Nodosilinea signiensis sp. nov. (Leptolyngbyaceae, Synechococcales), a new terrestrial cyanobacterium isolated from mats collected on Signy Island, South Orkney Islands, Antarctica

Ranina Radzi¹, Narongrit Muangmai², Paul Broady³, Wan Maznah Wan Omar¹, Sebastien Lavoue¹, Peter Convey⁴, Faradina Merican*¹

¹ School of Biological Sciences, Universiti Sains Malaysia, Minden, Penang, Malaysia, ² Department of Fishery Biology, Faculty of Fisheries, Kasetsart University, Chatuchak, Bangkok, Thailand, ³ School of Biological Sciences, University of Canterbury, Christchurch, New Zealand, ⁴ British Antarctic Survey, Cambridge, United Kingdom

* faradina@usm.my

Abstract

Terrestrial cyanobacteria are very diverse and widely distributed in Antarctica, where they can form macroscopically visible biofilms on the surfaces of soils and rocks, and on benthic surfaces in fresh waters. We recently isolated several terrestrial cyanobacteria from soils collected on Signy Island, South Orkney Islands, Antarctica. Among them, we found a novel species of Nodosilinea, named here as Nodosilinea signiensis sp. nov. This new species is morphologically and genetically distinct from other described species. Morphological examination indicated that the new species is differentiated from others in the genus by cell size, cell shape, filament attenuation, sheath morphology and granulation. 16S rDNA phylogenetic analyses clearly confirmed that N. signiensis belongs to the genus Nodosilinea, but that it is genetically distinct from other known species of Nodosilinea. The D1–D1´ helix of the 16S–23S ITS region of the new species was also different from previously described Nodosilinea species. This is the first detailed characterization of a member of the genus Nodosilinea from Antarctica as well as being a newly described species.

Introduction

Cyanobacteria are a widely distributed group of oxygenic photosynthetic prokaryotes that possess chlorophyll a and phycobiliproteins [1]. Despite their widespread occurrence and ecological importance, the taxonomy of cyanobacteria remains problematic. Cyanobacterial classification has recently undergone rapid revision based on the use of polyphasic approaches to define new taxa [2]. These approaches combine molecular characterization with cytomorphological and ecological characteristics and have been used to erect and describe new taxa and for the validation of classically described taxa [3, 4].
Amongst the different cyanobacteria groups, Leptolyngbyaceae (Synechococcales) is one of the most taxonomically challenging. Representatives of this family are morphologically simple, often with ambiguous and indistinct morphological features that lead to uncertainty in identification [5]. However, species within the group can be distinguished based on molecular evaluation. Ongoing taxonomic revisionary work to date has resulted in over 21 new genera being erected from the original genus Leptolyngbya [6–23]. The new erected genus were listed in various literature [6–11] and others are described [12–23].

The genus Nodosilinea [8] shows a clear phylogenetic separation from Leptolyngbya based on molecular assessment. The genus has been reported to possess a unique capability to form short convoluted lengths of trichomes within the bounding sheath, termed nodules. This was first documented in L. nodulosa [24]. The generic assignment of L. nodulosa was subsequently revised with the support of molecular phylogenetics and this species became the first member of the newly erected genus Nodosilinea, as N. nodulosa [24]. Subsequent studies have reported the ability to form nodules in all members of the genus when exposed to low-light conditions of < 4 μmol m−2 s−1 for 4 weeks. Some species have also been reported to perform nitrogen fixation [8, 24, 25]. Apart from nodule formation, the characteristics of the genus resemble those of Leptolyngbya, with uniseriate trichomes ranging from 0.5 to 3.5 μm wide within a thin sheath, similar vegetative and apical cell shapes, constriction at each cross wall and containing granules [26].

Representatives of the genus have been recorded from various habitats. Nodosilinea sp. LEGE 13457 and Nodosilinea sp. TM-3.1 were both isolated from the McMurdo Dry Valleys, Antarctica [27]. N. ramsarensis and N. radiophila were recorded from a thermal spring in Iran [28] and N. nodulosa from marine phytoplankton in the China Sea [24]. Other species are sub-aerial on walls (N. epilithica [8], N. chupicuarensis [25]) and occur in soils (N. conica [8]) and freshwater (N. bijugata [8, 29]).

In Antarctica, cyanobacteria are the most important primary colonizers of soils [30]. They play vital roles in soil stabilisation, photosynthetic carbon fixation, and the release of fixed nitrogen, whilst forming the base of the terrestrial food web [31]. Cyanobacteria are found across all geographical regions of Antarctica where they can form macroscopically visible mats, crusts or thin biofilms on the surfaces of soils and rocks and in streams, ponds and lakes, as well as occupying endolithic niches [32, 33].

