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Global distribution of cyanobacterial ecotypes in the cold biosphere

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Perennially cold habitats are diminishing as a result of climate change; however, little is known of the diversity or biogeography of microbes that thrive in such environments. Here we use targeted 16S rRNA gene surveys to evaluate the global affinities of cold-dwelling cyanobacteria from lake, stream and ice communities living at the northern limit of High Arctic Canada. Pigment signature analysis by HPLC confirmed the dominance of cyanobacteria in the phototrophic communities of these High Arctic microbial mats, with associated populations of chlorophytes and chromophytes. Microscopic analysis of the cyanobacteria revealed a diverse assemblage of morphospecies grouping into orders Oscillatoriales, Nostocales and Chroococcales. The 16S rRNA gene sequences from six clone libraries grouped into a total of 24 ribotypes, with a diversity in each mat ranging from five ribotypes in ice-based communities to 14 in land-based pond communities. However, no significant differences in composition were observed between these two microbial mat systems. Based on clone-library and phylogenetic analysis, several of the High Arctic ribotypes were found to be >99% similar to Antarctic and alpine sequences, including to taxa previously considered endemic to Antarctica. Among the latter, one High Arctic sequence was found 99.8% similar to Leptolyngbya antarctica sequenced from the Larsemann Hills, Antarctica. More than 68% of all identified ribotypes at each site matched only cyanobacterial sequences from perennially cold terrestrial ecosystems, and were <97.5% similar to sequences from warmer environments. These results imply the global distribution of low-temperature cyanobacterial ecotypes throughout the cold terrestrial biosphere.

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Introduction

Recently attention has been focused on how the Earth's atmosphere has rapidly warmed over the last decade; however, vast regions of the planet remain at temperatures near or below freezing. Extreme cold is a defining feature of High Arctic, Antarctic and high alpine sites, which are separated by large distances and climatic barriers. The ecology of these cryoenvironments is mostly microbial, and existence of a perennially cold terrestrial biosphere has implications for microbial speciation, dispersal, biogeography and gene exchange at a planetary scale. Globally dispersed microbial ecotypes have

been described from hot springs and other geothermal environments (Papke *et al.*, 2003; Bhaya *et al.*, 2007; Ward *et al.*, 2008), but microbiota at the opposite thermal extreme, cold-dwelling taxa, have received little attention.

Cyanobacteria are common throughout the terrestrial North and South Polar Regions, where they form benthic mats and films at the bottom of lakes, ponds and streams (Zakhia et al., 2007). These communities often dominate total ecosystem biomass and productivity, and must contend with persistent low temperatures, repeated freeze-thaw cycles and highly variable light, nutrient and osmotic regimes (Vincent, 2000). Filamentous, mucilage-producing Oscillatoriales are responsible for much of the biomass and three-dimensional structure of these polar mat consortia. They have been shown to tolerate a wide range of conditions and to maintain slow net growth despite the frigid ambient temperatures (Tang et al., 1997).

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Previous work on polar cyanobacteria using both morphological and molecular methods in the Polar Regions, has mostly been performed in the Antarctic, where cosmopolitan and endemic taxa are reported (Komárek, 1999; Taton et al., 2003, 2006a,b; Jungblut et al., 2005; Comte et al., 2007). By comparison, little is known about Arctic cyanobacteria, which although inhabiting a similar environment, are potentially more connected to temperate latitudes than Antarctica cyanobacteria, which are isolated by the Southern Ocean.

In the present study we evaluated the global distribution of cyanobacteria by comparing communities from the most northern reaches of North America (High Arctic Canada) with those from analogous sites in Antarctica. We determined the diversity and community structure of cyanobacterial mats collected from lakes, ponds and streams on land, and from meltwater lakes on ice shelves, at the northern limit of the North American Arctic, specifically Ward Hunt Island (latitude 83.1°N) and its vicinity in Quttinirpaaq ('top of the world' in Inuktitut) National Park, Nunavut, Canada. Cyanobacterial diversity was determined in the microbial mats by way of morphological characters, 16S rRNA gene similarity and pigment biomarkers.

Materials and methods

Study sites

The samples from Ellesmere Island in Quttinirpaaq National Park (Supplementary Figure S1), Canadian High Arctic, were taken between 8 and 15 July 2007 from the following sites: Ward Hunt Lake (WH-Lake) 83°N 05.289, 74°W 10.048; Quttinirpaag Lagoon (Q-Lagoon) 83°N 05.843, 74°W 15.018; Markham Ice Shelf (MIS) 83°N 01.898, 71°W 30.812; Ward Hunt Ice Shelf (WIS) 83°N 04.949, 74°W 26.281; Antoniades Pond (Pond-A) 82°N 58.957, 75°W 24.161 and the inflow from Lake B into Lake A (Inflow-A) 82°N 58.801, 75°W 25.372. All environmental measurements and samples were from 10- to 20-cm water depths. WH-Lake has a maximum depth of 5.5 m and a total area of 0.37 km². and is the most northern lake of North America (Villeneuve et al., 2001). The ice-free littoral zones were completely covered by cohesive microbial mats. Q-Lagoon is located between the northern coastline of Ward Hunt Island and an ice rise (thick ice on land). It has an area of 3 km², with meltwater inflows from the island and the ice rise. The shallow inshore region of the lagoon along Ward Hunt Island was covered with thick accumulation of microbial mat flakes. WIS (400 km²) is a floating mass of landfast ice approximately 40 m thick, with a ridge and trough morphology. The troughs were filled with meltwater ponds up to 15 km long, approximately 3 m deep and 10-20 m wide, with localized accumulations of sediments and loose aggregates of microbial mats ('matlets'; Mueller and Vincent, 2006). In July 2007, MIS had an area of 50 km² and a third of its surface was covered with sediment. Mat-containing sediments occurred on raised mounds of the ice and in meltwater ponds in the form of microbial matlets (Mueller *et al.*, 2006). Inflow A was approximately 10 m wide and 30 cm deep, and its submerged berm was coated with mucilaginous benthic microbial mats. Pond-A, with an area of approximately 300 m², contained thick mucilaginous orange pigmented microbial mats that covered the littoral zone.

