Bryophyte-Cyanobacteria Associations during Primary Succession in Recently Deglaciated Areas of Tierra del Fuego (Chile)

María Arróniz-Crespo¹,², Sergio Pérez-Ortega³, Asunción De los Ríos⁴, T. G. Allan Green¹,⁴, Raúl Ochoa-Hueso⁵, Miguel Ángel Casermeiro⁶, María Teresa de la Cruz⁶, Ana Pintado¹, David Palacios⁷, Ricardo Rozzi⁸,⁹, Niklas Tysklind¹⁰, Leopoldo G. Sancho¹

¹ Dept. de Biología Vegetal II, Universidad Complutense de Madrid, Madrid, Spain, ² School of Environment Natural Resources and Geography, Bangor University, Bangor, Wales, United Kingdom, ³ Dept. de Biología Ambiental, Museo Nacional de Ciencias Naturales, MNCN-CSIC, Madrid, Spain, ⁴ Biological Sciences, Waikato University, Hamilton, New Zealand, ⁵ Dept. de Biogeografía y Cambio Global, Museo Nacional de Ciencias Naturales, Madrid, Spain, ⁶ Departamento Edafología, Universidad Complutense de Madrid, Madrid, Spain, ⁷ Departamento Geografía Física, Universidad Complutense de Madrid, Madrid, Spain, ⁸ Sub-Antarctic Biocultural Conservation Program, University of North Texas, Denton, Texas, United States of America, ⁹ Institute of Ecology and Biodiversity, Universidad de Magallanes, Puerto Williams, Chile, 10 School of Biological Sciences, Bangor University, Bangor, Wales, United Kingdom

Abstract

Bryophyte establishment represents a positive feedback process that enhances soil development in newly exposed terrain. Further, biological nitrogen (N) fixation by cyanobacteria in association with mosses can be an important supply of N to terrestrial ecosystems, however the role of these associations during post-glacial primary succession is not yet fully understood. Here, we analyzed chronosequences in front of two receding glaciers with contrasting climatic conditions (wetter vs drier) at Cordillera Darwin (Tierra del Fuego) and found that most mosses had the capacity to support an epiphytic flora of cyanobacteria and exhibited high rates of N₂ fixation. Pioneer moss-cyanobacteria associations showed the highest N₂ fixation rates (4.60 and 4.96 µg N g⁻¹ bryo. d⁻¹) very early after glacier retreat (4 and 7 years) which may help accelerate soil development under wetter conditions. In drier climate, N₂ fixation on bryophyte-cyanobacteria associations was also high (0.94 and 1.42 µg N g⁻¹ bryo. d⁻¹) but peaked at intermediate-aged sites (26 and 66 years). N₂ fixation capacity on bryophytes was primarily driven by epiphytic cyanobacteria abundance rather than community composition. Most liverworts showed low colonization and N₂ fixation rates, and mosses did not exhibit consistent differences across life forms and habitat (saxicolous vs terricolous). We also found a clear relationship between cyanobacteria genera and the new fixation rates, and mosses did not exhibit consistent differences across life

Introduction

Cryptogamic covers account for nearly half of the biological nitrogen (N) fixation on land [1], representing an important supply of N to terrestrial ecosystems. In particular, dinitrogen (N₂) fixation by cyanobacteria-moss associations has been found to contribute significantly to the N economy in high-latitude ecosystems [2–6] where it can represent the main source of new N [7–10]. For example, moss-associated cyanobacteria can provide 2–58% of the N input to Arctic ecosystems and in Antarctica the percentage can reach up to 84% [11]. In boreal forests, up to 60% of N can come from N₂-fixation in mosses [12]. In early primary succession, however, the role of cyanobacteria-moss associations on N inputs has received considerably less attention, despite bryophytes being often the first plants to colonize newly exposed terrain [13]. The lack of studies is particularly marked in temperate and subpolar regions of the Southern Hemisphere, where bryophytes account for more than 50% of the plant species [14]. Rates of N₂ fixation in moss associated with cyanobacteria can be highly variable. Previous studies in the 1980s and early 1990s have shown that N₂ fixation varies greatly among bryophyte species [8,15–17] which may be due to differences in the microhabitat characteristics where the mosses grow [2]. Moderate increases in temperature leads to a general increase in N₂ fixation [18] whereas shading [19] and N fertilization (e.g. atmospheric N
deposition or throughfall) results in reduced rates of N₂ fixation in these associations [4,12]. Moisture is also considered a major environmental factor influencing N₂-fixation on moss-cyanobacteria associations, with a positive relationship between water availability and N-fixation rates [3,20]. The diversity and density of cyanobacteria species associated with mosses have also been described as important factors explaining the variability in N₂ fixation rates [21,22]. However, little is still known about the existence of species-specific association between bryophyte and cyanobacteria species [23], although some studies have shown that host species identity could be a stronger driver of epiphytic cyanobacteria diversity and composition [22]. While these studies have already started to elucidate the factors that drive N₂-fixation rates, the influence of bryophyte-cyanobacteria associations on the biogeochemical cycling of N and, in particular, during early primary succession still needs further assessment.

In Tierra del Fuego, glacial retreat has been occurring since the end of the Little Ice Age (around 1750 to 1830) [24], exposing new terrain for ecosystem development by the process of primary succession. Glacier forelands are typically nutrient-poor [25]. However, our preliminary studies at the Cordillera Darwin (Tierra del Fuego) found rapid rates of vegetation growth and quick succession to stages dominated by Nothofagus tree species [26]. These tree-dominated stages were present on the southern side of the mountain range after 34 years of soil surface exposure [26], compared to 80 years on the northern side of the mountain range (Palacios et al., unpublished). As inputs of N through atmospheric deposition are very low in such high latitude region, possibly < 0.7 kg N ha⁻¹ yr⁻¹ [27], biological fixation can greatly exceed bulk precipitation. For example, biological N fixation inputs from bryophytes during early succession, e.g. [28] reached 22 kg N ha⁻¹ yr⁻¹ in a New Zealand glacier foreland. Therefore, we considered that N₂ fixation and its interaction with other biotic and abiotic factors could be a key driver of early plant succession in recently deglaciated areas of Tierra del Fuego. The combination of three attributes: 1) the difference in the speed of ecological succession found between the two glaciers on southern and northern slopes; 2) the strong climatic gradients characteristic of this area: with high rates of precipitation on the southwestern side and dry conditions on the northern side of the mountain range [29]; and 3) the low rates of anthropogenic N deposition, make the Cordillera Darwin an unique site to further elucidate the potential drivers of N₂ fixation rates on bryophyte-cyanobacteria associations and its impact on C and N accumulation during post-glacial primary succession.

