Effects of nitrogen fertilization on diazotrophic activity of microorganisms associated with *Sphagnum magellanicum*

Martine A. R. Kox · Claudia Lüke · Christian Fritz · Eva van den Elzen · Theo van Alen · Huub J. M. Op den Camp · Leon P. M. Lamers · Mike S. M. Jetten · Katharina F. Ettwig

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**Abstract**

**Background and aims** In pristine ombrotrophic *Sphagnum*-dominated peatland ecosystems nitrogen (N) is often a limiting nutrient, which is replenished by biological N$_2$ fixation and atmospheric N deposition. It is, however, unclear which impact long-term N deposition has on microbial N$_2$ fixing activity and diazotrophic diversity, and whether phosphorus (P) modulates the response. Therefore, we studied the impact of increased N deposition and N depletion on microbial N$_2$ fixation and diazotrophic diversity associated with the peat moss *S. magellanicum*, and their interaction with P availability.

**Methods** Nitrogenase activities of *S. magellanicum*-associated microorganisms were determined by acetylene reduction assays (ARA) and $^{15}$N$_2$ tracer methods on mosses from two geographically distinct locations with different N deposition histories, high or low N deposition, and in samples depleted in N (grown 3 years in the greenhouse) versus recent field samples. The short-term response to increased N deposition was tested for mosses differing in N and P fertilization histories. In addition, diversity of diazotrophic microorganisms was assessed by *nifH* gene amplicon sequencing of N-depleted mosses.

**Results** We showed distinct and persistent differences in diazotrophic communities and their activities associated with *S. magellanicum* from sites with high versus low N deposition. Initially, diazotrophic activity was six times higher for the low N site. During incubation and repeated ARA, however, this activity strongly decreased, while it remained stable for the high N site. Activity for the high N site could not be increased by long-term experimental N deprivation. Short-term, experimental N application had an inhibitory effect on N$_2$ fixation for both sites, which was not observed in mosses with high indirect P availability.

**Conclusions** We conclude that although N deposition negatively affects N$_2$ fixation as also shown in previous studies, long-term effects of N deprivation on the diazotrophic activity and community are more complex. Furthermore, our results indicated that P availability might be an important factor in modulating the response of *Sphagnum*-associated diazotrophs to N deposition.
Keywords Peatland · Nitrogen fixation · Diazotroph diversity · Phosphorus · $^{15}$N$_2$ · Acetylene reduction · nifH · Nitrogen deposition

Introduction

Although nitrogen (N) in the form of dinitrogen gas (N$_2$) is the most abundant element in the Earth’s atmosphere, low availability of this essential element for plants is often limiting primary production in pristine ecosystems (Canfield et al. 2010). The rates of atmospheric N deposition have increased at record pace over the last decades in many regions of the world, due to industrialization, increased traffic and the intensive use of N-fertilizers and manure in agriculture (Gruber and Galloway 2008). Nowadays, biological N$_2$ fixation is estimated to contribute about 200–250 Tg N year$^{-1}$ (Tg = 10$^{12}$ g) to the biosphere, including N$_2$ fixation associated with agricultural crops (Gruber and Galloway 2008; Fowler et al. 2013), while anthropogenic input from fertilizers and increased atmospheric deposition of reactive nitrogen (NH$_x$, NO$_y$) nearly matches this value (up to 195 Tg N year$^{-1}$; Gruber and Galloway 2008). This has massively disturbed the N balance in many (pristine) ecosystems, and is expected to possibly change even N-limited sites to sites with excess N availability (Bobbink et al. 1998; Granath et al. 2014).

The effects of high N deposition are most severe in nutrient-poor ecosystems such as Sphagnum-dominated peatlands, which are very sensitive to increased nutrient influx (Lamers et al. 2000; Tomassen et al. 2004; Verhoeven et al. 2011; Novak et al. 2015). Here, additional N supply might lead to higher biomass production and therefore carbon sequestration, but an opposite or mixed effect can also be envisaged: N in the form of nitrate/nitrite could serve as electron acceptor for peat degradation, and increase degradability of otherwise very recalcitrant peat bog biomass (via an “improved” C:N stoichiometry and/or induced shifts from Sphagnum to vascular plants (Bragazza et al. 2012)), with greenhouse gases carbon dioxide (CO$_2$) and methane (CH$_4$) being the products of this oxygen-limited degradation. Since peatlands are estimated to store one-third of all terrestrial carbon (Gorham 1991), it is crucial to understand these feedback effects of increased N supply (Granath et al. 2014).

Many studies in peatlands have focused on the effects of high N deposition (i.e. >10 kg N ha$^{-1}$ year$^{-1}$; Lamers et al. 2000) on plants, since plants are assumed to be the main consumers of NH$_4^+$ and NO$_3^-$ in terrestrial ecosystems. Sphagnum mosses have been shown to be very efficient in retaining nitrogenous compounds from N deposition, preferentially taking up N in the form of NH$_4^+$ (Fritz et al. 2014). The effects of high N deposition on Sphagnum moss productivity are dependent on the level, where at low deposition levels, up to about 10 kg N ha$^{-1}$ year$^{-1}$, mosses might still benefit from the additional N input (Lamers et al. 2000; Berg et al. 2012). At deposition levels higher than 18 kg N ha$^{-1}$ year$^{-1}$, Sphagnum spp. become saturated with N and excess N leaches deeper into the system, which facilitates growth of vascular plants, changes nitrogen cycling and decomposition rates, and may result in degradation of peatlands (Lamers et al. 2000; Bragazza 2006; Wu and Blodau 2013; Novak et al. 2015). Moreover, high N availability in the form of NH$_4^+$ ultimately could become toxic to wetland plants, especially under P-limited conditions (Lamers et al. 2012). Yet, when phosphorus (P) availability is increased simultaneously, it is thought that the negative physiological effects that plants experience under high N loads can be alleviated, due to stimulation of primary production (Lamers et al. 2012; Fritz et al. 2012), and that also their microbial community might benefit directly from non-limiting P-supply.

