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OPEN Habitat heterogeneity and **connectivity shape microbial communities in South American peatlands**

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Bacteria play critical roles in peatland ecosystems. However, very little is known of how habitat heterogeneity afects the structure of the bacterial communities in these ecosystems. Here, we used amplicon sequencing of the 16S rRNA and *nifH* **genes to investigate phylogenetic diversity and bacterial community composition in three diferent sub-Antarctic peat bog aquatic habitats:** *Sphagnum magellanicum* **interstitial water, and water from vegetated and non-vegetated pools. Total and putative nitrogen-fxing bacterial communities from** *Sphagnum* **interstitial water difered signifcantly from vegetated and non-vegetated pool communities (which were colonized by the same bacterial populations), probably as a result of diferences in water chemistry and biotic interactions. Total bacterial communities from pools contained typically aquatic taxa, and were more dissimilar in composition and less species rich than those from** *Sphagnum* **interstitial waters (which were enriched in taxa typically from soils), probably refecting the reduced connectivity between the former habitats. These results show that bacterial communities in peatland water habitats are highly diverse and structured by multiple concurrent factors.**

Bacterial communities in peat bog ecosystems contribute signifcantly to nutrient cycling, to carbon sequestration and to greenhouse gas emissions¹⁻⁴. However, we still have a limited knowledge of the diversity and spatial distribution of bacterial assemblages in peat bogs. As a result, we do not know, for example, how the composition of bacterial communities and their traits difer across peatland water bodies; although recent work has shown that bacterioplankton communities in peat bog pools are diverse and variable⁵. Understanding the diversity and spatial distribution of microbial communities across diferent peatland aquatic habitats is important, because these habitats are highly heterogeneous^{[6](#page-6-2)}; and habitat heterogeneity can lead to changes in biodiversity patterns, which provide information to many ecological and evolutionary questions, as well as to conservation planning^{[7](#page-6-3)}. The latter is especially crucial, as peat bog ecosystems may be highly vulnerable to climate change^{[8](#page-6-4)}.

Habitat diferences between peatland water bodies may arise for a number of reasons. For example, peatlands can have contrasting nutrient conditions⁹, and it is well known that microbes often differ in their nutritional requirements¹⁰. Copiotrophic microbes have high nutritional requirements, while oligotrophic microbes are likely to outcompete copiotrophs in conditions of low nutrient availability¹⁰. Environmental parameters such as $\rm pH^5$ and $\rm DOC^{11}$ $\rm DOC^{11}$ $\rm DOC^{11}$ may also be important in determining microbial community composition in peatlands. The role of aquatic vegetation cannot be ignored either. The lack of surface vegetation allows the penetration of high levels of photosynthetically active radiation (PAR) and ultraviolet radiation, with consequences for nutrient cycling and primary production, which in turn, can promote compositional differences in microbial communities¹² Furthermore, peatland vegetation is ofen dominated by *Sphagnum* mosses, and diferent *Sphagnum* species har-bour different bacterial populations^{[13](#page-6-9)}.

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Peatland water bodies also differ in connectivity, which influences microbial diversity patterns^{14–16}. For instance, the total number of species in habitat patches connected by moderate dispersal should be higher than in a single large patch^{[17](#page-6-11)}. In addition to their effects on species richness, interconnected habitats may favour species with generalist traits. This should lead to a decline in community turnover because of increased homogenization of the metacommunity[18.](#page-6-12) Consequently, understanding how habitat heterogeneity and diferences in connectivity afect the distribution of bacterial communities and their traits is important to better understand the structure-function relationship in peat bog ecosystems and for conservation planning.

