Global phylogeography of pelagic *Polynucleobacter* bacteria: Restricted geographic distribution of subgroups, isolation by distance and influence of climate†

Martin W. Hahn,1* Ulrike Koll,1 Jitka Jezberová1,2 and Antonio Camacho3

1Research Institute for Limnology, University of Innsbruck, Mondseestrasse 9, A-5310 Mondsee, Austria.
2Biology Centre of the ASCR, v.v.i., Institute of Hydrobiology, České Budějovice, Czech Republic.
3Cavanilles Institute for Biodiversity and Evolutionary Biology, University of Valencia, Burjassot, Spain.

Summary

The free-living planktonic freshwater bacterium *Polynucleobacter necessarius* subspecies *asymbioticus* (>99% 16S rRNA similarity) represents a taxon with a cosmopolitan distribution and apparently ubiquitous occurrence in lentic freshwater habitats. We tested for intra-taxon biogeographic patterns by combining cultivation-independent and cultivation methods. A culture collection of 204 strains isolated from globally distributed freshwater habitats (Arctic to Antarctica) was investigated for phylogeographic patterns based on sequences of two markers, the 16S–23S internal transcribed spacers and the glutamine synthetase gene (*glnA*). Genetic distance between isolates showed significant geographic distance-decay patterns for both markers, suggesting that an isolation-by-distance mechanism influences the global phylogeography. Furthermore, a couple of subgroups showed restricted geographic distributions. Strains of one subgroup were exclusively obtained from tropical sites on four continents (pantropical subgroup). Cultivation-independent methods were used to confirm the restricted geographic distributions of two subgroups. The pantropical taxon could be detected in 63% of investigated tropical habitats but not in any of 121 European freshwater samples. Physiological tests indicated that almost all strains of the pantropical subgroup failed to grow at temperatures of 4°C, while strains affiliated with other subgroups showed good growth at this temperature. This suggests that thermal adaptation is involved in phylogeographic structuring of the global *Polynucleobacter* population.

Introduction

An increasing number of studies suggests that the distribution of many microbial taxa shows biogeographic patterns (Bass and Boenigk, 2011; Fontaneto and Brodie, 2011; Hanson et al., 2012), but not much is known about the underlying mechanisms. Basically, such biogeographic patterns can result from barriers to dispersal of organisms (i.e. limited gene flow in combination with mutation, selection and/or drift, which causes divergence of populations) (Hanson et al., 2012). Alternatively, these patterns may simply reflect geographic distribution patterns of site-specific environmental conditions (Tamames et al., 2010), such as climate-determined features, habitats or host species suitable for maintaining populations of microbial taxa. It is not doubted that such environmental selection (filtering) of taxa by site-specific ecological conditions is significant in shaping microbial distribution patterns (Baas Becking, 1934; Crump et al., 2004; Gray et al., 2007). The role of dispersal barriers in biogeography of microorganisms is, however, controversial. A school of microbiologists argues that at least free-living (not host-associated) microorganisms differ from macroorganisms in the lack of any dispersal barriers or limitations (Baas Becking, 1934; Finlay, 2002; de Wit and Bouvier, 2006). This hypothesis is partially supported, for instance, by observations of long-distance or even intercontinental transport of microorganisms (Schlichting et al., 1978; Gorbushina et al., 2007; Hervás et al., 2009; Pearce et al., 2009; DeLeon-Rodriguez et al., 2013). However, successful colonization of new habitats by microorganisms is a two-step procedure consisting of dispersal to and establishment of a population in the new...
habitat. The latter requires viable cells to be able to arrive (Gorbushina et al., 2007; Hervás et al., 2009), to grow under the local environmental conditions (Hervás et al., 2009) and to compete successfully with potential local inhabitants. Successful dispersal of microorganisms may thus be limited by pre- and post-colonization barriers (Marshall et al., 2010). Several investigations suggested that pre-colonization (dispersal) barriers exist for some microbial taxa (Papke et al., 2003; Whitaker et al., 2003), while other taxa seem not to be limited by dispersal (Gorbushina et al., 2007; van Grembergh et al., 2011). On the other hand, knowledge on potential abiotic or biotic post-colonization barriers is very limited.

Biogeography studies on microorganisms tend either to compare entire communities or to focus on particular taxa (Hanson et al., 2012). Most community studies are based on a single locus and employ partial small subunit rRNA (16S or 18S rRNA) sequences as phylogenetic marker, resulting in a pronounced limitation in phylogenetic resolution due to the conserved nature of these ribosomal sequences. This limitation in phylogenetic resolution may result in a failure to reveal existing biogeography (Bass and Boenigk, 2011; Hanson et al., 2012). By contrast, investigations focusing on particular taxa can potentially employ one or more loci with higher phylogenetic resolution, for instance 16S–23S internal transcribed spacers (ITS) (e.g. van Grembergh et al., 2011) or protein-encoding sequences (e.g. Whitaker et al., 2003).

A general problem in biogeography studies on both macro- and microorganisms is the confirmation of the absence of a studied taxon at a particular site or in a particular region. Cultivation-based investigations on microorganisms may easily overlook the presence of a particular taxon in samples because of the large diversity and high cell numbers of microbial communities. In order to lessen this problem, we studied in a first step the distribution of taxa by a cultivation-based approach, and tested in a second step for absence of taxa in certain regions by using environmental DNA samples and polymerase chain reaction (PCR)-driven detection methods, i.e. cultivation-independent methods. This combination of approaches and the fact that the studied organisms usually form persistent and well-detectable populations (Hahn et al., 2005; 2012) strongly decreased the probability of false negative detection results.

Here we study the biogeography of planktonic freshwater bacteria currently classified as Polynucleobacter necessarius subspecies asymbioticus (Hahn et al., 2009), which are also known as free-living PnecC bacteria (Hahn, 2003). This taxon represents a phylogenetically narrow group characterized by 16S rRNA gene sequence similarities of ≥ 99.2%. Ecologically, this taxon is characterized by an apparently ubiquitous presence in standing freshwater systems (Jezberová et al., 2010; Jezbera et al., 2012) and a cosmopolitan distribution (Hahn, 2003; Pearce et al., 2003; Crump et al., 2004; Hahn et al., 2009; Ghai et al., 2011). They contribute on average by 10% and 20% (extrapolated) to bacterioplankton cell numbers in Central European lentic freshwater systems (Jezberová et al., 2010; Jezbera et al., 2012) and in freshwater systems globally (Jezberová et al., 2010) respectively. Investigations employing phylogenetic markers with resolution beyond the resolution of 16S rRNA sequences suggested that ubiquity of the entire group in freshwater systems results from the presence of specialized lineages in systems differing in ecological conditions (Jezbera et al., 2011). Thus, the ecologically wide distribution of the subspecies is explained by an ecological radiation into differently specialized lineages, for instance adapted to different pH conditions. This explanation for the ubiquitous occurrence of the taxon in freshwater systems is summarized by the ubiquity-by-diversification hypothesis (Jezbera et al., 2011).

