



The role of mycorrhizal type and plant dominance in regulating nitrogen cycling in Oroarctic soils

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Abstract. Mycorrhizal fungi enhance plant access to nitrogen (N) in nutrient-poor environments like the Arctic tundra by
15 depolymerizing N-rich organic compounds into forms available to plants and microbes. As climate change reshapes plant
communities and mycorrhizal associations, shifting dominance from herbaceous species to shrubs, changes in mycorrhizal
type and plant species dominance may differentially stimulate N cycling. Both dominant and rare species, along with
mycorrhizal associations, contribute to ecosystem processes and stability, though the specific roles of these components in
nitrogen cycling and overall ecosystem functioning remain uncertain. We investigated how mycorrhizal associations and plant
20 diversity affect gross N mineralization and nitrification rates in an Oroarctic ecosystem using a plant removal experiment, *in situ* ¹⁵N labelling, and quantification of select nitrification genes. Treatments plots included (1) unmanipulated (Control); or
the removal of: (2) ectomycorrhizal and ericoid mycorrhizal (EcM/ErM) plants, letting arbuscular mycorrhizal and non-
mycorrhizal (AM/NM) plants dominate; (3) AM/NM plants, letting EcM/ErM plants dominate; (4) low-abundance species
(Dominant); and (5) high-abundance species (Rare). Gross N mineralization rates were 73 % and 78 % higher in EcM/ErM
25 and Dominant, respectively, compared to Control, while AM/NM and Rare showed more moderate increases of 30 % and 46
%. Gross nitrification was also highest in EcM/ErM, with a 26 % increase over Control. Gene abundances did not mirror
nitrification patterns. Archaeal ammonia oxidizers (AOA), Nitrospira-type nitrite oxidizers (NIS), and comammox clade A
(ComaA) were consistently more abundant than bacterial ammonia oxidizers (AOB), Nitrobacter-type nitrite oxidizers (NIB),
and comammox clade B (ComaB), suggesting a stable site-level nitrifier community. Dominant had the lowest gene copy
30 numbers overall, except for AOB which was highest. In addition, AOA gene abundance was significantly lower in Dominant
compared to Control, with a marginal reduction observed for NIS. Our findings highlight the key role of EcM/ErM fungi in
accelerating N cycling in Oroarctic soils, challenging traditional assumptions that N transformation rates are slow in EcM/ErM
dominated ecosystems. These insights underscore the need to consider mycorrhizal associations and plant community
composition when predicting tundra ecosystem responses to environmental change.



35 1 Introduction

The availability of soil nutrients plays a pivotal role in shaping tundra plant productivity and the composition of plant communities (Chapin et al., 1995; Shaver et al., 2001), as well as their responses to climate change (Aerts, 2009; Riley et al., 2021; Stow et al., 2004; Sturm et al., 2001). As climate change is particularly pronounced in the Arctic, shifts in plant growth and community composition are occurring (Bjorkman et al., 2020; Hollister et al., 2015), including increased plant productivity ("arctic greening") and shrub expansion ("shrubification") (Bjorkman et al., 2019; Mekonnen et al., 2021; Myers-Smith et al., 2011; Sistla et al., 2013; Tape et al., 2006). Changes in plant community composition contribute to shifts in biodiversity above- and below-ground (Mod and Luoto, 2016; Parker et al., 2018, 2021), with mycorrhizal fungi mediating these changes by influencing soil microbial community composition and activity, impacting soil carbon (C) content, and nutrient cycling ((Andresen et al., 2022; Bahram et al., 2020; Eagar et al., 2022; Hawkins et al., 2023; Hobbie and Högberg, 2012; Hobbie and Hobbie, 2006; Netherway et al., 2021; Phillips et al., 2013; Read, 1991; Sun et al., 2023; Tedersoo et al., 2020). Thus, changes in plant identity or functional diversity can alter nitrogen (N) availability through indirect effects on N mineralization, nitrification, and other N transformations (Isobe et al., 2018; Robertson and Groffman, 2015). These alterations can feed back to plant growth and enhance ecosystem C cycling (Hicks et al., 2020a, 2022; Mekonnen et al., 2021; Parker et al., 2021). Therefore, understanding the links between plant community composition, soil microorganisms, and N cycling is vital for predicting climate change impacts on tundra ecosystems, yet these interactions remain poorly understood (Dobbert et al., 2022).

Ecological communities are typically composed of a few abundant species and many rarer ones (Gaston, 2011; McGill et al., 2007). Traditionally, research has focused on the role of dominant species in ecosystem functioning, but both dominant and rare species contribute to ecosystem stability and processes (Avolio et al., 2019; Jain et al., 2014; Lyons et al., 2005; Lyons and Schwartz, 2001; Richardson et al., 2012; Säterberg et al., 2019; Smith and Knapp, 2003). Dominant plant species influence primary production, nutrient cycling, and soil microbial composition, due to their high biomass and resource use (Grime, 1998; Tedersoo et al., 2020). Their functional traits affect soil N availability by regulating N mineralization and nitrification (Clemmensen et al., 2021; Kielland, 1995; Liu et al., 2018; Michelsen et al., 1996; Ramm et al., 2022; Rozmoš et al., 2022; Tunlid et al., 2022). Rare species, in contrast, often exhibit higher functional diversity and may fill ecological roles not occupied by dominant species, facilitating niche differentiation and promoting ecosystem resilience (Dee et al., 2019; Hooper et al., 2005; Leuzinger and Rewald, 2021; Mouillot et al., 2013; Soliveres et al., 2016; Tang et al., 2023). While their overall biomass contribution is lower, their diverse traits and microbial interactions could play an important role in nutrient partitioning. Both dominant and rare species can form mycorrhizal associations, but differences in mycorrhizal types and plant-microbe interactions may drive variation in N cycling at the community level (Knops et al., 2002; Van der Krift and Berendse, 2001; Moreau et al., 2015, 2019).



Arbuscular mycorrhizal (AM) fungi are considered less common in Arctic ecosystems due to low cold tolerance (Kilpeläinen et al., 2016; Kytöviita, 2005; Ruotsalainen and Kytöviita, 2004; Wang et al., 2002), while ectomycorrhizal (EcM) and ericoid mycorrhizal (ErM) fungi dominate (Michelsen et al., 1998; Soudzilovskaia et al., 2017; Steidinger et al., 2019). Different mycorrhizal types differ in their influence on N mineralization rates and inorganic N availability (Björk et al., 2007; Phillips et al., 2013; Tedersoo et al., 2020). AM fungi facilitate rapid N turnover by promoting inorganic N uptake (Govindarajulu et al., 2005; Hodge and Storer, 2015; Savolainen and Kytöviita, 2022), EcM fungi access both organic and inorganic N, leading to intermediate N turnover rates (Hobbie et al., 2009; Kohler et al., 2015; Miyauchi et al., 2020; Orwin et al., 2011; Pelltier and Zak, 2018), and ErM fungi specialize in mobilizing N from complex organic compounds, contributing to slower N cycling (Bending and Read, 1996; Clemmensen et al., 2021, 2024; Fanin et al., 2022; Tybirk et al., 2000; Wurzbarger and Hendrick, 2009). In ecosystems dominated by a single mycorrhizal type, nutrient cycling may become increasingly constrained by that symbiosis, leading to homogenized soil N dynamics. For example, EcM fungi effectively access organic-N, stabilizing it in less labile forms and reducing N losses, whereas AM fungi promote greater N mobility, potentially increasing N loss (Hobbie and Ouimette, 2009). In contrast, communities composed of less abundant, locally rare species may support different or complementary N cycling functions compared to those dominated by the most abundant species, potentially enhancing functional redundancy and buffering against environmental fluctuations — even when species richness is held constant. To understand these dynamics, it is essential to disentangle the effects of plant dominance, species diversity, and mycorrhizal associations on N cycling.