Here, we analysed the total and nitrogen-fxing bacterial communities found in three diferent aquatic habitats (vegetated and clear pools, and *Sphagnum magellanicum* interstitial water) within two peat bog complexes (Rancho Hambre and Valle de Andorra, Tierra del Fuego, Argentina), using high-throughput DNA sequencing of 16S rRNA and *nifH* genes. Biological nitrogen fxation of atmospheric nitrogen is an important process in peatlands, because the growth of *Sphagnum* mosses in peatlands is ofen N limited[19](#page-6-13). Vegetated and clear pools can be regarded as nutrient-poor island-like habitats embedded in a landscape of *Sphagnum* and fed essentially by rainfall, whereas *Sphagnum* interstitial waters are less nutrient-poor and exhibit a major degree of connectivity and are further linked to the surrounding landscape. Tus, we predict that the structure of both total bacterial and putative nitrogen-fxing communities will refect the striking diferences in nutrient status and connectivity between these habitats.

Results and Discussion

Rarefaction plots, Chao1 and Good's coverage (95–100%) estimates indicated reasonable sequencing coverage, especially for *nifH* genes, (Supplementary Table S1, Supplementary Fig. S1); although more sequencing depth would be required to assess the true diversity of the samples. Overall, total bacteria and nitrogen-fxing taxa from *Sphagnum* interstitial waters were more diverse than those from pools (Supplementary Table S1, Supplementary Fig. S2). However, the diferences were not statistically signifcant for putative nitrogen-fxing taxa. Most of the bacterial taxa found were members of the *Proteobacteria*, *Verrucomicrobia*, *Actinobacteria*, *Bacteroidetes*, *Acidobacteria* and *Planctomycetes* [\(Fig.](#page-1-0) 1), all of which are commonly found in peatland ecosystems[1,](#page-6-0)[5,](#page-6-1)[20,](#page-6-14)[21.](#page-6-15) Nevertheless, *Betaproteobacteria* and *Actinobacteria* dominated pool communities, while *Sphagnum* interstitial water contained a larger proportion of *Alphaproteobacteria*, *Acidobacteria and Planctomycetes*. Tis is in agreement with previous studies showing that members of these phyla are commonly found in association with *Sphagnum* mosses^{[13](#page-6-9),[22](#page-6-16)}.

The differences in composition were also evident at the genus level. LEfSe analysis identified 26 bacterial genera as biomarkers [\(Fig.](#page-2-0) 2). *Sphagnum* interstitial water samples were enriched in genera such as *Aciditerrimonas* (*Actinobacteria*), *Mucilaginibacter* (*Bacteroidetes*) and *Acidisoma* (*Alphaproteobacteria*), whereas *Propionibacterium* (*Actinobacteria*), *Polynucleobacter* (*Betaproteobacteria*) and *Methylomonas* (*Gammaproteobacteria*) were overrepresented in pools. We do not have a complete understanding about the specifc ecology of these microorganisms in peatland ecosystems. However, members of the genus *Aciditerrimonas* and *Methylomonas* have been reported as *Sphagnum*-associated methanotrophs^{[4](#page-6-17),[23](#page-6-18)}, which are an important sink for the methane produced in peatland[s24](#page-6-19). *Mucilaginibacter* spp. has the ability to use a broad range of heteropoly-saccharides, such as xylan, laminarin, or pectin, in acidic and cold conditions^{[1](#page-6-0)}. Several bacterial strains affiliated

Figure 2. Least discriminant analysis (LDA) efect size taxonomic cladogram, based on 16S rRNA gene sequences, comparing all samples for the three habitats. Signifcantly discriminant taxon nodes are coloured and branch areas are shaded according to the highest ranked group for that taxon. If the taxon is not signifcantly diferentially represented among sample habitats, the corresponding node is coloured yellow.