The study presented here is based on a globally assembled collection of P.n. ssp. asymbioticus strains currently consisting of 204 isolates from 130 different sites. The collection contains strains from all six continents and all major climatic zones. We searched for phylogeography patterns by analysing sequences of two phylogenetic markers. The most important phylogeographic patterns suggested by the analysis of cultivated strains were tested and confirmed by cultivation-independent investigations.

Results
The habitats from which the 204 P.n. ssp. asymbioticus strains (Supporting Information Table S1) were isolated represent an ecologically broad spectrum of lakes and ponds distributed over all six continents and all major climatic zones (Fig. 1). A minor fraction of strains were isolated from running waters (5 strains), a reservoir (2 strains) and puddles on forestry roads (3 strains). Besides large differences in size, these habitats also vary strongly in physico-chemical conditions. For instance, pH and conductivity values ranged from 3.5 to 9.0 and 4 to 10 010 μS cm⁻¹ respectively. The latitudinal distribution of sites ranges from 78°N (Arctic region, Spitsbergen, Svalbard Islands, Norway) to 63°S (Antarctic Peninsula) (Fig. 1A). While the continent-specific origin of strains is strongly biased towards Eurasia (80% of strains) a more even climate-specific origin of strains can be noted (Table 1). Polair/alpine sites were located in the Arctic (i.e. north of 10°C July isotherm; 5 habitats, 7 strains), the maritime Antarctic (6 habitats, 10 strains) and at high altitudes in the European Alps (13 habitats, 20 strains).
The genetic markers used in the current investigation differed in sequence diversity. The nucleotide diversity of the 16S–23S ITS (543 alignment positions) and glnA sequences (603 bp) were 0.022 and 0.114 respectively. While the average similarity of ITS sequences was 97.5% (range 92.8–100%), the partial glnA sequences showed average sequence similarity of 88.7% (range 82.1–100%) and 97.0% (range 92.0–100%) on the nucleotide and amino acid levels respectively.

**Global phylogeography of P.n. ssp. asymbioticus**

Mantel tests were performed to search for a potential isolation by distance. All strains were included in this analyses, although (despite of the fact that) nine of the ten Antarctic strains were isolated at incubation temperatures of 15°C, and all other at room temperature (-20°C). A cultivation bias due to the different isolation temperatures cannot be ruled out; however, Antarctic strains, which were isolated at room temperature and 15°C, respectively, shared identical ITS and glnA sequences. Furthermore, cultivation-independent investigation with *Polynucleobacter*-specific primers (Hahn *et al.*, 2005) on the *Polynucleobacter* diversity in one of the investigated Antarctic lakes did not reveal the presence of any *Polynucleobacter* lineage missed by the cultivation approach (data not shown). Both phylogenetic markers analysed indicated a significant correlation between genetic and geographic distance (Table 2), which suggests isolation-by-distance pattern on a global scale. Significant but weak correlations between climate parameters (air temperature) and genetic distance of the glnA marker were also revealed, whereas the ITS marker showed no significant correlations with these climate parameters. By contrast, no significant correlation between differences in pH values of habitats from which strains were isolated and the genetic distance between strains could be observed for any of the phylogenetic markers assayed.

A closer look at the reconstructed phylogeny of strains revealed phylogeographic patterns of restricted geographic ranges of subgroups (Fig. 1A and B). One clade representing 10% of strains is exclusively represented by organisms originating from habitats at sites with a warm, mainly tropical climate (temperatures above 18°C in all 12 months; Peel *et al.*, 2007) or almost tropical climate (three strains from two sites in Brazil with minimum monthly mean temperatures of 17°C). As the sites of origin of these 21 strains are spread within tropical areas from South and North America, Asia and Africa, the clade formed by these organisms was named pantropical clade (Fig. 1B). Mantel tests on correlations of the genetic distance between strains of this clade and geographic distance, or selected environmental or climatic parameters, revealed a pronounced correlation with geographic distance ($r = 0.42$, $P < 0.01$, and $r = 0.39$, $P < 0.01$ for ITS and glnA markers respectively) but no significant correlation between genetic distance and any of the tested environmental or climatic parameters (data not shown). Thus, compared with the entire set of strains, a much stronger spatial influence on genetic divergence was suggested for the pantropical clade.

**Testing of range restrictions of the pantropical clade**

The pantropical clade is almost identical with the previously characterized group F13n (Jezbera *et al.*, 2011), which was defined based on the presence of a diagnostic oligonucleotide sequence in the 16S–23S ITS sequences of member strains. Only one strain of the pantropical clade lacks this diagnostic sequence (Supporting Information Fig. S1), thus 95% of the known clade members could be detected by a previously developed reverse line blot hybridization (RLBH) probe (Jezbera *et al.*, 2011).
Importantly, previous investigation of 121 samples from non-tropical habitats in Europe by RLBH resulted in a complete lack of detection of the pantropical clade, while other *Polynucleobacter* groups were frequently detected in these samples (Jezbera et al., 2011 and Table 3). By contrast, investigation of environmental DNA extracted from 30 freshwater habitats in tropical Uganda (Supporting Information Table S2) by RLBH resulted in 63% of habitats being positive for the pantropical clade (82% of habitats yielding PCR products were positive). Presence of strains affiliated with this clade was also confirmed by an analysis of a metagenomic data set representing microbial communities of the Amazon River (Ghai et al., 2011). Our analysis revealed that this metagenomic data
set contains a couple of glnA sequences clustering within the pantropical clade (data not shown).