Sphagnum mosses and their microbial associates have been demonstrated to be specific throughout the moss’s life cycle, with a higher microbial diversity observed in Sphagnum species from more eutrophic habitats (Opelt et al. 2007; Bragina et al. 2011, 2014). The mosses consume carbon dioxide (CO$_2$) produced during decomposition in lower peat layers, while epi- and endophytic microorganisms of Sphagnum contribute additional CO$_2$ by converting methane (CH$_4$) to CO$_2$ (Raghoebarsing et al. 2005; Kip et al. 2011). Microbial analyses of peat mosses demonstrated the presence of a wide variety of diazotrophs, which might serve as nutrient suppliers to the moss (Andersen et al. 2012; Berg et al. 2012; Bragina et al. 2014). Nitrogen is an important nutrient limiting or co-limiting primary production in boreal peatlands. The only known biological process that is able to convert N$_2$ into a reactive form of N replenishing the ecosystems’ N-pool is N$_2$ fixation, which is known to be performed by a large variety of prokaryotes called diazotrophs (Robertson and Vitousek 2009). In peatlands, diazotrophic activity (N$_2$ fixation)
has been shown to account for up to 35% of the N assimilated by *Sphagnum* species (Berg et al. 2012; Larmola et al. 2014). Biological N\textsubscript{2} fixation is catalyzed by the enzyme nitrogenase, consisting of an $\alpha_2\beta_2$ heterotetramer (NifD and NifK respectively) and a homodimer dinitrogen reductase $\gamma_2$ (NifH; Rees et al. 2005). The *nifH* gene encoding for the NifH homodimer is most widely used as biomarker to study the diversity of diazotrophs (Gaby and Buckley 2012; Levy-Booth et al. 2014).

Moreover, diazotrophy has been suggested to be an important process in the accumulation of N during succession of peatlands (Larmola et al. 2014), where methanotrophic diazotrophs rather than cyanobacteria may account for a significant part of the N\textsubscript{2} fixation activity. In a Canadian ombrotrophic bog, investigation of the active diazotrophic community showed higher relative abundance of *nifH* transcripts originating from methanotrophs than those from cyanobacteria (Vile et al. 2014). Nevertheless, many questions on the relative importance of different diazotrophic microorganisms in peat mosses remain and in contrast to plant communities, the effect of N deposition on microbial assemblages in peatlands has been less well explored (Bodelier 2011; Andersen et al. 2012; Ho and Bodelier 2015).

The expression of the nitrogenase enzyme is down regulated by ammonium availability. Also it is known that high concentrations of nitrate might result in formation of nitrite, which irreversibly inactivates the nitrogenase enzyme (Postgate 1982). Multiple studies have shown that N deposition affects diazotrophic activity in a dose-dependent way. Recently, Rousk et al. (2013b) showed that N\textsubscript{2} fixation rates in the forest moss *Pleurozium schreberi*, mainly colonized by cyanobacteria, were not sensitive to N amendments up to 10 kg N ha$^{-1}$ year$^{-1}$. This result was similar to the upper limit of N deposition that *Sphagnum* mosses could adsorb before N compounds started to leach into deeper peat layers, as calculated in Lamers et al. (2000). However, when the *Pleurozium* mosses were experimentally deprived of N, after having been exposed to high N deposition rates, their microbial community reacted with an increase in N\textsubscript{2} fixation rates (Rousk et al. 2013b). Similarly, along a natural gradient of N deposition levels, the diazotrophic activity (mainly of cyanobacteria) associated with the forest mosses *Hylocomium splendens* and *P. schreberi* was shown to decrease with increasing N deposition levels (Leppänen et al. 2013). Studies on the effect of N fertilization on *S. fuscum* and *S. angustifolium*-associated diazotrophs in a pristine peat bog showed that the diazotrophic activity was negatively affected by high N input, although moss productivity did not change (Vile et al. 2014). The difference in response pattern to N deposition between the study of Rousk et al. (2013b) and the others indicates that increased N deposition may not have a uniform effect on diazotrophs in all moss species. Rather, it suggests that the response may be influenced by diazotrophic community composition, as well as by differential plant–microbe interactions.

For the globally occurring peat moss species *Sphagnum magellanicum*, N deposition effects with respect to its microbial diazotrophic community have not been studied yet. Because of its abundance in ombrotrophic and oligotrophic sites from boreal to temperate regions (Laine et al. 2009), it is an attractive species to study the effects of habitat difference. The main question in this study was whether *S. magellanicum*-associated N\textsubscript{2} fixation was affected by high N deposition, and whether P availability modulated this effect. We compared, for the first time, the same moss species (*S. magellanicum*) from a pristine versus an N-loaded site to study whether N limitation stimulates N\textsubscript{2} fixation. Along this line, long-term effects of N deposition and of N deposition history on N\textsubscript{2} fixation in *Sphagnum* mosses had not been studied before. Like in forest mosses, we expected (1) *Sphagnum* associated N\textsubscript{2} fixation rates to decrease upon N fertilization, since N will than be taken up directly from the environment rather than costly being produced via N\textsubscript{2} fixation. Secondly, it was expected (2) that N\textsubscript{2} fixation rates increase upon N deprivation. Next, P is known to be a directly limiting factor for diazotrophy and often becomes the limiting factor for primary production with excess N input (Vitousek et al. 2002), however, it is not known whether indirect P availability, in the form of high P-content in plant-tissue, has the same effect. Therefore, we hypothesized that (3) high indirect P availability to stimulate N\textsubscript{2} fixation, and to alleviate the negative effects of N deposition (Smith and Bennett 1999; Fritz et al. 2012). We expected (4) N\textsubscript{2} fixation to be induced by limited N availability, either by absolute N limitation or by relative N limitation, (i.e. relative to the P limitation that plants and microorganisms experience). It was expected (5) that high indirect P availability would result in a higher N demand for both plants and microorganisms, ultimately stimulating N\textsubscript{2} fixation.
fixation. The role of P as potential stimulator (direct or indirect) of N₂ fixation in Sphagnum mosses has not yet been experimentally tested.

Our hypotheses were investigated in two microcosm fertilization experiments. First, to test the effect of N deposition on Sphagnum associated N₂ fixation activity, N fertilization was applied to S. magellanicum collected from field sites exposed to either low or high N deposition over a long time period. Similarly, S. magellanicum from these sites that had been deprived of N for 3 years, were tested in the same set up and their diazotrophic community of the N-deprived mosses was compared. In a second experiment, it was investigated whether Sphagnum mosses with a high P content due to former P fertilization, could modulate the effect of N fertilization.