The aim was to determine how the functional diversity in plant-mycorrhizal associations (AM/NM vs EcM/ErM) and structural diversity (dominant vs. rare species) regulate soil N cycling, and which diversity component is the primary driver. To address this, we conducted a plant removal experiment and in-situ ¹⁵N labelling to determine gross N mineralization and nitrification rates, processes central to N availability in soils as they regulate N supply and loss. Additionally, we used quantitative PCR (qPCR) to quantify six microbial genes related to nitrification, assessing the genetic potential for this process. Thus, we hypothesize that (1) gross N mineralization rates will be highest in EcM/ErM-dominated plots, as these fungi are particularly efficient at accessing organic N sources, breaking down complex organic compounds, and mobilizing N from organic matter. In addition, (2) gross nitrification rates will be highest in AM/NM-dominated plots, since AM fungi and NM plants promote rapid N turnover and enhance the activity of nitrifying microorganisms. (3) The higher gross nitrification rates will also correspond with greater genetic potential for ammonia- and nitrite oxidation in AM/NM-dominated plots. Finally (4) mycorrhizal type will have a stronger influence on N processes than plant community structure (i.e. Rare vs Dominant), as mycorrhizal fungi directly affect N acquisition and cycling through their symbiotic relationships with plants, whereas the influence of plant community structure is more indirect.



2 Methods

2.1 Study site and design

This study was conducted at the Tarfala Research Station in the Tarfala valley of the Kebnekaise Mountains, northern Sweden, at elevations ranging from 1098 to 1114 m a.s.l. (Latitudes: 67°54'14.16"N to 67°54'15.16"N, Longitudes: 18°37'23.80"E to 18°37'29.39"E). The geomorphology of the valley reflects its glacial history, with landforms shaped by retreating ice masses and a substrate dominated by rocky debris. The study area is situated near the terminal moraines marking the maximum extent of Storglaciären during the Little Ice Age (~1910) (Holmlund, 1987) and is characterized by shallow soils developed on till, classified as Leptosols and Regosols (Fuchs et al., 2015). Prominent plant species are the graminoids *Carex bigelowii*, *Carex nigra*, *Deschampsia flexuosa*, *Festuca vivipara*, and *Juncus trifidus*; the deciduous dwarf shrubs *Salix polaris* and *Vaccinium uliginosum*; the evergreen dwarf shrubs *Dryas octopetala* and *Empetrum nigrum*; and the forbs *Bistorta vivipara* and *Silene acaulis*. The mean annual air temperature from 1995 to 2019 was -2.6 ± 1.8 °C with the coldest month in February (-10.5 ± 5.5 °C) and the warmest month in July (8.4 ± 3.7 °C) (SMHI 1995-2019; raw data retrieved from www.smhi.se). The summer mean precipitation is 458 ± 201 mm (Dahlke et al., 2012, Tarfala Research Station 1980-2011; available at <https://bolin.su.se/data/tarfala/climate.php>).

We established a plant removal experiment in 2016 with one unmanipulated control and four treatments designed to manipulate plant community structure: 1) Control, where no plant species were removed; 2) AM/NM, where all plants with EcM or ErM associations were removed, leaving only plants with AM associations or no mycorrhizal association (NM); 3) EcM/ErM, where all plants with AM or NM associations were removed, leaving only plants with EcM or ErM associations; 4) Dominant, where rare plant species were removed, leaving the eight most dominant species (9 rare species removed; Table S1); and 5) Rare, where dominant plant species were removed, retaining the eight rarest species (7–11 dominant species removed; Table S1). Rare and dominant plant species were determined by reducing species richness to approximately 50 % of the community species pool while maintaining a mixed mycorrhizal plant community. Species removal was performed by clipping vegetation at the soil surface, with treatments maintained from 2016 to 2019 by removing regrowth of undesired species each growing season. The treatments were distributed across 32 plots arranged into eight blocks, each containing four plots (one for each treatment group, except for Rare and Dominant, which were represented in four blocks each). There were eight replicates for the AM/NM-dominant, EcM/ErM-dominant, and control treatments, and four replicates for the Rare and Dominant plant community treatments. Each consisted of a smaller survey area (1 m²) to exclude edge effect of the trenching (4 m²) designed to exclude external mycorrhizal colonization. Trenches were dug around the 4 m² perimeter and lined with 1 µm mesh to a depth of 0.3 m, allowing water movement but preventing root and mycorrhizal penetration.



2.2 Plot-level plant diversity

130 To determine plot-level plant community structure, we conducted two vegetation surveys (pre-clipping, July 2015 and post-clipping, July 2019) using point intercept measurements (Molau and Mølgaard, 1996) on the central 1 m² quadrats for each plot. In addition, all species within the 4 m² plot not registered by point intercept were noted. We estimated the cover and counts of each species to determine species richness. We also calculated the transient changes in community dynamics initiated by altered plant interactions and estimated changes in above-ground biomass (Molau, 2010).

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2.3 Nitrogen dynamics

Gross soil N dynamics were investigated in the field using the virtual soil core ¹⁵N tracing approach (Rütting et al., 2011) and a mirror ¹⁵N labelling approach, allowing investigation of N transformations in the intact mycorrhizosphere. Within each plot, we set up two groups of four injection locations in opposing corners: one corner for ¹⁵N-labelled ammonium (NH₄⁺) and the
140 other for ¹⁵N-labelled nitrate (NO₃⁻), to avoid cross-contamination. We conducted the ¹⁵N labelling by injecting ¹⁵(NH₄)₂SO₄ (Cambridge Isotope Laboratories) to quantify gross N mineralization or K¹⁵NO₃ (Cambridge Isotope Laboratories) to quantify nitrification, both labels with a ¹⁵N fraction of 99 %, to a soil depth of 6 cm, both treatments also receiving the unlabelled other moiety. There were 11 injection points per location using a template for guidance (Rütting et al., 2011), each point receiving 1.14 mL of solution containing 15.0 µg NH₄-N mL⁻¹ and 4.5 µg NO₃-N mL⁻¹, which is equivalent to c.a. 3.8 µg NH₄-N g⁻¹ dry
145 soil and 1.2 µg NO₃-N g⁻¹ dry soil. These amounts were calculated based on soil concentrations measured in tundra soil in the nearby Latnjajaure Field Station (Björk et al., 2007). We destructively harvested soil cores at 2, 25, 49, and 97 hours after labelling, using sharpened PVC tubes (3 cm in diameter) inserted down to 6 cm depth at the four respective locations.

Soil cores were immediately processed at the Tarfala Research Station to extract inorganic N (i.e., NO₃⁻ and NH₄⁺) following
150 initial sieving (mesh size 2 mm). 10 g of field moist soil were extracted using 20 ml of 1 M KCl and placed on a shaker for 60 min at 250 rpm before filtration with Whatman 1 G/F filter paper (11 µm). The extracts were stored frozen at -20°C until further analyses. Concentrations and the ¹⁵N fraction of NH₄⁺ were determined from soil KCl extracts using the micro diffusion technique (Biasi et al., 2022; Brooks et al., 1989), followed by ¹⁵N analysis on an elemental analyzer (Europa EA-GSL, Sercon Ltd., UK) coupled to an Isotope Ratio Mass Spectrometer (Sercon 20-22, Sercon Ltd., UK). NO₃⁻ concentrations and the ¹⁵N
155 fraction in all samples were determined from soil KCl extracts using the SPINMAS technique (Stange et al., 2007). The TN, TC, C:N ratio, and bulk ¹⁵N in soil were measured using the EA-IRMS described above. Dried soil was first ground (Retsch MM400, frequency 23.0 1/s, for 2 min) and around 15 mg from each sample was placed into a tin capsule.



2.4 Soil characteristics

160 Samples for abiotic and biotic soil characteristics were taken from the top 6 cm of soil (organic layer only) to match the soil
depth used during the ^{15}N labelling. Four soil samples ($10 \times 10 \times 6$ cm 250 cm^3 each) were collected from each plot after the ^{15}N
labelling experiment to avoid destructive sampling within the plot during the experiment. Stones, plant shoots and roots were
removed from the collected samples immediately after sampling at the Tarfala Research Station, which were then sieved
through a 2 mm mesh. The four sieved soil samples from each plot were combined and homogenized. Subsamples were
165 separated from the homogenized soil for various analyses elsewhere, including pH, gravimetric soil water content (GWC, g/g),
soil organic matter (SOM), and DNA extraction for abundance of microbial communities. Subsamples for DNA extraction
were stored frozen until further analyses.