Canonical correspondence analysis (CCA) indicated a significant influence of climate proxies (air temperature) and geographic (latitude) parameters on distribution of the pantropical clade (Fig. 2) compared with the other clades. Both of which suggest the influence of thermal conditions on the range of this taxon. As suggested by climate data, as well as by the available data on water temperatures in habitats of origin of the Polynucleobacter strains, freshwater systems inhabited by the pantropical clade differ from the other habitats mainly in the absence of low winter temperatures and weaker seasonality (Fig. 1C). Even in several habitats northern than 60° latitude surface water temperatures > 25°C were measured, which are comparable with typical surface temperatures of large tropical lakes such as Tanganyika or Malawi (World Lake Database, http://wldb.ilec.or.jp/LakeDB2/). By contrast, typical winter temperatures of non-tropical lakes, i.e. < 10°C, cannot be expected in lowland tropical freshwater systems. We tested if this basic difference between tropical and non-tropical habitats is reflected by the thermal adaptation of Polynucleobacter strains (Fig. 3). Only one of the 20 tested pantropical clade strains grew at 4°C, but showed only weak growth (colony size) in comparison with the control culture. In contrast, all 20 tested non-tropical strains affiliated with diverse lineages grew well at this low temperature, while only three out of five strains of the subtropical–tropical clade were able to grow.

**Endemicity of the Antarctic clade?**

In total, 37 strains were isolated from cold environments, i.e. polar and alpine lakes and ponds (Table 1). Strains isolated from southern (Antarctic) and northern hemisphere (Arctic and alpine) habitats clustered separately in all phylogenetic reconstructions performed. While the

| Table 2. Results of Mantel tests with data on pairwise genetic distance (Y) of isolates determined either for their 16S–23S ITS sequences or their partial glnA gene sequences against pairwise distance/differences of various parameters (X). |
|------------------------------------------------------------------------------|
| glnA (nt, 202 Sequences) | Geographical distance | Annual temperature | Warmest month | Coldest month | Warmest quarter | Coldest quarter | pH |
|---------------------------|----------------------|------------------|--------------|--------------|----------------|----------------|----|
| Correlation coefficient (rY) | 0.210 | 0.098 | 0.126 | 0.079 | 0.115 | 0.078 | -0.027 |
| Determination of Y by X (%) | 4.4 | 1.0 | 1.6 | 0.6 | 1.3 | 0.6 | - |
| P (rYrand ≥ rYobs) | <0.0001 | 0.0001 | <0.0001 | 0.0016 | <0.0001 | 0.0015 | 0.86 |
| ITS (nt, 204 sequences) | Correlation coefficient (rY) | 0.276 | -0.035 | -0.004 | -0.046 | -0.011 | -0.049 | -0.082 |
| Determination of Y by X (%) | 7.6 | - | - | - | - | - | - |
| P (rYrand ≥ rYobs) | <0.0001 | 0.90 | 0.54 | 0.95 | 0.65 | 0.97 | 1.00 |

*a. P < 0.01.*

These parameters included geographic distance of sites of origin of the strains, pH measured on site and several climate parameters predicted by a climate model for the sites of origin of the strains. Note that the climate data refer to air temperatures, which represent only rough proxies for thermal conditions in aquatic systems.

© 2014 The Authors. Environmental Microbiology published by Society for Applied Microbiology and John Wiley & Sons Ltd, Environmental Microbiology, 17, 829–840.
Antarctic strains formed one (ITS sequences, GlnA AA sequences) or two monophyletic clusters (glnA nt sequences) (Fig. 1B), most of the Arctic and alpine strains appeared to be more closely related to strains obtained from temperate sites.

We intended to search for the Antarctic genotypes in the tropics and the northern hemisphere. Due to the lack of a diagnostic ITS sequence, a RLBH probe suitable for detecting the Antarctic clade could not be developed. Instead, a primer pair for specific detection of the larger branch of the Antarctic clade (Supporting Information Fig. S1) was developed and applied in a nested PCR approach to 16 environmental DNA samples from Antarctic lakes on Livingston Island and at Hope Bay, and 57 samples from lakes and ponds in Uganda (13 habitats) and Europe (44 habitats). The European sites included 10 subalpine and 7 alpine habitats, all located in the Austrian Alps. In almost all cases, nested PCR with environmental DNA of non-Antarctic habitats yielded in the first PCR step a PCR product. This confirmed the presence of *Polynucleobacter* bacteria. In the second PCR step, however, only one non-Antarctic sample resulted in a PCR product. Sequencing of this sole PCR product resulted in a sequence clustering in phylogenetic trees outside of the Antarctic clade. Consequently, this PCR product was not scored as detection of the clade (Table 3). Thirteen of the 16 samples of Antarctic lakes resulted in detection of the Antarctic clade either by cultivation or nested PCR or both (Supporting Information Table S3).

**Discussion**

Previous studies on diversity and environmental distribution of *Polynucleobacter* bacteria focused on sites in Central Europe (Jezbera *et al.*, 2011). Here we enlarged the focus to a global scope and revealed a phylogeographic pattern influenced by geographic distance and climate. Importantly, pH representing the most important environmental parameter strongly influencing *Polynucleobacter* community composition on a regional scale (Jezbera *et al.*, 2011) was not found to influence genetic distance between strains on a global scale (Table 2). Instead, climate-related parameters and isolation-by-distance mechanisms are suggested to structure *Polynucleobacter* communities on this scale.

The used climate data are only weak but are the only available proxies for year-round thermal conditions in the

**Table 3.** Detection of the pantropical, the F1 and the Antarctic clade in various regions by cultivation and cultivation-independent methods.