to *Acidisoma* (e.g. *A. sibiricum* and *A. tundrae*) are aerobic, acidophilic, peat-inhabiting chemo-organotrophs that utilize a variety of sugars, polyalcohols and polysaccharides[25](#page-6-20). Fermenting bacteria such as *Propionibacterium* can degrade carbohydrates and polymeric compounds, therefore participating in carbon turnover 26 . Members of the genus *Polynucleobacter* can perform the assimilatory reduction of nitrate and assimilate sulphur and sulphat[e27.](#page-6-22) In regard to putative nitrogen-fxing microbes, most sequences were related to *Proteobacteria*, *Firmicutes*, *Cyanobacteria* and *Verrucomicrobia* (Supplementary Fig. S3). At the genus level, sequences closely related to *Methylobacter* (*Gammaproteobacteria*) and *Desulfobulbus* (*Deltaproteobacteria*) species dominated in pools, while sequences related to *Bradyrhizobium* (*Alphaproteobacteria*) and *Burkholderia* (*Betaproteobacteria*) species were overrepresented in *Sphagnum* interstitial waters. The *nifH* deduced amino acid sequences showed ≥96% similarity in BLAST analysis in all cases. Noteworthy, both *Bradyrhizobium* and *Burkholderia* species have been previously reported to associate with *Sphagnum* mosses²⁸. In fact, *Burkholderia* strains have been found to be transmitted by *Sphagnum* mosses over their life cycle, highlighting the relevance of this association^{[29](#page-6-24)}. In all, our fndings as well as other[s19,](#page-6-13)[28,](#page-6-23)[30](#page-6-25), suggest that nitrogen fxation may be important in facilitating plant growth under ombrotrophic, nitrogen-limited conditions in bog ecosystems.

Relatively few bacterial OTUs were shared between all three communities ([Fig.](#page-3-0) 3), but these shared OTUs accounted for 89% and 37% of the sequences for total and nitrogen-fxing bacterial communities, respectively. Interestingly, a large proportion of the taxa abundant in *Sphagnum* interstitial waters were rare in pools and *vice versa* [\(Fig.](#page-3-1) 4), suggesting a signifcant degree of habitat preference. Consequently, total bacterial communities, using Bray-Curtis dissimilarities, were found to difer in composition between *Sphagnum* interstitial water and pool samples (PERMANOVA: $F_{2,23} = 3.43$, $R^2 = 0.25$, $P = 0.001$; [Fig.](#page-4-0) 5a), but not between vegetated and clear pools (P>0.5). Similar fndings were obtained using weighted UniFrac dissimilarities (Supplementary Fig. S4a). Putative nitrogen-fxing bacterial communities gave comparable results to those of the total bacteria (Mantel $r= 0.65$, $P= 0.001$) ([Fig.](#page-4-0) 5b, Supplementary Fig. S4b). Interestingly, total bacterial communities were considerably less variable in composition in *Sphagnum* interstitial water samples than in pool samples ([Fig.](#page-4-0) 5a, Supplementary Figs S4a and S5a), which is interpreted as a sign of biotic homogenization. In contrast, nitrogen-fxing bacterial communities were equally variable between the three habitats ([Fig.](#page-4-0) 5b, Supplementary Fig. S4b).

When diferences in water chemistry were included in a non-metric multidimensional scaling ordination plot, *Sphagnum* interstitial water and pool samples grouped separately (PERMANOVA: $F_{2,23} = 8.9$, $R^2 = 0.45$, $P = 0.001$; [Fig.](#page-4-0) 5c). Conversely, no differences were found between vegetated and clear pools ($P > 0.05$). In addition, permutation dispersion showed that vegetated pools were more similar in chemical composition than *Sphagnum* interstitial water and clear pools (Supplementary Fig. S5b). A *post-hoc* ANOVA analysis found *Sphagnum* interstitial water samples having higher values of, phosphate, DOC and Chl *a* than pools (Supplementary Fig. 6). Using distance-based redundancy analysis we found that DOC and Conductivity explained 16% and 6% ($P=0.001$) of the variation in total [\(Fig.](#page-4-0) 5d) and nitrogen-fxing (not shown) bacterial communities, respectively.

We propose three non-mutually exclusive explanations for the observed patterns in community composition, related to diferences in (1) abiotic environmental conditions, (2) biotic interactions and (3) connectivity. Firstly,

Figure 3. Venn diagram showing the number of (**a**) shared total bacterial and (**b**) nitrogen-fxing OTUs. Te percentage of sequences associated with OTUs is shown in parentheses. CP, clear pools; VP, vegetated pools; SM, *S. magellanicum* interstitial waters.