Leptolyngbya sensu stricto is one of the most widely recorded genera in Antarctica [34–41]. However, detailed characterization combining morphological, ultrastructural and molecular approaches has been performed to date on only three named species (L. bijugata [42], L. borchgrevinkii and L. frigida [43]) and two unidentified species [44]. Three species of Leptolyngbya (i.e. L. foveolarum, L. notata and L. perelegans) were identified in a broad-scale floristic survey conducted on Signy Island, South Orkney Islands, in the 1970s [34]. Their taxonomic assignments, however, were based solely on morphological traits and have not subsequently been subjected to molecular assessment.

In this study, we report the polyphasic characterization of a cyanobacterial strain superficially resembling Leptolyngbya, recently isolated from soils collected on Signy Island.

## Materials and methods

### Ethics statement

New algal material was collected from Signy Island in the 2015/16 austral summer season under permit number 46/2015 issued under the United Kingdom Antarctic Act. The Department of Quarantine and Inspection Services Malaysia (MAQIS) provided import permit JP1412016065849 to permit the samples to be imported to Malaysia (https://www.maqis.gov.
Sample origin and culture conditions
Mat samples were collected under permit as listed above from cracks and crevices in rocks and beneath loose fragments of stone on a west-facing slope below Robin Peak (60.6833˚ S, 45.6333˚ W) on Signy Island during the expedition of British Antarctic Survey in austral summer of 2015/16. All apparatus used for obtaining the samples was sterile and samples were stored and subsequently transported frozen (-20˚C) in sterile containers to the Universiti Sains Malaysia. Cyanobacterial strains were then brought into culture by inoculation on 1% agarised full strength BG-11 medium in Petri dishes [45]. Culture media were supplemented with 100 μg mL⁻¹ cycloheximide to prevent growth of eukaryotes [46]. Cultures were incubated at 15 ± 2˚C with 24 h light supplied by cool white fluorescent lamps at < 4 μmol m⁻² s⁻¹. After 4 weeks of incubation, algal growth developing on the plates was examined microscopically.

Morphological characterization
Morphological examination was conducted using an Olympus BX-53 light microscope (Olympus America Inc., Center Valley, PA, USA) at 100 – 2000X magnification. Photomicrographs were taken. Illustrations were made with the aid of a camera lucida. Specimens were analysed based on morphological characteristics, particularly filament and trichome width, sheath morphology, cell colour, shape of intercalary and apical cells and presence of granules. Size measurements were made on 30 randomly chosen replicate specimens for each morphospecies. The characteristics of the strain studied were compared with descriptions in the literature [8, 25, 26].

Transmission electron microscopy
Transmission electron microscopy (TEM) samples were fixed in McDowell-Trump fixative solution [47] prepared in 0.1 M phosphate buffer and later post-fixed with 1% osmium tetroxide. The fixed material was dehydrated in an ethanol series (50%, 75%, 95%, and 100%) and embedded in Spurr’s resin [48] mixed in a rotator overnight. The ultrathin sections were treated with uranyl acetate and lead citrate to improve contrast [49]. Thin sections were collected on copper grids and were observed using a JEM 2000FX (JEOL, Tokyo, Japan) operating at 100 kV.

Molecular analyses
DNA was extracted using the G-spin for bacteria genomic DNA extraction kit (iNtRON Biotechnology, Korea) following the manufacturer’s protocol. DNA sample concentration was measured using a Thermo Scientific NanoDrop instrument. The 16S rDNA gene and the 16S–23S internal transcribed spacer (ITS) region were amplified using the polymerase chain reaction (PCR) and the combination of primers 2 (5’–GGG GGA TTT TCC GCA ATG GG–3’) and 3 (5’–CGC TCT ACC AAC TGA GCT A– 3’) for the 16S rRNA gene and primers 1 (5’– CTC TGT GTG CCT AGG TAT CC– 3’) and 5 (5’–TGT AGC TCA GGT GGT TAG– 3’) for the ITS region [50]. This resulted in products of approximately 1,600 bp for the 16S rRNA gene and 600 bp for the ITS region. The reaction mix comprised 2 μL of extracted DNA used in 50 μL reactions containing 1 μL of each forward and reverse primer, 21 μL of ultrapure water and 25 μL of MyTaq™ Red Mix, which is a pre-prepared
mixture of buffer, dNTPs and Taq polymerase (Bioline, United Kingdom). PCR was carried out using a Bio-Rad Thermal Cycler with standard parameters set as follows: 95°C for 2 min, 95°C for 15 sec, 55°C for 15 sec (30 cycles), and 7 min 20 sec extension at 72°C. Once the reaction was completed, the integrity of the PCR product was verified using a 2% agarose gel.