| Taxon          | Origin of Samples | Method                  | Total number of investigated samples or strains | Positive samples or strains (#) | Positive samples or strains (%) | Reference               |
|----------------|-------------------|-------------------------|-----------------------------------------------|--------------------------------|--------------------------------|-------------------------|
| Pantropical clade | Europe            | RLBH (probe F13n), env. DNA | 121                                           | 0                              | 0.0                            | Jezbera *et al.*, 2011  |
| Pantropical clade | Uganda            | RLBH (probe F13n), env. DNA | 30 (22)                                       | 18                             | 60.0 (81.8)                    | This study              |
| Pantropical clade | Outside tropics   | Cultivation, strains    | 178                                           | 0                              | 0.0                            | This study              |
| Pantropical clade | Tropics           | Cultivation, strains    | 26                                            | 21                             | 80.6                          | This study              |
| Antarctic clade  | Outside Antarctica| Cultivation, strains    | 194                                           | 0                              | 0.0                            | This study              |
| Antarctic clade  | Antarctica        | Cultivation, strains    | 10                                            | 10                             | 100.0                         | This study              |
| Antarctic clade  | Europe            | PCR, env. DNA           | 57                                            | 0                              | 0.0                            | This study              |
| Antarctic clade  | Antarctica        | PCR, env. DNA           | 16                                            | 12                             | 75.0                          | This study              |
| Clade F1*       | Europe            | RLBH (probe F1), env. DNA | 121                                           | 0                              | 0.0                            | Jezbera *et al.*, 2011  |
| Clade F1*       | Uganda            | RLBH (probe F1), env. DNA | 30 (22)                                       | 16                             | 53.3 (72.7)                   | This study              |

a. Clade F1 is only represented by two strain, one isolated from a tropical site and one isolated from a subtropical site (see Suppl. Materials Fig. S1).

b. Only 22 of 30 samples resulted in PCR products suitable for analysis by RLBH.

RLBH, reverse line blot hybridization; env. DNA, environmental DNA; #, number.

Fig. 2. Canonical correspondence analysis (CCA) of the relationship of particular clades (triangles) to environmental and climatic conditions (arrows) characterizing the habitat from which the clade members were isolated. This two-dimensional model explains 63% of the observed variability. Only clades that showed significant correlations in this model are depicted. Conduct, conductivity (μS per centimetre); WarmQuart, mean air temperature of warmest quarter; ColdQuart, mean air temperature of coldest quarter. For data on clade F15 see Suppl. Materials Figs S1 and S2.
freshwater systems inhabited by the isolated bacterial strains. During warmer seasons, surface water temperatures of lakes and ponds are usually above average air temperatures (Fig. 1C) due to the heat storage capacity of standing water bodies. At sites with winterly average air temperature below 0°C bacteria and other organisms do not experience growth conditions below freezing point. Vertical differences of water temperatures in stratified lakes are only a minor point here, because almost all (one exception) bacterial strains contained in the culture collection were obtained from surface waters. Due to the limitations of climate data as proxies for thermal conditions in standing aquatic systems, the observed weak correlation between glnA genetic data and climate data may underestimate the influence of thermal conditions on the global structuring of the Polynucleobacter population. Lack of correlation with ITS data could, in addition, have been influenced by the lower genetic diversity of this marker (Hahn et al., 2012) and by the presence of hypervariable regions potentially masking phylogenetic signals due to extensive homoplasy (van Gremberghe et al., 2011).

Do range restrictions of subgroups really exist?

Of course all the results by cultivation and molecular detection experiments (Table 3) do not provide evidence for complete absence of taxa in the entire tropical or non-tropical zones respectively. Complete absence is simply a non-testable hypothesis. Even ultra-deep sequencing of environmental DNA cannot prove the absence of a specific taxon in a certain area. Another potential reason for overlooking microorganism in a certain habitat is sampling of the system during an inappropriate period characterized by only low numbers of the taxon of interest. This is especially a problem in bloom-forming taxa, while Polynucleobacter bacteria rather tend to establish persistent populations fluctuating in numbers much less than typical bloom-forming microorganisms (Hahn et al., 2005; 2012; Wu and Hahn, 2006a,b). Although, as stated above, a perfect prove of complete absence of a certain taxon in a certain natural system is basically impossible, the observed patterns of detection/no detection obtained by two independent methods are at least for some subgroups very strong (Table 3). It is important to note that a previous study demonstrated that cultivation and cultivation-independent (cloning and sequencing) investigations on the Polynucleobacter population of a humic pond revealed the same diversity of 16S–23S ITS genotypes and also suggested a low genetic diversity of Polynucleobacter populations (Hahn and Pöckl, 2005; Hahn et al., 2005). Thus, it is not expected that cultivation-based investigation results in a strongly biased picture of the Polynucleobacter diversity.

The phenomenon of restricted ranges seems not to be limited to the pantropical and the Antarctic subgroup. The currently available distribution data (based on both cultivation and RLBH) also suggest for some other Polynucleobacter clades geographic range restrictions. This includes the subtropical–tropical clade (Fig. 1), which is partially represented by the previously defined group F2 (Jezbera et al., 2011), clade F1 (Table 3; Jezbera et al., 2011), clade F10 (Jezbera et al., 2011) and clade F15 (Jezbera et al., 2011), which is represented by 10% of the isolated strains (Table 4, Supporting Information Fig. S1). For groups F1, F10 and F15, suitable RLBH probes were available for testing presence/absence in Central European and African habitats located in Uganda. Similar to the pantropical clade, group F1 could not be detected in Europe, but it was detected in several Ugandan habitats (Table 3). Whereas for the other two groups, an opposite
Table 4. Genetic and ecological characteristics of three putative endemic Polynucleobacter taxa. All data are based on isolated representatives.

| Taxon                | Number of strains | Number of haplotypes | Minimum similarity (%) | Nucleotide diversity Pi | pH range | Maximum distance (km) |
|----------------------|-------------------|----------------------|------------------------|-------------------------|----------|----------------------|
| Pantropical clade    | 21                | 19                   | 97.3                   | 0.01005                 | 5.7–8.2  | 17234                |
| F15                  | 20                | 8                    | 99.2                   | 0.00272                 | 4.2–7.2  | 3264                 |
| Antarctic clade      | 10                | 5                    | 97.9                   | 0.00881                 | 6.8–8.5  | 10 (222)*            |

a. Maximum distance of sites of origin of cultivated strains is 10 km, maximum distance between sites of detection is 222 km.

Maximum distance and pH range refer to habitats from which strains affiliated with the taxa were obtained. See Supporting Information Fig. S2 for estimated range limits of each of the three taxa.

A trend was observed. In one case very weak (compared with detections in European samples), most likely false positive, detections in Ugandan samples were noted, though (Supporting Information Table S2). Further investigations are required for confirming these trends, however, these observations suggest that range restrictions are not exceptional among subgroups of Polynucleobacter bacteria.

