Fungal community and functional responses to soil warming are greater than for soil nitrogen enrichment

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Soil fungi are key regulators of forest carbon cycling and their responses to global change have effects that ripple throughout ecosystems. Global changes are expected to push many fungi beyond their environmental niches, but there are relatively few studies involving multiple, simultaneous global change factors. Here, we studied soil fungal diversity, community composition, co-occurrence patterns, and decomposition gene responses to 10 years of soil warming and nitrogen addition, alone and in combination. We specifically examined whether there were fungal community characteristics that could explain changes in soil carbon storage and organic matter chemistry in chronically warmed and fertilized soil. We found that fungal communities in warmed soils are less diverse and shift in composition. Warming also favored hyperdominance by a few mycorrhizal fungal species and lowered manganese peroxidase but increased hydrolytic enzyme encoding gene potentials. Nitrogen addition did not significantly affect fungal community composition but, like warming, did reduce fungal diversity and favored overdominance by a unique set of mycorrhizal taxa. Warming alone and in combination with nitrogen addition also reduced negative but increased positive fungal co-occurrence probabilities, promoting species coexistence. Negative fungal co-occurrence was positively correlated to soil carbon content, while the proportion of fungal hydrolytic enzyme encoding genes was negatively correlated with soil carbon content. This may reflect fungal life history trade-offs between competition (e.g., reduced negative co-occurrence) and resource acquisition (e.g., higher abundance of hydrolytic enzyme encoding genes) with implications for carbon storage.

Keywords: Arbuscular mycorrhizae, Climate change, Ectomycorrhizae, Fungi, Global change, Nitrogen deposition, Soil carbon storage, Soil warming

Introduction

Forest soils are an important and large carbon (C) sink (Pan et al., 2011). They harbor hyperdiverse fungal communities that act as key decomposers (Schneider et al., 2012; Peay et al., 2016) and, in turn, are regulators of forest soil C storage (Averill and Hawkes, 2016; Frey, 2019; Lindahl et al., 2021). Increasing evidence shows that fungal communities are sensitive to environmental changes, including anthropogenic nitrogen (N) deposition (Lilleskov et al., 2011; Morrison et al., 2016; Van der Linde et al., 2018; Moore et al., 2021), climate warming (Treseder et al., 2016; Fernandez et al., 2017; Morrison et al., 2019; Cao et al., 2020), nonnative invasive species (Lekberg et al., 2013; Gibbons et al., 2017; Anthony et al., 2019), and myriad other global changes (see review by Zhou et al., 2020). How fungi respond to global changes at the local level can have cascading effects throughout forests, influencing tree growth and mortality and soil carbon storage. However, and until recently, it has been technically challenging to trace how fungal community compositional changes are linked to explicit fungal functional shifts which impact processes such as soil C storage.

Ectomycorrhizal fungi (EMF) are tree symbionts that are especially important mediators of organic N acquisition from soil organic matter (SOM) and can strongly affect overall SOM chemistry (Read and Perez-Moreno, 2003; Lindahl and Tunlid, 2015). EMF relative abundances in forest soil are positively correlated with proteolytic enzyme activities (Averill and Hawkes, 2016; Morrison et al., 2019), soil C:N ratio (Anthony et al., 2017), and soil N...
immobilization (Franklin et al., 2014; Clemmensen et al.,
2015). There is also evidence for a narrow set of EMF
terrestrial fungi that drive SOM cycling within forests (Clemmensen et al., 2015; Heinonsalo et al., 2015; Lindahl et al.,
2021). For example, the relative abundance of Cortinarius acutus, an EMF species capable of producing manganese peroxi-
dases, was linked to a 33% decrease in C stocks in the
organic soil horizon of a Swedish boreal forest (Lindahl et al.,
2021). Thus, predicting how forest SOM will vary with
future global changes can be improved with concomitant
knowledge of the fungal community, including ectomycorrhizal
species.

Climate warming and atmospheric N deposition are
particularly important environmental factors that can
influence soil fungi, SOM cycling, and soil C storage.
Warming often decreases fungal biomass (DeAngelis et al.,
2015; Morrison et al., 2019) and has been shown to
shift fungal community structure (Morrison et al., 2019;
Cao et al., 2020) and the relative abundances of key fungal
taxa (Fernandez et al., 2017; Mucha et al., 2018). Warming
can also alter SOM chemistry by accelerating decomposi-
tion, resulting in lower soil C storage (Pisani et al., 2015;
Melillo et al., 2017). Simulated N deposition also reduces
fungal biomass at high N addition levels (Moore et al.,
2021), shifts fungal community structure (Lilleskov et al.,
2011; Morrison et al., 2016), and alters SOM chemistry by
inhibiting decomposition. Most previous studies have
focused on one or the other of these global change drivers,
yet these are happening simultaneously in the real world.
Because fungal communities often exhibit high degrees of
functional redundancy (Rineau and Courty, 2011; Banerjee et
al., 2016), it is also uncertain whether shifts in fungal
diversity and community composition in response to
warming and N enrichment ultimately translate into func-
tional changes which impact SOM decomposition.