Sampling and water analysis

Two or more samples were obtained from all the sites, except for Pond-A and Inflow-A where only single samples were taken. For the ice shelf sites, adjacent meltwater ponds were considered replicates and replicates for the lakes were from sites up to several hundred meters apart along the shorelines. After collection the samples were divided into subsamples for pigment, DNA and morphological analysis. Mat material for DNA and pigments was stored at $-80\,^{\circ}\text{C}$ and the subsamples for morphological characterization were kept in the dark at $4\,^{\circ}\text{C}$ until examination by microscopy.

Water temperature, pH and conductivity were determined at each site using a portable instrument (pH/Con 10 Series; Oakton Instruments, Vernon Hills, IL, USA). Water samples for nutrient analysis were collected from just above the microbial mats, in acid-washed bottles, and stored at 4 °C until analysis. Total nitrogen and total phosphorus were determined by standard methods (Strainton *et al.*, 1977; QuikChem 10-107-06-2-K) at Institut National de la Recherche Scientifique (Quebec City, QC, Canada).

Microscopic characterization

Cyanobacteria in the mats were examined at × 1000 magnification using an Olympus inverted light microscope (model IX71) equipped with DIC and phase contrast. Images were taken and measurements were taken using an ocular micrometer. Separation of taxa was based on morphological descriptions (Geitler, 1932; Anagnostidis and Komárek, 1988, 1990; Komárek and Anagnostidis, 1989, 1998; Villeneuve et al., 2001; Taton et al., 2008).

Pigment analysis

Total pigments from subsamples were extracted in the dark by grinding the frozen material for 2 min followed by sonication (3 \times 20 s at 20 W) in 4 or 6 ml 90% acetone:water (vol/vol) mixture, and left overnight at $-20\,^{\circ}\text{C}$ under an argon gas atmosphere. The extracts were recovered following centrifugation at 4150 r.p.m. for 15 min at 4 $^{\circ}\text{C}$. The supernatant was then filtered through a 0.2 μ m pore size PTFE Acrodisc filter (Pall Corporation, Ann Arbor, MI, USA) and stored in the dark at $-70\,^{\circ}\text{C}$ under an argon atmosphere until high-performance liquid chromatography (HPLC) analysis. This extraction procedure was repeated for the residual material

until no further coloration was detected in the extract solution. HPLC analysis was performed on 50 µl of injected sample using a ProStar HPLC system (Varian, Palo Alto, CA, USA) equipped with a Symmetry C8 column (3.5 μ m pore size, 4.6 \times 150 mm; Waters Corporation, Milford, MA, USA) at 25 °C with a C8 guard column (5 μm pore size, $3.9 \times 20 \,\mathrm{mm}$; Waters Corporation). The HPLC peaks were detected by diode-array spectroscopy (350-750 nm). Absorbance chromatograms at 384 nm (for scytonemin), 440 nm (for chlorophylls) and 450 nm (for carotenoids) were recorded. Chlorophylls were also detected by fluorescence (excitation, 40 nm; emission, 650 nm). The HPLC solvent protocol was based on gradient dilution with two solvent mixtures (Zapata et al., 2000). The flow rate was $1 \,\mathrm{ml}\,\mathrm{min}^{-1}$, with an equilibrium time of $5 \,\mathrm{min}$. Standards (chlorophyll (chl.)-a, b and c_2 ; β , β carotene, canthaxanthin, diadinoxanthin, echinenone, fucoxanthin, lutein, myxoxanthophyll and zeaxanthin) were obtained for identification and quantification of detected pigments (Sigma Inc., St Louis, MO, USA; DHI Water & Environments, Hørsholm, Denmark; Bonilla et al., 2005). Other carotenoids were quantified based on the published extinction coefficients of related pigments: HFUlike, $142 \,\mathrm{lg^{-1}\,cm^{-1}}$; uriolide-like, $166 \,\mathrm{lg^{-1}\,cm^{-1}}$; 19'-hexanoylofucoxanthin-like, 142 lg⁻¹ cm⁻¹; peridinin-like, $136 \lg^{-1} cm^{-1}$; astaxanthin, $210 \lg^{-1} cm^{-1}$; antheraxanthin-like, 244 l g⁻¹ cm⁻¹; diatoxanthin, 210 lg⁻¹ cm⁻¹ and monadoxanthin-like, 250 lg⁻¹ cm⁻¹. The extinction coefficient of β , β -carotene (250 l g^{-1} cm⁻¹) was used for unknown carotenoids (Jeffrey et al., 1997) and scytonemin was quantified as described by Garcia-Pichel et al. (1992).