Based on the current data, it is not possible to establish detailed estimations on the range boarders of subgroups. The geographic pattern of detection of the pantropical clade, the correlation of the distribution with climatic data and the revealed inability of strains to grow at low temperatures suggest that this clade is more or less restricted to the tropical climatic zone. However, whether and how far the range extends into the subtropical zones of both hemispheres is currently unknown (Supporting Information Fig. S2).

The Antarctic clade was observed to be widely distributed in lakes on Byers Peninsula (Camacho, 2006; Toro et al., 2007; Villaescusa et al., 2010; 2013) and at Hope Bay, 222 km distant (Supporting Information Table S3), while there was no hint of the presence of this clade in European and African lakes and ponds. Consequently, the Antarctic clade is currently known exclusively from several freshwater lakes located at the Antarctic Peninsula (Supporting Information Fig. S2). Whether the clade is restricted to Antarctic freshwater habitats or it is also present on other continents, is currently not known. The presence of Polynucleobacter in other freshwater lakes of maritime Antarctica was demonstrated previously (Pearce et al., 2003; Villaescusa et al., 2013), but these detections were exclusively based on 16S rRNA sequences as opposed to the analysis of 16S–23S ITS, and so whether those Polynucleobacter bacteria belong to the Antarctic clade cannot be analysed. Currently, we cannot exclude that at least alpine Andean lakes may represent habitats outside Antarctica potentially colonized by the cold-adapted Antarctic clade (Supporting Information Fig. S2).

Thermal adaptation of strains

The pantropical subgroup showed thermal adaptations different from strains isolated from habitats located in the temperate climatic zone (Fig. 3). Tropical strains are characterized by a lack or very weak growth at low temperatures, which fits well to the thermal conditions in their home habitats. Climate-linked differences in thermal adaptation of closely related strains of planktonic freshwater bacteria affiliated with another phylum were demonstrated previously (Hahn and Pöckl, 2005). This previous study compared optimal and maximum growth temperatures of actinobacterial strains isolated from habitats located in three climatic zones. The seven investigated strains shared identical 16S rRNA sequences and the 16S–23S ITS sequences possessed sequence similarities in the range of 99.0–100%. Thus, sequence diversity of both markers was smaller than sequence diversity revealed for the Polynucleobacter strains investigated in the current study. However, even these closely related strains of Actinobacteria showed pronounced differences in thermal adaptations, which reflected differences in thermal characteristics of their home habitats.

Intercontinental distribution of the pantropical clade

The range of the pantropical clade was found to be longitudinally unlimited within the tropical climatic zone (Fig. 1A). This could either be explained by a historical distribution already present before the break-up of the supercontinent Pangaea about 150–140 Ma before present or alternatively by later colonization of parts of the distribution area by natural or anthropogenically assisted intercontinental dispersal. Such successful intercontinental dispersal was demonstrated for spore-forming bacteria (Gorbushina et al., 2007). Interestingly, Polynucleobacter bacteria were detected in the troposphere (DeLeon-Rodriguez et al., 2013), but whether Polynucleobacter bacteria, which are unable to form spores, are transported over large distances without loss of viability, remains unknown. If gene flow between continents was absent or limited, separation of palaeotropical and neotropical sub-
groups of the clade should have resulted in genetic divergence following the break-up of Pangaea. By assuming isolation over a period of 140 Ma and an assumed divergence rate of the 16S rRNA genes of 0.00625–0.0125% per Ma, which represent conservative estimates of 25–50% of the slowest rate estimated for obligately endosymbiotic bacteria (Kuo and Ochman, 2009), sequence divergence within the pantropical clade of 0.9–1.8% could be expected. The lower divergence value is similar to the maximum divergence (0.8%) observed for the entire Pn. ssp. asymbioticus group (64 16S rRNA sequences ≥ 1477 bp). However, the maximum palaeo-neotropical divergence of almost full-length 16S rRNA sequences affiliated with the pantropical clade was only 0.2%, while a palaeotropical (Asia) and two neotropical strains (South America) even shared identical 16S rRNA sequences. Furthermore, in contrast to other bacterial taxa (Bahl et al., 2011), the phylogenetic reconstructions with both markers revealed no phylogenetic structure separating palaeo- and neotropical strains (Fig. 1B). Altogether, these findings suggest successful intercontinental dispersal of pantropical clade strains. The observed distance-decay pattern hints at rare but natural intercontinental dispersal events too weak to balance divergence between distantly located populations. On the other hand, this pattern argues against a recent pantropical range expansion driven by anthropogenic activities, such as intercontinental transport of freshwater as ballast water (Seiden et al., 2010).

Phylogenetic resolution and intra-taxon ecological diversification

As stressed previously, phylogenetic resolution of methods applied in phylogeographic investigations on microorganisms is the crucial factor for depths of analysis (Bass and Boenigk, 2011; Hanson et al., 2012). While the 16S rRNA gene of Polynucleobacter bacteria lacks almost any intraspecific phylogeographic signal (Hahn, 2003), the two markers employed in this study revealed a deep phylogeographic structure in Pn. ssp. asymbioticus. The presented analysis suggests that this structure results from both environmental selection of ecotypes and restrictions in gene flow between distant sites. Weighting of the proportional impact of these two mechanisms on the phylogeographic structuring will require larger data sets on ecologically more homogenous subgroups.

The analysed strain collection is currently geographically biased due to overrepresentation of strains obtained from European sites. This geographic bias and the small number of phylogenetic markers included in the study are potentially limiting the potential for revealing intercontinental structuring of Polynucleobacter communities. For instance, genetic differentiation between North American and European populations potentially caused by geographic isolation could not be revealed due to the lack of a suitable number of strains originating from North America. Future isolation efforts should focus on sites outside of Europe. Furthermore, efforts for establishment of a multilocus sequence typing scheme for Polynucleobacter bacteria are required. Both improvements will enable analysis of biogeographic structures with an increased resolution.

Apart from geographic reasons, the environmental filtering of ecotypes seems to take place at two different levels. On a regional level, climatic conditions seem to act as a filter, sorting out ecotypes with unsuitable thermal adaptation (Fig. 3). At a local level, habitat-specific environmental conditions such as pH, conductivity, oxygen concentration, and quantity and quality of dissolved organic carbon further select for suitably adapted genotypes (Jezbera et al., 2011). This two-level environmental filtering results in a patchy phylogeographic structure, which is more complex than revealed so far.