GWC was measured by oven-drying 10 g of wet soil at 100°C for 24 hours. SOM content was determined using the loss-on-
170 ignition method by heating the soil at 550°C for 6 hours. Soil pH was measured in water (10 g soil, 1:1 deionized water) and
in 1 M KCl (10 g soil, 1:4). Field measurements of soil temperature at a depth of - 5 cm ($T_{\text{soil}}^\circ\text{C}$) and soil water content at 0 –
6 cm (volumetric soil moisture content; VWC) were recorded on four days corresponding to the ^{15}N injection time points.
These measurements were taken at four locations within each plot using a hand-held thermometer and an ML3 ThetaProbe
(Delta-T Devices, Cambridge, U.K.), respectively. Bulk density was determined using intact soil cores (5 cm length, 7.2 cm
175 diameter, 203.6 cm^3 volume), collected by block ($N = 8$), and oven dried at 100°C for 24 hours.

2.5 DNA extraction and qPCR of the ITS region and 16S rRNA and nitrification-associated genes

The frozen, sieved soil was freeze-dried and then ground for 2 minutes using a ball mill. The DNA was extracted from 0.25 g
of the milled soil using the NucleoSpin soil kit (Macherey-Nagel, Duren, Germany), with SL2 buffer with enhancer and
180 according to the manufacturer's protocol. DNA was quantified using the Qubit 2.0 fluorometer (Invitrogen, Thermo Scientific).
qPCR was used to determine the size of total bacterial and fungal communities in the soil by quantifying 16S rRNA gene and
ITS, respectively. Additionally, nitrification-associated functional genes were quantified, including *amoA* (encoding ammonia
monooxygenase from archaeal (AOA) and bacterial (AOB) ammonia oxidizers, and clade A (comA) and clade B (comB)
complete ammonia oxidizers in the *Nitrospira* genus) and *nxrB* (encoding nitrite oxidoreductase from either *Nitrospira*-type
185 (NIS) or *Nitrobacter*-type (NIB) nitrite oxidizing bacteria) (Table S2). Quantification was done using the C1000TM Thermal
Cycler CFX96TM Real-Time System, and CFX ConnectTM Real-Time System (BioRad, CA, USA). All reactions were carried
out in duplicate with a 15 μL reaction volume containing 0.1 mg/mL BSA, 1x SYBR Green Supermix (BioRad), 0.2-1.0 μM
of each primer (Table S2), and 6 ng of template DNA. Standard curves were generated for each gene using serial dilutions
(10^2 - 10^8 copies/ μL) of linearized plasmids containing the target genes. The cycling conditions, primer sequences, and
190 concentrations for each gene are available in Table S3. The amplifications were validated by melting curve analyses and



agarose gel electrophoresis. Prior to quantification, potential inhibition of PCR reactions was checked by amplifying a known amount of the pGEM-T plasmid (Promega, Madison, WI, USA) using plasmid specific M13 primers and addition of soil DNA or non-template controls for each sample. No inhibition was detected with the amount of DNA used. Gene copy numbers were adjusted for the amount and concentration of extracted DNA and normalized per gram of dry soil.

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2.6 Data analysis

All statistical analyses were conducted in R (R Core Team, 2024) with RStudio interface (Posit team, 2024), except for the Isotope tracing model described below. R packages used included *tidyverse* (v2.0.0, Wickham et al., 2019), *rstatix* (v0.7.2, (Kassambara, 2023b)), *knitr* (v1.45, (Xie, 2023)), *kableExtra* (v1.4.0, (Zhu, 2024)), *ggpubr* (v0.6.0, (Kassambara, 2023a)), *sjPlot* (v2.8.15, Lüdtke, 2023). Additional R packages are described within the methods below.

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2.6.1 Vegetation diversity

Vegetation data from both surveys were analyzed using Correspondence Analysis (CA) to explore the relationships between time and treatment. One EcM/ErM plot was removed from the analysis because it had a vascular plant species richness of zero based on the point-framing survey in 2019. This plot had 88 % bryophyte cover and although *Salix herbacea*, *S. polaris*, and *Empetrum nigrum* were still present there were no direct hits from the point-framing survey, indicating a presence of less than 1 % coverage.

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2.6.2 Isotope (^{15}N) tracing model

Process-specific gross N transformation rates were quantified using the ^{15}N tracing model *Ntrace* (Müller et al., 2007; Rütting and Müller, 2007). We used a model setup, including three N pools (organic N, NH_4^+ and NO_3^-) and four N transformation processes: mineralization of organic N (M_{Norg}), immobilization of NH_4^+ and NO_3^- (I_{NH_4} and I_{NO_3}) and NH_4^+ oxidation (O_{NH_4} , i.e. nitrification). The N transformations were described by first-order kinetics, except for M_{Norg} which followed zero-order kinetics. The kinetic parameters of the N transformations were approximated numerically for each treatment separately with Monte Carlo sampling through a random walk aiming to minimize a misfit function (quadratic weighted error) between the modelled and observed values. Model inputs were mean values and standard deviations of NH_4^+ and NO_3^- content and their respective ^{15}N abundances. The initial ^{15}N content of the organic N pool was not measured at the plots and was instead assumed to be at natural abundance (0.366 %). Iterative approximation of the N cycle rates creates normally distributed probability density functions, for which the mean values and standard deviations were calculated (Müller et al., 2007). For pathways described by first-order kinetics, gross N rates were calculated as the product of the kinetic factor and substrate content. *Ntrace* and the optimization algorithm were set up in Matlab version R2023b and Simulink version 23.2 (The MathWorks Inc.). Rates are reported per gram of C to account for differences in organic matter content across soils and to facilitate better comparison.

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The *Ntrace* provides robust estimates of gross N transformation rates but was here applied to treatment averages, hence did not allow investigation of potential block effects. To do so, we additionally quantified gross N mineralization and nitrification for each plot based on the isotope pool dilution (IPD) principle and the analytical tracing model by (Kirkham and Bartholomew, 1954) using the first two timesteps of the ^{15}N tracing experiment. All gross N transformation rates are normalized for the soil C content. To assess potential block effects on gross N mineralization and nitrification rates, we fitted generalized linear models (GLMs) with Block as a fixed effect using the *glmmTMB* package (v1.1.8, Brooks et al., 2017). Given the right-skewed distribution of the data, a zero-inflated Gamma distribution with a log link function was used for mineralization rates, while a standard Gamma distribution was applied for nitrification rates. Model significance was assessed using Type II Wald chi-square tests.

2.6.3 Soil characteristics and microbial genes

To analyse the impacts of mycorrhizal status and vegetation composition on soil characteristics and microbial genes, we fitted Generalized Linear Mixed Models (GLMMs) with *glmmTMB* (v1.1.8, (Brooks et al., 2017)). Each model included Treatment as a fixed effect and Block as a random effect. Given that Block showed significant effects for several response variables, additional GLMs were fitted with Block as a fixed effect to explore its specific influence. These results are presented in the supplementary material for completeness, though Block was not originally intended as a primary focus of the experimental design. We validated models using the *DHARMA* package (v0.4.6, (Hartig, 2022)). Pairwise comparisons between treatments were conducted with *emmeans* (v1.10.0, (Lenth, 2024)).

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We conducted paired samples Wilcoxon signed-rank tests with the *wilcox.test* function within the *stats* package (R Core Team, 2024) to assess differences in log-transformed gene abundances between the sample groups ITS and 16S rRNA, AOA and AOB, ComaA and ComaB, as well as NIB and NIS.