In this study, we measured how the diversity and com-
position of dominant fungal functional groups (i.e., EMF,
arbuscular mycorrhizal fungi [AMF], saprotrophs, pathogens)
responded to soil warming and N addition, alone and in combination. We also measured fungal functional
genes using a gene capture approach which allowed us to
target specific aspects of fungal decomposition which may
be sensitive to environmental change and linked to SOM
chemistry at the molecular level. Warming at our study
site has enhanced soil respiration by 30%–50% (Contosta et al., 2011) and reduced C storage by 35% (Anthony et al.,
2020). Nitrogen additions have had ephemeral effects on
soil respiration and no significant effect on soil C storage.
Warming in combination with nitrogen has had the great-
est positive effect on soil respiration and intermediate
effects on soil C storage (unpublished data). As the primary
decomposers in this ecosystem, we expected shifts in the
fungal community to mirror changes in soil respiration
and C storage. Specifically, we hypothesized that warming
and warming × N would have larger effects on fungal
communities and decomposition genes compared with
N addition alone. All of the treatments have shifted SOM
chemistry to varying degrees (Pisani et al., 2015), with
a general increase in the decomposition of most SOM
compounds in the heated plots and a suppression of SOM
decomposition in the N addition plots (vandenEnden et
al., 2021). Because warming in combination with N has
shifted SOM chemistry to be more similar to the heated
compared to the N addition plots (vandenEnden et al.,
2021), we expected fungal functional gene changes in the
heating × N plots to be more similar to heated plots and
distinct from the N addition plots. Earlier studies on fungi
at this site were performed on a subset of the experimental
plots (i.e., control vs. heated only plots) during the
initial study phase (Pec et al., 2021) and in the context of
an encroaching nonnative plant (Wheeler et al., 2017;
Anthony et al., 2020), but the sensitivities of soil fungi
to long-term warming, N additions, and their combination
independent of plant invasion have not been explicitly
investigated.

Materials and methods
Site description and experimental design
The Soil Warming × Nitrogen Addition experiment at the
Harvard Forest Long-Term Ecological Research site (42°50’
N, 72’18’ W) was established in 2006 and is described in
detail in Contosta et al. (2011). It is located in a mixed
deciduous stand with Quercus rubra, Quercus velutina,
Acer rubrum, Acer pensylvanicum, Fagus grandifolia, and
Betula papyrifera canopy trees. Mean annual temperature
is 8.3°C and mean annual precipitation is 1,247 mm (Boose, 2021). The full experiment includes 3 × 3 m plots
replicated 6 times for each of 4 treatments: control,
heated, N addition, and heated × N addition. Heated plots
are elevated 5°C above ambient soil temperatures using
buried heating cables installed at 10 cm depth and spaced
20 cm apart; N addition plots receive equal monthly doses
of aqueous ammonium nitrate (0.83 g N m⁻²) for a total of
5 g N m⁻² yr⁻¹ throughout the growing season (May–
October).

Soil sampling, processing, and analyses
We collected soil samples in July 2016 from 5 of the 6
replicate plots from each treatment. Five rather than 6
replicates were sampled because 1 heated plot is no lon-
ger functional due to an electrical issue, and we chose to
have a balanced design. Within each plot, we collected
a 10 × 10 cm block from the organic horizon to the depth
of the mineral soil, followed by collection of a cylindrical
core of mineral soil (5 cm width × 10 cm depth). Two
paired organic and mineral soil samples were collected
within each plot and homogenized into composites by
depth increment (i.e., organic horizon and 0–10 cm min-
eral soil). Samples were stored on ice in the field and then
at 4°C within 12 h of sampling. Within 24 h, soil was
passed through a 4 mm sieve, and a subsample for molec-
ular analysis was placed at −80°C. Additionally, a subsam-
ple for phospholipid and neutral lipid fatty acid analyses
to measure fungal and AMF biomass, respectively, was
frozen at −20°C and then freeze dried. All remaining soil
was dried at 105°C for 48 h for analysis of soil pH (10 g
soil: 20 mL deionized water) and total soil organic C and N
using dry combustion (Perkin Elmer 2400 Series II CHN,
Waltham, MA). A summary of the edaphic responses to the
treatments can be found in Anthony et al. (2020).
**Molecular analyses of fungal composition and functional genes**