Material and methods

Sample collection

Sphagnum magellanicum samples were collected from two geographical locations: (1) a low N deposition site located in the Andorra valley mire, Tierra del Fuego, Argentina (<0.5 kg N ha⁻¹ year⁻¹, Fritz et al. 2012) and (2) a high N deposition site, the Dwingelderveld, Dwingeloo, The Netherlands (>25 kg N ha⁻¹ year⁻¹ for more than 30 years; Lamers et al. 2000; Patberg 2011; for details see Table 1). The pristine bog vegetation of the Tierra del Fuego site was dominated by S. magellanicum interspersed with few vascular plants (cover <2 %) such as Empetrum rubrum, Nothofagus antarctica, Carex magellanica, Marsipospermum grandiflorum. The Dutch high N deposition site Dwingelderveld consisted of several patches of bog vegetation that differed in successional state, containing species such as S. magellanicum, S. fallax, S. squarrosum, Andromeda polifolia and Eriophorum vaginatum (Patberg 2011). Mosses from both sites were collected in February 2010 and February 2013.

Experimental design and maintenance

Mosses collected from both sites in 2010 were maintained in the greenhouse facilities at Radboud University until 2013 (in total 37.5 months until incubation period) and were experimentally deprived of N by watering with artificial rainwater without added N-compounds (see Table 2 time table and Table S1 for
Mosses collected in 2013 were collected from the field and together with the N-deprived mosses from the greenhouse transferred to the climate rooms to acclimatize for 8 weeks prior to N fertilization experiments. From each site (high versus low N deposition) and treatment history (N-deprived or fresh from the field) 6 replicates were used in the experiments, except for mosses from the low N site collected in 2013, which had only 4 replicates due to limited sample size.

In a second experiment we tested whether P modulates N deposition effects on N$_2$ fixation. From the Argentinean field site (low N deposition), $S$. magellanicum was collected from former experimental plots (Fritz et al. 2012). The plots had been treated from 2008 to 2010 with increased levels of either P (P; 10 kg P ha$^{-1}$ year$^{-1}$, as Na$_2$HPO$_4$ dissolved in bog water) or nitrogen (N; 40 kg N ha$^{-1}$ year$^{-1}$, as NH$_4$NO$_3$ dissolved in bog water; see Table 2 for timeline and Fritz et al. (2012) for further details). As a control, unfertilized mosses from the same site were collected at the same time. Microcosm treatments are referred to as 10P, 40N and control, 0NP (Table 1; Fritz et al. 2012). From each plot, $S$. magellanicum was collected and divided into 4 replicates, yielding a total of 12 samples. All samples collected from the field sites were transported in plastic bags, which were kept in the dark during transport.

### N fertilization treatments

For clarity, throughout this manuscript deposition is defined as being exposed to the increased anthropogenic load in the field (increased atmospheric deposition), whereas experimental fertilization both in the field and in the lab is defined as watering with NH$_4$NO$_3$-supplemented water. Prior to incubation in microcosms, fresh weights of mosses were determined, after which they were incubated in glass microcosms that were closed with metal clamps and a rubber O-ring (Bormioli Rocco, Fidenza, Italy). The microcosms consisted of glass jars that had been modified by the addition of a standard crimp-cap serum bottle opening to the glass lid, creating a vessel that was gastight (tested) and accessible through a standard butyl rubber stopper. Most incubations were performed in 1.2 L glass microcosms containing multiple $S$. magellanicum individuals (>10 individual shoots, ±8 cm length) to mimic natural density. Due to smaller sample sizes, mosses from the previously fertilized plots (40N, 10P, and 0NP/low N-site mosses from Argentina) were incubated in similar density in microcosms of 0.28 L.

Incubation microcosms were kept in a cooled water basin (16 °C; cryostat) within a climate chamber with artificial light (10 h dark, 14 h light, 150 µmol m$^{-2}$ s$^{-1}$ PAR, provided by 2 × 400 W lamps (Philips, Master Son-T PiaPlus, Belgium)). To minimize site-specific effects other than N deposition, mosses were acclimatized in this climate room for 8 weeks prior to N fertilization treatments, during which water in the microcosms was replaced three times a week by fresh N-free artificial rainwater (composition see Table S1), keeping the water level stable at 1 cm below the capitula (300 ml for large microcosms; 60 ml for small microcosms; pH range 4.0–4.5). Hereafter, half of all replicates of each experiment were randomly assigned to receive N

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**Table 2** Overview of fertilization measures and moss sampling in time, also see Table 1 for more info on sample locations

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Methane (CH₄) is detectable with this system. Gas sampling was performed over time in a non-destructive manner. Nitrogenase activity was determined two times at the end of the acclimatization period (t = 5) and 5 days (t = −5) prior to N fertilization) and 8 (t = +8) and 16 days (t = +16) after start of the N fertilization treatment (t = 0). Only nitrogenase activity rates of the different treatments from day +8 and +16 were used for further comparisons and correlation analysis.

To convert ethylene production rates to N₂ fixation rates, ¹⁵N₂ labelling experiments were performed on incubations that still showed nitrogenase activity at the end of the experiment (field control samples from both high-N and low-N sites and 10P samples). These experiments could only be performed at the end of the incubation, as they require destructive sampling. Incubations were split and mosses were incubated for 48 h either with ¹⁵N₂ (5 % v/v headspace) or with both ¹⁰N₂ (5 % v/v headspace) and acetylene (5 % v/v headspace) to check for potential inhibition by acetylene, and a sample was kept as ¹⁰N₂ background control. Ethylene production was measured as described above. To determine ¹⁵N₂ uptake, mosses were collected for δN determination of the background and the enriched plant material after 48 h, segmented into three parts (0–2 cm from top, 2–4 cm and 4–6 cm) and dried (72 h at 70 °C). Moss samples were ground for 2.5 min at 30 Hz with a stainless steel bead (ø 5 mm) using a Retsch MM200 ball mill (Retsch, Haan, Germany). 12 to 15 mg of each sample was transferred into a 5 × 8 mm tin cup and combusted by flash combustion (1800 °C) on a CNS analyzer (EA 1110 Carlo Erba - Thermo Fisher Scientific) coupled to an Isotopic Ratio Mass Spectrometer (Finnigan DeltaPlus, interface Conflo III). Conversion factors were calculated according to Leppänen et al. (2013). Background ¹⁰N₂ levels were uniform, for the 10P mosses average Δ15N/14N (%) was 0.797 ± 0.396. Since the results of the heavy isotope labelling did not generate one stable conversion factor due to large variations, nitrogenase activity rates were not converted to corresponding N₂ fixation, but are expressed as the directly measured ethylene production rates throughout this study.

