Sa21 (=NNIBR2021 642BA7) has been isolated from a freshwater sample, Gangneung-si, Korea.

Description of Bacillus cytotoxicus MMS21-DH2R7

Cells are Gram-staining-positive, flagellate and rodshaped. Colonies are rhizoid, raised, undulate and white color after 3 days on marine agar 2216 at 30°C. In API 20NE, positive for arginine dihydrolase, urease, glucose fermentation, gelatinase and β -galactosidase, but negative for nitrate reduction, indole production and esculin hydrolysis. Utilize D-glucose, L-arabinose, D-mannose, D-mannitol and potassium gluconate. Does not utilize capric acid, malic acid, trisodium citrate, *N*-acetyl-glucosamine, D-maltose, adipic acid and phenylacetic acid. Strain MMS21-DH2R7 (=NNIBR2021642BA1) has been isolated from a freshwater sample, Gongju-si, Korea.

Description of *Sphingobacterium canadense* MMS21-DH2R2

Cells are Gram-staining-negative, non-flagellated and

rod-shaped. Colonies are circular, convex, entire and yellow color after 2 days on R2A at 30°C. In API 20NE, positive for nitrate reduction, glucose fermentation, arginine dihydrolase, urease, esculin hydrolysis and β -galactosidase, but negative for indole production and gelatinase. Utilize D-glucose, L-arabinose, D-mannose, *N*-acetyl-glucosamine and D-maltose. Does not utilize D-mannitol, potassium gluconate, capric acid, malic acid, trisodium citrate, adipic acid and phenylacetic acid. Strain MMS21-DH2R2 (=NNIBR2021642BA2) has been isolated from a freshwater sample, Gongju-si, Korea.

Description of *Metabacillus mangrovi* MMS21-DH3MA7

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are circular, convex, entire and white color after 3 days on marine agar 2216 at 30°C. In API 20NE, positive for arginine dihydrolase, esculin hydrolysis, urease, gelatinase and β -galactosidase, but negative for nitrate reduction, indole production and glucose fermentation. Utilize D-glucose, L-arabinose, D-mannose, D-mannitol, *N*-acetyl-glucosamine, D-maltose and potassium gluconate. Does not utilize capric acid, malic acid, trisodium citrate, adipic acid and phenylacetic acid. Strain MMS21-DH3MA7 (=NNIBR2021642BA3) has been isolated from a freshwater sample, Gongju-si, Korea.

Description of *Fredinandcohnia onubensis* MMS21-DH2MA11

Cells are Gram-staining-positive, flagellate and rodshaped. Colonies are irregular, flat, entire and white color after 3 days on marine agar 2216 at 30°C. In API 20NE, positive for nitrate reduction, esculin hydrolysis, gelatinase and β -galactosidase, but negative for indole production, glucose fermentation, arginine dihydrolase and urease. Utilize L-arabinose, potassium gluconate, *N*-acetyl-glucosamine and D-maltose. Does not utilize D-glucose, D-mannose, D-mannitol, capric acid, malic acid, trisodium citrate, adipic acid and phenylacetic acid. Strain MMS21-DH2MA11 (=NNIBR2021642BA49) has been isolated from a freshwater sample, Gongju-si, Korea.

Description of Aequorivita sinensis MMS21-MC1

Cells are Gram-staining-negative, non-flagellated and rod-shaped. Colonies are beige color after 3 days on marine agar 2216 at 30°C. In API 20NE, positive for urease and esculin hydrolysis, but negative for nitrate reduction, indole production, β -galactosidase, glucose fermentation, arginine dihydrolase and gelatinase. Utilize D-mannose, D-mannitol and trisodium citrate. Does not utilize D-glucose, L-arabinose, *N*-acetyl-glucosamine, potassium gluconate, capric acid, malic acid, D-maltose, adipic acid and phenylacetic acid. Strain MMS21-MC1 (=NNIBR2021 642BA11) has been isolated from a freshwater sample, Asan-si, Korea.

Description of *Pectobacterium aroidearum* MMS21-Er7

Cells are Gram-staining-negative, flagellate and rodshaped. Colonies are circular, flat, undulate and white color after 3 days on R2A at 30°C. In API 20NE, positive for glucose fermentation, but negative for esculin hydrolysis, β -galactosidase, nitrate reduction, indole production, arginine dihydrolase, urease and gelatinase. Utilize D-glucose, L-arabinose, D-mannose, D-mannitol, *N*-acetyl-glucosamine and D-maltose. Does not utilize, adipic acid, phenylacetic acid, potassium gluconate, capric acid, malic acid and trisodium citrate. Strain MMS21-Er7 (=NNIBR 2021642BA10) has been isolated from a freshwater sample, Daejeon, Korea.