We characterized fungal diversity, community composition, the relative abundance of key functional groups, and fungal functional genes related to decomposition. Genomic DNA was extracted from 250 mg of soil (Qiagen PowerSoil Kit; Qiagen, Hilden, Germany). Total fungi, including saprotrophs, EMF, and pathogenic species were analyzed using ITS2 DNA metabarcoding with the primer pair fITS7-ITS4 (White et al., 1990; Ihrmark et al., 2012; BioProject: PRJNA522440), and AMF were analyzed using 18S DNA metabarcoding with the primer pair NS31-AML2 (Simon et al., 1992; Lee et al., 2008; PRJNA522442). A description of the polymerase chain reaction (PCR) and thermocycler conditions is found in Anthony et al. (2020). Fungal functional genes were targeted and enriched using biotinylated RNA probe capture. A set of 20,005 probes were designed by Arbor Biosciences from 2,322 fungal gene sequences (a list of the gene sequences can be found in Moore et al., 2021). Each probe was 100 bp and varied with 3× tiling density. Probes were hybridized with sheared template DNA (350 bp) on a thermocycler using the manufacturer protocol and the MyBaits kit (Arbor Biosciences, Ann Arbor, MI, USA). Hybridized DNA was isolated and amplified using PCR. Amplified libraries were prepared for sequencing using the PCR-free TruSeq Prep Kit (Illumina, San Diego, CA, USA) and were sequenced on an Illumina NextSeq 150 platform (PRJNA63326). A thorough description including additional details on the probe designs and thermocycler conditions is in Moore et al. (2021).

**Bioinformatics**

All sequences (ITS, 18S, functional genes) were passed through quality control measures, removing poor quality reads (Phred scores <2 at a 20 bp sliding window) using Trimmomatic (Bolger et al., 2014). Forward and reverse reads were then merged at a 20 bp overlap with a 5% mismatch (ITS2) and at a 10 bp overlap allowing 10% mismatch (18S) using the join_paired_ends.py function in QIME (Caporaso et al., 2010). The entire ITS2 region was then excised from flanking 5.8S and 28S regions using a custom database made from the NCBI template DNA (Simon et al., 1992; Xin v7 (Li, 2013). The database was made using the index algorithm, and forward and reverse reads were aligned using the bwa mem algorithm. We only aligned forward sequences since forward and reverse sequences did not pair well, and we removed alignments with <90% sequence similarity using SAMtools. Sequences were then sorted by genes encoding for hydrolitic enzymes (beta-glucosidase, cellobiohydrolase, and cellobiidoxygenase) or oxidative enzymes (lignin peroxidase, manganese peroxidase, laccase, and laccase-like multicopper oxidases).

**SOM chemistry**

Molecular components of SOM were analyzed in soil extracts using gas chromatography-mass spectrometry (GC-MS) with an Agilent 7890B GC with a 5977B MS with electron impact ionization. See vandenEnden et al. (2021) for a complete description of the methods. In short, we quantified n-alkanes, n-alkenoic acids, sugars, curcin, suberin, and lignin-derived (vanillyl, syringyl, and cinnamyl phenols) compounds. Compounds were identified using a library of MS spectra (Wiley registry v9), National Institute of Standards and Technology (2008), and a custom mass spectral library. Quantification was based on standards specific to each compound class and normalized by soil C content (mg g⁻¹ C).

**Statistical analyses**

All statistical analyses were conducted in R (v3.6.1; R Development Core Team, 2019) and significance was set to a P value ≤ 0.05. We used analysis of variance (ANOVA) to test the effects of warming and N additions plus their interaction on univariate response variables using the base R aov function. We used the Anova function in the car package (Fox et al., 2007) to calculate type III (sequential) ANOVA tables. Multiple comparisons were evaluated by computing estimated marginal means using the emmeans function in the lsmeans package (Lenth, 2018). All models were inspected to meet the assumptions of ANOVA based on the distribution of residuals and QQnorm plots. All molecular data sets were normalized to the lowest sequencing depth using the rarefy function in vegan (Oksanen et al., 2013). We computed species richness and diversity (Shannon-Index) using the specnumber and diversity functions in vegan, respectively. We calculated Pielou’s evenness index J as Shannon/log(richness). We analyzed differences in community composition using permutational ANOVA (PERMANOVA) and the adonis function in vegan and visualized differences by treatments using non-metric multidimensional scaling (NMDS) and the metaMDS function in vegan. OTUs contributing most to
Table 1. Permutational analysis of variance summarizing changes in fungal community composition in the organic horizon across treatments. DOI: https://doi.org/10.1525/elementa.2021.000059.t1