Plant and porewater analyses

To check for potential abiotic changes and nutrient contents of the respective porewater and peat moss, peat moss samples were collected in duplicate before N fertilization and 1 week after the start of N fertilization, each time directly before acetylene reduction assays were performed. Samples were oven-dried, milled and combusted as described above. Combustion products from 5 mg dry biomass were separated on a Porapak QS-column and analyzed for total C and N content. One week after the start of the N treatments, porewater samples were taken before watering from the microcosms, using rhizons (pore size 2 µm; Rhizosphere Research Products, Wageningen the Netherlands). Element concentrations of Na, K, P in the porewater (10 mL, acidified with 0.1 mL 65 % HNO₃) were analyzed in duplicate by inductively coupled plasma-optical emission spectroscopy (ICP-OES iCAP 6000, Thermo Fischer Scientific, Carlsbad, USA).
Molecular analyses

**DNA extraction and nifH amplification**

Total DNA was extracted only from intact *S. magellanicum* plants that were deprived of N by growth in the greenhouse (originating from both the low and the high N deposition site). These samples were collected 4 weeks after the start of the acclimatization period, before acetylene reduction assays. Three moss individuals were grinded using pestle, mortar and liquid N₂, of which 0.5 g fresh weight was used for DNA extraction using the Powersoil DNA isolation kit (MO BIO laboratories, Carlsbad, USA), with minor adaptation: in order to increase the final DNA concentration the elution buffer was added in three portions (in total 100 μL), instead of at once. After each addition (33.3 μL) the sample was briefly centrifuged.

A fragment of the *nifH* gene (360 bp) was amplified by polymerase chain reaction (PCR) in a nested approach: the primary step included the forward primer 19F (5′-GCIWTYAYGGIAARGGIGG-3′; Ueda et al. 1995) and the reverse primer *nifH*3 (5′-ATRTTRTTNGCNGCRTA-3; Zani et al. 2000); in the second step, the forward primer *nifH*1 (5′-TGYGAYCCNAARGCNGA-3) and the reverse primer *nifH*2 (5′-ANDGCCATCATYTCNCC-3; Zehr and McReynolds 1989) were used. For PCR, PerfCTA Quanta master mix (Quanta Biosciences, Gaithersburg, USA) was used. Both PCR protocols started with an initial denaturation step of 98 °C for 5 min, followed by 30 cycles of denaturation at 95 °C for 30 s, annealing at 55 °C for 1 min, and elongation at 72 °C for 1 min; completed by a final elongation step at 72 °C for 10 min. Nested PCR was performed with 1 μL PCR product of the first PCR. Correct length of both PCR products was checked with gel electrophoreses (GE) on an agarose gel (1 %, runtime of 1 h at 60 mV). Products of the nested PCR with correct lengths (360 base pairs (bp)) were used for amplicon sequencing.

**Amplicon sequencing**

Amplicon sequencing of nested PCR products was done using the Ion Torrent Personal Genome Machine™ (Life technologies, Carlsbad, USA) and the Ion Xpress™ Barcode Adapter Kits 1–2 (Life technologies, Carlsbad, USA). DNA products of several PCR-reactions of the same sample were pooled to obtain sufficient amounts of DNA. First, PCR products were purified using magnetic beads (Agencourt® AMPure® XP Kit, Beckman Coulter, Brac, USA) and magnetic rack (DynaMag™-2 magnet). After each following step purification of the DNA fragments was performed using this method.

After determination of purity and concentration using the Bioanalyzer® 2100 and the High sensitivity DNA kit (Agilent Technologies, Santa Clara, USA), DNA fragments of the same sample were pooled in equimolar amounts, to a final concentration of 50 ng DNA μL⁻¹. Next, DNA fragments were end-repaired, adapters containing specific barcodes (for each sample a different barcode) were ligated, followed by a nick-repair. Finally, amplification (6 cycles, according to manufacturer’s protocol) was performed, followed by quantification and a purity check of the product using the Bioanalyzer® 2100 and the High sensitivity DNA kit (Agilent Technologies, Santa Clara, USA). All barcoded samples for sequencing were pooled in equimolar amounts to a final concentration of 26 pM DNA. Pooled fragments were amplified on Ion Sphere particles using the Ion One Touch™ Instrument and Ion One Touch™ 400 Template Kit (Life Technologies, Carlsbad, USA) according to the manufacturer’s instructions. After enrichment of the Template-Positive Ion Sphere™ Particles using the Ion One Touch™ ES (Life Technologies, Carlsbad, USA), the spheres were loaded on an Ion 314™ v2 Chip. Subsequently, DNA fragments were sequenced using the Ion PGM™ 400 Sequencing Kit according the manufacturer’s instructions (MAN0007242 Revision 2.0).

**Amplicon sequence analysis**

The amplicon library (NCBI SRA database project number PRJNA280035, samples SAMN03454249, SAMN03454250) yielded 188,517 reads with an average read length of 266 basepairs (bp) from the low N deposition site and 197,813 with an average length of 281 bp from the high N deposition site. Prior to clustering, the reverse complements were generated for reads starting with the reverse primer, using the ‘reverse.seqs’ function in mothur (1.22.2; Schloss et al. 2009). Reads were quality-filtered, so that reads shorter than 320 bp and having more than two expected errors were
removed (12.6 %; Fastq_filter command USEARCH; Edgar 2010). Next, amplicon sequences were dereplicated, sorted according to number of replicate sequences belonging to the representative, followed by length sorting, after which the remainder of 164,235 reads were clustered using UCLUST (Edgar 2010) at a distance level of 10 % (nucleotides). Resulting OTU-representatives (centroids) were tested for chimeras using UCHIME (Edgar et al. 2011). All OTU-representative sequences were subsequently imported into ARB (Ludwig et al. 2004), translated into protein sequences in the correct reading frame, aligned and used for phylogenetic analysis using the nifH database described in Gaby and Buckley (2014). Neighbour-joining trees were calculated based on the amino acid alignment (alignment position 179–368) using the neighbour-joining algorithm and Kimura correction implemented in ARB.