Specifically, we aimed to evaluate the role of bryophyte-cyanobacteria associations during post-glacial primary succession because (i) moss establishment is a positive feedback process that enhance N inputs in newly exposed terrain [13] and (ii) moss-associated cyanobacteria can be an important contributor to N inputs through the process of N₂ fixation [28,30]. Our aims were (1) to describe how soil properties and N₂ fixation related parameters change along two pro-glacial chronosequences at the Cordillera Darwin (Tierra del Fuego), (2) how species identity (both bryophytes and cyanobacteria), climatic conditions (wetter vs. drier), and the stage of ecological succession, influence N₂ fixation rates of bryophyte-cyanobacteria associations, and thereby ecosystem N accumulation, and (3) whether any of these associations are species specific.

Materials and Methods

Ethics Statement

The Chilean National Forestry Corporation (CONAF) issued the scientific collecting permit for each location considered in this study. This study did not involve endangered or protected species.

Site description and chronosequence establishment

The study area is located at the Cordillera Darwin in the southwester part of Tierra del Fuego (Chile) (Fig. 1a). Around 80% of the mountain range is glaciated. One glacier was selected on each side of the mountain range for the present study; both glaciers have been receding steadily for several decades and present a well-defined sequence of moraine bands. The south-side glacier, SG (Pia Glacier, 54°46′S 69°40′W) flows into the east arm of Bahia Pia; plant succession goes from bare ground to full forest (stage of succession dominated by Nothofagus spp., >10 m high) in 34 years [26]. At the north-side glacier, NG (Parry Glacier, 54°41′S 69°23′W) the succession is slower and the equivalent stage of succession dominated by Nothofagus appears after 80 years of soil exposure (Palacios et al., unpublished data). A characteristic feature of the area is the steep meteorological gradients across the Cordillera Darwin with heavy precipitation and solid cloud cover being typical over the southern and western fjords, whilst to the north and east conditions are much drier [29]. This results in a stark contrast in annual precipitation between both glaciers. Precipitation rates where SG flows, Bahia Pia (1600 mm) are twice as high as where NG flows, Seno Almirantazgo fjord (800 mm) [29]. Mean annual air temperature, however, is relatively constant over the entire region, with only a slight drop from 5.9 °C at Punta Arenas to 4.5 °C at Bahia Pia [31].

The chronosequence in each glacier has been established by using several independent techniques to date surfaces: aerial photographs, dencrochronology, (sampling tree rings), and lichenometry (dating of rock surface from lichens growth) utilizing the growth rate and thallus size of the lichen Rhizocarpon geographicum (L.) DC [26].

Forest vegetation in the area consists of Nothofagus antarctica (G. Forst.) Oerst., and N. betuloides (Mirb.) as dominant tree species. After glacier retreat, the successional sequence at NG is characterized by pioneer lichens (Placopsis perragusa (Nyl.) Nyl., Rhizocarpon geographicum (L.) DC., Xanthoria candelaria (L.) Th. Fr.) and mosses (Dendroligotrichum squamosum (Hook. f. & Wilson) Cardot and Andreaea spp.), followed by dwarf shrubs (Empetrum rubrum Wahl. Ex Willd., Gaultheria mucronata (L. f.) Hook. & Arn.), perennial herbs (Gunnera magellanica Lam) and moss (Racomitrium spp.) vegetation assemblage. At SG the successional sequence goes from pioneer lichen (same species as in NG plus Placopsis stenophylla (Hue) J.M. Lamb and Placopsis perragusa L.M. Lamb) and the moss Ditrichum cylindricarpum (Mull. Hal., F. Muell, followed by perennial herbs vegetation assemblage (Gunnera magellanica Lam and Uncinia tenax Poeppl. ex Kunth) and the development of Nothofagus spp. forest (Fig. 1b).

Sample collection and transport

Field work took place in 2009 during the summer season which corresponds to the active period for N₂ fixation at high southern latitudes [32]. Sampling sites were established at increasing surface ages in front of both glaciers. There were six sites at SG with surface ages of 1, 4, 7, 10, 19 and 34 yrs and six sites at NG with surface ages of 5, 18, 20, 26, 66 and 80 yrs. At both glaciers, the younger site corresponds to bare ground close to the glacier front whereas the older site was covered within Nothofagus forests. Three transects were randomly established along the chronosequence in each glacier foreland and one independent sample was collected in each transect per site (surface age) so that there were three replicates per surface age for soil, each bryophyte species and cyanobacteria measurements.

Soil sampling. 200 g of soil samples were collected using the core method, samples were collected from the upper 0–4 cm of the soil. Samples were placed into, and sealed in, suitably sized plastic
bags. Collected samples were immediately frozen at −20 °C and returned frozen to the Universidad Complutense de Madrid for soil characterization.

**Bryophyte sampling.** The dominant bryophyte species at each site (surface age) along the chronosequence were collected. In total 16 bryophyte species (including both mosses and liverworts) with varying morphologies, life forms and habitats (saxicolous and terricolous species) were selected. These included 10 mosses *Dendroligotrichum squamosum* (Hook. f. & Wilson) Cardot, *Andreaea laxifolia* (Hook. f. & Wilson), *A. alpina* (Hedw.), *Acroschisma wilsonii* (Hook. f.) A. Jaeger, *Racomitrium didymum* (Mont.) Lorentz, *R. laevigatum* (A. Jaeger), *R. lanuginosum* (Schimp. ex R.S. Williams), *R.*

Figure 1. Study area and chronosequence description. (a) Map of the study area at the Cordillera Darwin in the southwestern part of Tierra del Fuego (Chile) showing the location of the south-side (SG) and north-side (NG) glaciers on each side of the mountain range. (b) Four representative sites along north-side glacier and south-side glacier chronosequences at Cordillera Darwin. At both glaciers, the younger site corresponds to bare ground close to the glacier front whereas the older site was covered within *Nothofagus* forests. Years since deglaciation per site are included in the picture.

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Cyanobacteria sampling. The diversity and distribution of cyanobacteria associated with bryophytes at each site (surface age) was determined by molecular analysis. A second set of bryophyte samples (around 10 shoots per species) were collected in the same bryophyte patches described above. Collectors used disposable laboratory latex gloves when taking samples. Samples were placed into, and sealed in suitably sized plastic bags. Collected samples were immediately frozen at −20°C and returned frozen to the Museo Nacional de Ciencias Naturales in Madrid (CSIC) where DNA from epiphytic cyanobacteria was extracted and analysed.

Soil chemical properties

Samples were stored at −20°C. Soils were air-dried, sieved (<2 mm) homogenized before conducting the following analysis:

Soil pH. Soil pH was measured in water suspension (soil-water, 1:2.5) [33].