Phylogenetic analyses and ITS folding

All sequences were edited and assembled using the Geneious 11.0 software package (Biomatters, http://www.geneious.com). Sequence alignments were prepared using the MUSCLE algorithm in Geneious 11.0 and then manually checked by eye. The dataset included 47 OTUs, consisting of sequences newly obtained in this study together with additional sequences retrieved from GenBank of closely related species of *Nodosilinea*, more distantly related species of *Leptolyngbya sensu stricto* and one outgroup taxon (*Gloeobacter violaceus* FR798924). Some *Leptolyngbya* sequence that were retrieved from GeneBank currently continue to be known as *Nodosilinea* and we identify these sequences with “*Leptolyngbya*” in the phylogenetic tree. All new sequences generated in this study have been deposited in GenBank under accession numbers USMFM MN585774 and USMFM MN585775. Phylogenetic analyses were performed using two different methods: maximum likelihood (ML) and Bayesian inference (BI). Before ML and BI analyses, the best-fit model of DNA substitution was determined using the program Kakusan4 [51]. ML analyses were performed using with RaxML v7 [52] in Geneious 11.0 using the general time-reversible invariant-sites (GTRI) nucleotide substitution model with the default parameters. The bootstrap probability of each branch was calculated using 1000 replications. BI analyses were performed with the program MrBayes v3.1.2 [53]. Two independent analyses, each consisting of four Markov chains, were run simultaneously for 3,000,000 generations, sampling every 100 generations. Log likelihood and parameter values were assessed with Tracers ver. 1.5 [54]. A burn-in of 25% of saved trees was removed, and the remaining trees were used to calculate the Bayesian posterior probability values. ML and BI trees were edited with the program FigTree v1.3.1 [53].

The 16S–23S ITS region was used for modelling of secondary structure folding. The tRNA genes were identified using tRNAscan-SE 2 [55]. The secondary structure of the D1–D1′ helix was modelled using the Mfold WebServer with default conditions.

Results

A single strain was successfully isolated from one sampling site and showed the diagnostic traits of the genus *Nodosilinea*. However, it has morphological characteristics that did not correspond to any previously described species. In addition, our phylogenetic study confirmed that this strain forms a distinct lineage within the genus with respect to the nucleotide sequences of both the 16S rDNA gene and the 16S-23S ITS region. We hereafter refer to this strain under the name *Nodosilinea signiensis* sp. nov.

*Class* Cyanophyceae  
*Order* Synechococcales  
*Family* Leptolyngbyaceae  
*Genus* Nodosilinea  
*Nodosilinea signiensis* sp. nov.

**Description.** (Culture conditions) Mat creeping on agar, pale blue-green to olive-green in colour. Filaments long, immotile, solitary, occasionally forming spirals under normal light conditions of 27 μmol m⁻² s⁻¹ (Fig 1A and 1C). Under low light intensity of <4 μmol m⁻² s⁻¹, uniseriate trichomes can lie parallel or twisted around one another within a common sheath.
(Fig 1D and 1E). This resembles nodule formation whereby filament width in these areas becomes wider, up to 5.0 μm wide (Fig 2C–2E). Cells 1.0 (1.5)–2.0 μm wide, 1.0–2.0 (2.3) μm long, shorter to longer than wide, discoid to barrel-shaped (Fig 2A–2G). Apical cells rounded, non–capitate, without calyptra and lacking granules (Fig 2G). Sheath colourless, thin.

**Etymology.** *Nodosilinea signiensis*, *Nodosilinea* = "Knotted line" Perkeson et al. (2011); *signiensis* (sig.nie’n.sis) adj. signiensis = originated from Signy Island.

**Habitat.** Mats in cracks and crevices in rocks and beneath loose fragments of stone collected from west-facing slope below Robin Peak (60.6833˚ S, 45.6333˚ W).

**Occurrence.** South Orkney Islands, Signy Island.

**Observations.** In cultures incubated for 4 weeks under low light, filaments are entangled, curved or very occasionally form a spiral. Cell division without lengthening of the enclosing sheath causes the trichomes to sometimes bend and fragment to form two or three twisted structures that resemble nodules, although this was rarely observed.

**Morphological assessment.** The strain displays similar nodule formation to that of the seven currently described species within the genus. *Nodosilinea* sp. LEGE 13457 and *Nodosilinea* sp. TM-3.1, both originating from Antarctica [27], are not included here in the comparison as both strains lack detailed taxonomic characterization. Cell width of *Nodosilinea signiensis* sp. nov. falls within the range of all previously recorded species of *Nodosilinea*.