DNA extraction

On return from the field, the mat material for DNA analysis was freeze-dried and suspended in 800 µl XSbuffer (1% potassium-methyl-xanthogenate; 800 mM ammonium acetate; 20 mm EDTA; 1% sodium dodel sulfate: 100 mm Tris-HCl (pH 7.4); Tillett and Neilan, 2000). The mixture was vortex-mixed and incubated at 65 °C for 4 h. The extracts were cooled overnight at −20 °C and cell debris were removed by centrifugation at 12000 g for 10 min. An equal volume of phenolchloroform-isoamyl alcohol (25:24:1) was then added to the removed aqueous phase and centrifuged at 3000 g for 3 min. The two steps were repeated twice. DNA was precipitated overnight by addition of 1 volume of isopropanol and 1/10 volume of 4M ammonium acetate at -20 °C. The precipitated DNA was pelleted by centrifugation at 12 000 g for 10 min and washed with 70% ethanol. The extracted DNA was then resuspended in 100 µl of sterile water.

PCR

All PCR reactions were performed using Advantage 2 PCR kits with proof-reading ability, in a 20 μl reaction mix using 10 × Advantage 2 SA PCR buffer and 0.2 mm dNTPs (Fermentas, Foster City, CA. USA), according to the manufacturer (Clontech, Mountain View, CA, USA). PCR amplification of cyanobacterial 16S rDNA was performed using 0.5 µM of each cyanobacteria-specific primer 27F1 (5'-AGAGTTTGATCCTGGCTCAG-3') and 809R (5'-G CTTCGGCACGGCTCGGGTCGATA-3'). As described by Jungblut et al. (2005), these primers provide broad coverage of cyanobacterial taxa.

Cloning, RFLP analysis and sequencing

Prior to cloning, the amplified PCR products were verified by gel electrophoresis and amplicons of the target size were purified with the Qiaquick PCR Purification kit (Qiagen, Mississauga, CA, USA). PCR products were cloned using the TOPO TA cloning kit (Invitrogen, Carlsbad, CA, USA). Ligation and transformation were performed according to the manufacturer's protocols. Positive clones were transferred to 96-well plates containing Luria Bertani medium with 7% glycerol. The inserted 16S rRNA sequences were amplified using vector-specific primers M13f and M13r, and subjected to restriction-fragment length polymorphism (RFLP) screening. Amplicons (4 µl) were digested (overnight in separate incubations with 5 U of restriction enzymes AluI and HpaII; Fermentas, Hanover, NH, USA) in a final reaction volume of 10 µl with the appropriate buffer at 37 °C. The resulting digests were run on 2.5%, low-melting point agarose gel and the generated RFLP patterns were visualized using the Bio-Rad Laboratories Gel Doc imaging system and Quantity One software (Bio-Rad Laboratories, Hercules, CA, USA, version 4.5.1). At least two clones for each unique RFLP pattern were sequenced using the vector-specific $\mathsf{T7}$ universal primer (single read) at the Centre Hospitalier de l'Université Laval (CHUL, QC, Canada), using an ABI 3730xl system (Applied Biosystems, Foster City, CA, USA), which included a purification step.

Total-mat-community RFLP

Communities were compared using direct RFLP analysis of PCR products from total genomic DNA using the same restriction enzymes as above. Five units of AluI and HpaII (Fermentas) were added to 6 μl of PCR product for a final volume of 10 μl. The generated RFLP patterns were run on 2.5%, lowmelting point agarose gel and analyzed as described above. The RFLP community patterns were compared using Restdist and Neighbor in PHYLIP version 3.67 (Felsenstein, 1989).

Phylogenetic analysis and diversity calculations All sequences were checked for chimeras using the Chimera check program at Ribosomal Data Project II (Maidak et al., 2001) and they were excluded from



further analysis. Sequences were edited and trimmed using 4Peaks (version 1.7). The approximately 750-nt sequences were aligned using ClustalX (version 1.8; Thompson et al., 1994) and were checked manually using BBEdit Lite (version 6.1). Reference sequences were from GenBank and for each phylotype the closest match based on a BLAST search (Altschul et al., 1990) of GenBank was selected as a reference sequence. If the closest match was an uncultured clone, we also included the closest isolated strain. For comparisons, we also searched for environmental 16S rDNA sequence data from other Arctic and Antarctic sites (Priscu et al., 1998; Nadeau et al., 2001; Taton et al., 2003, 2006a, b; Jungblut et al., 2005).

The genetic differences between the cyanobacterial communities from the clone-libraries were calculated using Unifrac (Lozupone *et al.*, 2006). Each pair of environments was compared with a weighted Unifrac matrix that takes abundances of different sequences into account using Unifrac Significance test.

Phylogenetic trees were constructed using neighbor-joining with the Kimura-2-Parameter distance matrix (DNAdist, Neighbor) and maximum-likelihood (DNAml) was computed with PHYLIP (version 3.67, 19). Aligned partial 16S rRNA gene sequences corresponding to Escherichia coli sequence positions 129-775 were used. Confidence levels were calculated for each method by bootstrapping with 1000 and 100 reassembly events for neighbor-joining and maximum-likelihood, respectively (Seqboot, Consense). One representative for each ribotype was included in the phylogenetic analysis, and individual ribotypes or Operational Taxonomic Units (OTUs) were defined as groups of sequences, which were at least 97.5% similar (Stackebrandt and Göbel, 1994; Taton et al., 2003).