Total soil N (TN) and total soil carbon (TC) were analysed in an autoanalyser system LECO CNS 2000II.

Soil extractable nitrogen. Soil nitrate (NO₃-N) and nitrite (NO₂-N) were extracted by shaking soil aliquots (30 g DW) in (1:5) water [33], filtered and analysed by ionic chromatography Metrohm 761 compact IC. Ammonium (NH₄-N) was extracted by shaking 10 g of soil in 30 ml of 0.5 M K₂SO₄ for 30 min, filtered and colorimetrically assessed [34].

Nitrogen-fixation measurements

Prior to N₂-fixation measurements, all bryophyte samples were reactivated for 48 h in a growth chamber, photosynthetic photon flux density (PPFD) of 100 µmol m⁻² s⁻¹ over the wavelength 400–700 nm, 10°C, with a 18 : 6 h photoperiod (light : dark) and maintained hydrated to ensure recovery (confirmed by chlorophyll fluorescence, data not shown). N₂-fixation rates of bryophyte-cyanobacteria associations were estimated under controlled conditions using the acetylene reduction assay (ARA) [35]. Revitalized shoot apices (around 10–15 shoots or 0.07–0.20 dry weight basis) were placed into 10 ml glass incubation vessels and sealed with a rubber septum. 1 ml of headspace was removed and replaced by 1 ml of acetylene (generated from CaC₂ and water), resulting in headspace enrichments of 10–20% (enrichment was calculated using a image analysis program. Eight species were sampled at more than one site and the other eight were sampled at a single site on the chronosequence. A full description of bryophyte species sampled by site are shown in (Table S1). Monospecific patches of fresh bryophyte species were collected, air dried and sent to the Universidad Complutense de Madrid in paper bags for analysis that included: (1) rates of N₂ fixation, (2) quantification of the abundance of epiphytic cyanobacteria, and (3) abundance of stable isotopes (¹⁵N and ¹⁵N) and shoot N and C contents.

Cyanobacteria abundance, composition and phylogenetic analysis

Cyanobacteria abundance by HPLC analysis of echinenone. Bryophyte samples (~3–10 cm² of fresh tissue per sample depending on the species) were deep-frozen until pigment analyses. Pigments were extracted in 95% cool acetone in the presence of sodium ascorbate. Echinenone (a cyanobacterial pigment used as a proxy of cyanobacteria abundance) was separated by HPLC according to the method of [39] slightly modified. After passing through a 0.45 µm nylon filter, 25 µL of the extract was injected into a C18 column. The mobile phase rate was 1.2 mL min⁻¹ and the elution time lasted 30 min. Solvents for HPLC analysis were degassed before use by bubbling helium. The HPLC system was equipped with a photodiode array detector. For peak identification and quantification, a pure commercial standard was used. The relative abundance of cyanobacteria was expressed as the amount of echinenone on a bryophyte surface area basis (nmol echinenone/cm² bryophyte). Digital pictures of each sample were taken before freezing and the surface area of each sample was calculated using a image analysis program.

DNA extraction and polymerase chain reaction (PCR), denaturing gradient gel electrophoresis (DGGE). For cyanobacteria DNA extraction, bryophyte samples (around 200 mg dry weight) were introduced into 15 ml falcon tubes with distilled water and vigorously shaken (maximum speed) for 10 over an 18-h period in 10 samples with known different cyanobacteria abundance (data not shown). Ethylene resulting from the reducing activity was measured by gas chromatography. N₂-fixation rates were calculated as nanomoles of acetylene reduced per gram bryophyte dry weight per hour (nmol C₂H₂ g⁻¹ dw h⁻¹). Theoretical stoichiometric ratio of acetylene to nitrogen (3:1) is recommended to be corrected by parallel uptake of labelled ¹⁵N₂ for each organism and also, possibly, for location [36,37]. Thus, to estimate N₂ fixation rates (¹⁵N₂ uptake), consecutive incubations of the same samples with ¹⁵N₂ after ARA was performed following [9]. 2 ml of headspace were replaced by 2 ml ¹⁵N₂ gas (98 atom % ¹⁵N enriched), resulting in headspace enrichments of 20–30% and, after 18 h incubation at the same conditions as ARA samples, were immediately oven dried (60°C for 48 h), ground in a ball mill and sent for ¹⁵N enrichment and total N analysis (UCDavis Stable Isotope Facility, University of California, Davis, USA). Three control samples for each bryophyte-cyanobacteria association were used to determine the natural abundance ¹⁵N. The amount of N fixed was calculated according to [6,38]. N₂-fixation rates were calculated as micrograms of ¹⁵N incorporated per hour per gram of bryophyte on a dry weight basis and expressed as micrograms of fixed N per day (µg N g⁻¹ dw bryophyte d⁻¹).

ARA was highly correlated with ¹⁵N₂ uptake (N= 89, R² = 0.689, P<0.001). ARA measurements were used to compare N₂ fixation rates between taxa and rates reported by other authors. ¹⁵N₂ uptake was used to evaluate the potential impact of N₂ fixation on ecosystem nitrogen (N) accumulation.

Stable isotope analysis and shoot N and C content

All isotopic analyses were conducted at the UCDavis Stable Isotope Facility, Department of Plant Sciences, University of California, Davis, USA. δ¹⁵N analyses were performed by elemental analyser/continuous flow isotope ratio mass spectrometry using a ANCA-GSL elemental analyser interfaced coupled with a PDZ Europa 20-20 isotope ratio mass spectrometer. Atom% was calculated from δ¹⁵N values. Total tissue C and N was also analysed.
minutes in a vortex using a falcon tube adapter to release the epiphytic cyanobacteria from bryophytes. The extracted cyanobacteria contained in the water were removed with a 5 ml pipette and cells were concentrated by centrifugation in an ultracentrifuge at 3000 rpm for 5 min. The concentrated cyanobacteria cells were used for DNA extraction.

Denaturing gradient gel electrophoresis (DGGE) of cyanobacterial PCR-amplified 16S rRNA gene fragments was used to survey the genetic diversity of isolated epiphytic cyanobacteria. Total genomic DNA for DGGE analysis was extracted in three replicates from the cyanobacteria isolates samples using CTAB extraction method described in [40]. A fragment of the cyanobacterial 16S rRNA gene suitable for DGGE analysis was amplified in each sample from total genomic DNA using the following primer pair: CYA359fGC and CY781r [41]. Each 25 μl volume of PCR mix [75 mM Tris pH 9.0/50 mM KCl/20 mM (NH₄)₂SO₄] contained 1 unit of Taq polymerase, 0.2 mM of each dNTPs, 0.4 mM of each primer, 100 mg of bovine serum albumin, 1.5 mM of MgCl₂, 5 μl de 5×Taq Master PCR enhancer (Prime) and ca 10–50 ng genomic DNA. Annealing conditions were 60°C. The DGGE system was used for the analysis of cyanobacterial fragments of 16S rRNA gene according to the manufacturer’s instructions. Acrylamide gels (6%) with a 30 to 60% urea-formamide denaturing gradient were prepared. Lanes were loaded with 22 μl of PCR product, run at a constant 200 V for 7 h at 60°C, and EtBr stained to visualize and photograph the resultant bands according to [42]. The most predominant bands were excised and incubated 1 h at 60°C in water before PCR amplification. The eluted DNA was reamplified as described above but using primers devoiding the GC clamp. The PCR products were cleaned on Quick Spin columns. Both complementary strands were sequenced separately at the MACROGEN sequencing company.