| Treatment | Sums of Sq. | F Value | R² | P Value |
|-----------|-------------|---------|----|---------|
| Total fungi |             |         |    |         |
| Heated     | .56         | 1.45    | .07 | 0.028   |
| Nitrogen (N) | .50       | 1.28    | .06 | 0.08    |
| Heated × N | .46         | 1.18    | .06 | 0.16    |
| AMF        |             |         |    |         |
| Heated     | .33         | 2.32    | .12 | 0.026   |
| Nitrogen   | .16         | 1.16    | .06 | 0.30    |
| Heated × N | .22         | 1.53    | .08 | 0.15    |
| EMF        |             |         |    |         |
| Heated     | .58         | 1.42    | .07 | 0.047   |
| Nitrogen   | .53         | 1.31    | .07 | 0.11    |
| Heated × N | .43         | 1.05    | .05 | 0.42    |
| Saprotrophs|             |         |    |         |
| Heated     | .61         | 1.85    | .09 | 0.01    |
| Nitrogen   | .42         | 1.25    | .06 | 0.143   |
| Heated × N | .53         | 1.6     | .08 | 0.019   |
| Pathotrophs|             |         |    |         |
| Heated     | .34         | 0.93    | .05 | 0.6     |
| Nitrogen   | .34         | 0.92    | .05 | 0.58    |
| Heated × N | .48         | 1.31    | .07 | 0.14    |

Significant differences (P ≤ 0.05) are bolded. The degrees of freedom for each factor is 1 and the total degrees of freedom for each model is 19. AMF = arbuscular mycorrhizal fungi; EMF = ectomycorrhizal fungi; Sq. = squares.

dissimilarity between control and treatment plots were identified using similarity percentage analysis and the simper function in vegan. All multivariate analyses were based on fungal relative abundances computed as Bray–Curtis dissimilarities using the vegdist function in vegan.

Fungal co-occurrence was computed using probabilistic models and the cooccur function in the cooccur package (Griffith et al., 2016). We separated the data set based on significant differences in community composition to avoid spurious negative co-occurrences due to differences in environmental conditions (Blanchet et al., 2020). We then removed any species with fewer than 2 occurrences in the data sets. We used a combinatorics approach to compute co-occurrence (Veech, 2013) by setting prob = “comb.” At the plot level, we summarized every nonrandom, positive and negative pairwise co-occurrence that was observed. We then calculated our 2 response variables, the mean probability of positive and negative species co-occurrences at the plot level in order to compare overall co-occurrence probabilities across the treatments.

We tested the correlation between fungal community characteristics and functional genes with SOM compound chemistry and soil organic C using a regression framework. We first examined overall differences in SOM compounds using PERMANOVA and NMDS as described above. We then correlated fungal predictor variables, including community composition, represented as NMDS1 and NMDS2 for total fungi, EMF, saprotrophs, and pathotrophs, richness and Shannon diversity (all groups listed above), co-occurrence (negative and positive), and the relative abundances of fungal functional genes. We used the envfit function in vegan to examine correlations between fungal characteristics and SOM composition and linear regression to test the relationship between SOM compounds and every fungal predictor variable. Separate linear regression models were then run for every combination and inspected for normality. We used the log concentration of SOM compounds where nonlinear relationships existed.

Results
Fungal biomass was not affected by the treatments (Table S1), but heating significantly altered fungal diversity and community composition while N additions only significantly affected fungal diversity. Fungal community composition in the organic horizon was distinct in ambient compared to heated plots (F_1,19 = 1.45, P = 0.03; Table 1, Figure 1), but this effect was not detected in the mineral horizon (Table S2). A similar response to heating in the organic horizon was observed for saprotrophic fungi (F_1,19 = 1.85, P = 0.01), EMF (F_1,19 = 1.42, P = 0.05), and AMF (F_1,19 = 2.32, P = 0.03; Table 1, Figure 1). In contrast to community composition, both heating and N additions reduced fungal diversity (F_1,19 = 4.97, P = 0.04) and EMF Shannon diversity (F_1,19 = 12.16, P = 0.003; Figure 2, Table S3) and community evenness, measured as Pielou’s evenness index J’ (Table S4). Saprotrophic diversity was not sensitive to the treatments. AMF diversity (F_1,19 = 7.77, P = 0.009) and community evenness (F_1,19 = 9.5, P = 0.004) were also reduced by heating but not by N additions. Overall, fungal communities in the control plots were significantly more even than the treatment plots, and evenness contributed more to differences in diversity than richness. Changes in community composition and diversity across the treatments were observed in the organic horizon and to a lesser extent or not at all in mineral soil; thus, all subsequent analyses report on communities found in the organic horizon where we see the greatest impact on soil C pools.