Conclusions

This study revealed a dimension of microbial diversity, which cannot be discovered by current ultra-deep sequencing projects employing next generation sequencing of 16S rRNA markers. Linking of genetic diversity data and phenotypic data suggested a phylogeographic structuring of the Pn. ssp. asymbioticus population, which is at least partially shaped by adaptation to different thermal niches. Future studies have to reveal the extent of the ecological differences between subgroups involved in this phylogeographic structuring, as well as the contribution of restrictions in gene flow to the revealed patterns.

Experimental procedures

Samples

The study is based on two different kinds of samples, i.e. bacterial strains representing the investigated taxon and environmental DNA directly extracted from water samples. The strains were isolated as pure cultures from various freshwater systems by using the acclimatization method (Hahn et al., 2004). The established culture collection consists of 204 Polynucleobacter necessarius subspecies asymbioticus (Hahn et al., 2009) strains. Samples of environmental DNA were obtained from many freshwater systems by sampling surface waters (up to 0.5 m depth), harvesting of planktonic organisms on 0.2 μm filters and subsequent DNA extraction as described previously (Jezbera et al., 2011).

Sequencing of genetic markers and phylogenetic analysis

16S–23S ITS and partial glutamine synthetase gene sequences (glnA) were amplified from gDNA of pure cultures.
of strains, sequenced and phylogenetically analysed as described previously (Jezbera et al., 2011). Both markers represent single copy genes/sequences in Polynucleobacter genomes (Hahn et al., 2012).

Detection of taxa by RLBH

16S–23S ITS sequences of P.n. ssp. asymbioticus were amplified from environmental DNA samples by using taxon-specific primers, as described previously (Jezbera et al., 2011). The obtained mixtures of amplified ITS sequences were investigated for the presence/absence of certain Polynucleobacter subgroups by RLBH by using a set of thirteen Polynucleobacter-subgroup-specific hybridization probes (Jezbera et al., 2011).

Detection of the Antarctic clade by PCR

A primer pair for specific detection of Antarctic Polynucleobacter genotypes in environmental DNA samples was developed. The primers Limnop-glnA-F (5′-CATCATT AAGCATGCGAAGAAC-3′) and Limnop-glnA-R (5′-CCA TCATCAATGCAGCAAAGCAA-3′) target the glnA genes of 8 of the 10 obtained Antarctic Polynucleobacter strains. PCR conditions were optimized by using gDNA extracted from target strains. A nested PCR approach was applied to environmental samples in order to increase the sensitivity of detection of Antarctic genotypes. The Polynucleobacter-specific primers glnA1212F and glnA1895R (Jezbera et al., 2011) were used in a first step for amplifying partial glnA sequences. PCR products were subjected to a second PCR with the new primers Limnop-glnA-F and Limnop-glnA-R.

Thermal adaptation of cultivated strains

The capability of selected strains to grow at low temperatures was tested by dropping 10 µl of culture grown in liquid NSY medium (Hahn et al., 2004) on NSY agar plates and subsequently incubating them (triplicates for each strain) at 4°C for up to 15 weeks. Positive controls were incubated at room temperature (> 23°C). Experiments resulting in lack of growth at 4°C were repeated.

Climate data

Climate data were used as proxies of thermal conditions of aquatic systems from which Polynucleobacter strains were isolated. Site-specific climate data were obtained from the WorldClim data set (v. 1.3, with a resolution of 2.5 min) (Hijmans et al., 2005a), covering the global land areas except Antarctica, using the DIVA-GIS software (Hijmans et al., 2005b). This included data on annual mean air temperature, mean maximum air temperature of the warmest month, mean minimum air temperature of the coldest month, mean air temperature of the warmest quarter and mean air temperature of the coldest quarter. Climate data at the site of origin of the Antarctic strains (Byers Peninsula, Livingston Island, South Shetlands Islands, and Hope Bay, Maritime Antarctica) were recorded on location (Villaescusa et al., 2010; Schiaffino et al., 2011; Bañon et al., 2013).

Statistical analyses

Using Mantel tests, isolation by distance was investigated as the correlation between matrices of pairwise genetic distances of strains and matrices of pairwise geographic distances of sites of origin of strains. In the same way, correlations of genetic distance and differences in climatic or pH conditions between sites of origin of strains were searched. Pairwise genetic distances between strains were calculated for both markers (nt sequences) by using the Tamura 3-parameter substitution model with non-uniformity of evolutionary rates among sites and by assuming that a certain fraction of sites are evolutionarily invariable (T92 + G + I substitution model) by using the MEGA software, version 5 (Tamura et al., 2011). Pairwise geographic distances (great-circle distance) were calculated with the Geographic Distance Matrix Generator (Peter Ersts, American Museum of Natural History, Center for Biodiversity and Conservation, http://biodiversityinformatics.amnh.org/open_source/gdmg) from geographic coordinates. Using the ARLEQUIN software (Excoffier et al., 2005), correlations between matrices were analysed by Mantel tests with 100,000 permutations.

Multivariable analyses were performed using the CANOCO software (Ter Braak and Šmilauer, 1998). CCA was carried out with centring and standardization by species norm. Forward selection was used to choose significant (P < 0.05) explanatory (environmental) variables by a Monte Carlo permutation test with 999 unrestricted permutations.

Nucleotide sequences

Accession numbers are provided in Supporting Information Table S1.

Acknowledgements

We are grateful to many colleagues for providing water samples, especially to Ondřej Komárek and Karel Šimek, to Martha Kaddumukasa for organizing and participating in sampling of habitats in Uganda, to Maria Gadermaier for excellent technical assistance and to Anton Baer for English text corrections. This study was largely supported by the Austrian Science Fund (project I482-B09 granted to MWH), as well as by the Grant Agency of the Czech Republic (project EEF/10/E011 granted to J. Jezbera). Antarctic work and some other sampling and sample-processing were supported by grants CGL2005-06549-C02-02/ANT from the Spanish Ministry of Education and Science and CGL2012-38909 (ECOLAKE) from the Spanish Ministry of Economy and Competitiveness, both given to AC and co-financed by European FEDER funds. We are indebted to colleagues from LIMNOPOLAR project for their help in Antarctic sampling.