For sequence analysis, we obtained closely related sequences with known taxonomy using Blast searches in GenBank. Later, sequences obtained from DGGE bands were reduced to OTUs in mothur v.1.28 [43] using a cutoff of 0.01% [44]. A reference sequence for each OTU was chosen for subsequent analysis by means of the command get_ottrep in mothur v.1.28. 20 sequences corresponding to each of the 0.01% cut-off OTUs were aligned with a total of 29 sequences retrieved from the GenBank (Table S2). Alignment of all sequences were carried out in Muscle 3.8.31 [45] and visually checked in Bioedit 7.0.5.2 [46]. Nucleotide substitution model was statistically selected with the help of jModelTest [47], program available at http://darwin.uvigo.es). Model selection was made according to the Akaike’s information criterion [AIC, [48]; the General Time Reversible substitution model [49], with estimation assuming a gamma distribution (GTR+G) had the lowest –lnL value. Phylogenetic relationships among OTUs and sequences with known taxonomy were studied by means of Maximum Likelihood (ML) and Bayesian phylogenetic inference. ML analysis were carried out in PHYML 3.0 [50] through the webserver http://www.atgc-montpellier.fr/phyml/ using 1,000 bootstrap repetitions in order to assess node support. Bayesian analyses were carried out in Mr. Bayes 3.1.2 [51] running eight different chains and 5,000K generations. 50% majority rule consensus tree is shown in (Fig. 2).

Sequences generated in this study were deposited in GenBank under accession numbers KJ576732-KJ576785.

Statistical and data analyses
We calculated the Shannon index \( H = -\sum (ni/N) \ln (ni/N) \) [52], where \( ni \) is the number of times we found each ith OTU and \( N \) is the total number of unique OTUs \((N = 20) \) found in the study area, in order to evaluate changes in diversity of epiphytic cyanobacteria along the chronosequence \((H_{\text{north}})\) and between bryophytes species \((H_{\text{specie}})\).

To be able to compare parameters between glacier chronosequences with different sampling site ages we first tested if the different stages of ecological succession, identified by the soil development transformation trends observed along the chronosequence, were comparable between the two glacier chronosequences; we used T; TN and soil pH as descriptors of soil development status at each site and extracted one general factor component of soil development using principal component analysis [53]. The resulting scores were tested with two-way ANOVA to detect differences on the stage of ecological succession along the chronosequence and between glaciers, with sampling sites (0 to 5) and glacier location (north vs. south) as fixed factors. These analyses indicated that the two chronosequences were comparable despite the different absolute soil ages where the samples were collected (Fig. S1). Differences between glaciers on soil chemical properties, N₂-fixation rates \(( ^{15} \text{N} \text{uptake})\), abundance of epiphytic cyanobacteria, and C : N ratio in bryophyte tissue were then tested with two-way ANOVAs. All data were tested for homoscedasticity (Levene’s test for equal variances) and normality (IBM SPSS statistics, Armonk NY, USA). When data did not meet these assumptions, they were rank or Log₁₀ +1 transformed prior to statistical analyses. Analyses were carried out with IBM SPSS Statistics 19 (IBM SPSS statistics, Armonk NY, USA).

A fundamental assumption about chronosequences is that the communities and ecosystems of the younger sites are developing in the same temporal sequence as the older sites developed (termed a space-for-time substitution). At any location on the chronosequence the vegetation present represents what will eventually form on younger sites and which will, itself, change to that found on the older sites present. As such, the assumption of independence is violated. We used non-parametric Kruskall-Wallis test to evaluate changes of parameters measured at the different stages of ecological succession (mean values per site) along each glacier chronosequence separately, using stage of ecological succession (sampling site) as the grouping factor. When significant differences were found, Mann-Whitney U-test post hoc tests were run to detect differences between sites. Bonferroni’s correction was used to determine the level of significance of the post hoc tests.

At species level, we used conditional inference trees (CTREE) [54] to assess and visualise the relative importance of bryophyte species identity (species effect), climatic conditions (glacier location: north vs. south), and stage of ecological succession (sampling site effect) in determining N₂ fixation capacity and cyanobacteria abundance on bryophytes. CTREE work by first testing the null hypothesis of independence between the predictors (here bryophyte species, glacier and site) and response variable (here N₂ fixation); if dependence is found, then determining the best split value for the predictor with the strongest effect on the response; and third, repeating the first two steps with each branch until independence cannot be rejected. Relationships are represented as dichotomous trees with nodes indicating split points for significant predictor variables, and terminal nodes for the final groups of responses. Being nonparametric, CTREE make no assumptions about the data, and are appropriate where nonlinear relationships between predictive and response variables and complex interactions between variables, such as covariance, may be expected [54]. CTREE were constructed in R 3.0.1 (R Development Core Team, 2013) with the package PARTY [55], with 10,000 resamples of Bonferroni-adjusted \( \alpha = 0.05 \) significance levels and with at least one
observation into each daughter node. N₂ fixation and cyanobacteria abundance data were log +1 transformed and site included as an ordered categorical variable.

Linear regression analysis was used to evaluate whether abundance of epiphytic cyanobacteria or its diversity on host bryophyte (H. speciosa) had a larger influence on the rate of N₂ fixation.

Correspondence analysis in each glacier separately were used to graphically examine the relationships between OTUs of cyanobacteria and the different stages of ecological succession and also to detect selectivity associations between cyanobacteria OTUs and bryophyte species (6 species with cyanobacteria colonies sampled at more than one site were used in this analyses). Analysis were carried out in R 3.0.1 (R Development Core Team, 2013) using the package ca [56].