Fig 1. *Nodosilinea signiensis* sp. nov. grown in culture. (A), filaments. (B), rounded apical cell (arrow). (C), spiral formation (arrow). (D) and (E), trichomes twisted within the enclosing sheath, young trichome forming nodules (arrow). Scale bars: 20 μm.

https://doi.org/10.1371/journal.pone.0224395.g001
Cell length closely resembles five of the seven described species (N. conica, N. chupicuarensis, N. nodulosa, N. radiophila and N. ramsarensis), while N. epilithica and N. bijugata have longer cells at 8 μm and 6.2 μm, respectively (Table 1). The vegetative cell shape of N. signiensis was distinct, being the only species showing a variety of cell shapes ranging from isodiametric to longer than wide or shorter than wide. Cell constriction changed from slightly constricted to distinctly constricted as trichomes matured. In all other species of Nodosilinea, the cell shape was reported to be stable throughout the development of the trichome (Table 1). Trichomes of the present strain showed no attenuation towards the apex, in contrast to the abrupt tapering in N. conica (Table 1). Nodosilinea signiensis sp. nov. is the only species lacking cell granulation. Sheath development was very similar to N. conica and N. chupicuarensis, with the thin, colourless sheath occasionally becoming wide in mature filaments (Table 1). Three species, N. conica, N. radiophila, and N. ramsarensis, have been recorded from extreme environments (Table 1).

**Transmission electron microscopy.** Thylakoids were parietal (4–5 per cell), visible in longitudinal section but poorly seen under cross section (Fig 3). Nodule formation was not observed under TEM. Cyanophycin granules were visible among the thylakoids but carboxysomes were absent. The cell wall was simple, similar to Leptolyngbya species, and the sheath was distinct.

**Phylogenetic analysis.** The sequence dataset consisted of 1,482 bp, including gaps. Our tree suggests that “Leptolyngbya antarctica” should be reclassified into the genus Nodosilinea, as with several other sequence of “Leptolyngbya” deposited in GenBank. Partial sequences of the 16S rDNA of N. signiensis were distinct from other Nodosilinea sequences including “Leptolyngbya antarctica” by at least 2% (≥ 29 differences). Both ML and BI analyses yielded an identical topology, and therefore only the ML tree with an L score of 11654.762 is presented (Fig 4). The 16S rDNA phylogeny clearly indicated that our strain was nested within the genus Nodosilinea (98% ML bootstrap percentage (BP) and 1.00 posterior probability (PP). The relationships within the Nodosilinea clade are complex and showed several low- to well-supported monophyletic groups. Nodosilinea signiensis forms a monophyletic group with two sequences of two specimens previously identified as “Leptolyngbya” sp. and one sequence of Nodosilinea sp. with high support from both ML (BP = 79%) and BI (PP = 93%).

---

**Fig 2. Morphological characteristics of Nodosilinea signiensis sp. nov.** (A)–(E), trichome variously curved, sometimes straight, tangled together or twisted, characteristic nodules (arrow). (F) and (G), trichomes lying together within a common sheath, mature rounded apical cells (arrow). Scale bar: 10 μm.

https://doi.org/10.1371/journal.pone.0224395.g002

(Table 1)
Table 1. Comparison of characteristics of the previously described eight species of *Nodosilinea* [8, 24, 25, 28] and *N. signiensis* sp. nov. (’?’ indicates feature unknown).

| Characters | Nodosilinea signiensis sp. nov. | Nodosilinea epilithica | Nodosilinea bijugata | Nodosilinea conica | Nodosilinea chupicuarensis | Nodosilinea nodulosa | Nodosilinea radiophila | Nodosilinea ramsarensis |
|------------|--------------------------------|------------------------|---------------------|------------------|--------------------------|---------------------|-----------------------|------------------------|
| Cell length (μm) | 1.0–2.0 (2.3) | 1.0–8.0 | 1.5–6.2 | 0.9–2.4 | 1.1–1.3 | 1.1–1.5 | 1.0–2.0 (0.8) | 1.0–1.5 |
| Cell width (μm) | 1.0 (1.5) | 1.5–2.5 | 1.5–1.7 | 2.5–2.7 | 1.2 | 1.2–2.4 | 2.0–5.0 | 1.0–2.0 |
| Cell shape | Isodiametric, longer than wide/ barrel shape | Barrel shaped, shorter to longer than wide | Isodiametric, longer than wide | Isodiametric, shorter than wide | Isodiametric | Isodiametric, longer than wide | Isodiametric, longer than wide | Isodiametric, longer than wide |
| Cross-wall | Slightly constricted to strongly constricted | Distinctly constricted | Slightly constricted | Constricted | Slightly constricted to strongly constricted | Distinctly constricted | Distinctly constricted |
| Filaments | Solitary, immotile, forming spiral | Forming nodules in low light | Rarely forming nodules | Rarely forming nodules | Multiseriate, motile, forming nodules | Forming nodules | No formation of nodules | Rarely forming nodules |
| Attenuated | Absent | ? | ? | Tapering abruptly | ? | ? | ? | ? |
| Apical cells | Rounded | Rounded | Rounded | Rounded | Dome-shaped | Rounded | ? | ? |
| Granule | Ungranulated | Granulated | Granulated | ? | Granulated at cross walls | Granulated at cross walls | Granulated at cross walls |
| Sheaths | Very thin, colourless | Thin, colourless, occasionally becoming wide and diffusent | Often absent, thin, colourless | Soft, thin, colourless | Thin, clear | Thin, colourless, occasionally becoming wide and diffusent | Thin, colourless | Thin, colourless |
| Special features | Uniseriate trichome lie parallel or twisted around one another within a common sheath that resembles nodules | Cells typically barrel shaped after cell division, cylindrical in nondividing trichomes | Cells typically barrel shaped to spherical; inflated sheaths | Abundant nodule formation | Filament forming a tight spiral | ? | ? | ? |
| Occurrence | Soil—Signy Island, Antarctica | House wall -Peninsula Gargano, town of Vieste (Foggia), Italy. | Littoral zone—Eutrophic Lake Piaseczno, Poland | Sevilleta Long Term Ecological Research, New Mexico Soil -Chihuahuan Desert, USA | Stone monument surface—Central Mexico | Marine—South China Sea | Benthic mat in thermal spring (27 <C)—Taleshy Mahalleh, Ramsar Iran | Soil around thermal spring (32 <C)—Khaksefid, Ramsar, Iran. |