References

Baas Becking, L.G.M. (1934) Geobiologie of inleiding tot de milieukunde. The Hague, the Netherlands: W.P. Van Stockum & Zoon (in Dutch).
Bahl, J., Lau, M.C.Y., Smith, G.J.D., Vijaykrishna, D., Cary, S.C., Lacap, D.C., et al. (2011) Ancient origins determine global biogeography of hot and cold desert Cyanobacteria. *Nat Commun* **2**: 163. doi:10.1038/ncomms1167.

Baahn, M., Justel, A., Velazquez, D., and Quesada, A. (2013) Regional weather survey on Byers Peninsula, Livingston Island, South Shetland Islands, Antarctica. *Antarct Sci* **25**: 146–156.

Bass, D., and Boenigk, J. (2011) Everything is everywhere: a twenty-first century de-/ reconstruction with respect to protists. In *Biogeography of Microorganisms. Is Everything Small Everywhere?*, Vol. 79. Fontaneto, D. (ed.). Cambridge: Cambridge University Press, pp. 88–110. Special.

Camacho, A. (2006) Planktonic microbial assemblages and the potential effects of metazooplankton predation on the food web of lakes from the maritime Antarctica and sub-Antarctic islands. *Rev Environ Sci Biotechnol* **5**: 167–185.

Crump, B.C., Hopkinson, C.S., Sogin, M.L., and Hobbie, J.E. (2004) Microbial biogeography along an estuarine salinity gradient: combined influences of bacterial growth and residence time. *Appl Environ Microbiol* **70**: 1494–1505.

DeLeon-Rodriguez, N., Lathem, T.L., Rodriguez-R, L.M., Barazesh, J.M., Anderson, B.E., Bayersdorf, A.J., et al. (2013) Microbiome of the upper troposphere: species composition and prevalence, effects of tropical storms, and atmospheric implications. *Proc Natl Acad Sci* **110**: 2575–2580.

de Wit, R., and Bouvier, T. (2006) ‘Everything is everywhere, but, the environment selects’ what did Baas Becking and Beijerinck really say? *Environ Microbiol* **8**: 755–758.

Excoffier, L., Laval, G., and Schneider, S. (2005) Arlequin ver. 3.0: an integrated software package for population genetics data analysis. *Evol Bioinform Online* **1**: 47–50.

Finlay, B.J. (2002) Global dispersal of free-living microbial eukaryote species. *Science* **296**: 1061–1063.

Fontaneto, D., and Brodie, J. (2011) Why biogeography of microorganisms? In *Biogeography of Microorganisms. Is Everything Small Everywhere?*, Vol. 79. Fontaneto, D. (ed.). Cambridge: Cambridge University Press, pp. 3–10. Special.

Ghai, R., Rodriguez-Valera, F., McMahon, K.D., Toyama, D., Rinke, R., de Oliveira, T.C.S., et al. (2011) Metagenomics of the water column in the pristine upper course of the Amazon River. *PLoS ONE* **6**: e23785.

Gorbushina, A.A., Cort, R., Schulte, A., Lazanus, D., Schnetger, B., Brumsack, H.J., et al. (2007) Life in Darwin’s dust: intercontinental transport and survival of microbes in the nineteenth century. *Environ Microbiol* **9**: 2911–2922.

Gray, N.D., Brown, A., Nelson, D.R., Pickup, R.W., Rowan, A.K., and Head, I.M. (2007) The biogeographical distribution of closely related freshwater sediment bacteria is determined by environmental selection. *ISME J* **1**: 596–605.

van Gremberghhe, I., Leliaert, F., Mergeay, J., Vanormelingen, P., Van der Gucht, K., Debeer, A.E., et al. (2011) Lack of phylogeographic structure in the freshwater cyanobacterium *Microcystis aeruginosa* suggests global dispersal. *PLoS ONE* **6**: e19561. doi:10.1371/journal.pone.0019561.

Hahn, M.W. (2003) Isolation of strains belonging to the cosmopolitan *Polynucleobacter necessarius* cluster from freshwater habitats located in three climatic zones. *Appl Environ Microbiol* **69**: 5248–5254.

Hahn, M.W., and Pöckl, M. (2005) Ecotypes of planktonic *Actinobacteria* with identical 16S rRNA genes adapted to thermal niches in temperate, subtropical, and tropical freshwater habitats. *Appl Environ Microbiol* **2**: 766–773.

Hahn, M.W., Stadler, P., Wu, Q.L., and Pöckl, M. (2004) The filtration-acclimatization-method for isolation of an important fraction of the not readily cultivable bacteria. *J Microbiol Methods* **57**: 379–390.

Hahn, M.W., Pöckl, M., and Wu, Q.L. (2005) Low intraspecific diversity in a *Polynucleobacter* subcluster population numerically dominating bacterioplankton of a freshwater pond. *Appl Environ Microbiol* **71**: 4539–4547.

Hahn, M.W., Lang, E., Brandt, U., Wu, Q.L., and Scheuerl, T. (2009) Emended description of the genus *Polynucleobacter* and the species *P. necessarius* and proposal of two subspecies, *P. necessarius* subsp. nov. and *P. necessarius* subsp. *asymbioticus* subsp. nov. *Int J Syst Evol Microbiol* **59**: 2002–2009.

Hahn, M.W., Scheuerl, T., Jezeberová, J., Koll, U., Jezebera, J., Šimek, K., et al. (2012) The passive yet successful way of planktonic life: genomic and experimental analysis of the ecology of a free-living *Polynucleobacter* population. *PLoS ONE* **7**: e32772.

Hanson, C.A., Fuhrman, J., Horner-Devine, M.C., and Martiny, J.B.H. (2012) Beyond biogeographic patterns: processes shaping the microbial landscape. *Nat Rev Microbiol* **10**: 497–506.

Hervás, A., Camarero, L., Reche, I., and Casamayor, E.O. (2009) Viability and potential for immigration of airborne bacteria from Africa that reach high mountain lakes in Europe. *Environ Microbiol* **11**: 1612–1623.

Hijmans, R.J., Cameron, S.E., Parra, J.L., Jones, P.G., and Jarvis, A. (2005a) Very high resolution interpolated climate surfaces for global land areas. *Int J Clim* **25**: 1965–1978.

Hijmans, R.J., Guarino, L., Jarvis, A., O’Brien, R., and Mathur, P. (2005b) *DIVA-GIS Version 5.2: Manual*. URL http://diva-gis.org/.