**Results**

**Genetic diversity of isolated epiphytic cyanobacteria**

A total of 52 bands (18 from NG and 34 from SG) were recovered from DGGE analyses and subsequently sequenced, producing 28 unique sequences. After their collapse into OTUs (0.01% cutoff), 20 cyanobacteria putative taxa were obtained. The phylogenetic analysis of the relationships between these OTUs and sequences with known taxonomy obtained from the GenBank is depicted in Fig. 2. Seven 0.01% cutoff OTUs could be assigned to the species aggregate *Nostoc commune*. Six OTUs (5, 7, 9, 12, 15, 17) were assigned to unidentified Nostocaeae because no clear relationship was found in the phylogenetic tree with any certain genus. A similar case is presented for OTUs 10, 13, 14, 16, and 18, which did not show a clear and well supported clade with sequences of *Stigonema* included in the phylogenetic analysis. Nevertheless, BLAST searches of these OTUs recovered sequences identified as *Stigonema*. In both cases, the most likely explanation for such lack of phylogenetic support in our tree for these two OTU groups is the shortness of fragment of the 16S gene used in the analyses. Finally, OTU 20 is clearly and statistically supported related to *Leptolyngbya* and OTU 19 is likewise related to *Microcoleus* (Fig. 2).

**Soil properties of glacier forelands**

Overall comparisons between both glaciers (two-way ANOVAs) showed no differences on pH, TC and TN between glaciers (NG vs SG) and higher concentration of inorganic N at SG (ANOVA P= 0.037). At NG, the concentration of inorganic N in young soils (younger than 26 years) was lower (from 0.16 to 8.23 mg kg⁻¹).
compared to concentrations observed in soils at SG very early after glacier retreated (15.42 mg kg⁻¹ after 4 years soil exposure and 34.03 mg kg⁻¹ after 10 years, Fig. 3d). Within the Nothofagus forests inorganic N concentration in the soil was nearly 3-fold higher at SG (13.05 mg kg⁻¹) compared to NG (4.52 mg kg⁻¹).

All soil parameters showed a significant overall site effect (ANOVA P<0.001) and no interaction between glacier location (northern vs. southern) and site (0 to 5). Within each chronosequence (non-parametric tests), soil pH decreases from 6.4 to 4.4 in NG (P<0.01) and from 6.1 to 4.5 in SG (P<0.005), with the higher values always found in the younger soils (Fig. 3a). TC increased from very low values (0.33% at NG and 0.24% at SG) to over 33% at NG (P<0.05) and 39% at SG (P<0.05) in the oldest soils (50 and 34 years of soil exposure respectively) inside the Nothofagus spp. forest (Nothofagus-dominated state, Fig. 3b). TN also increased along the chronosequence from 0.02% in NG and undetectable levels in SG to 0.99% in NG (P<0.05) and 1.51% in SG (P<0.005) inside the forest site (Fig. 3c). Soil extractable inorganic N (NO₃⁻ + NO₂⁻ + NH₄⁺) followed a similar trend increasing with time since deglaciation at both glaciers (P<0.05 for NG and P<0.01 for SG, Fig. 3d).

**Nitrogen-fixation, cyanobacteria abundance and shoot C : N contents**

N₂ fixation rates (¹⁵N₂ uptake) and epiphytic cyanobacteria relative abundance, when averaged across all bryophyte species per site, were higher at SG compared to NG (ANOVA P<0.001 and P<0.01 respectively). A significant overall interaction (ANOVA P<0.001) between glacier locations (northern vs. southern) and the stage of ecological succession (sampling sites: 1 to 5) suggested that variation of N₂ fixation rates and cyanobacteria abundance along each chronosequence differed between glaciers. At NG, N₂ fixation rate (Fig. 3e) followed an optimum increasing from the lowest rates at 18 and 20 year old sites (0.52 and 0.3 μg N g⁻¹ DW bryo. d⁻¹, respectively) to rates that were 2-fold higher in intermediate-aged sites (0.94 and 1.42 μg N g⁻¹ DW bryo. d⁻¹) at 10 and 66 years old sites respectively) and then declined in the 80 years old forest site (0.00 μg N g⁻¹ DW bryo. d⁻¹). Epiphytic cyanobacteria abundance (Fig. 3f) followed a similar trend with the highest concentration at 66 years old site (42.99 mmol echin. cm⁻²) and lowest concentration at 80 and 20 years old soil (4.5 and 7.94 mmol echin. cm⁻²). In contrast, at SG the highest N₂ fixation rate and cyanobacteria abundance was found in young soils (4.60 and 4.96 μg N g⁻¹ DW bryo. d⁻¹ and 51.30 and 53.33 mmol echin. cm⁻²) at 4 and 7 year old sites respectively) compared to older sites (ranging from 0.64 to 1.34 μg N g⁻¹ DW bryo. d⁻¹ and 6.46 to 23.96 mmol echin. cm⁻²), at soil age 10 to 34 years old (Fig. 3g).

Shoot C : N ratios of bryophyte tissue, when averaged across all bryophyte species per site, did not differ between glacier (ANOVA F=0.165) but change with soil age (ANOVA P<0.001). Shoot C : N increased with time since deglaciation at NG (P<0.001) with the lower values always found in the younger soil (Fig. 3g). At SG shoot C : N ratios showed the same pattern but the effect of soil age was not significant (Fig. 3g).

**Differences on bryophyte N₂ fixation capacity and cyanobacteria abundance**

The conditional inference tree (Fig. 4a) revealed that both species identity and the stage of ecological succession (sampling site) were important variables in the N₂ fixation rates at species level (ARA, mmol C₂H₂ g⁻¹ dm bryo. h⁻¹). Three species of mosses (Ditrichum cylindricarpum, Racomitrium suberuptipulum and R. laevigatum) showed the highest rates of N₂ fixation (Table 1 and node 3 in Fig. 4a); Intermediate N₂ fixation rates were found for a second group of species (node 4), namely Racomitrium didymum, R. lanuginosum, Archaeisella walloni and the liverwort Cryptochaeta grandiflora. An increment of N₂ fixation rates was detected between younger sites (node 5: sites 1–3) and older sites (node 6: sites 4 and 5) for node 4. However, a more detailed analysis by individual species, revealed that there was an effect of the stage of ecological succession on N₂ fixation rates only on the two most common species sampled in this study, R. didymum and A. walloni (statistical data not shown). The remainder of the species (node 7) had either low N₂ fixation rates as detected in Dicranuloma chilense, Andreae laxifolia, Andreae alpina and Anastrophyton udowratulphen, or zero N₂ fixation rates, as in the liverworts Cladomatosus undatus, Hypnodendryphyllum densifolium, Chloroscyphus leptanthus, Heteroscyphus magellanicus and the endohydric moss Dendrogyrichtrum squammosum (Table 1 and Fig. 4a). With our dataset, no effect was detected for glacier location (northern vs. southern) at species level.