Statistical analyses

For the high and low N-deposition samples, assumptions of normality for all data (acetylene reduction (ARA), C and N content, Porewater concentrations, pH) were tested using the Shapiro-Wilk test to check for normal distribution, and homogeneity of variances was tested with Levene’s test. To analyze the data from 0NP/10P/40N plots, the Fligner-Killeen test was used, because of the small number of replicates, since this test is more robust with respect to departures from normality. Non-normal distributed data was transformed to approach a normal distribution, either by square root transformations or by ln-transformations. When parametrical conditions were not met, a non-parametric Kruskal-Wallis test was applied (i.e. for the N content data).

Differences in nitrogen fixation activity, C content and porewater data were all analyzed by three-way ANOVA, followed by a Tukey Honest Significant Differences (HSD) test. To test differences in nitrogenase activity measurements and N-fertilization effects, a two-way ANOVA was performed. Type I was applied to the formerly fertilized plots (0NP, 10P, 40N); factors used were origin (0NP, 10P or 40N), and treatment (N-fertilized or control). A type III three-way ANOVA was applied to test for differences between the different samples in the high versus low N-deposition site experiment, correcting for unbalanced design in the comparison. For both experiments test were carried out separately for the two periods: acclimatization period ($t = 13$ and $t= 5$ days) and N fertilization period ($t = +8$ and $t= +16$ days). Factors used in the ANOVA for the high-low N deposition site were: origin (high or low N site), N deprivation (N-deprived or field), and treatment (N-fertilized or control; also see Table S8). Since the ANOVA tests showed high numbers of interactions on the ARA data, the N-fertilization treatment effect was also tested using a paired t-test on a priori paired data of the N-fertilization period.

Data shown here are the averages of the relevant replicates (mean ± standard error). Correlations between ARA data and N contents, C contents or porewater nutrient levels of the formerly fertilized plots were analyzed using Spearman’s test, which is more robust for small number of replicates. All statistical analyses were performed using R version 2.15.2 (R Development Core Team 2012), and significance levels were indicated as follows: $\cdot = p < 0.1$, $*= p < 0.05$, $** = p < 0.01$, $*** = p < 0.001$.

Results

N-accumulation in Sphagnum

Sphagnum N content was 2.5 times lower for mosses from the low N site than for the high N site ($\chi^2 = 6.86$, $p < 0.01$; Fig. 1). This strong difference in N contents had disappeared in N-deprived mosses. For the formerly fertilized mosses from Argentina, N content also

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Fig. 1 N content (%) in Sphagnum samples freshly collected from the field and from Sphagnum samples after 3 years of N-deprived greenhouse incubations originating from the high- and low N deposition sites, and its change upon treatment with high N fertilization. Bars represent the mean ± SE (3 replicates per sample). Effects of N fertilization and of the N deposition history were tested with Kruskal-Wallis test (resp. $\chi^2 = 11.1368$, $p < 0.001$; $\chi^2 = 6.8612$, $p < 0.01$)
differed, being lowest for the 0NP-plot mosses and highest for the 40N-plot mosses ($p < 0.001$; Fig. 2). Overall, both N and P fertilization increased the N content of *Sphagnum* tissue compared to the unfertilized mosses, while C content was not affected ($F_{1,20} = 21.15$, $p < 0.001$). After short-term laboratory N fertilization, the N content increased by 25% in mosses from both low- and high-N deposition sites ($\chi^2 = 11.14$, $p < 0.001$; Fig. 1), and by 35% in mosses from the formerly fertilized plots ($\chi^2 = 4.44$, $p < 0.05$; Fig. 2). For the formerly fertilized plots, it was apparent that the increase in N-content in the 10P mosses was much higher than for the 40N or 0NP mosses (Fig. 2). In none of the experiments did the pattern observed in N content of the mosses correlate with nitrogenase activity.

Porewater Na, K and P concentrations

Experimental N fertilization led to increased porewater concentrations of potassium (K) and sodium (Na) in both experiments (Table S2 and Table S3). Both K and Na concentrations were generally three times lower in microcosms with mosses from the field than those with N-deprived mosses (resp. $F_{1,14} = 22.25$, $p < 0.001$ $F_{1,14} = 11.39$, $p < 0.01$). Porewater P concentrations of mosses from the low N deposition field site were very low, only 10% of the P concentration in all other microcosms ($F_{1,14} = 9.55$, $p < 0.01$). Moreover, in these microcosms the P concentrations did not further decrease upon N fertilization, unlike in the other treatments, where P concentrations decreased by 33%. In microcosms with mosses from the formerly N-fertilized plot (40N), P concentrations became remarkably higher upon N fertilization ($F_{2,8} = 3.38$, $p = 0.086$; supplementary table S3), when compared to the other treatments. Furthermore, nitrogenase activities in this treatment were negatively correlated with P concentrations in porewater (Spearman $c = -0.71$, $p = 0.01$, $n = 12$), indicating that higher P uptake from pore water was linked to higher N$_2$ fixation rates.

Nitrogenase activity in mosses derived from low and high N deposition sites

The nitrogenase activity was seven times higher for the low N site than for the high N site (Fig. 3a). During incubation and repeated ARA, however, activity strongly decreased for the low N site, while it remained stable for the high N site (Table S5). Furthermore, during acclimatization nitrogenase activity was on average two times lower in N-deprived samples ($71 \pm 21$ C$_2$H$_4$ g$^{-1}$ dw day$^{-1}$). Prior to N fertilization, nitrogenase activity was determined twice, showing an overall average of $116 \pm 25$ C$_2$H$_4$ g$^{-1}$ dw day$^{-1}$. A week before start ($t = -5$) of N-fertilization, nitrogenase activity measurements showed no difference between replicates of the different mesocosms or between the two sites ($F_{1,18} = 3.96$, $p = 0.06$). At $t = +8$ the activity was 40 times higher in the mosses from the high N deposition site compared to the low N deposition site (see Fig. 3b) as a result of the stronger decrease. Throughout the experiment, no CH$_4$ formation was detected in any of the incubations.