Library coverage, the Shannon-Wiener diversity index (H'), Chao1 non-parametric richness estimates and rarefaction curves were calculated using DO-TUR (Schloss and Handelsman, 2005) on a Juke-Cantor distance matrix. 16S rRNA gene sequences are available under GenBank accession numbers FJ977098-FJ977164 (Supplementary Table S1).

Results

Environmental properties

The six collection sites spanned a range of environmental conditions, with overlying water temperatures from 0.9 °C in the WIS meltwater ponds to 6 °C in Pond-A (Supplementary Table S2). Pond-A had the highest pH among all the sites (8.28), and the lowest pH values were recorded in the meltwater ponds of WIS (6.45) and MIS (6.53). Highest conductivities were at the ice-based sites, with 637, 384.8 and 269.0 $\mu S \, cm^{-1}$ on MIS, WIS and Q-Lagoon, respectively. Land-based sites had conductivities of 137 $\mu S \, cm^{-1}$ or less. Nutrient concentrations were highest in Pond-A, with 0.961 mg l $^{-1}$ total nitrogen

and $0.016\,\mathrm{mg}\,l^{-1}$ total phosphorus. Total nitrogen concentrations were similar between the two iceshelf sites ($0.149\,\mathrm{mg}\,l^{-1}$ for MIS and $0.156\,\mathrm{mg}\,l^{-1}$ for WIS), whereas total phosphorus concentrations were 0.009 and $0.014\,\mathrm{mg}\,l^{-1}$, respectively. The lowest nutrient concentrations were in Q-Lagoon and WH-Lake, with 0.033 and $0.089\,\mathrm{mg}\,l^{-1}$ total nitrogen, and 0.004 and $0.003\,\mathrm{mg}\,l^{-1}$ of total phosphorus, respectively.

Pigment diversity

Each microbial community contained diverse pigments, including chlorophylls, scytonemins, carotenoids and their degradation products (Supplementary Table S3). Chl.-a concentrations ranged from $3.9 \,\mu \text{g cm}^{-2}$ (Pond-A) to $42.6 \,\mu \text{g cm}^{-2}$ (WIS). Chl.-b was identified in all the sites except WH-Lake, with concentrations of $3.5 \,\mu\mathrm{g}\,\mathrm{cm}^{-2}$ or less, whereas chl.-c was only identified in Pond-A (0.5 µg cm⁻²). The cyanobacterial pigment scytonemin and its reduced derivative, red-scytonemin, were the most abundant pigments in mats from WH-Lake, WIS and MIS, with concentrations of up to 474.8 µg cm⁻² (WIS). Low concentrations of scytonemin were detected in Pond-A, with 0.28 µg cm⁻², and none in Q-Lagoon. High concentrations of the carotenoids zeaxanthin, echinenone, β-carotene and a lutein-like carotenoid were present in all mat samples. The pigments canthaxanthin, fucoxanthin, 19'-hexanoylofucoxanthin and 4ketomyxol-like carotenoid were only separated in some of the microbial mats, with diatoxanthin; astaxanthin and diadinoxanthin-like, antheraxanthinlike, monadoxanthin-like, peridinin-like carotenoids identified only in Pond-A mats.

Morphological classification

Microscopic analyses confirmed that cyanobacteria constituted the greatest proportion of biomass in all of the High Arctic communities. Based on morphological criteria, they were found to be composed of taxa within orders Chroococcales, Nostocales and Oscillatoriales (Supplementary Table S4). Five known Chroococcales genera were identified (Gloeocapsa cf. alpina, Chroococcus cf. prescotii, Chlorogloea, Aphanocapsa cf. hyalina and Merismopedia cf. angularis), along with one unclassified coccoid morphotype. Genera within order Nostocales included Nostoc, Dichothrix and Tolypothrix, and within the Oscillatoriales the identified genera were Leptolyngbya (Leptolyngbya cf. frigida), Pseudanabaena (Pseudanabaena cf. amphigranulata), Phormidium (Phormidium autumnale) and Oscillatoria (Oscillatoria sancta). Overall we distinguished six Chroococcales, five Nostocales and 13 Oscillatoriales based on classical morphological characters.

Cyanobacterial 16S rRNA gene analysis

We constructed targeted 16S rRNA gene clone libraries using genomic environmental DNA from

all the sites, vielding a total of 426 clones with the correct insert. Initial RFLP analysis showed that there were much larger differences among sites than between duplicate samples from the same site (Supplementary Figure S2). The highest diversities were from the three land-based sites, with 12-14 OTUs, defined as >97.5\% similarity, per site. Similarly, Chao statistics of the land-based sites ranged from 21.7 to 34.7, while the WIS mats contained only five OTUs (Table 1). However, the

six cyanobacterial communities did not differ in

pairwise comparisons in a weighted Unifrac matrix

(Lozupone *et al.*, 2006).

Six OTUs were from order Chroococcales (Figure 1 and Table 2). Three of these were most similar to cultured representatives: clone ArC22 was up to 99.1% similar to Synechococcus sp. PCC 7502 (AF448080); ArC20 98.5% to Chamaesiphon subglobosus PCC 7430 (AY170472) and ArC19 98.8% to Snowella litoralis 1LT47S05, AJ781041). The novel ribotype, ArcC21, had highest similarity of 93.5-93.8% to Gloeothece sp. SK40 (AB067576). Two other ribotypes had 93.1-93.4% (ArC23) and 95.8% (ArC24) similarity to uncultured Gloeobacter sp. HAVOmat17 (EF032784).