The conditional inference tree (Fig. 4b) also showed a significant effect of bryophytes species identity on cyanobacteria relative abundance (mmol echinomone/cm² bryophyte) with the highest colonization found in the mosses Ditrichum cylindricarpum, Racomitrium suberuptipulum, R. laevigatum, R. lanuginosum and the liverwort Cryptochaeta grandiflora (Table 1 and node 2 in Fig. 4b). Variability of N₂ fixation rates between bryophyte species was best explained by changes in epiphytic cyanobacteria abundance (positive linear regression: $R^2 = 0.44, P < 0.001$, Fig. 5a) rather than the Shannon diversity index of cyanobacteria on host bryophytes (positive linear regression: $R^2 = 0.19, P = 0.012$, Fig. 5b).

**Community structure of epiphytic cyanobacteria**

The Shannon index of epiphytic cyanobacteria OTUs per site ($H'_{otu}$) at NG rose from 0.29 to a maximum of 0.96 at 26 years followed by a steady decline to original level at the oldest site. At SG, $H'_{otu}$ increased sharply from 0.38 and 0.32 at the two youngest sites to 1.37 at the oldest site within Nothofagus forest (Fig. 3b). Correspondence analysis showed a clear relationship between cyanobacteria genera and the stage of ecological succession at both glaciers. At NG (Fig. 6a), sites 2 and 3 share similar cyanobacteria communities, Nostoc commune was present at site 2 (OTU 1 & 3) and site 3 (OTU 6) whereas unidentified Nostocaceae was confined to site 3 (OTU 5, 7 & 15) and site 4 (OTU 17); Stigonema spp was found just at site 4 (OTU 16) and Microcoleus spp. (OTU 19) at site 5. At SG (Fig. 6b), all sites differed in their cyanobacteria communities, Nostoc commune was present at sites 1 (OTU 2), site 2 (OTU 1), site 3 (OTU 4 & 8) and site 5 (OTU 11); unidentified Nostocaceae at site 4 (OTU 9) and site 5 (OTU 15 & 12) and Stigonema spp. at site 4 (OTU 18) and site 5 (OTU 16, 13, 10 & 14). Leptolyngbya spp. (OTU20) was found at site 1. No relationship was observed between the genus of cyanobacteria and bryophyte species (Table 1 and Fig. S2).

**Discussion**

Patterns of soil properties and N₂ fixation related parameters along chronosequences

The trend observed in soil development followed the classic sequence reported for glacier foreland chronosequences [57,58]. Soil pH fell and total C and N contents increased with surface age along both chronosequences. However, the time required differs significantly to rates reported in other proglacial areas. Soils developed from barren rocks to Histosol, an organic soil, with 39% C in SG and 33% C in NG after 34 and 80 years of soil exposure, respectively (Fig. 3b). This contrasts with 2% C reported in other
studies reached 80 years after deglaciation [57]. Total soil N increased from very low or undetectable values at the youngest sites to 1.5% in SG and 0.99% in NG in the forest soil over this short period, again contrasting with concentration of 0.2% TN after 80 years exposure observed in the Central Alps [57]. These results suggest that conditions for vegetation succession are more favourable under the oceanic climatic conditions of Southern Hemisphere high latitude regions, such as Tierra del Fuego.

Figure 3. Patterns of soil properties and N₂ fixation related parameters along chronosequences. Site characteristics along the chronosequence in the north-side glacier (NG: black circles) with site soil (surface) age of 5, 18, 20, 26, 66 and 80 yrs and south-side glacier (SG: white circles) with site soil (surface) age of 1, 4, 7, 10, 19 and 34 yrs. At both glaciers, the younger site corresponds to bare ground close to the glacier front whereas the older site was covered within Nothofagus forests. Significant changes of parameters along each glacier chronosequence is indicated by *, p<0.05, **, p<0.01***, p<0.001*** (Krukall-Wallis test). Different letters indicate significant differences between site ages after Mann–Whitney U-test post-hoc (Bonferroni corrected), bold letters are used for NG. Statistical differences on soil properties and N₂ fixation related parameters between the two glacier chronosequences are indicated in the result section.

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Soil development along the two chronosequences, from bare
ground to full forest, showed similarities in soil pH and total C and
N concentration, indicating similar stages of soil development at
both sites (Fig. 3a,b,c and Fig. S1). However, the time-required for
C and N accumulation was much less at SG (Fig. 3d). In addition,
after 4 and 7 years of soil exposure extractable inorganic N
concentration was already much higher in the soil at SG
(13.42 mg kg\(^{-1}\) and 12.51 mg kg\(^{-1}\) respectably, Fig. 3d) than at
NG, where values remained low (0.16 to 8.23 mg kg\(^{-1}\)) even after
26 years of soil exposure. We also found the highest rates of N\(_2\)
fixation and cyanobacteria abundances at SG.
fixation and cyanobacteria abundance in the pioneer terricolous moss *Ditrichum cylindricarpum* at 4 and 7 years soil exposure in SG (Fig 3e,f Table 1), where it forms a dense carpet in the soil (Fig. 1b). By contrast, in NG biological N$_2$ fixation and epiphytic cyanobacteria abundance on bryophytes was very low during the early stages of succession (Fig. 3e,f). It is likely that higher annual rainfall on SG is associated with higher rates of N$_2$ fixation on bryophyte-cyanobacteria associations in young soils, since fixation of N is strongly controlled by moisture availability [30]. Earlier studies have shown the importance of free living colonies of diazotrophs for C and N inputs during early stages of succession in glacial forelands [59–62] which could contribute to nutrient accumulation at our sites. Fixation of N by cyanobacteria associated with bryophytes that colonize soil surfaces very early after glacier retreat, as shown by our results, could also help accelerate soil development in recently deglaciated areas under wetter conditions. As succession proceeds in SG, terricolous mosses are replaced by perennial herbs like *Gunnera magellanica*, which forms symbiotic associations with the cyanobacterial genus *Nostoc* [63], and becomes the dominant species together with *Uncinia tenuis*, *Nothofagus* forests.

### Table 1. Mean (N = 3) N$_2$ fixation rates (nmol C$_2$H$_2$ g$^{-1}$ dm bryo. h$^{-1}$) cyanobacteria abundance (nmol echinenone/bryophyte cm$^2$) and cyanobacteria genera associated with each individual bryophyte species along both glacier chronosequences.