Short-term N fertilization effects on nitrogenase activity in mosses from low- and high N deposition sites

N fertilization reduced nitrogenase activity in mosses from both low and high N deposition sites, and also for N-deprived mosses. Already 8 days after the start of N fertilization ($t = +8$), all N fertilized microcosms showed lower nitrogenase activity rates than controls (Fig. 3b). For the field mosses, 8 days of N fertilization led to an 80 times lower N$_2$ fixation activity rate ($1.4 \pm 0.55$ nmol C$_2$H$_4$ g$^{-1}$ dw day$^{-1}$) compared to their controls ($113 \pm 30$ nmol C$_2$H$_4$ g$^{-1}$ dw day$^{-1}$; $t(9.0) = 3.42$, $p < 0.01$). A similar trend of N fertilization was observed for the N-deprived moss incubations ($t(13.1) = 1.72$, $p = 0.055$).
Short-term N fertilization effects on nitrogenase activity in mosses with a history of P or N fertilization

Similar to the results for low and high N deposition sites, nitrogenase activity decreased over time for all incubations with mosses from former fertilization plots 10P, 40N and 0NP. At the very beginning of the acclimatization period, Sphagnum-associated nitrogenase activity in the 0NP-microcosms showed almost three times higher activity (1012 ± 101 nmol C 2H4 g−1 dw day−1), than for incubations from the fertilized plots 10P and 40N (respectively 368 ± 14 and 340 ± 53 nmol C 2H4 g−1 dw day−1; $F_{2,18} = 17.55, p < 0.0001$). Five days prior to start of the N-fertilization treatment, nitrogenase activities had decreased compared to levels at which nitrogenase activities started, but the activities were similar between the different microcosms (0NP/10P/40N $F_{2,9} = 1.59, p = 0.26$).

During N-fertilization, nitrogenase activities in all but the 10P samples decreased, and even became undetectable in the 40N-microcosms ($F_{2,8} = 24.84, p < 0.001$; Fig. 4). In contrast, 10P-microcosms showed on average two times higher nitrogenase activity after N fertilization (118 ± 7 C2H4 g−1 dw day−1) than their unfertilized controls (50 ± 17 nmol C 2H4 g−1 dw day−1; $t_{4.04} = 3.63, p < 0.05$).

15N label incorporation

The 15N incorporation method to assess N2 fixation rates could only be used at the end of the incubation experiments, as it required destructive sampling. Only in mosses from the 10P-plot, both a clear 15N incorporation could be observed.

Fig. 3 C2H4 production rates of mosses (mean ± SE) from low- and high N deposition sites, during acclimatization period (a, n = 6), where different letters indicate significant differences as tested by ANOVA followed by TukeyHSD ($F_{1,40} = 24.06, p < 0.001$), and C2H4 production rates during the fertilization period (b, n = 3), where different letters indicate significant differences as tested by ANOVA analysis followed by TukeyHSD, with significant interactions between treatment and N-deposition site ($F_{1,36} = 9.79, p < 0.01$) and between treatment and N-deprivation ($F_{1,36} = 9.0381, p < 0.01$). The treatment effect was tested separately on a priori paired data using a paired t-test for field ($t_{(9.04)} = 3.42, p < 0.01$) and N-deprived samples ($t_{(12.3)} = 1.72, p = 0.055$)

Fig. 4 Effect of N fertilization on C2H4 production rates of mosses from past long-term in situ fertilization experiments with either P (10P), N (40N) or treated with unamended bog water (0NP). Displayed values indicate mean C2H4 production ± SE (n = 3). The treatment effect was tested on a priori paired data using a paired t-test ($t_{(3,13)} = 2.220, p = 0.055$)
incorporation signal and acetylene reduction rate was obtained. Here, uptake of $^{15}$N$_2$ was lower in incubations with both acetylene and $^{15}$N$_2$ compared to incubations with only $^{15}$N$_2$ (Fig. 5; $F_{1,12} = 11.92$, $p < 0.01$), indicating competition and/or inhibition by acetylene. The weighted average of the $^{15}$N$_2$ incorporation in mosses incubated with both acetylene and $^{15}$N$_2$ generated a conversion factor of 3.11 (mol N$_2$ fixed/mol C$_2$H$_2$ reduced) similar to those found in other studies (Leppänen et al. 2013), although showing high variation (SEM = 1.1).

Molecular analysis of microbial diazotrophic communities

Clustering of the nifH amplicon sequences derived from the N-deprived mosses of the low- and high N sites was performed at 10 % nucleotide-distance level. The resulting 73 nifH OTU-representatives were analyzed at amino-acid level (Suppl. table S4). Ninety-five percent of all reads were covered by 20 OTU-representatives, with the remaining 53 OTU-representatives had only a low coverage (Fig. 6). The OTU-representatives clustered in three of the four main groups of the nifH gene diversity (Zehr et al. 2003) and the diversity considerably differed between mosses from the two sites.

Overall the diazotrophic community consisted mostly of proteobacteria, but showed distinct diversity for the mosses originating from two different geographical regions. Cyanobacterial nitrogenases were only present in S. magellanicum from the high N deposition site (Fig. 6, Cluster I; OTU-representatives 15, 33, 41, 43, 62; related to Calothrix, Leptolyngbya, Trichormus). Furthermore, the diazotrophic community within the high N deposition differed from the low N deposition site in that many OTU-representatives showed high similarity to uncultured, environmental nifH representatives. Within cluster I and III, several OTUs, mostly from the high N deposition site, were related to uncultured groups: Within cluster II groups such as from a Russian peat bog (OTU 17 and 18), from peat-rhizosphere-soil environments (OTU 13, 21, 26 and 72), estuarine seawater (OTU 9, 27 and 37); Within cluster I the nifH representatives were related to clusters of uncultured nifH sequences derived from peat, soil, rhizosphere and glaciers (Cluster I - OTU 34, 40, 44, 48, 56, 58, 60 and 68).

The diazotrophic community in S. magellanicum from the low N deposition was dominated by reads associated to Bradyrhizobiaceae, Rhizobiaceae, Rhodospirillales and Chlorobiales. The difference with the high N deposition site was apparent in that low N deposition site had the highest number of reads clustering closely to Verrucomicrobial nitrogenases (cluster III; OTU 3, 4, 7, 8, 14, 16, 19, 20, 23 and 28). Furthermore, three highly abundant OTU-representatives (OTU 5, 32 and 36) that were only present in mosses from the low N deposition site formed a new distinct group within cluster III, in between Desulfobacterales and environmental nifH sequences previously found in peat.