Three of the OTUs were within order Nostocales, and one within Stigonematales (Figure 2 and Table 2). Nostocales had the highest sequence similarity to cultured *Nostoc* spp., including *Nostoc* commune KU002 (ArcC17, 98.2-98.4% similarity to AB088375) and *Nostoc* sp. PCC 7906 (ArcC18, 97.7% similarity to AB325908), and therefore were conservatively classified as cosmopolitan ribotypes. OTU ArcC16 had less than 94.9% sequence similarity to Stigonema ocellatum SAG 48.90 (AJ544082) and appears to be a novel phylotype within order Stigonematales.

Fifteen of the OTUs grouped within order Oscillatoriales (Figure 2 and Table 2). Eleven of these had highest similarities (97.5% or usually >99) to sequences from cold environments and seven OTUs (ArcC04–07 and ArcC11–13) were grouped with clones or strains previously identified solely from Antarctic microbial communities, including the Vestfold and Larsemann Hills (East Antarctica), McMurdo Ice Shelf, Lake Fryxell and Lake Bonney (McMurdo Dry Valleys) (Priscu et al., 1998; Taton et al., 2003; Jungblut et al., 2005). At five of the Arctic sites, these clones made up between 20.5 and 70% of the total diversity (Figure 3). The second set of OTUs included ArcCo1, 08 and 15, and was grouped with sequences from Antarctic and other cold environments (glaciers and glacial surface snow, Kuytun Glacier 51, Tian Shan Mountains, China; AY493581, AY151728, DQ181742). This set accounted for 12.0–94.4% of the total diversity. The oscillatorian ribotype ArcC02 shared the highest similarity with an environmental sequence from Kuytun Glacier 51 (surface snow, China; EU263766) and had a relative abundance of 1.3% in one of the High Arctic sites. All of these three categories of phylotypes were classified as cold ecotypes since they have only been reported from cold habitats to date.

Discussion

Phototrophic community diversity

Polar cyanobacteria withstand the extremes of their environment through production of photoprotective screening and quenching pigments, as well as by their highly efficient light-capturing systems, nutrient storage ability and freeze-thaw tolerance (Hawes and Schwarz, 2001; Zakhia et al., 2007). The HPLC pigment signatures of the High Arctic assemblages that we sampled in the present study provided semi-quantitative information on phototrophic community composition. Chl.-a concentrations in WH-Lake and WIS were similar to that in earlier reports (Bonilla et al., 2005; Mueller et al., 2005, 2006). Previous studies reported higher concentrations of chl.-a in mats from WIS and MIS than

Table 1 Diversity indices, coverage, number of clones and OTUs for the six microbial mat communities

	WH-Lake	A-Pond	Inflow-A	Q-Lagoon	MIS	WIS
Number of OTUs	12	14	12	7	10	5
Shannon index	1.6	2.0	1.8	1.3	1.8	0.7
Chao1 (max.)	21.7	21.7	34.7	ND	15.3	ND
Coverage	91.3	94.9	88	98.6	98.5	98.9
Number of clones	69	78	50	74	66	89
Ribotype assemblage						
Antarctic ribotypes (%)	40.6	20.5	70.0	62.2	51.5	4.5
Antarctic and non-polar ribotypes (%)	27.5	60.3	12.0	18.9	39.4	94.4
Non-polar, cold ribotypes (%)	0.0	1.3	0.0	0.0	0.0	0.0
Total cold ecotypes (%)	68.1	83.3	82.0	81.1	90.9	98.9
Non-polar ribotypes (%)	2.9	11.5	10.0	18.9	9.1	1.1
Novel ribotypes (%)	29.0	6.4	8.0	0.0	0.0	0.0

Abbreviations: Inflow-A, inflow from Lake B into Lake A; MIS, Markham Ice Shelf; OUT, Operational Taxonomic Unit; Pond-A, Antoniades Pond; Q-Lagoon, Quttinirpaaq Lagoon; WH-Lake, Ward Hunt Lake; WIS, Ward Hunt Ice Shelf. ND: Chao1 could not be determined due to low OTU diversity.