![Fig 3. TEM of *Nodosilinea signiensis*. (A), cross-section of a cell within its surrounding sheath. (B) longitudinal section of a trichome. Scale bar: 1 μm. *s* = sheath; *t* = thylakoids; *n* = nucleoplasm. Thylakoids are arranged more or less parallel in a parietal position.](https://doi.org/10.1371/journal.pone.0224395.g003)
ITS Secondary structure. The D1–D1´ helix, a semi-conserved subregion of the 16S–23S ITS region, was examined in the genus *Nodosilinea*. The putative secondary structures of the D1–D1´ helix of *N. signiensis* sp. nov., together with those of other species of *Nodosilinea*, are presented in Fig 5. The D1–D1´ helix of eight *Nodosilinea* species contained 62–64 nucleotides, and seven of them (*N. signiensis* sp. nov, *N. radiophila* TMS2B, *N. ramsarensis* KH-S S2.6, *N. chupicuarensis* PCA471, *N. epilithica* Kovacik 1998/7, *N. nodulosa* UTEX 2910, *N. bijugata* Kovacik 1986/5a) possessed a 6 nucleotide unilateral bulge with highly conserved sequence (CACUCU), and shared the same basal stem structure (GACC–GGUC) (Fig 5). Despite their similar D1–D1´ helix structures, the sequences of *N. signiensis* sp. nov. differed from those of the seven previously described species.

Discussion

Our study on a cyanobacterial strain superficially resembling *Leptolyngbya*, isolated from soils obtained on Signy Island, has resulted in the identification of a new species of *Nodosilinea*. *Nodosilinea signiensis* sp. nov. is differentiated from other previously described species based on cell size, cell shape, filament attenuation, sheath morphology, granulation and geographical distribution. Due to the simple morphology of this genus, some characteristics overlap between the species. However, the distinct cell shapes and the lack of filament attenuation separate *N. signiensis* sp. nov. from all previously described species. Cultivation clearly showed that nodule formation is only facilitated in low light conditions. Therefore, strains assigned to similar genera that have not been exposed to low light could have been misidentified. The occurrence of this species in an Antarctic terrestrial habitat further supports its separation from previously described species, based on the strikingly different biotope.
Our genetic data further indicate that *N. signiensis* sp. nov. is distinct from other species of *Nodosilinea*. The 16S rDNA phylogenetic results are congruent with morphological examination in confirming that *N. signiensis* sp. nov. is a member of the genus *Nodosilinea*. Within *Nodosilinea*, *N. signiensis* sp. nov. showed similarity values of < 98% compared with species from the basal subclade. Stackbrandt and Goebel [56] recommended that the similarity cut-off for bacterial species recognition should be 97%. Recently, Stackbrandt and Ebers [57] raised the cut-off to 98.7%, a value later supported by Yarza et al. [58]. On this basis, *N. signiensis* sp. nov. represents a species distinct from the seven previously described *Nodosilinea* species. *Leptolyngbya antarctica* is recognised as the sister clade within the genus *Nodosilinea*. Morphological study of *L. antarctica* [59] and the genetic evidence confirm that *N. signiensis* sp. nov. and *L. antarctica* are clearly distinct from each other. The evolutionary relationships of *N. signiensis* sp. nov. within the genus remain largely unclear, possibly due to the limited number of variable sites in the 16S rDNA gene. Future use of multiple genetic makers is required to fully clarify phylogenetic relationships within *Nodosilinea*.
Modelling of the secondary structure of the ITS region indicated that the structure of the D1-D1’region of N. signiensis sp. nov. was closely similar to that of other species of Nodosilinea, even though the genetic sequences were distinctly different. Previous studies have shown that analysis of 16S rDNA gene phylogeny coupled with the secondary structure of the ITS region provides a suitable tool for separation of cyanobacteria species [8, 60, 61, 62, 63]. We note that the secondary structure of the ITS region in Nodosilinea is conserved among species. This study demonstrates that the integration of other lines of evidence, from morphological and genetic sequence data, provides an effective means to improve the taxonomy of species of Nodosilinea.