The only OTU-representatives with similarity to cluster II, which contained functional alternative nitrogenases (anfH) and nitrogenases of archaeal origin, were OTU 12 and 64. OTU 12 was overall the most abundant OTU-representative with most reads originating from the low N deposition site. OTU-representatives 12 and 64 clustered in between nifH groups from Alphaproteobacteria and Firmicutes of Cluster II, but not close to alternative nitrogenases.

Discussion

We studied the effect of N deposition on the activity of the diazotrophic community in Sphagnum magellanicum. Sphagnum mosses with a history of either low or high N deposition in the field were investigated with respect to their nitrogenase activities, and this

![Fig. 5](https://example.com/fig5.png)
Fig. 6 Phylogenetic relationship of partial nifH sequences, based on the deduced amino acid alignment. The Neighbor joining tree was generated in ARB, showing nifH OTU-representatives generated with USEARCH (dist < 0.90) and relevant publically available nifH sequences. OTU-representative IDs are indicated on the right, as well as relative read abundance of the OTU per sample (low N-site in light grey, high N-site in dark grey). The size of the pie charts indicates read abundance for the OTU-representatives.
was complemented by short-term N fertilization experiments. In addition, similar experiments were performed on mosses from the same locations that had been deprived for N for 3 years.

Long- and short-term effects of increased N deposition

Nitrogenase activity was expected (hypothesis 4, H4) to be higher in mosses receiving low N deposition (pristine site), which was initially indeed the case (Fig. 3a), while P contents were similar for both sites (0.27 vs. 0.33 mg g⁻¹; Fritz et al. 2012; H.B.M. Tomassen, unpublished results). These results are comparable to observations for nitrogenase activity in forest mosses (Pleurozium spp.) along an N deposition gradient, where sites with higher N deposition showed lower N₂ fixation rates (Leppänen et al. 2013). Surprisingly however, nitrogenase activity in Sphagnum mosses with a long-term history of low N deposition strongly decreased upon incubation and repeated ARA, while activity remained equally for the high N site (see “Methodological considerations” below). Furthermore, experimental, 3-year N deprivation in the greenhouse did not result in increased nitrogenase activity, unlike what was hypothesized (H2). Changes in the environmental conditions (such as temperature and duration of light) could have contributed to that. When testing short-term effects of N fertilization, nitrogenase activity decreased in mosses from both sites, as was expected (H1). These negative effects in the N-fertilization period were, however, not observed for mosses with a high P content (around 1.8 mg g⁻¹ (Fritz et al. 2012)), which is in line with our hypothesis (H3). This lack of negative effects on nitrogenase activity may either indicate a direct regulatory role of P on N₂ fixation rates, or an indirect role due to higher N demands as a result of higher P availability. Competition for N between the plant and different guilds of microorganisms might explain why we observe an increase in nitrogen fixation activity when both P and N are highly available. In this scenario, diazotrophs would not be able to successfully compete with the moss and/or other microbes for direct uptake of N from the environment.

The lack of stimulation of N₂ fixation by N deprivation, and the interaction with P availability indicated that the effect of N deposition on N₂ fixation is not as straightforward as initially assumed. It indicates that rather than forming a closely regulated symbiosis with their Sphagnum host, N₂ fixing microorganisms might live relatively independently within the mosses’ hyaline cells. In this case, the plants would only indirectly benefit from diazotrophic activities of the microorganisms they harbor, via increased turnover of N in the plant-microbe system, as was also postulated by Ho and Bodelier (2015).

Comparing our N-content data of the mosses that were kept as controls in the short-term N fertilization experiment, with N-content data in Fritz et al. (2012), we observed that the previously fertilized mosses were not N-saturated at start of the N-fertilization treatments and could still increase their N-contents (see supplementary material table 7). Even though in the field porewater N concentrations were 7 times higher for the high N site than in the low N site (20 vs. 2.8 μmol L⁻¹; H.B.M. Tomassen, unpublished results and Fritz et al. 2012), the N content of all mosses in this study increased after short-term N fertilization and diazotrophic activity decreased, with the highest increase in N contents seen in the mosses with a high P-content, as hypothesized (5).

The negative feedback of N fertilization might be regulated at two different levels: at the bacterial cell level, where high N availability may lead to a down-regulation of energetically costly N₂ fixation by negative feedback of NH₄⁺ (Dixon and Kahn 2004) or due to nitrite damage (Postgate 1982), or at the level of plant-microbe interactions, e.g. by changing the release of plant exudates as energy-rich substrates for heterotrophic diazotrophs. In a meta-analysis of 92 studies the second mechanism has been demonstrated for rhizospheric communities of vascular plants: higher nutrient availability (including N) resulted in lower C-loss via root exudation (Fernández-Martínez et al. 2014). The latter explanation would imply that saturation of the N requirement in Sphagnum would lead to lower diazotrophic activity by limiting the energy supply to endophytes. It warrants further investigation to precisely determine the mechanisms that control the N₂ fixing activity of Sphagnum-associated diazotrophs.

Methodological considerations

Apart from the N deposition effects, an overall decrease in nitrogenase activity over time was observed for the low N site, unlike for the high N site. Besides an influence of incubation conditions such as light, nutrient and possibly oxygen availability that are different from field conditions, the repeated exposure to acetylene may well have contributed to the decrease. ARA was
measured four times during 40 days in the same microcosms. Acetylene is known to inhibit the (methane and ammonium) monooxygenases of methanotrophs and other alkane- and ammonium-utilizing bacteria (De Bont and Mulder 1976), as well as the nitrous oxide reductace of denitrifiers (Yoshinari et al. 1977).

In our experimental set-up, most nitrogenase activity with the $^{15}$N$_2$ tracer method was observed in capitula of the moss, a site where previous studies hardly detected methanotrophic activity (Raghoebarsing et al. 2005). Moreover, selective inhibition of methanotrophic diazotrophs by acetylene is unlikely, since no methane and therefore substrate for methanotrophs was provided during the experiments or the acclimatization period of this study, nor was there any formation of methane observed. Therefore, deprivation of methane during the complete experimental phase might have contributed to the observed decrease in N$_2$ fixation activity. Especially for mosses from the low N-deposition site, where the strongest decrease in N$_2$ fixation activity was observed, since for similar pristine sites methanotrophic diazotrophs have been shown to contribute to N$_2$ fixation significantly (Vile et al. 2014).