<table>
<thead>
<tr>
<th>Glacier</th>
<th>Site</th>
<th>Bryophyte species</th>
<th>Mean reduction rate (nmol C$_2$H$_2$ g$^{-1}$ dm bryo. h$^{-1}$)</th>
<th>Cyanobacteria abundance (nmol echin. cm$^{-2}$)</th>
<th>Cyanobacteria identification</th>
</tr>
</thead>
<tbody>
<tr>
<td>SG 1</td>
<td>Ditrichum cylindricarpum</td>
<td>283.4</td>
<td>55–520</td>
<td>51.5 (3.2)</td>
<td>N. commune. Leptolymania sp</td>
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<tr>
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<td>Ditrichum cylindricarpum</td>
<td>270.1</td>
<td>70–455</td>
<td>53.3 (8.2)</td>
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<tr>
<td>SG 4</td>
<td>Racotritium subciriplus</td>
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<td>43–144</td>
<td>30.7 (12.1)</td>
<td>N. commune</td>
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<td>SG 5</td>
<td>Racotritium subciriplus</td>
<td>81.6</td>
<td>49–142</td>
<td>40.3 (36.2)</td>
<td>N. commune. Unidentified Nostocaceae Stigonema</td>
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<tr>
<td>NG 3</td>
<td>Racotritium laevigatum</td>
<td>63.5</td>
<td>33–125</td>
<td>8.5 (6.9)</td>
<td>N. commune</td>
</tr>
<tr>
<td>NG 4</td>
<td>Racotritium laevigatum</td>
<td>59.9</td>
<td>30–101</td>
<td>37.2 (9.4)</td>
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<td>NG 2</td>
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<td>7.7</td>
<td>3–16</td>
<td>14.5 (14.5)</td>
<td>N. commune</td>
</tr>
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<td>NG 3</td>
<td>Racotritium didymum</td>
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<td>12–56</td>
<td>8.5 (5.3)</td>
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<td>NG 4</td>
<td>Racotritium didymum</td>
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<td>22–80</td>
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<td>0</td>
<td>0.3 (0.3)</td>
<td>N. commune</td>
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<tr>
<td>NG 5</td>
<td>Racotritium didymum</td>
<td>20.7</td>
<td>7–34</td>
<td>21.5 (10.7)</td>
<td>Unidentified Nostocaceae</td>
</tr>
<tr>
<td>NG 1</td>
<td>Andreaea laxifolia</td>
<td>66.6</td>
<td>53–75</td>
<td>39.4 (19.6)</td>
<td>N. commune. Stigonema</td>
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<tr>
<td>NG 3</td>
<td>Andreaea laxifolia</td>
<td>8.7</td>
<td>0–26</td>
<td>15.8 (15.5)</td>
<td>Non found</td>
</tr>
<tr>
<td>NG 2</td>
<td>Cryptochila grandiflora</td>
<td>7.7</td>
<td>3–16</td>
<td>14.5 (14.5)</td>
<td>N. commune</td>
</tr>
<tr>
<td>NG 5</td>
<td>Cryptochila grandiflora</td>
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<td>3–124</td>
<td>17.1 (4.5)</td>
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<td>NG 2</td>
<td>Acroschisma wilsonii</td>
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<td>0–3</td>
<td>11.8 (6.0)</td>
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</tr>
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<td>Acroschisma wilsonii</td>
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<td>1–14</td>
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<tr>
<td>SG 3</td>
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<td>25–40</td>
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<td>SG 4</td>
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<td>18.4</td>
<td>9–28</td>
<td>19.5 (6.3)</td>
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<tr>
<td>SG 5</td>
<td>Acroschisma wilsonii</td>
<td>39.8</td>
<td>8–72</td>
<td>9.9 (4.9)</td>
<td>Unidentified Nostocaceae</td>
</tr>
<tr>
<td>NG 4</td>
<td>Racotritium lanuginosum</td>
<td>39.2</td>
<td>10–98</td>
<td>76.5 (22.0)</td>
<td>Stigonema</td>
</tr>
<tr>
<td>NG 3</td>
<td>Dicranoloma chilense</td>
<td>19.2</td>
<td>9–33</td>
<td>2.7 (2.7)</td>
<td>Unidentified Nostocaceae</td>
</tr>
<tr>
<td>NG 5</td>
<td>Dicranoloma chilense</td>
<td>2.9</td>
<td>0–9</td>
<td>5.9 (5.9)</td>
<td>Non found</td>
</tr>
<tr>
<td>NG 5</td>
<td>Anastrophyllum involutifolium</td>
<td>3.5</td>
<td>0–11</td>
<td>5.7 (5.1)</td>
<td>Non found</td>
</tr>
<tr>
<td>SG 4</td>
<td>Clasmatocolea humilis</td>
<td>1.5</td>
<td>0–3</td>
<td>15.2 (4.5)</td>
<td>Stigonema</td>
</tr>
<tr>
<td>NG 2</td>
<td>Andreaea alpina</td>
<td>0.9</td>
<td>0–3</td>
<td>2.7 (2.7)</td>
<td>Non found</td>
</tr>
<tr>
<td>NG 5</td>
<td>Blepharidophyllum densifolium</td>
<td>0.0</td>
<td>0</td>
<td>6.1 (6.1)</td>
<td>Microcoleus</td>
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<tr>
<td>SG 5</td>
<td>Chlomyces leptanthus</td>
<td>0.0</td>
<td>0</td>
<td>4.8 (4.8)</td>
<td>Non found</td>
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<tr>
<td>NG 1</td>
<td>Dendroligotrichum squamosum</td>
<td>0.0</td>
<td>0</td>
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<td>Non found</td>
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<tr>
<td>NG 2</td>
<td>Dendroligotrichum squamosum</td>
<td>0.0</td>
<td>0</td>
<td>2.5 (2.5)</td>
<td>Non found</td>
</tr>
<tr>
<td>NG 5</td>
<td>Heterococcus magellanicus</td>
<td>0.0</td>
<td>0</td>
<td>0.1 (0.1)</td>
<td>Non found</td>
</tr>
</tbody>
</table>

In both glaciers the younger site (site 1) correspond to early succession site close to the glacier front whereas the older site (site 5) was settled within Nothofagus forests. Sites 3–4 represent intermediate stages of ecological succession. No species were sampled at site 0. SE for cyanobacteria abundances is shown in brackets. NG: north-side glacier; SG: south-side glacier of the mountain range.
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N₂-Fixation in Bryophytes from Deglaciated Areas