 Table 2
 OTUs defined from the six clone libraries of Arctic microbial mats

OTU ribotype	OTU ribotype Similarity (%) Highest match (accession nur	Highest match (accession number)	Similarity %	Highest cultured match (accession number)	Ecotype
ArcC 01 ArcC 02 ArcC 03 ArcC 03 ArcC 05 ArcC 06 ArcC 09 ArcC 10 ArcC 11 ArcC 11 ArcC 11 ArcC 12 ArcC 14 ArcC 15 ArcC 17 ArcC 12 ArcC 12 ArcC 21 ArcC 22 ArcC 22 ArcC 22 ArcC 22	99.6–99.9 98.7 99.2 97.9 97.5–98.7 97.5–97.7 99.1–99.9 94.1–94.8 98.9–99.2 ————————————————————————————————————	Uncultured bacterium clone KuyT-water-120 (EU263787) Uncultured bacterium clone KuyT-ice-23 (EU263766) Uncultured bacterium clone KuyT-ice-32 (EU263774) Uncultured cyanobacterium clone RJ088 (DQ181681) Uncultured cyanobacterium clone Fr121 (AY151728) Uncultured cyanobacterium clone Fr121 (AY151728) Uncultured cyanobacterium clone RD107 (DQ181677) Uncultured cyanobacterium clone H-B02 (DQ181686) Uncultured cyanobacterium clone H-B02 (DQ181686) Uncultured cyanobacterium clone H-B02 (DQ181686) Uncultured cyanobacterium clone H-D15 (DQ181742)	97.9–98.2 96.3 90.9 97.6–98.3 98.8–99.6 91.6 99.3 93.5–94.2 95.7–97.0 91.7–92.1 99.5–99.8 91.7–92.1 99.5–99.8 91.7–91.8 91.7–92.1 99.5–99.8 91.7–91.8 91.7–91.8 91.7–91.8 91.7–91.8 91.7–91.8 91.7–91.8 91.7–91.8 91.7–91.8 91.7–91.8 91.7–91.8	Phormidium priestleyi ANT.LH66.1 (AY493581) Limnothrix redekei CCAP 1443/1 (LRE580007) P. priestleyi ANT.LH52.6 (AY493579) P. priestleyi ANT.LH52.6 (AY493579) P. priestleyi ANT.PROGRESS2.6 (AY493585) P. priestleyi ANT.PROGRESS2.6 (AY493585) P. priestleyi ANT.PROGRESS2.6 (AY493585) Phormidium sp. SAG 37.90 (EF654082) L. frigida ANT.LH52.2 (AY493575) cf. Leptolyngbya sp. Greenland 9 (DQ431004) Pseudanabaenaceae cyanobacterium DPG1-KK5 (EF654067) Pseudanabaenaceae cyanobacterium DPG1-KK5 (EF654067) Pseudanabaenaceae cyanobacterium DPG1-KK5 (EF654067) Pseudanabaenaceae cyanobacterium DPG1-KK5 (EF654067) L. frigida ANT.LH0 (AY493607) L. frigida ANT.LH18.1 (AY493607) Cf. Leptolyngbya antarctica ANT.LH18.1 (AY493607) Cf. Leptolyngbya sp. Greenland 9 (DQ431004) Stigonema ocellatum SAG 48.90 (SOC544082) Nostoc commune KU002 (AB088375) Nostoc sp. PCC 7906 (AB325908) Snowella litoralis 1LT47805 (AJ781041) Chamcesiphon subglobosus PCC 7430 (AY170472) Gloeothece sp. SK40 (AB067576) Synechococcus sp. PCC 7502 (AF448080) Gloeobacter violaceus PCC 7421 (BA000045)	Cold Cold Cold Cold Cold Cold Cold Cold

Abbreviation: OTU, Operational Taxonomic Unit. Highest cultured match was also included when the highest BLAST match was an uncultured clone from GenBank.



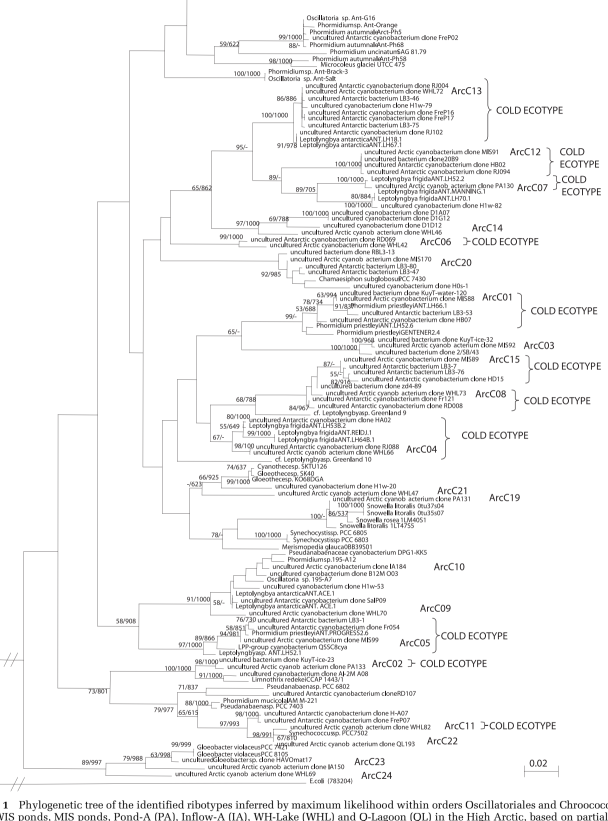


Figure 1 Phylogenetic tree of the identified ribotypes inferred by maximum likelihood within orders Oscillatoriales and Chroococcales from WIS ponds, MIS ponds, Pond-A (PA), Inflow-A (IA), WH-Lake (WHL) and Q-Lagoon (QL) in the High Arctic, based on partial 16S rRNA gene analysis. Bootstrap values based on maximum likelihood (bold) and neighbor-joining methods are indicated at the nodes when equal or greater than 50%; * indicates where Figure 2 joins Figure 3. Inflow-A, inflow from Lake B into Lake A; MIS, Markham Ice Shelf; Pond-A, Antoniades Pond; Q-Lagoon, Quttinirpaaq Lagoon; WH-Lake, Ward Hunt Lake; WIS, Ward Hunt Ice Shelf.