245 We utilized the *corr.test* function within the *psych* package (v2.4.1, (Revelle, 2024)) to conduct correlation analyses to explore the relationships between gene abundances and environmental variables. We calculated Spearman rank-order correlation coefficients to quantify the strength and direction of these relationships. To address the issue of multiple testing and control the family-wise error rate, we applied a Holm correction. We categorized correlation coefficients based on their strength: weak ($0 < |r| < 0.4$), moderate ($0.4 < |r| < 0.7$), and strong ($|r| > 0.7$).

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Principal Component Analysis (PCA) was employed for dimensionality reduction. The first three Principal Components (PCs) were retained, and ANOVAs were performed on them, incorporating Treatment and Block as fixed effects. The ANOVA outputs provided adjusted p-values, which were further examined using Tukey tests to identify significant differences between treatment groups and blocks.

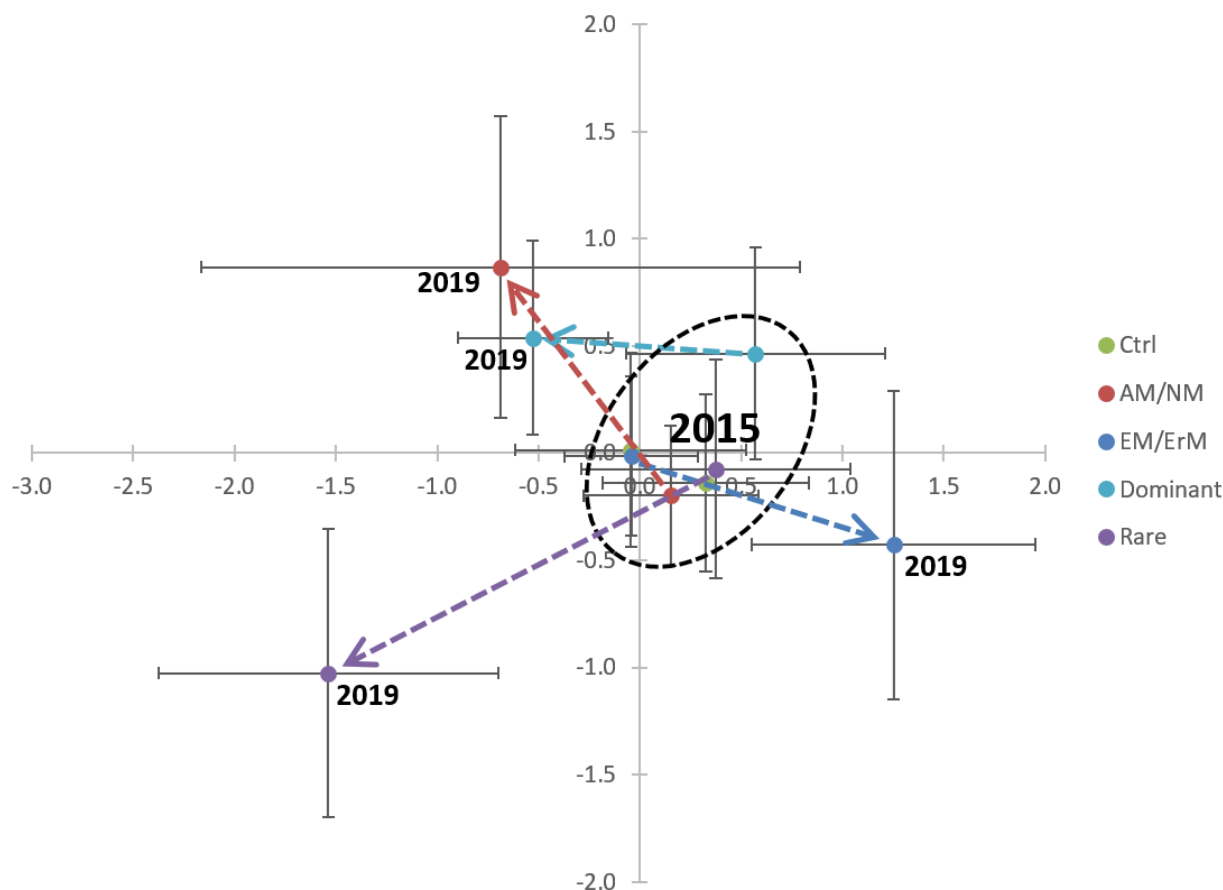


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3 Results

3.1 Vegetation diversity treatment effect

The treatments clearly shifted the plant community in three directions within the ordination space from its original structure in 2015 (Fig. 1). The AM/NM community and the Dominant community clustered together, whereas the EcM/ErM community and the Rare community formed their own distinct clusters after clipping treatment. The control plots in 2019 remained similar to the plant communities recorded in 2015 before the experiment was established.



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Figure 1: Changes in plant communities over the course of the experiment. Mean values ($\pm 85\%$ confidence interval corresponding to an $\alpha = 0.05$ test; see (Payton et al., 2000, 2003) of sample scores from the correspondence analysis (CA), comparing the abundances of plant species before treatment in 2015 and four years after treatment in 2019. The eigenvalues are 0.499 for Axis 1 and 0.469 for Axis 2. Axis 1 explains 10.68 % of the total variance, and Axis 2 explains 10.04 %, together accounting for 20.72 % of the total variance. Treatments: Ctrl = control; AM/NM = plants with



270 arbuscular mycorrhizal association or no mycorrhizal association; EcM/ErM = plants with ectomycorrhizal and ericoid
mycorrhizal associations; Dominant = rare plant species removed allowing the eight most dominant plant species to grow
in the plots; and Rare = dominant species removed keeping the eight rarest plant species.

3.2 Soil characteristics

275 During the labelling period, VWC was significantly higher in EcM/ErM-dominated plots compared to the control ($z = 2.19$, $p = 0.029$). T_{soil} was significantly lower in Dominant plots relative to the control ($z = -2.44$, $p = 0.015$) and marginally lower in
EcM/ErM plots relative to the control ($z = -1.88$, $p = 0.06$). Pairwise comparisons showed that T_{soil} in AM/NM was significantly
higher than EcM/ErM (estimate = 0.02, SE = 0.006, $p = 0.007$) and Dominant (estimate = 0.03, SE = 0.008, $p = 0.003$). SOM
was significantly lower in AM/NM-dominated plots compared to control ($z = -2.35$, $p = 0.019$). No other significant differences
280 were found for the remaining soil characteristics (Table 1, Table S4).

285 **Table 1: Soil properties at the Tarfala study site (Sweden). Variables: soil moisture (VWC), soil temperature (T_{soil}),
laboratory gravimetric soil water content (GWC), soil organic matter (SOM), pH, C:N ratio, TN. Values represent mean
± standard error (N = 32). VWC and T_{soil} are averaged values taken over four days of measurements, while all other
properties are based on one measurement per soil sample collected from each plot. Treatments: only ecto- and ericoid
mycorrhiza plant associations present (EcM/ErM), only arbuscular and non-mycorrhiza associations present (AM/NM);
removal of dominant plant species (Rare); removal of rare plant species (Dominant). Significant differences from control
are bolded (* = $p < 0.05$, # = $p < 0.1$) based on general linear mixed-effects models (GLMMs) (Table S4).**

Treatment	n	VWC (%)	T_{soil} (°C)	GWC (g/g)	SOM (%)	pH	C/N	TN (%)
Control	8	26.0±1.6	10.7±0.1	55.7±1.9	37.2±2.5	4.9±0.1	15.3±0.5	1.1±0.2
AM/NM	8	26.6±1.1	10.9±0.1	52.0±2.3	29.9±2.7*	4.9±0.1	15.6±0.6	0.9±0.1
EcM/ErM	8	29.8±1.9*	10.4±0.2#	54.9±1.4	33.7±1.6	5.0±0.1	14.2±0.6	1.4±0.4
Dominant	4	27.4±2.8	10.4±0.2*	57.6±4.4	38.1±5.9	5.0±0.0	16.7±0.7	0.9±0.1
Rare	4	31.4±2.8	10.5±0.3	53.8±1.8	31.4±3.6	5.0±0.1	14.9±0.7	1.0±0.2