**Differences on bryophyte N₂ fixation capacity and cyanobacteria abundance**
Few studies have investigated interspecific or intraspecific variation in N₂ fixation capacity on bryophytes, particularly in the context of post-glacial primary succession. In our study, most bryophyte species have the ability to support an epiphytic flora of cyanobacteria. However, N₂ fixation rates varied between and within bryophyte species (Table 1). In line with results reported in previous studies [5], we detected differences in N₂ fixation rates between mosses and liverworts. Five of the six studied liverworts showed no or very low rate of N₂ fixation and cyanobacteria abundance (Table 1), and only one species, the terricolous liverwort _Cryptochila grandiflora_, showed high fixation rates (52.6 nmol C₂H₂ g⁻¹ dm bryo. h⁻¹) and cyanobacteria colonization. Among moss species, _Dendroligotrichum squamosum_ also showed no N₂ fixation or cyanobacteria colonization, possibly due to their rigid, water repellent leaves which may confer less favorable moisture conditions. However, morphology or habitat could not explain variations found among other moss species. Both terricolous (_Ditrichum cylindricarpum_, _Racomitrium laevigatum_, _R. lanuginosum_) and saxicolous species (_Racomitrium subcrispipilum_, _R. didymium_, _Acroschisma wilsonii_) showed high N₂ fixation rates (over 40 nmol C₂H₂ g⁻¹ dm bryo. h⁻¹) compared to rates reported in Arctic [8] and sub-Antarctic [16] ecosystems. Regarding intraspecific variations, we observed that the rates of N₂ fixation of the saxicolous mosses _Racomitrium didymum_ and _Acroschisma wilsonii_ (the most frequently sampled species in the study area) were affected by the stages of ecological succession. The rate of N₂ fixation significantly increases on _R. didymum_ and _Acroschisma wilsonii_ populations that grow at sites that have been longer deglaciated, reflecting growth of the cyanobacterial population as vegetation develops when growing epiphytically on these mosses.

**Community structure of epiphytic cyanobacteria**
In the present study, we also aimed to evaluate the influence of cyanobacteria community structure on bryophyte N₂ fixation rates. We found that abundance of epiphytic cyanobacteria appears to be a better factor explaining differences in N₂ fixation rates (Fig 5) than epiphytic cyanobacteria diversity on host bryophyte species (Fig. 5). It is also noteworthy that _Nostoc commune_-associated bryophyte species, both mosses and liverworts, showed the highest rate of N₂ fixation (Table 1), which is in agreement...
Figure 6. Relationships between cyanobacteria and the stages of ecological succession. Relationships between OTUs of epiphytic cyanobacteria obtained at a 0.01% cut-off and the different stages of ecological succession (triangles) - the younger site (site 1) correspond to bare ground close to the glacier front whereas the older site (site 5) correspond to Nothofagus forests - in the north-side glacier (a) and south-side glacier (b) after correspondence analysis. Site 1 is not included in (a) because cyanobacteria were not found epiphytically on bryophytes at that site. For OTUs identification please see Fig. 2.

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with the suggestion that *Nostoc commune*, in particular, is an important contributor of soil N in both polar regions [32]. Specific affinities of individual cyanobacteria genera with particular bryophyte species were not apparent, as the same bryophyte species can support the three main cyanobacteria genera found here (*Nostoc*, unidentified Nostocaceae group and *Stigonema*) (Table 1 and Fig. S2). However, a relationship between cyanobacteria genera and the different stages of succession was apparent in both glaciers. With increasing age along the chronosequences, cyanobacteria shifted from: *Nostoc commune*-dominated communities at young sites; to unidentified Nostocaceae-dominated communities at intermediate sites; and finally to *Stigonema* spp-dominated communities at older sites (Fig. 6). Such procession correlated with the soil development trends found here and contrasts with results observed in mature boreal forests where moss species identity serves as an important determinant of cyanobacterial communities that inhabit mosses [22]. With our data set, our results suggest that the stage of succession acts as a strong driver in determining the composition of epiphytic cyanobacteria on bryophytes during primary succession. Despite this similarity between the two glaciers, diversity of epiphytic cyanobacteria on bryophytes during primary succession. Despite this similarity between the two glaciers, diversity of epiphytic cyanobacteria at SG was higher than at NG and increased with time since deglaciation (Fig. 3b), reflecting more favourable conditions for cyanobacteria growth on the wetter southern side of Cordillera Darwin.

Conclusions

We have compared, from a multidisciplinary perspective, chronosequences in front of two receding glaciers with contrasting climatic conditions (wetter vs drier) and differing rates of ecological succession in Tierra del Fuego (Chile). In doing so, we have shown that bryophyte-cyanobacteria associations have the potential to contribute to N accumulation very early after glacier retreat, a process that could help promote soil development. At intermediate-aged sites, and under drier conditions, bryophytes can grow among dwarf shrub plants, forming dense mats that host cyanobacteria species and exhibit high N₂ fixation rates, contributing to N accumulation and cycling. We also found that differences in N₂ fixation capacity between bryophyte species were primarily driven by the abundance of epiphytic cyanobacteria rather than cyanobacterial community composition. Most liverworts showed low colonization, whilst in mosses rates of N₂ fixation did not exhibit consistent differences across life forms and habitat (saxicolous vs terricolous). Finally, we identified that the main factor structuring community composition of epiphytic cyanobacteria was the stage of ecological succession, with no relationship to host species identity.

Supporting Information

Figure S1 Scores variation of the general factor component of soil development along the glacier chronosequence in the north-side (black bars) and south-side (white bars) of Cordillera Darwin. Sites with higher loadings reflect a more developed soil. The general factor component of soil development was obtained by reduction of soil pH and total soil carbon and nitrogen by principal component analysis. Mean values (± SE) of three replicate transect are shown.

**Figure S2** Correspondence analysis between OTUs of cyanobacteria obtained at a 0.01% cut-off and each individual bryophyte species, just the six most abundance bryophyte species with cyanobacteria colonies were used in this analysis, identification of OTUs can be see in Fig.2.

Table S1 Changes in bryophyte species along two proglacial chronosequence at the Cordillera Darwin (Tierra del Fuego, Chile).

Table S2 Gen Bank accession numbers corresponding to species and specimens used for tree inference.

Appendix S1 Calculation of different vial headspaces for ARA and ^15^N₂ uptake essays.

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Author Contributions

Conceived and designed the experiments: MA-C SPO TGAG LGS. Performed the experiments: MA-C SPO. Analyzed the data: MA-C SPO. Conceived and designed the experiments: MA-C SPO TGAG LGS. Performed the experiments: MA-C SPO. Analyzed the data: MA-C SPO. Performed the experiments: MA-C SPO. Analyzed the data: MA-C SPO. Conceived and designed the experiments: MA-C SPO TGAG LGS. Performed the experiments: MA-C SPO. Analyzed the data: MA-C SPO. Performed the experiments: MA-C SPO. Analyzed the data: MA-C SPO. Performed the experiments: MA-C SPO. Analyzed the data: MA-C SPO. Carried out sampling field work: AP. Organised field logistics: AP RR LGS. Selected field sites: DP RR. Arranged required permissions and permits: RR. Discussed the results and commented on the manuscript: MA-C SPO TGA LGS ADLR ROH RR NT MC MTDLC AP.

References