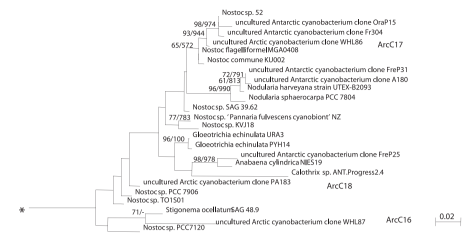


Figure 2 Phylogenetic tree of the identified ribotypes inferred by maximum likelihood within orders Nostocales and Stigonematales from WIS, MIS, Pond-A (PA), Inflow-A (IA), WH-Lake (WHL) and Q-Lagoon (QL) in the High Arctic, based on partial 16S rRNA gene analysis. Bootstrap values based on maximum likelihood (bold) and neighbor-joining methods are indicated at the nodes when equal or greater than 50%; * indicates where Figure 3 joins this figure. Inflow-A, inflow from Lake B into Lake A; MIS, Markham Ice Shelf; Pond-A, Antoniades Pond; Q-Lagoon, Quttinirpaaq Lagoon; WH-Lake, Ward Hunt Lake; WIS, Ward Hunt Ice Shelf.

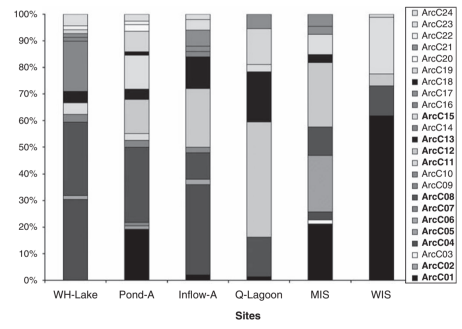


Figure 3 Percentage abundance of ribotypes in the cyanobacterial communities of WH-Lake, Pond-A, Inflow-A, Q-Lagoon, MIS and WIS. Ribotypes are highlighted in bold. Inflow-A, inflow from Lake B into Lake A; MIS, Markham Ice Shelf; Pond-A, Antoniades Pond; Q-Lagoon, Quttinirpaaq Lagoon; WH-Lake, Ward Hunt Lake; WIS, Ward Hunt Ice Shelf.

from Antarctic microbial ice-shelf mats (Howard-Williams et al., 1989, 1990; Mueller et al., 2005). Cyanobacteria-specific markers, such as scytonemin, echinenone, zeaxanthin and canthaxanthin, dominated the microbial mats in WIS, MIS, WH-Lake and Q-Lagoon, consistent with cyanobacterial dominance of the total phototrophic biomass (Vincent et al., 2004; Bonilla et al., 2005; Mueller et al., 2005), and similar markers have been reported for Antarctic microbial mats (Vincent et al., 1993). In

addition, in some of the mats high concentrations of red-scytonemin, a reduced product of scytonemin, were detected; scytonemin and red-scytonemin are sheath pigments, which protect cyanobacterial cells against UV-A radiation.

Chl.-b and lutein, pigments specific to Chlorophyta, were identified in most microbial mats. These pigments could potentially be associated with the genera Mougeotia, Zygnema and Cosmarium, which were identified morphologically in samples



from WH-Lake by Villeneuve et al. (2001). Similarly the Chlorophytes Chlorosarcinopsis, Chlamydomonas, Chlamydocapsa and Chlorella have been reported previously from WIS (Mueller et al., 2005). In Pond-A, the most abundant and diverse pigments were specific for Chromophyta, including chl.-c2, diadinoxanthin-like, diatoxanthin, astaxanthin, monadoxanthin-like and fucoxanthin. The second highest concentrations of pigments were characteristic for Chlorophyta in Pond-A, and cyanobacteriaspecific signatures were also identified in Pond-A, however at lower concentrations. Diatoxanthin and fucoxanthin are common in diatoms in particular (Jeffrey et al., 1997). Chromophyta-specific pigments were also identified in the other microbial mats, however at lower concentrations. Pond-A temperatures were high compared with that in the other sites, and nutrient concentrations were elevated, potentially due to enrichment by a population of aquatic birds (red-throated loons, Gavia stellata) that we observed at this site.

Morphological diversity

As in other Arctic and Antarctic freshwater ecosystems, mat-forming cyanobacteria were the most conspicuous members of the well-developed benthic communities. Light-microscopy results were similar to that of previous studies of microbial mat communities from the High Arctic (Bonilla et al., 2005; Mueller et al., 2005). The microbial mat communities were made up of morphospecies within orders Oscillatoriales, Nostocales and Chroococcales, and were similar to Antarctic microbial mats (Howard-Williams et al., 1989; Taton et al., 2003, 2006a,b; Jungblut et al., 2005). Morphospecies in Oscillatoriales were the most abundant taxa at all the sites, followed by those in Chroococcales and Nostocales. In particular, morphospecies related to *Leptolyngbya*, Pseudanabaena, Phormidium, Oscillatoria and Nostoc are characteristic of polar mats and form their overall structure (Vincent, 2000). The morphological diversity of Chroococcales was similar to freshwater ponds in the Larsemann and Vestfold Hills region: Antarctica, however analogous communities on the McMurdo Ice Shelf and in the McMurdo Dry Valleys, lacked any Chroococcalean morphotypes (Taton et al., 2003; Jungblut et al., 2005).

Interestingly, we did not find *Nodularia* at any of the Arctic sites, even though it has been described regularly for Antarctic microbial mats, in particular in the McMurdo Ice Shelf, McMurdo Dry Valleys and Larsemann and Vestfold Hills (Taton et al., 2003, 2006a; Jungblut et al., 2005). In contrast, sequences related to Gloeobacter, as found here in the High Arctic, have never been reported from Antarctica. All of these closest matches are to ribotypes from temperate climatic zones, suggesting the connectivity of Arctic environments to lower latitudes. These findings contrast with data on Antarctic mats from the McMurdo Region, which are conspicuously lacking in Chroococcales (Taton et al., 2003; Jungblut et al., 2005).