Figure 4. Taxonomic composition and abundance patterns of the two groups of 16S rRNA-derived OTUs that showed the largest changes in abundance between *Sphagnum* **interstitial water and pools samples.** Te group Other encompasses unclassifed sequences together with orders representing ≤1% of total sequences. (**a**,**b**) OTUs that dominated pools (n=169). (**c**,**d**) OTUs that dominated *S. magellanicum* interstitial waters $(n=200)$. Taxonomic data are presented as per cent contribution of each bacterial order to total sequences. OTUs were selected following the procedure described by Ruiz-González *et al.*[15](#page-6-26) using a mean distance >15. CP, clear pools; VP, vegetated pools; SM, *S. magellanicum* interstitial waters.

the fact that bacterial communities were diferent in the *Sphagnum* interstitial waters and the pools indicates a strong habitat efect, which is consistent with the concept of species sorting (e.g., ref. [31\)](#page-6-27). Conductivity, a proxy for salinity, and DOC were the dominant drivers of the total bacterial communities ([Fig.](#page-4-0) 5d). Previous studies have identified DOC to be a key driver for aquatic microorganisms³², and it is well established that different aquatic bacterial taxa have different preferences in terms of carbon utilization³³. Similarly, salinity has been found to greatly influence bacterial diversity in aquatic ecosystems³⁴, although to our knowledge, this is the first time it has been reported in peatland water bodies.

Secondly, only 6–16% of the total variation in bacterial community composition could be explained by con-ductivity and DOC, suggesting that other abiotic or biotic factors may be of greater importance^{[35](#page-7-3)}. For example, it is well known that *Sphagnum* mosses are colonized by species-specifc microbial populations, which fulfl important functions (e.g. nutrient supply and pathogen defence) for moss growth and health^{[13,](#page-6-9)30}. For instance, *Sphagnum fallax* is colonized mainly by *Verrucromicrobia* and *Planctomycetes*, while *Sphagnum magellanicum* is dominated by *Alphaproteobacteria* and *Gammaprotobacteria*[13.](#page-6-9) Both phyla were well represented in *Sphagnum* interstitial water samples [\(Fig.](#page-1-0) 1). Zooplankton, protists and viruses could also have a role in structuring these communities^{[36](#page-7-4)}. Different bacterial taxa are known to vary in their resistance to both grazing and viral lysis^{[37](#page-7-5)} and mesocosm experiments have shown that predation by fagellates and ciliates can structure the composition of bacterial communitie[s38.](#page-7-6) Interestingly, a number of studies have reported that *Alphaproteobacteria* are resistant to predation (e.g., ref. [39](#page-7-7)), whereas others have indicated that members of the *Betaproteobacteria* are vulnerable to grazing (ref. [33](#page-7-1) and references therein). We note that *Alphaproteobacteria* dominated *Sphagnum* interstitial water samples, while *Betaproteobacteria* dominated pool communities ([Fig.](#page-1-0) 1).

Tirdly, we suggest that the distinction between *Sphagnum* interstitial water and pools communities may be linked to diferences in connectivity within these two habitats. *Sphagnum* interstitial water communities are more likely to be interconnected and further linked to terrestrial communities through hydrological networks, favouring microbial dispersal. Dispersal can lead to increased local species richness because it allows new species to enter communities¹⁸, leading to higher gamma diversity and reduced community dissimilarity¹⁷, patterns we observed in *Sphagnum* interstitial water samples. Similar fndings have been reported in previous studies for invertebrates^{[40](#page-7-8)}, plants¹⁷, and bacteria and viruses⁴¹, suggesting that this may be a universal phenomenon. In addition, *Sphagnum* interstitial water samples showed a greater proportion of *Acidobacteria* and *Gammaproteobacteria*, phyla commonly found in terrestrial communities¹⁶. In contrast, clear and vegetated pools can be seen as discrete, unconnected patches isolated from the surrounding soils, which allows for the development of more dissimilar communities, even if environmental conditions are similar, as those of vegetated pools ([Fig.](#page-4-0) 5c). Tis concept may be supported by the increase in abundance and diversity of common freshwater phyla such as *Betaproteobacteria* and *Actinobacteria*[15](#page-6-26),[16](#page-6-28),[42](#page-7-10) in pool communities relative to *Sphagnum* interstitial water communities. Indirect mechanisms such as dispersal-mediated trophic interactions can also generate apparent patterns of dispersal limitation in aquatic metacommunities⁴³.