Acknowledgments
We thank Dr Japareng Lalung for collection of the original soil sample on Signy Island. British Antarctic Survey staff at Signy research station are thanked for logistical and other practical support. This paper also contributes to the international SCAR ‘State of the Antarctic Ecosystem’ (AntEco) research programme. Peter Convey is supported by NERC core funding to the BAS ‘Biodiversity, Evolution and Adaptation’ Team.

Author Contributions
Conceptualization: Ranina Radzi, Faradina Merican.
Data curation: Ranina Radzi, Narongrit Muangmai, Paul Broady, Peter Convey.
Formal analysis: Ranina Radzi, Narongrit Muangmai, Paul Broady, Peter Convey.
Funding acquisition: Faradina Merican.
Investigation: Ranina Radzi.
Methodology: Ranina Radzi, Paul Broady, Peter Convey, Faradina Merican.
Resources: Wan Maznah Wan Omar.
Software: Narongrit Muangmai.
Supervision: Paul Broady, Peter Convey, Faradina Merican.
Validation: Narongrit Muangmai, Peter Convey, Faradina Merican.
Visualization: Narongrit Muangmai, Paul Broady.
Writing – original draft: Ranina Radzi.
Writing – review & editing: Narongrit Muangmai, Paul Broady, Wan Maznah Wan Omar, Sebastien Lavoue, Peter Convey, Faradina Merican.

References
1. Castenholz RW, Waterbury JB. Taxa of the cyanobacteria. Bergey’s Manual of Systematic Bacteriology. 1989; 3: 1727–1728.
2. Komárek J. A polyphasic approach for the taxonomy of cyanobacteria: principles and applications. European Journal of Phycolology. 2016; 51: 346–353.
3. Komárek J, Kaštovský J, Mareš J, Johansen JR. Taxonomic classification of cyanoprokaryotes (cyanobacterial genera) using a polyphasic approach. Preslia. 2014; 86: 295–335.
4. Mareš J. Multilocus and SSU rRNA gene phylogenetic analyses of available cyanobacterial genomes, and their relation to the current taxonomic system. Hydrobiologia. 2018; 811: 19–34.
5. Komárek J. Several problems of the polyphasic approach in the modern cyanobacterial system. Hydrobiologia. 2018; 811:7–17.
6. Abed RM, Garcia-Pichel F, Hernández-Mariné M. Polyphasic characterization of benthic, moderately halophilic, moderately thermophilic cyanobacteria with very thin trichomes and the proposal of *Halomicrocena eccentricum* gen. nov., sp. nov. Archives of Microbiology. 2002; 177: 36–70. https://doi.org/10.1007/s00203-001-0390-2 PMID: 11976745

7. Turicchia S, Ventura S, Komářková J, Komárek J. Taxonomic evaluation of cyanobacterial microflora from alkaline marshes of northern Belize. 2. Diversity of oscillatoriaceous genera. Nova Hedwigia. 2009; 89: 165–200.

8. Perkerson RB III, Johansen JR, Kovácík L, Brand J, Kaštovský J, Casamatta DA. A unique pseudanaenalean (Cyanobacteria) genus *Nodosilinea* gen. nov. based on morphological and molecular data. Journal of Phycology. 2011; 47: 1397–1412. https://doi.org/10.1111/j.1529-8817.2011.01077.x PMID: 27020364

9. Tatone A, Wilmotte A, Šmarda J, Elster J, Komarek J. *Plectolyngbya hodgsonii*: a novel filamentous cyanobacterium from Antarctic lakes. Polar Biology. 2011; 34: 181–191.

10. Dadheech PK, Mahmoud H, Kotut K, Krienitz L. *Zammitia*, *Billia*, & *Albertanoia* P. The subaerophytic cyanobacterium *Haloleptolyngbya alcalis* gen. et sp. nov., a new filamentous cyanobacterium from the soda lake Nakuru, Kenya. Hydrobiologia. 2012; 691: 269–283.

11. Zammit G, Billi D, Albertano P. The subaerophytic cyanobacterium *Oculatella subterranea* (Oscillatoriales, Cyanophyceae) gen. sp. nov.: a cytological and morphological description. European Journal of Phycology. 2012; 47: 341–354.

12. Dvořák PE, Hindák FR, Hašler PE, Hindaková A, Pouličková AL. Morphological and molecular studies of *Neosynechococcus sphagnicola*, gen. et sp. nov. (Cyanobacteria, Synechococcales). Phytotaxa. 2014; 170: 24–34.