Biogeography of polar cyanobacteria

The Polar Regions offer ideal sites for testing microbial endemism since they contain parallel environments separated by vast geographical distances and potential barriers to dispersal (Staley and Gosink, 1999). Many bacteria and microbial eukaryotes have been identified as possibly endemic to Antarctica, including several cyanobacterial species (Komárek, 1999; Taton et al., 2006b). However, our clone-library analyses indicate that three taxa previously identified as Antarctic endemics (Phormidum priestleyi Fritsch, L. frigida (Fritsch) Anagn. and Kom., and Leptolyngbya antarctica (West and West) Anagn. and Kom.; Komárek, 1999; Taton et al., 2006b) were more than 99% similar to sequences from the Canadian High Arctic (Table 2); for example, ArC05 is 99.6% similar to P. priestlevi (ANT.PROGRESS2.6; AY493585) and ArC13 is 99.8% similar to *L. antarctica* (ANT.LH18.1; AY493607). Furthermore, several of the uncultured cvanobacterial clones from East Antarctica and the McMurdo Dry Valleys identified as endemic, had the highest percentage match, up to 99.9%, to some of our High Arctic sequences. Similarly, clonelibrary analysis of high-altitude saline wetland mats included a 99% match to L. frigida (ANT.LH701, and L. antarctica (ANT.LH18.1, AY493574) AY493607) based on partial 16S rRNA gene analyses (Dorador et al., 2008). Nadeau et al. (2001) previously reported that within another clade of Antarctic Oscillatoriales, there was an 11-bp insertion earlier found in a Svalbard soil isolate, which implied a shared evolutionary history.

In sum, these findings suggest the presence of cold-habitat-specific cyanobacterial assemblages, with individual ribotypes that are up to 99.9% similar in the Arctic and Antarctic, and conspicuously absent from other climate zones. Molecularclock analysis of several bacterial taxa suggests that a 1% divergence in 16S rRNA gene sequence corresponds to an evolutionary time span of approximately 50 million years (Moran et al., 1993; Ochman et al., 1999). This would imply that the Arctic and Antarctic ribotypes described here have been isolated or subject to reduced genetic exchange for less than 10 million years. Cyanobacteria isolated from cold environments all have temperature optima growth rates in the range 15-20 °C, suggesting that they likely had their evolutionary origins within temperate latitudes (Tang et al., 1997; Nadeau et al., 2001) and subsequently colonized perennial cold habitats.

Additional analyses using the ITS region (Comte al., 2007), multi-locus sequence analyses (Whitaker et al., 2003) and broader genomic and metagenomic analyses are needed to determine whether cold-dwelling oscillatorians belong to



narrow ecotypes, analogous to the Synechococcus ecotypes from geothermal springs (Bhaya et al., 2007) and N. commune in Antarctica (Novis and Smissen, 2006). An ecotype may be defined as a group of ecologically similar cyanobacteria, with genetic diversity within the ecotype limited by a cohesive force, either periodic selection or genetic drift, or both (Cohan and Perry, 2007), where in our case the environmental force is extreme cold. This corresponds to high-latitude and high-altitude regions where growth of higher plants is severely limited and temperatures are near-zero in summer (Thomas et al., 2008). At least on a 16S rRNA gene level, cyanobacteria from these cold regions are more related to each other than to those in the temperate groups.

Our molecular findings suggest that microbiota of the cryosphere have been globally distributed with local habitat selection (Baas-Becking, 1934; Finlay and Fenchel, 2004). This could occur via mechanisms of long-range transport, similar to atmospheric studies documented for microbes, such as bacteria in Saharan dust transported over the Atlantic (Griffin et al., 2002; Gorbushina et al., 2007), and across Antarctica and the Southern Hemisphere (Hughes et al., 2004; Muñoz et al., 2004). Short-term exchange between the Arctic and Antarctica may be favored by seasonal oscillation of the Hadley cells, which contributes to inter-hemisphere mixing in the troposphere, as revealed by model analysis of long-lived tracers (Bowman and Cohen, 1997).

The present day distribution may be accentuated over longer time scales (Cermeño and Falkowski, 2009) via global freeze-up events such as the Precambrian glaciations (Kirschvink *et al.*, 2000) and dispersal of microbiota throughout the cold biosphere. More recent glacial events may have also favored genetic exchange between the Polar Regions, as suggested for cold-water foraminifers (Darling *et al.*, 2000), although such cooling could also lead to isolation and divergence of some populations (Darling *et al.*, 2004), and dinoflagellates (Montresor *et al.*, 2003).

The dispersal of low-temperature ecotypes may differ from those in other extremes, for example geothermal hot-spring cyanobacteria (Papke et al., 2003; Souza et al., 2008) and hyperthermophiles such as Sulfolobus (Whitaker et al., 2003) that occupy much more localized as well as distantly separated habitats. Furthermore, cold-adapted cyanobacteria are well equipped to withstand potential nutrient limitations, temperature fluctuation, dehydration and elevated UV radiation during longdistance aerial transport. As a result, cold-adapted cyanobacteria may show much reduced genetic divergence in comparison with the known degree of diversification of microbial taxa at the other thermal extreme; for example, Sulfolobus endemism in hot springs (Whitaker et al., 2003). Global circulation models currently predict accelerated

warming and massive contraction of glacial environments over the next few hundred years (IPCC, 2007), which may force the cold ecotypes identified here into similarly localized habitats or extinction.

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