Jezebera, J., Jezeberová, J., Brandt, U., and Hahn, M.W. (2011) Ubiquity of *Polynucleobacter necessarius* subspecies *asymbioticus* results from ecological diversification. *Environ Microbiol* **13**: 922–931.

Jezebera, J., Jezeberová, J., Koll, U., Hornák, K., Šimek, K., and Hahn, M.W. (2012) Contrasting trends in distribution of four major planktonic betaproteobacterial groups along a pH gradient of epilimnia of 72 freshwater habitats. *FEMS Microbiol Ecol* **81**: 467–479.

Jezeberová, J., Jezebera, J., Brandt, U., Lindström, E.S., Langenheder, S., and Hahn, M.W. (2010) Ubiquity of *Polynucleobacter necessarius* subsp. *asymbioticus* in lentic freshwater habitats of a heterogeneous 2000 km² area. *Environ Microbiol* **12**: 658–669.

Kuo, C.H., and Ochman, H. (2009) Inferring clocks when lacking rocks: the variable rates of molecular evolution in bacteria. *Biol Direct* **4**: 35.
Marshall, D.J., Monro, K., Bode, M., Keough, M.J., and Swearer, S. (2010) Phenotype-environment mismatches reduce connectivity in the sea. *Ecol Lett* **13**: 128–140.

Papke, R.T., Ramsing, N.B., Bateson, M.M., and Ward, D.M. (2003) Geographical isolation in hotspring cyanobacteria. *Environ Microbiol* **5**: 650–659.

Pearce, D.A., van der Gast, C.J., Lawley, B., and Ellis-Evans, Schiaffino, M.R., Unrein, F., Gasol, J.M., Massana, R., Papke, R.T., Ramsing, N.B., Bateson, M.M., and Ward, D.M. (2003) Bacterioplankton community diversity in a maritime Antarctic lake, determined by culture-dependent and culture-independent techniques. *FEMS Microbiol Ecol* **45**: 59–70.

Pearce, D.A., Bridge, P.D., Hughes, K.A., Sattler, B., Marshall, D.J., Monro, K., Bode, M., Keough, M.J., and M. W. Hahn, U. Koll, J. Jezberová and A. Camacho (2011) Bacterial community structure in a latitudinal gradient of lakes: the roles of spatial versus environmental factors. *Freshw Biol* **56**: 1973–1991.

Schlichting, H.E., Speziale, B.J., and Zink, R.M. (1978) Dispersal of algae and protozoa by Antarctic flying birds. *Antarct J US* **13**: 147–149.

Seiden, J.M., Way, C., and Rivkin, R.B. (2010) Microbial hitchhikers: dynamics of bacterial populations in ballast water during a trans-Pacific voyage of a bulk carrier. *Aquat Invasions* **5**: 13–22.

Strahler, A.H., and Strahler, A.N. (2005) *Physical Geography: Science and Systems of the Human Environment*. New York: Wiley. 794pp.

Tamames, J., Abellan, J.J., Pignatelli, M., Camacho, A., and Moya, A. (2010) Environmental distribution of prokaryotic taxa. *BMC Microbiol* **10**: 85.

Tamura, K., Peterson, D., Peterson, N., Stecher, G., Nei, M., and Kumar, S. (2011) MEGAS: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. *Mol Biol Evol* **28**: 2731–2739.

Ter Braak, C.J.F., and Šmilauer, P. (1998) *CANOCO for Windows Version 4.02*. Wageningen, The Netherlands: Centre for Biometry Wageningen, CPBO-DLO.

Toro, M., Camacho, A., Rochera, C., Rico, E., Bahón, M., Justel, A., et al. (2007) Limnological characteristics of the freshwater ecosystems of Byers Peninsula, Livingston Island, in maritime Antarctica. *Polar Biol* **30**: 635–649.

Villaescusa, J.A., Casamayor, E.O., Rochera, C., Velázquez, D., Chicote, A., Quesada, A., et al. (2010) A close link between bacterial community composition and environmental heterogeneity in maritime Antarctic lakes. *Int Microbiol* **13**: 67–77.

Villaescusa, J.A., Casamayor, E.O., Rochera, C., Quesada, A., Michaud, L., and Camacho, A. (2013) Heterogeneous vertical structure of the bacterioplankton community in a non-stratified Antarctic lake. *Antarct Sci* **25**: 229–238.

Whitaker, R.J., Grogan, D.W., and Taylor, J.W. (2003) Geographic barriers isolate endemic populations of hyperthermophilic archaea. *Science* **301**: 976–978.

Wu, Q.L., and Hahn, M.W. (2006a) Differences in structure and dynamics of *Polynucleobacter* communities in a temperate and a subtropical lake revealed at three phylogenetic levels. *FEMS Microbiol Ecol* **57**: 67–79.

Wu, Q.L., and Hahn, M.W. (2006b) High predictability of the seasonal dynamics of a species-like *Polynucleobacter* population in a freshwater lake. *Environ Microbiol* **8**: 1660–1666.

Zwolinski, Z.B., Rachiewicz, G., Mazurek, M., and Paluszkiwicz, R. (2007) The geoecological model for small tundra lakes, Spitsbergen. *Landform Anal* **5**: 113–118.

**Supporting information**

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

**Fig. S1.** Complete phylogenetic trees (NJ) of the 16S–23S ITS and the glnA sequences (compare Fig. 1B). Subgroups mentioned in the paper are highlighted by different colours. Strains matching to the RLBH probe F13n targeting the Antarctic clade (Jezbera et al., 2011) and strains matching to PCR primers used for targeting the major branch of the Antarctic clade (this study) are indicated by brackets.

**Fig. S2.** Putative range limits of the Pantropical (red), the Antarctic (blue) and the F15 clade (yellow). Range limits deduced from detections by cultivation and cultivation-independent methods are shown as solid lines of a dot (Antarctic clade, region represented by Byers Peninsula and Hope Bay). Hypothetical range limits are shown as dashed lines.

**Table S1.** *Polynucleobacter* strains, geographic origin and Genbank accession numbers of 16S–23S ITS and glnA sequences.

**Table S2.** Detection of *Polynucleobacter necessarius* ssp. *asymbioticus* subgroups by reverse line blot hybridization (RLBH) in habitats located in Uganda.

**Table S3.** Antarctic lakes investigated for the presence of the Antarctic clade.