13. Vázquez-Martínez J, Gutierrez-Villagomez JM, Fonseca-García C, Ramírez-Chavez E, Mondragón-Sánchez M. Partida-Martínez L, Molina-Torres J. *Nodosilinea chupicuarensis* sp. nov. (Leptolyngbyaceae, Synechococcales): a subaeriel cyanobacterium isolated from a stone monument in central Mexico. Phytotaxa. 2018; 334: 167–182.

14. Anagnostidis K, Komarek J. *Cyanoprokarota*. Teil 2: Oscillatoriales. *Subwasserflora von Mitteleuropa*; Band 19/2 Spektrum Akademischer Verlag Heidelberg. 2005.
27. Costa MS, Rego A, Ramos V, Afonso TB, Freitas S, Preto M, Lopes V, Vasconcelos V, Magalhaes C, Leao PN. The conifer biomarkers dehydroabietic and abietic acids are widespread in Cyanobacteria. Scientific Reports. 2016; 6: 23436. https://doi.org/10.1038/srep23436 PMID: 26996104

28. Heidari F, Hauer T, Zima J, Riahi H. New simple trichal cyanobacterial taxa isolated from radioactive thermal springs. Fottea. 2018; 18:137–49.

29. Kongisser RA. On the Methods of Studying Phytoplankton [In Russian]. Trav. Soc. Nat. Leningrad. 1925.

30. Wynn-Williams DD. Television image analysis of microbial communities in Antarctic fellfields. Polarforschung. 1988; 58: 239–249.

31. Gaydon DS, Probert ME, Buresh RJ, Meinke H, Timsina J. Modelling the role of algae in rice crop nutrition and soil organic carbon maintenance. European Journal of Agronomy. 2012; 39: 35–43.

32. Friedmann EI. Endolithic microorganisms in the Antarctic cold desert. Science. 1982; 215: 1045–53. https://doi.org/10.1126/science.215.4536.1045 PMID: 17771821

33. Quesada A, Vincent WF (2012). Cyanobacteria in the cryosphere: snow, ice and extreme cold. In Ecology of cyanobacteria II. Dordrecht: Springer; 2012. pp. 387–399.

34. Broady PA. The terrestrial algae of Signy Island, South Orkney Islands. British Antarctic Survey; 1979.

35. Friedmann EI, Hua M, Ocampo-Friedmann R. Cryptoendolithic lichen and cyanobacterial communities of the Ross Desert, Antarctica. Polarforschung. 1988; 58: 251–259. PMID: 11538357

36. Priscu JC, Fritsen CH, Adams EE, Giovannoni SJ, Paepl HK, Gordon DA. Lanolin BD, Pinckney J. Perennial Antarctic lake ice: an oasis for life in a polar desert. Science. 1998; 280: 2095–2098. https://doi.org/10.1126/science.280.5372.2095 PMID: 9641910

37. Mataloni G, Tell G, Wynn-Williams DD. Structure and diversity of soil algal communities from Cierva Point (Antarctic Peninsula). Polar Biology. 2000; 23: 205–211.

38. Cavacini P. Soil algae from northern Victoria Land (Antarctica). Polar Bioscience 2001; 14: 45–60.

39. Mueller DR, Pollard WH. Gradient analysis of cryoconite ecosystems from two polar glaciers. Polar Biology. 2004; 27: 66–74.

40. Brinkmann M, Pearce DA, Convey P, Ott S. The cyanobacterial community of polygon soils at an inland Antarctic nunatak. Polar Biology. 2007; 30: 1505–1511.

41. Vishnivetskaya TA. Viable cyanobacteria and green algae from the permafrost darkness. In Permafrost soils. Berlin, Heidelberg: Springer; 2009. pp. 73–84.

42. Stoyanov P, Moten D, Mladenov R, Dzhambazov B, Teneva I. Phylogenetic relationships of some filamentous cyanoprykaryotic species. Evolutionary Bioinformatics. 2014; EBO-S13748.

43. Jancusova M, Kovacik L, Pereira AB, Dusinsky R, Wilmotte A. Polyphasic characterisation of 10 selected ecologically relevant filamentous cyanobacterial strains from the South Shetland Islands, Maritime Antarctica. FEMS Microbiology Ecology. 2016; 92: 215001.Stoyanov P, Moten D, Mladenov R, Dzhambazov B, Teneva I. Phylogenetic relationships of some filamentous cyanoprykaryotic species. Evolutionary Bioinformatics. 2014; EBO-S13748.