290 Bulk density (g cm^{-3}) was measured by block, not by treatment: A) 0.18, B) 0.22, C) 0.21, D) 0.22, E) 1.01, F) 0.37, G) 0.41, H) 0.36.
Natural abundance $\delta^{15}\text{N}$ was measured by block, not by treatment: A) 0.95, B) 1.76, C) -0.08, D) 0.92, E) 2.62, F) 2.16

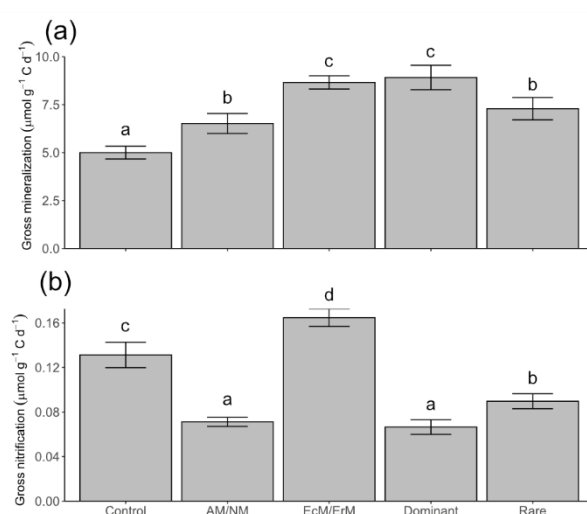
3.3 Gross nitrogen dynamics

Compared to the control, all treatments showed significantly higher gross N mineralization rates (Fig. 2a). EcM/ErM and
Dominant showed the largest increases, at 73 % and 78 % above the Control, respectively, while AM/NM and Rare had more
295 moderate increases of 30 % and 46 %. Gross nitrification rates were 1-2 orders of magnitude lower than gross N mineralization
rates (Fig. 2b). Significant differences in gross nitrification rates were also observed, with EcM/ErM showing a 26 % increase,



while Rare, AM/NM, and Dominant exhibited reductions of 32 %, 46 %, and 49 %, respectively, compared to the Control (Fig. 2b).

300 Average gross N mineralization and nitrification rates, calculated using the IPD approach, showed a similar pattern to those obtained through *Ntrace* (Table S5). However, the IPD based rates had much higher variability. In some instances, we even observed implausible negative rates. Block effects on gross N transformation rates were not statistically significant for mineralization ($X^2(7) = 9.19$, $p = 0.24$), but marginally significant for nitrification ($X^2(7) = 13.08$, $p = 0.07$).



305 **Figure 2: Gross N mineralization and nitrification rates (mean and 85 % confidence interval). Rates were quantified by the *Ntrace* model with different manipulated vegetation (Control = no manipulation; AM/NM = plants with arbuscular mycorrhizal association or no mycorrhizal association; EcM/ErM = plants with ectomycorrhizal and ericoid mycorrhizal associations; Dominant = rare plant species removed allowing the eight most dominant plant species to grow in the plots; and Rare = dominant species removed keeping the eight rarest plant species). Different lowercase letters above the bars**
 310 **indicate significant differences based on whether the 85 % confidence intervals overlap.**

3.4 Abundance of bacteria and fungi

The bacterial 16S rRNA gene copy numbers were consistently higher than fungal ITS rRNA gene copy numbers across all treatments (EcM/ErM, AM/NM, Rare, Dominant), ranging from 2.07×10^9 to 2.75×10^9 and 1.05×10^8 to 1.68×10^8 copies
 315 g^{-1} dry soil respectively ($V = 528$, $p < 0.001$, $n = 32$). In the AM/NM, fungal abundances were marginally lower ($z = -1.67$, $p = 0.094$) and the ITS:16S rRNA gene copy ratio were lower ($z = -2.13$, $p = 0.033$) compared to the Control (Fig. 3a). No other treatments significantly affected the bacterial or fungal abundance (Table S6, S7) or the ITS:16S ratio (Fig. 3a, Table S8, S9).

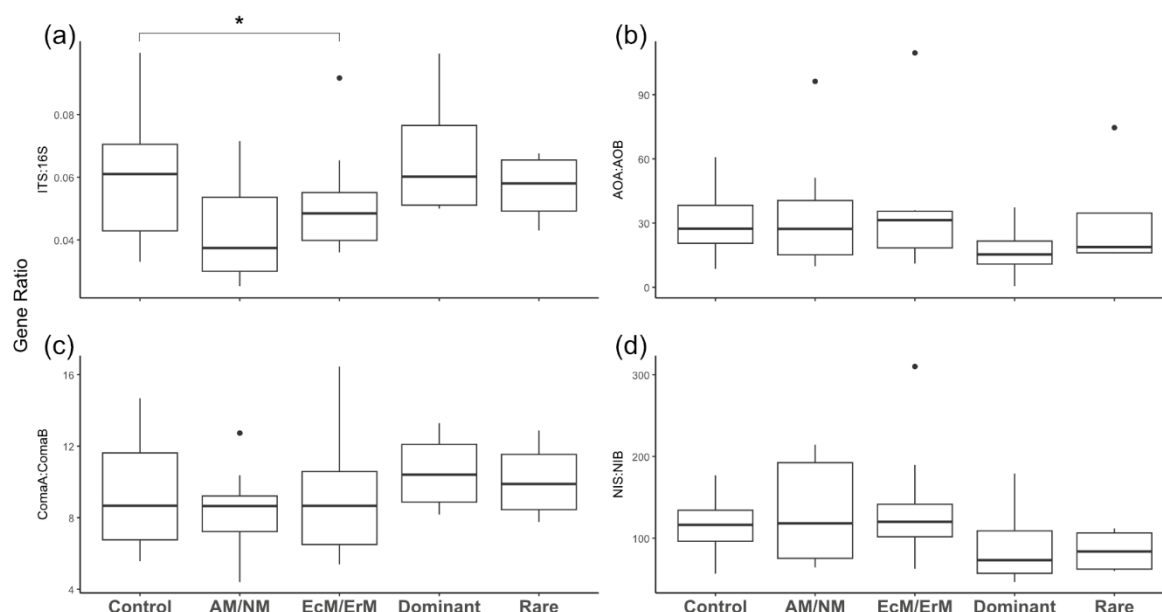


Figure 3: Soil gene abundance ratios in response to plant removal treatment. Gene abundance ratios for A) fungi (ITS) vs bacteria (16S rRNA gene); B) Archaeal ammonia oxidizer (AOA) vs bacterial ammonia oxidizer (AOB); C) comammox bacteria, clade ComaA vs comammox clade ComaB; and D) nitrite-oxidizing Nitropira (NIS) vs Nitrospira (NIB). Treatments: no manipulation = Control; removal of plants with ecto and ericoid mycorrhiza associations = AM/NM; removal of plants with arbuscular mycorrhiza & no mycorrhiza associations = EcM/ErM; removal of rare plant species = Dominant; removal of dominant plant species = Rare. Symbols above the boxplots denote significant differences for each group relative to a control group as determined through Generalized Linear Mixed Models (GLMMs) (* < 0.05) (Table S9).

3.5 Nitrifier gene abundance

We observed the most notable variations in nitrification gene copy numbers between functional groups capable of the same transformation step in nitrification. Gene abundances exhibited distinct differences between functional group pairs: AOA > AOB ($V = 527$, $p < 0.001$, $n = 32$); NIS > NIB ($V = 0$, $p < 0.001$, $n = 32$); and ComaA > ComaB ($V = 528$, $p < 0.001$, $n = 32$). Of the six genes, *amoA* in ComaA and *nxrB* NIS, both representing the bacterial genus *Nitrospira*, were consistently the most abundant genes (Table S10).