Description of Aquabacterium lacunae MMS21-SJ6-8

Cells are Gram-staining-negative, flagellate and rodshaped. Colonies are circular, convex, entire and white color after 3 days on marine agar 2216 at 30°C. In API 20NE, positive for esculin hydrolysis, but negative for nitrate reduction, indole production, glucose fermentation, arginine dihydrolase, urease, gelatinase and β -galactosidase. Utilize D-glucose. Does not utilize L-arabinose, D-mannose, D-mannitol, potassium gluconate, capric acid, malic acid, trisodium citrate, *N*-acetyl-glucosamine, D-maltose, adipic acid and phenylacetic acid. Strain MMS21-SJ6-8 (=NNIBR2021642BA47) has been isolated from a freshwater sample, Okcheon-gun, Korea.

Description of *Streptomyces shenzhenensis* MMS21-OR1

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are circular, convex, entire and brown color after 3 days on R2A at 30°C. In API 20NE, positive for urease and esculin hydrolysis, but negative for nitrate reduction, β -galactosidase, indole production, glucose fermentation, arginine dihydrolase, and gelatinase. Utilize D-mannose, D-mannitol and trisodium citrate. Does not utilize *N*-acetyl-glucosamine, potassium gluconate, capric acid, malic acid, D-glucose, L-arabinose, Dmaltose, adipic acid and phenylacetic acid. Strain MMS21-OR1 (=NNIBR2021642BA9) has been isolated from a freshwater sample, Cheongju-si, Korea.

Description of Nocardia africana MMS21-JYBA42

Cells are Gram-staining-positive, non-flagellated and rod-shaped. Colonies are punctiform, convex, entire and peach color after 3 days on R2A at 30°C. In API 20NE, positive for nitrate reduction, but negative for esculin hydrolysis, β -galactosidase, indole production, glucose fermentation, arginine dihydrolase, urease and gelatinase. Does not utilize D-glucose, L-arabinose, D-mannose, D-mannitol, potassium gluconate, capric acid, malic acid, trisodium citrate, *N*-acetyl-glucosamine, D-maltose, adipic acid and phenylacetic acid. Strain MMS21-JYBA42 (=NNIBR2021642BA8) has been isolated from a freshwater sample, Gangneung-si, Korea.

CONFLICTS OF INTEREST

The author of this paper has no affiliation with any interests and is solely responsible for the paper.

ACKNOWLEDGEMENTS

This research was supported by the "Survey of freshwater organisms (Prokaryotes)" (NNIBR202101204) research program of the Nakdonggang National Institute of Biological Resources.

References

- Felsenstein, J. 1985. Confidence limit on phylogenies: an approach using the bootstrap. Evolution 39:783-791.
- Hall, T.A. 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/ NT. Nucleic Acids Symp. Ser. 41:95-98.
- Kimura, M. 1983. The Neutral Theory of Molecular Evolution. Cambridge: Cambridge University Press, Cambridge, New York.
- Kumar, S., G. Stecher, M. Li, C. Knyaz and K. Tamura. 2018. MEGA X: molecular evolutionary genetics analysis across computing platforms. Mol. Biol. Evol .35:1547-1549.
- Lane, D.J. 1991. 16S/23S rRNA sequencing. In: Stackebrandt, E. and M. Goodfellow (eds.), Nucleic acid techniques in bacterial systematics. Wiley, New York, USA.
- Mukherjee, S., D. Stamatis, C.T. Li, G. Ovchinnikova, J. Bertsch, J.C. Sundaramurthi, M. Kandimalla, P.A. Nicolopoulos, A. Favognano, I.A. Chen, N.C. Kyrpides and T.B.K. Reddy. 2022. Twenty-five years of Genomes OnLine Database (GOLD): data updates and new features in v.9. Nucleic Acids Res. https://doi.org/10.1093/nar/gkac974
- Newton, R.J., S.E. Jones, A. Eiler, K.D. McMahon and S. Bertilsson. 2011. A guide to the natural history of freshwater lake bacteria. Microbiol. Mol. Biol. Rev. 75:14-49.
- Saitou, N. and M. Nei. 1987. The neighbor-joining method: a new method for reconstructing phylogenetic trees. Molecular Biology and Evolution 4(4):406-425.
- Thompson, J.D., T.J. Gibson, F. Plewniak, F. Jeanmougin and D.G. Higgins. 1997. The CLUSTAL_X windows interface:

flexible strategies for multiple sequence alignment aided by quality analysis tools. Nucleic Acids Res. 25:4876-4882.

Wetzel, R.G. 2001. Limnology: lake and river ecosystems: Gulf Professional Publishing.

Yoon, S.H., S.M. Ha, S. Kwon, J. Lim, Y. Kim, H. Seo and J. Chun. 2017. Introducing EzBioCloud: A taxonomically united database of 16S rRNA and whole genome assemblies. Int. J. Syst. Evol. Microbiol. 67:1613-1617.

Submitted: December 19, 2022 Revised: February 19, 2024 Accepted: February 22, 2024