In summary, this study showed that species sorting, as a result of both abiotic and biotic interactions, plays a pivotal role in explaining diferences in microbial (general and putative nitrogen-fxing bacteria) community diversity and composition across peatland water habitats. In contrast, within habitat diferences were better explained by the degree of microbial dispersal, which is higher in *Sphagnum* interstitial water communities. Tis has implications for conservation planning, as the high turnover in species composition in pool communities suggests that maximizing protected area is the key to maintaining diversity in the long term. It remains to be elucidated how these changes in microbial structure and composition will afect ecosystem function.

Materials and Methods

Study site, sampling and chemical analysis. Two peatlands from Tierra del Fuego Province (Argentina) separated by a distance of 50km were sampled in February 2014: Andorra peat bog (AN), located in the Andorra Valley (54°45′ S; 68°20′ W) and Rancho Hambre peat bog (RH), in Tierra Mayor Valley (54°44′ S; 67°49′ W) (Supplementary Fig. 7). Both peatlands are raised, nutrient-poor ombrotrophic peat bogs[44](#page-7-12). Samples were collected at twelve sampling points along a transect crossing each peat bog dome $(n = 24)$. The sampling points included three diferent habitat types (four replicates each): interstitial water from *Sphagnum magellanicum* matrix (SM), clear pools (CP) and vegetated pools (VP), with the latter two representing patches within the *Sphagnum* matrix. Vegetated pools showed prolifc growth of *Sphagnum* cf. f*mbriatum* and in some instances *Drepanocladus uncinatus*, while clear pools showed a mud bottom mainly formed by decaying *S. magellanicum*. Pools were sampled at the margins, while interstitial water from the *Sphagnum* matrix was collected by aseptically squeezing the mosses *in situ*. Sampled water (250 ml) was filtered through 0.22-₁ m sterile nitrocellulose membranes (Nalgene, Rochester, NY, USA), and membranes placed in RNAlater (Sigma-Aldrich, St. Louis, Mo, USA) and stored at 4 °C until further processing.

pH and conductivity were measured *in situ* using a Hach Sension 156multiparametric probe (Hach, Loveland, CO, USA). Nutrient (ammonium, phosphate, total N and total P) concentrations were measured using a Hach DR2800 spectrophotometer and their corresponding reagent kits, following standard methods described in the Hach DR2800 spectrophotometer procedures manual [\(www.hach.com\)](http://www.hach.com). Ammonium and phosphate concentrations were determined according to the salicylate (No. 8155) and ascorbic acid (No. 8048) Hach methods, respectively. Total N and total P were determined by performing an acid digestion with potassium persulphate and boric acid⁴⁵ followed by nitrate and phosphate determinations by the cadmium reduction (No. 8192) and ascorbic acid (No. 8048) Hach methods, respecrively. Dissolved organic carbon (DOC) was determined using the high temperature Pt catalyst oxidation method (Shimadzu analyser TOC-5000A, SM5310B technique) following the recommendations of Sharp *et al.*[46](#page-7-14). Chlorophyll *a* (Chl *a*) concentration corrected for phaeopigments was determined spectrophotometrically (for details see ref. [14](#page-6-10)). Metadata and water chemistry values are shown in Supplementary Table S2.