Overall, gene abundances were minimally affected by treatment, except for Dominant treatment having lower AOA abundance ($z = -2.66$, $p = 0.008$; Fig. 4a), and marginally lower NIS abundance ($z = -1.89$, $p = 0.058$; Fig. 4f) compared to the control (Table S11). Gene copy abundance ratios were unaffected by treatment (Fig 3b-d, Table S9).

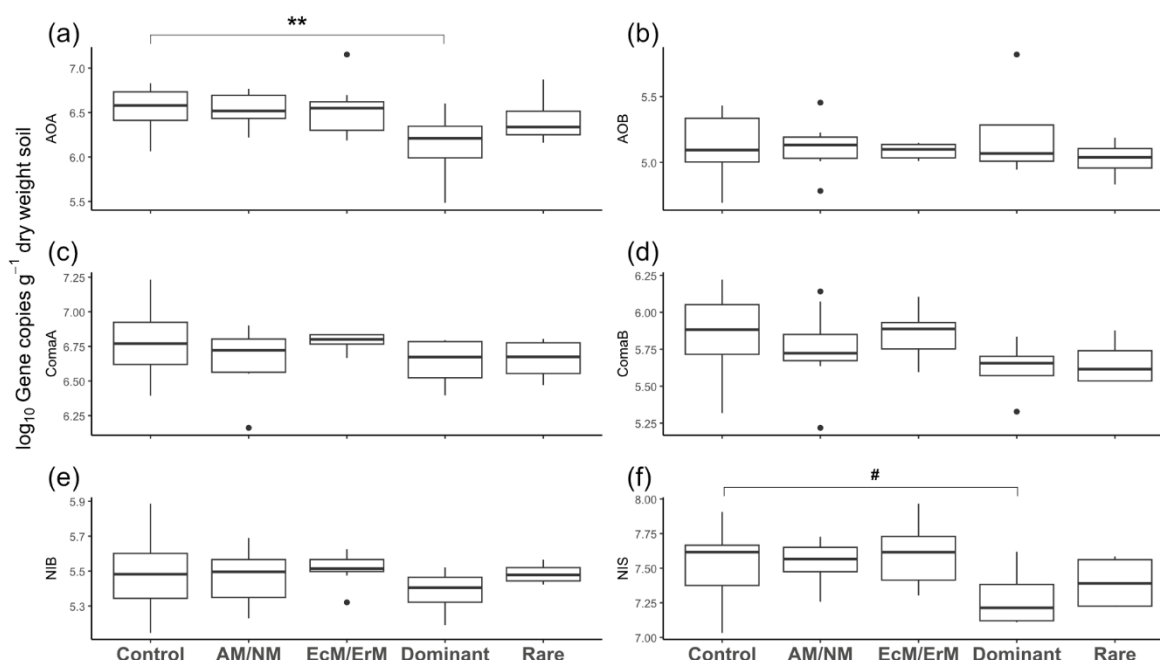


Figure 4: Gene abundances representing six functional groups involved in nitrification in response to plant removal treatment. A) Ammonia-oxidizing archaea (AOA), B) ammonia-oxidizing bacteria (AOB), C) complete ammonia oxidizers (comammox) clade A (comaA), D) complete ammonia oxidizers (comammox) clade B (comaB), E) nitrite-oxidizing Nitrobacter (NIB), and F) nitrite-oxidizing Nitrospira (NIS). Treatments: no manipulation (Control); removal of plants with ecto and ericoid mycorrhiza associations = AM/NM; removal of plants with arbuscular mycorrhiza & no mycorrhiza associations = EcM/ErM; removal of rare plant species = Dominant; removal of dominant plant species = Rare. Symbols above the boxplots denote significant differences for each group relative to a control group as determined through Generalized Linear Mixed Models (GLMMs) (< 0.01, # < 0.1) (Table S11).**

3.6 Relationships between gene abundances, vegetation, and edaphic factors

We found no significant correlations between gene abundances and Simpson's diversity index of plants, VWC, GWC, T_{soil} , SOM, pH, TN, C/N, and BD after adjusting for multiple testing (Table S12). However, we observed a strong positive correlations between abundance of 16S rRNA genes and ITS ($r = 0.73$, $p < 0.01$), AOB ($r = 0.73$, $p < 0.01$), ComaA ($r = 0.77$, $p < 0.01$), NIB ($r = 0.75$, $p < 0.01$), AOA and NIS ($r = 0.81$, adj. $p < 0.01$), ComaA and ComaB ($r = 0.72$, $p < 0.01$), and moderate positive correlations between 16S rRNA genes and ComaB ($r = 0.67$, adj. $p < 0.01$), ITS and AOB ($r = 0.69$, $p < 0.01$), AOB and ComaA ($r = 0.63$, $p = 0.02$), ComaA and NIB ($r = 0.63$, adj. $p = 0.02$), and ComaB and NIB ($r = 0.60$, $p = 0.04$) gene abundances. We also observed moderate positive correlations between Simpson's diversity index of plants and VWC ($r = 0.63$, adj. $p = 0.02$), VWC and BD ($r = 0.60$, adj. $p = 0.04$), and BD and elevation ($r = 0.60$, adj. $p = 0.04$). There was a marginally moderate negative correlation between T_{soil} and BD ($r = -0.58$, adj. $p = 0.08$).

When considering the combined effects of vegetation diversity, soil characteristics, and the abundance of bacterial, fungal, and nitrifier genes, the first three principal components accounted for 58.2 % of the total variance, with PC1, PC2, and PC3



explaining 25.9 %, 18.5 %, and 13.7 %, respectively (Fig. S1, S2). The loadings for each component indicate that no single variable drives the variance (Table S13). The strongest positive loadings on PC2 were vegetation diversity, VWC, AOA abundance, and NIS abundance, while the strongest negative loading was C/N ratio. For PC3, T_{soil} was the strongest negative loading, and elevation, VWC, GWC, and BD were the strongest positive loadings. However, some variables show relatively stronger contributions to certain components. For PC1, neither Treatment ($F_{(4,20)} = 0.81$, $p = 0.53$) nor Block ($F_{(7,20)} = 1.28$, $p = 0.31$) had a significant effect on the PC1 scores. For PC2, Treatment showed a significant effect on PC2 scores ($F_{(4,20)} = 3.40$, $p = 0.028$), while Block was not significant ($F_{(7,20)} = 1.52$, $p = 0.22$). Tukey's test indicated a significant difference between the EcM/ErM and Dominant (adj. $p = 0.015$), and a notable difference between Rare and Dominant (adj. $p = 0.079$) treatment groups. For PC3, Treatment ($F_{(4,20)} = 3.71$, $p = 0.02$) and Block ($F_{(7,20)} = 12.01$, $p = 0.00$) showed a significant effect on PC3 scores. Tukey's test indicated a significant difference between the EcM/ErM and AM/NM ($p = 0.039$), and between Rare and AM/NM treatment groups ($p = 0.038$). Tukey's test also showed significant differences between Blocks with Blocks A-D showing negative PC scores and Blocks E-H showing positive scores (Fig. S2). Elevation, which increased from Block A to H (Table S14, Fig. S3a), influenced multiple properties despite the subtle 15-meter gradient. The proportional cover of EcM/ErM plots decreased from Block A-G but deviated in Block H, where the cover resembled that of Blocks B-D. Additionally, Blocks A-D were drier than Blocks E-F, and soil temperature decreased with elevation.