44. Taton A, Grubisic S, Brambilla E, De Wit R, Wilmott A. Cyanobacterial diversity in natural and artificial microbial mats of Lake Fryxell (McMurdo Dry Valleys, Antarctica): a morphological and molecular approach. Applied and Environmental Microbiology. 2003; 69: 5157–69. https://doi.org/10.1128/AEM.69.9.5157-5169.2003 PMID: 12957897

45. Rippka R, Deruelles J, Waterbury JB, Herdman M, Stanier RY. The Journal of cell biology. 1963; 17: 208. https://doi.org/10.1083/jcb.17.1.208 PMID: 13986422

46. McDowell EM, Trump BF. Histologic fixatives suitable for diagnostic light and electron microscopy. Archives of pathology & laboratory medicine. 1976; 100: 405–414.

47. Spurr AR. A low-viscosity epoxy resin-embedding medium for electron microscopy. Journal of ultrastructure research. 1969; 26: 31–43. https://doi.org/10.1016/s0022-5320(69)90033-1 PMID: 4867011

48. Reynolds ES. The use of lead citrate at high pH as an electron-opaque stain in electron microscopy. The Journal of cell biology. 1963; 17: 208. https://doi.org/10.1083/jcb.17.1.208 PMID: 13986422

49. Boyer SL, Flechtner VR, Johansen JR. Is the 16S–23S rRNA internal transcribed spacer region a good tool for use in molecular systematics and population genetics? A case study in cyanobacteria. Molecular Biology and Evolution. 2001; 18: 1057–1069. https://doi.org/10.1093/oxfordjournals.molbev.a003877 PMID: 11371594

50. Kanehisa M, Goto S, Yato F, Furumichi M, Tanabe M. KEGG for integration and interpretation of large-scale molecular data sets. Nucleic Acids Research. 2011; 40: D109–D114. https://doi.org/10.1093/nar/gkr988 PMID: 22080510

51. McDowell EM, Trump BF. Histologic fixatives suitable for diagnostic light and electron microscopy. Archives of pathology & laboratory medicine. 1976; 100: 405–414.
52. Stamatakis A. RAxML-VI-HPC: maximum likelihood-based phylogenetic analyses with thousands of taxa and mixed models. Bioinformatics. 2006; 22: 2688–2690. https://doi.org/10.1093/bioinformatics/btl446 PMID: 16928733

53. Ronquist F, Huelsenbeck JP. MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics. 2003; 19: 1572–1574. https://doi.org/10.1093/bioinformatics/btg180 PMID: 12912839

54. Rambaut A. FigTree, version 1.3. 1. Computer program distributed by the author; 2009, http://tree.bio.ed.ac.uk/software/figtree.

55. Lowe TM, Chan PP. tRNAscan-SE Online: integrating search and context for analysis of transfer RNA genes. Nucleic Acids Research. 2016; 44: W54–7. https://doi.org/10.1093/nar/gkw413 PMID: 27174935

56. Stackebrandt E, Goebel BM. Taxonomic note: a place for DNA-DNA reassociation and 16S rRNA sequence analysis in the present species definition in bacteriology. International Journal of Systemic Evolutionary Microbiology. 1994; 44: 846–849.

57. Stackebrandt E. Taxonomic parameters revisited tarnished gold standards. Microbiology Today. 2006; 33: 152–155.

58. Yarza P, Yilmaz P, Pruesse E, Glöckner FO, Ludwig W, Schleifer KH, Rosselló-Mora R. Uniting the classification of cultured and uncultured bacteria and archaea using 16S rRNA gene sequences. Nature Reviews Microbiology. 2014; 12: 635. https://doi.org/10.1038/nrmicro3330 PMID: 25118885

59. Komárek J. Phenotype diversity of the cyanobacterial genus Leptolyngbya in the maritime Antarctic. Pol Polar Res. 2007; 28(3): 211–31.

60. Řeháková K, Johansen JR, Casamatta DA, Xuesong L, Vincent J. Morphological and molecular characterization of selected desert soil cyanobacteria: three species new to science including Mojavia pulchra gen. et sp. nov. Phycologia. 2007; 46: 481–502.

61. Johansen JR, Kovacik L, Casamatta DA, Iková KF, Kuďařovský J. Utility of 16S-23S ITS sequence and secondary structure for recognition of intrageneric and intergeneric limits within cyanobacterial taxa: Leptolyngbya corticola sp. nov. (Pseudanabaenaceae, Cyanobacteria). Nova Hedwigia. 2011; 92: 283–302.

62. Vaccarino MA, Johansen JR, Brasilolena angustatum sp. nov (Nostocales), a new filamentous cyanobacterial species from the Hawaiian Islands. Journal of Phycology. 2012; 48: 1178–1186. https://doi.org/10.1111/j.1529-8817.2012.01203.x PMID: 27011277

63. Osorio-Santos K, Pietrasik N, Bohunická M, Miscoe LH, Kováčik L, Martin MP, Johansen JR. Seven new species of Oculatella (Pseudanabaenales, Cyanobacteria): taxonomically recognizing cryptic diversification. European Journal of Phycology. 2014; 49: 450–470.