DNA extraction and amplicon sequencing. DNA was extracted from half of each flter, cut into small pieces with a sterile blade, using a PowerSoil DNA isolation kit (MoBio Laboratories, Carlsbad, CA, USA). PCR was performed in a single-step PCR using HotStarTaq Plus Master Mix Kit (Qiagen, Valencia, CA) with primer pairs 515F (5′-GTGYCAGCMGCCGCGGRA-3′) and 909R (5′-CCCCGYCAATTCMTTTRAG-3′) for the 16S rRNA gene[s47,](#page-7-15) and nifH1F (5′-TGYGAYCCNAARGCNGA-3′) and nifH2R (5′-ADNGCCATCATYTCNCC-3′) for the nifH genes⁴⁸. Triplicate PCR products were pooled after amplification, mixed in equal concentrations with samples targeting a specifc gene and purifed using Agencourt Ampure beads (Agencourt Bioscience Corporation, MA, USA). Sequencing was carried out on an Illumina MiSeq2000 using a paired-end approach⁴⁹ at the Molecular Research LP next generation sequencing service (<http://www.mrdnalab.com>).

16S rRNA gene sequences were analysed in MOTHUR^{[50](#page-7-18)}, following a previously established pipeline⁵¹, using the Silva core set (<http://www.arb-silva.de>) for alignment. Reads were removed from further analysis if at least one of the following criteria was met: (i) reads shorter than 200bp, (ii) number of ambiguous bases greater than 5, and (iii) presence of homopolymers with more than 8bp. One mismatch to the sample-specifc barcode and to the target-specifc primer were allowed. Afer removal of chimeras and poor quality reads, a total of 3625057 sequences were obtained. Sequences were grouped into OTUs, defned using a 97% sequence similarity cut-of. Taxonomic affiliation was determined using the naive Bayesian rRNA classifier in MOTHUR; 80% confidence level. Singletons were removed and each sample was rarefed to 39908 sequences (the lowest number of sequences in any sample). A total of 21221 OTUs (957792 sequences) were retained for analysis.

Amplicons of the *nifH* genes were analysed using the FunGene pipeline [\(http://fungene.cme.msu.edu/](http://fungene.cme.msu.edu/FunGenePipeline/) [FunGenePipeline/](http://fungene.cme.msu.edu/FunGenePipeline/))⁵². Filtered reads were translated into amino acid sequences and clipped at 60 aa. Further analyses were carried out on amino acid sequences clustered at 95% similarity[53.](#page-7-21) Representative sequences for each cluster were classifed using the NCBI algorithm BLASTP. Singletons were removed and each sample was rarefed to 911 sequences. 327OTUs (18220 sequences) from 20 samples, four samples did not give any amplifcation product, were kept for further analysis.

Statistical analysis. OTU richness and diversity indices (Shannon, Inverse Simpson and Pielou's evenness), together with rarefaction curves, Chao1 and Good's coverage estimates were calculated using MOTHUR. We applied mixed model ANOVA to determine signifcant diferences in diversity and water chemistry between habitat types using the phia package⁵⁴. In these analyses, we specified peat bog ID as a random factor. Abiotic data were standardized and pair-wise distances computed based on Euclidean distances. The community data matrices was Hellinger-transformed and the Bray-Curtis distance measure was used to generate a dissimilarity matrix. Weighted UniFrac dissimilarities were also obtained^{[55](#page-7-23)}. The structure of the bacterial community and the environmental variables were visualized using non-metric multidimensional scaling. The effect of abiotic data in explaining variation in bacterial community structure was assessed by distance-based redundancy analysis afer forward selection of the best set of parameters that could explain the variation in community composition. A permutational analysis of variance was used to test for diferences in composition between habitats, 'adonis' function (strata = peat bog ID) in vegan for R; whereas permutation dispersion was used to test for differences in their within-habitat dissimilarity, 'betadisper' function.

We used LEfSe analysis^{[56](#page-7-24)} to explore the presence of taxonomic groups that can serve as biomarkers for different classes (habitats). Statistically signifcant groups are reported with high LDA (Linear Discriminant Analysis) scores, which characterize the degree of consistency in relative abundance between features (genera) together with their efect relevance in each class. Correlations between the two biotic distance matrices were tested using the 'mantel' function in the ecodist package for R.

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Author Contributions

A.V., M.V.Q., D.C. and G.M. designed research. F.O. and M.V.Q. performed research. F.O. and A.V. analysed the data. A.V. wrote the manuscript. All authors commented on the manuscript at all stages.

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