4 Discussion

4.1 EcM/ErM communities enhance both gross mineralization and nitrification in a conservative tundra N cycle

As hypothesized, we found the highest gross N mineralization rates in the EcM/ErM treatment, but unexpectedly, the treatment with only dominant species in the plant communities also exhibited high rates. Notably, all treatments showed elevated mineralization compared to the unmanipulated control. By contrast, our hypothesis was not supported for gross nitrification. The EcM/ErM treatment was the only one showing higher nitrification rates compared to control, while all other treatments exhibited decreased rates. A previous study in a hemiboreal forest found that the presence of EcM increased gross N mineralization threefold, while gross nitrification remained largely unaffected (Holz et al., 2016). EcM-dominated ecosystems are commonly assumed to cycle N more slowly because EcM fungi promote organic N retention and decomposition of more recalcitrant substrates, whereas AM-dominated ecosystems exhibit faster N cycling due to greater reliance on inorganic N uptake and relatively fast N mineralization rates (Averill et al., 2019). However, our small-scale experimental study does not support this hypothesis, as we found significantly higher gross N cycling in the presence of EcM/ErM compared to the plots with AM/NM. This is consistent with a recent meta-analysis on rhizosphere effects on gross N mineralization (Gan et al., 2022), demonstrating that EcM-associated plant species enhanced gross N mineralization more than AM-associated species. EcM/ErM mycorrhizal treatments circulated N faster than the other treatments, also indicated by the lower gross mineralization-to-nitrification ratio in the EcM/ErM mycorrhizal treatments (53 for EcM/ErM mycorrhizal treatments vs. 92,



81, and 134 for AM/NM and the diversity treatments). According to the mass ratio hypothesis, the plant functional traits and
395 relative abundances of dominant species within a community are highly influential for ecosystem processes (Grime, 1998).
Our study partly supports the mass ratio hypothesis by demonstrating that mycorrhizal type, particularly EcM/ErM, can be
regarded as a key functional plant trait influencing N cycling. However, we found that dominant species were not necessarily
associated with faster or more open N cycling overall, despite high mineralization rates. The high gross mineralization-to-
nitrification ratio (134) in the Dominant treatment suggests a more conservative, ammonium-driven N cycle. This may reflect
400 competitive dynamics, where dominant species more effectively acquire NH_4^+ , thereby reducing substrate availability for
nitrifiers. In this way, dominant species could exert a strong influence on the N cycle by both enhancing mineralization and
constraining nitrification, resulting in a faster but tighter cycle that favours internal N recycling. By contrast, rare species
communities exhibited lower mineralization but relatively higher nitrification (gross mineralization-to-nitrification ratio of
81), potentially indicating a more open N cycle and increased risk of N losses via leaching or gaseous pathways. These
405 differences may arise from functional similarity and resource monopolization in dominant communities (Eisenhauer et al.,
2023), versus greater functional complementarity and microbial interactions in rare communities (Niklaus et al., 2006). Thus,
our findings suggest that mycorrhizal status, particularly EcM/ErM associations, plays a more significant role in shaping gross
N cycling dynamics than species dominance alone.

410 The observed increase in gross N mineralization in all manipulation treatments compared to control could be due to the
increased carbon input from decaying roots of plants that were removed by clipping. Although treatments were initiated four
years before our study, clipping also occurred during the growing season leading up to it. After clipping, roots remain in the
soil and decompose, triggering a priming effect on the microbial community, which increases N mineralization and
rhizodeposition (Bengtson et al., 2012; Dijkstra et al., 2013). Early-stage decomposition generally progresses the fastest due
415 to the rapid loss of soluble carbon (Aber et al., 1990). However, root decay rates significantly decline in the second year
compared to the first (McLaren et al., 2017). Thus, the observed gross N mineralization rates are likely slightly elevated, which
suggests that the gross N mineralization rates are affected by the experimental manipulation. However, the significantly higher
mineralization rates in the Dominant treatment, despite its similar community composition to the AM/NM, indicate that species
identity and associated functions in inherent plant species drive the pattern we observe and play an important role in shaping
420 nitrogen cycling dynamics.

Moreover, we found that gross N mineralization rates were 1-2 orders of magnitude faster than the gross nitrification rate, and
the ratio of gross nitrification to NH_4^+ immobilization was low. This is a strong indicator of a conservative N cycle with
minimal N losses to the environment, which is typical in N-limited ecosystems (Schimel and Bennett, 2004; Tietema and
425 Wessel, 1992). Nitrogen limitation is further supported by our $\delta^{15}\text{N}$ data for SOM. The $\delta^{15}\text{N}$ values of SOM depend mainly
on external N sources and ecosystem N losses. In N-rich ecosystems with high denitrification, N with low $\delta^{15}\text{N}$ is lost, resulting
in higher soil $\delta^{15}\text{N}$ values (Bai and Houlton, 2009). Conversely, in N-limited ecosystems, the primary input is via biological



N fixation, which has minimal fractionation, resulting in soil $\delta^{15}\text{N}$ values close to 0 (Amundson et al., 2003), as we observe. Few studies have investigated gross N cycling rates *in situ* in tundra ecosystems (Ramm et al., 2022), but our gross N
430 mineralization rates in the control plots ($5.0 \pm 0.3 \mu\text{mol g}^{-1} \text{C d}^{-1}$) is similar to *in situ* rates obtained in other low Arctic and
oroarctic ecosystems (Buckeridge et al., 2010; Gil et al., 2022; Paré and Bedard-Haughn, 2012). Rates on *in situ* gross
nitrification is even more scarce for tundra ecosystems. The global average gross nitrification rate in mineral soils has been
estimated to $0.56 \mu\text{mol g}^{-1} \text{C d}^{-1}$ (Elrys et al., 2021), whereas in permafrost mineral soils it is about half this rate, $0.27 \mu\text{mol g}^{-1} \text{C d}^{-1}$ (Ramm et al., 2022). Our control plot nitrification rates are lower ($0.13 \pm 0.01 \mu\text{mol g}^{-1} \text{C d}^{-1}$), and also in the lower end
435 of what been observed in alpine grasslands, 0.16 and $0.27 \mu\text{mol g}^{-1} \text{C d}^{-1}$ (Jin et al., 2023; Shaw and Harte, 2001). High soil
C content ($> 5\%$) can decouple N mineralization and nitrification (Gill et al., 2023) by increasing heterotrophic N demand
and intensifying competition for ammonium between heterotrophs and autotrophs (Booth et al., 2005; Keiser et al., 2016; Silva
et al., 2005). Hence, our gross rates suggest that N availability in the Fennoscandian oroarctic tundra is low and low enough
for the ecosystem to operate with a conservative N cycle. This leads to reduced N losses and further reinforces that N is a
440 limiting factor controlling ecosystem productivity.

4.2 Distinct soil nitrifier community within an otherwise stable microbial community

Despite the distinct roles of mycorrhizal fungi in N cycling (Castaño et al., 2023; Hobbie and Högberg, 2012; Tedersoo et al.,
2020), the AM/NM and EcM/ErM plots did not differ in N-cycling gene abundances. However, altering plant composition
445 revealed differences. Prior studies show AM-dominated soils contain more inorganic N and up to five times more N-cycling
gene copies than EcM soils ((Mushinski et al., 2021; Zhang et al., 2022), while EcM is linked to greater gross N mineralization
but lower nitrification rates than AM (Seyfried et al., 2023). Our Dominant community plots had lower abundance of AOA
and NIS functional groups, coinciding with reduced gross nitrification rates, likely because plants outcompete nitrifiers for
 NH_4^+ (Hayashi et al., 2016) or from decreased microbial reliance on NH_4^+ (Hobbie and Hobbie, 2006; Schimel and Chapin,
450 1996). In Rare community plots, nitrification gene abundances were comparable to Control plots despite lower gross
nitrification rates. Since plant species richness was similar across Dominant and Rare treatments, our results suggest that
dominant species traits may drive ecosystem function, echoing findings in the ecological literature (Grime, 1998; MacGillivray
et al., 1995). Interestingly, these traits seem to differ or be suppressed in the AM/NM community, which showed a similar
plant composition to Dominant plots. Thus, the presence of EcM/ErM plants in an AM/NM-dominated community, even as a
455 minor component, can shift how plant community traits influence N dynamics in Arctic soils.

There were limited treatment effects on the abundance of nitrification genes, and bacterial and fungal communities, except in
the AM/NM community, which showed reduced fungal abundance. This reduction led to a significantly lower ITS:16S rRNA
gene ratio compared to the Control. Although Arctic soils contain a high abundance of AM fungi, they are typically dominated
by EcM/ErM fungi (Brachmann et al., 2025), likely explaining the decrease in ITS:16S rRNA gene in the AM/NM community.