Lastly, some bacteria have been reported to use the ethylene produced as a substrate for growth (Capone 1993), and ethylene is a plant hormone, which might affect the moss performance (Groen and Whiteman 2014). Together these effects may have caused a shift in the microbial community or in the plant-microbe interaction, resulting in a decrease of nitrogenase activity.

In several other studies, conversion factors between acetylene reduction and N$_2$ fixation rates were calculated by complementing the AR method with $^{15}$N$_2$ incorporation, resulting in a wide range of values for conversion factors (Leppänen et al. 2013; Vile et al. 2014; Knorr et al. 2014). Unlike in these studies, repeated measurements of the same microcosms in the present study made the use of the otherwise preferable, but destructive $^{15}$N$_2$ uptake method impossible. Comparisons of ethylene production rates from acetylene with N$_2$ fixation rates indicated inhibition of N$_2$ fixation by acetylene (Fig. 5). The variation in conversion factors to calculate N$_2$ fixation rates based on ethylene production rates emphasized the uncertainty when using standard conversion factors. Finally, although endogenous ethylene production is generally not observed in peatlands (Schwintzer 1983; Weber et al. 1983), we cannot exclude it as a confounding factor.

Comparison of diazotrophic communities

The molecular analysis of the diazotrophic community, which was performed only on N-deprived Sphagnum magellanicum mosses, showed clearly distinguishable communities for the mosses of two different origins, still persisting after 3 years of identical growth conditions in the greenhouse. Even though the community of both moss samples may no longer be identical to the original community in the field, it demonstrates a remarkable persistence, or ‘memory effect’ (Gunderson 2000). Key distinguishing phylotypes were Cyanobacteria, Verrucomicrobia, and several environmental nifH clusters. While Cyanobacteria and OTUs 17 & 18 were almost exclusively present in mosses from the high N site, Verrucomicrobia and OTUs 5, 32 and 36 were most prominently present in mosses with a history of low N deposition. This finding underlined the need for complementary studies focusing on microbial culturing, as there are no cultured representatives of this entire branch of commonly found Verrucomicrobia yet, and therefore their physiological characteristics are unknown. Studies dedicated to the enrichment and isolation of this group would complement molecular analyses, and yield valuable information on their physiological characteristics that are so far poorly understood.

Several groups of the Sphagnum-associated diazotrophic community abundant in both sites, especially belonging to the Alpha- and Betaproteobacteria, have also been observed in Sphagnum plants from alpine sites and Russian bogs (Zadorina et al. 2009; Bragina et al. 2014). Whereas these may be part of a basic Sphagnum microbiome, specific environmental factors apart from N availability (i.e. water table dynamics, O$_2$, P and CH$_4$ availability), are likely to also have contributed to the differences in the diazotrophic communities. Our findings are in contrast to a study comparing the microbial diversity of two Sphagnum species from three sampling sites in the Alps (Bragina et al. 2011). There it was observed that the same Sphagnum species from different sites harboured similar microbial communities. The clear community differences we found within the same species from Europe and South America questions if this is valid for distances on a global scale. Although long-distance spore (and attached community) dispersal has been documented, it probably nevertheless is so rare that species effects are dominated by geographical effects on community diversity (Shaw et al. 2014).
Interactions with P-availability

When focusing on the N-fertilization period of this study, the formerly P-fertilized mosses demonstrated higher nitrogenase activity upon N fertilization, compared to their non-fertilized controls, in stark contrast to all other samples. It is remarkable that even 3 years after cessation of P fertilization, moss P content was still higher (Fritz et al. 2012). Since nitrogenase activity only increased in those P-enriched mosses, it is likely that all other samples were P limited. This may particularly affect N₂-fixing microorganisms, which generally require more P than non-N₂-fixers (Vitousek et al. 2002). When looking at the effect of N-fertilization on N-content, the steeper increase in N-content in the 10P mosses indicated that 10P mosses may better retain the applied N, probably because compared to the 0NP and 40N mosses they have less relative P-limitation. Since P had been incorporated into plant tissue, it was most probably no longer directly available to diazotrophs. Therefore we concluded that in case of P- or other relative nutrient limitation in the plants, N fertilization may not be of benefit to plants, but only lead to a shift in their microbial community and activity. The most probable mechanism explaining the increased nitrogenase activity is increased exudation of energy-rich substances by Sphagnum, which could then enable heterotrophic diazotrophs to increase their N fixation rates. Furthermore, since nitrogenase activity of the 10P mosses overall was higher than in the 40N or 0NP mosses, our results also confirmed findings by Larmola et al. (2014), who showed that P availability exerted a strong control on N₂ fixation activity.

It is likely that N₂ fixation can match the increased N-demand of Sphagnum at higher P availability, as has been discussed for P addition experiments with feather mosses occurring in boreal forests (Rousk et al. 2013a). P addition experiments generally did show a stimulating effect on Sphagnum growth (Limpens et al. 2004; Fritz et al. 2012) and in P. schreberi on N₂ fixation (Rousk et al. 2013a). In accordance with these studies, the P-fertilized mosses in this study demonstrated an increase of both the C and N content upon N fertilization, which indicated that these mosses, in contrast to all others, could make use of the additional N-load for growth (Fritz et al. 2012). As increased P availability will directly affect both the plant and its microbial community, future experiments are needed to disentangle the capability of Sphagnum to actively regulate diazotrophic activity versus regulation on the microbial community level only.

Conclusions and outlook

Our study demonstrated that the nitrogenase activity of Sphagnum-associated diazotrophs was clearly negatively affected by N fertilization. Saturation of the N demand of the diazotrophic community in combination with the energy costs associated with N₂ fixation, and potentially also N saturation of Sphagnum, most likely resulted in a decrease of nitrogenase activity. Our results also showed that indirect P availability is able to modulate this response. However, the fact that nitrogenase activity was not stimulated by experimental N deprivation indicated that the impact of N deposition on Sphagnum-associated N₂ fixation was much less apparent than assumed before, and far from understood. This study warranted further investigation of how Sphagnum-associated diazotrophs might control N₂ fixation in relation to the environmental N availability and how this might be influenced by P availability. In addition, the ‘memory effect’ of the diazotrophic community associated with one Sphagnum species with respect to long-term N deposition, questioned the specificity of endo- and epiphytic diazotrophs for Sphagnum species and pointed at an important role of environmental parameters.

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