460 Despite the limited treatment effects, distinct communities for ammonia oxidation and nitrite oxidation emerged. For ammonia
oxidation, AOA was more abundant than AOB, consistent with findings from other Arctic soils (Alves et al., 2013; Banerjee
et al., 2011; Lamb et al., 2011). AOA, with their flexible ammonia requirements (Verhamme et al., 2011), often dominate in
low-N (Di et al., 2010; Erguder et al., 2009) and acidic soils (Gubry-Rangin et al., 2010; Prosser and Nicol, 2012). Their
metabolic and physiological versatility (Alves et al., 2019), along with cold-tolerance genes (Pessi et al., 2022), highlights
465 their important role in N cycling in Arctic soils. Further differentiation was observed among comammox clades (ComaA and
ComaB), with ComaA being more abundant. Although comammox is underexplored in Arctic soils (Guo et al., 2024), studies
from coastal Antarctica indicate clade B dominates nitrification there (Han et al., 2024). Biochemical differences, such as
hydrogen oxidation capabilities (Han et al., 2024; Palomo et al., 2018), may contribute to niche specialization. Arctic soils,
experiencing periodic anoxia during thawing, likely favour ComaA due to its adaptation to fluctuating oxygen conditions,
470 including its ability to switch metabolic pathways and efficiently manage oxygen consumption through hydrogen oxidation.
Among nitrite oxidizers, NIS was more abundant than NIB. NIS thrives under low-nitrite conditions, where its periplasmic
localization provides a competitive advantage (Nowka et al., 2015). However, the periplasmic location also makes NIS more
sensitive to environmental fluctuations (Wilks and Slonczewski, 2007), whereas NIB, with its cytoplasmic NXR complex, can
outcompete NIS under less stable conditions and at higher nitrite concentrations (Bartosch et al., 2002; Taylor and Mellbye,
475 2022). Our results show a distinct nitrifier community and suggest that Arctic soils favour a more resource-efficient, yet
environmentally responsive, ammonia and nitrite oxidation strategy, supporting our findings of a conservative N cycle.
Moreover, we observed correlations between nitrification genes (Table S12), including a strong positive correlation between
AOA and NIS, suggesting potential synergistic interactions (Jones and Hallin, 2019; Ke et al., 2013; Stempfhuber et al., 2016)
within the microbial community. This reinforces the idea that N cycling in these soils is structured by microbial traits and
480 environmental pressures rather than competitive interaction with plants and mycorrhizal fungi.

4.3 Mismatch between gene abundances and *in-situ* activity

We found a mismatch between genetic potential for nitrification and *in situ* activity (gross nitrification rates) in the mycorrhizal
manipulated plots. While higher gene abundances sometimes correlate with nitrification potential and rates (Ke et al., 2013;
485 Laffite et al., 2020; Ribbons et al., 2016; Rocca et al., 2015), similar inconsistencies as in our study have been observed in
high-Arctic soils, where the abundances of ammonia-oxidizing archaea (AOA) and ammonia-oxidizing bacteria (AOB) do not
always correlate with ammonia oxidation potential (Hayashi et al., 2016). Thus, gene abundance alone does not necessarily
predict nitrification rates, as environmental factors (Avrahami and Conrad, 2003; Hicks et al., 2020b; Hu et al., 2014; Li et al.,
2020a; Oshiki et al., 2016; Rousk et al., 2010; Stempfhuber et al., 2016; Taylor and Mellbye, 2022; Wright and Lehtovirta-
490 Morley, 2023), and competition (Huang et al., 2024; Jung et al., 2022; Yang et al., 2022) likely play an interacting role.
Additionally, our gene targets did not encompass alternative N sources, for example nitrogen fixation (Castaño et al., 2023) or
the full nitrification potential of the soil. For example, *Nitrotoga*, a cold-adapted genus of nitrite oxidizing bacteria (NOB)



(Alawi et al., 2007), competes with our targeted groups of NOB (NIB and NIS) (Alawi et al., 2009; Karkman et al., 2011; Nowka et al., 2015), but was not considered in our study.

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4.4 Limited impact of environmental factors

Overall, neither mycorrhizal type nor plant species richness treatments had a strong influence on soil properties, nor did soil properties affect nitrification gene abundances. This result was unexpected, as above- and below-ground processes are often considered interconnected (Wardle et al., 2004). Changes in mycorrhizal type and vegetation typically influence soil properties (Netherway et al., 2021; Welker et al., 2024; Wurzbürger and Brookshire, 2017), and shifts in soil conditions, management practices, or environmental conditions can affect N dynamics (Björk et al., 2007; Li et al., 2020b) and nitrification gene abundances (Zhan et al., 2023). However, vegetation is not always the primary driver of N dynamics; other environmental factors, like soil moisture, can play a more important role (Fisk et al., 1998). Recent studies suggest that below-ground communities and functions can resist changes in vegetation cover and diversity (Fanin et al., 2019; Kirchhoff et al., 2024). Consistent with this, we found no clear environmental drivers of gene abundance (Table S12). However, we observed relationships among environmental factors: vegetation diversity was positively correlated with VWC measured during the week of the labelling, while soil bulk density was positively related to elevation and VWC but negatively related to soil temperature. These relationships may be temporally dynamic, as soil moisture can strongly influence N transformation rates earlier in the growing season, with its effect diminishing later in the season (Steltzer and Bowman, 1998). Notably, our soil samples were collected during the late growing season. When analyzing vegetation diversity, soil characteristics, and gene abundances together, clear treatment differences emerged. Differences were observed between EcM/ErM and Dominant, and to a lesser extent between Dominant and Rare, driven by vegetation diversity, VWC, AOA and NIS abundances, and C:N ratio. Differences between AM/NM and EcM/ErM, and AM/NM and Rare were driven by soil temperature, elevation, VWC, GWC, and soil bulk density. Block effects also emerged as a key factor. There were distinct and subtle environmental gradients represented in elevation change (over a short 15-meter gradient), vegetation cover, and soil characteristics (Table S14, Fig. S3a,b). Similar block effects were observed in another plant removal study involving plant-mycorrhizal associations (Kirchhoff et al., 2024) even after two years of treatment. Notably, our study spanned four years, further highlighting the persistence of these spatial influences.

520 5 Conclusions

Our study reveals that EcM/ErM mycorrhizal associations significantly enhance N cycling in Oroarctic tundra, challenging the conventional view that EcM-dominated ecosystems cycle N more slowly. Elevated gross N mineralization rates in EcM/ErM plots suggest that these fungi are more efficient at accessing and mobilizing N from organic matter. Despite stable



microbial communities, the AM/NM plots showed reduced fungal abundance, reflecting the dominance of EcM/ErM fungi in
525 Arctic soils. Distinct communities for ammonia and nitrite oxidation emerged, with AOA being more abundant than AOB and
NIS more abundant than NIB. This supports a resource-efficient, yet environmentally responsive, N cycling strategy in these
soils. However, a mismatch between gene abundances and nitrification rates suggests that environmental factors and biological
competition play significant roles. Altering plant diversity revealed differences in nitrification gene abundances, with dominant
plots showing lower AOA and NIS gene abundances, indicating that dominant plant species may suppress or outcompete
530 nitrifiers. Our findings emphasize the importance of EcM/ErM in N cycling and provide a deeper understanding of ecosystem
processes in tundra environments. Future research should focus on long-term experiments and monitoring to better understand
how changing plant diversity and mycorrhizal associations under varying climatic conditions affect ecosystem functioning.

Author contribution

535 CRediT authorship contribution statement

Aurora Patchett: investigation, data curation, formal analysis, visualization, original draft preparation, review and editing.
Louise Rütting: conceptualization, investigation, data curation, validation, supervision, original draft preparation, review and
editing. Tobias Rütting: conceptualization, investigation, data curation, formal analysis, visualization, validation, supervision,
original draft preparation, review and editing. Samuel Bodé: investigation, validation, review and editing. Sara Hallin:
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