EMF were the dominant fungal guild (approximately 70% relative abundance; Figure S1), and there were no treatment-level differences in EMF relative abundances as a group, nor that of any other fungal guild (i.e., saprotrophs, pathotrophs; Table S5). Rather, there were particular ectomycorrhizal, and to a lesser extent, saprotrophic OTUs that drove dissimilarity in fungal community composition between treatment and control plots (Table S6). Heated plots were hyperdominated by ectomycorrhizal Russula laurocerasi, while N addition plots were hyperdominated by Russula subsulphurea compared to control.
plots (i.e., these OTUs were at an order of magnitude higher relative abundances in treatment versus control plots; **Figure 3**; Table S7). In the 2-factor heated × N plots, ectomycorrhizal *Boletus rubropunctus* had 6 times greater relative abundance compared to control plots. Representing the core sensitive taxa, ectomycorrhizal *Amanita fulva*, *Cortinarius sp. 171*, and *Cortinarius sp. 1565* and the non-mycorrhizal *Eurotiales sp. 18*, *Herpotrichiellaceae sp. 143*, and *Mortierella pulchella* were all negatively affected by the treatments.

Fungal community co-occurrence patterns were altered by heating but not by N additions. In both the heated and the heated × N addition plots, there was lower negative co-occurrence among fungal community members compared to unheated plots ($F_{1,19} = 4.4$, $P = 0.05$; **Figure 4A**). Conversely, there was higher positive co-occurrence among fungi in the heated and heated × N addition plots ($F_{1,19} = 6.9$, $P = 0.02$; **Figure 4B**). Taxa that most negatively co-occurred with other species in the heated plots included saprotrophic, EMF, and pathotrophic fungal species. There were no changes in the co-occurrence of AMF communities.

Overall fungal functional gene composition was not affected by the treatments, but the relative abundances of individual decomposition genes and groups of genes were altered by heating and N additions (Table S8). Fungal genes encoding for manganese peroxidases were at reduced proportions in the heated plots (**Figure 5A**), particularly in the single factor heated (i.e., heating only) treatment. In contrast, the total relative abundance of
hydrolytic enzyme encoding genes was higher in the heated plots (Figure 5B). Although there was no change in the relative abundance of hydrolytic enzyme encoding genes in the N addition plots, cellobiohydrolase encoding genes were at reduced proportions (Figure S2).

Finally, we examined which fungal community attributes were associated with changes in SOM compounds and soil C storage in the treatment plots. SOM chemistry was altered by heating (PERMANOVA: $F_{1,19} = 4.5, P = 0.005$) but not by N additions ($F_{1,19} = 1.3, P = 0.28$), and the heated plots were enriched in suberin- and cutin-derived compounds compared to unheated plots (Figure 6A). The relative abundance of fungal hydrolytic enzyme encoding genes was positively associated with both suberin- and cutin-derived compounds. In contrast, lignin-derived compounds, short- and long-chain aliphatic acids, n-alkanes, and total sugars were associated with the unheated plots. Neither fungal community composition nor diversity (total or that of any group) were correlated with total soil C or any particular group of SOM compounds; however, fungal negative co-occurrence was strongly and negatively correlated with suberin- and cutin-derived compounds (Figure S3). Fungal negative co-occurrence was also strongly and positively correlated with overall soil organic C contents (Figure 6B).
Discussion

Few studies have considered how soil fungi in forests will be affected by future warming conditions, atmospheric N deposition, and their combination (Rillig et al., 2019; Zhou et al., 2020). But as the main decomposers of SOM in forests (Schneider et al., 2012), fungal responses to warming and N additions may be linked to soil C storage in the face of climate change. Here, we show that fungal communities are less diverse (i.e., lower evenness), shift in composition, and have lower manganese peroxidase and higher hydrolytic enzyme production potentials in response to heating in comparison to N additions. This indicates that fungal potentials to decompose cellulolytic versus ligninolytic substrates responded in opposite directions to heating. The novel gene capture and enrichment technique we applied to soil systems made it uniquely possible for us to measure these specific fungal-mediated decomposition pathways. Changes in fungal functional genes and co-occurrence were also associated with distinct SOM compound concentrations and soil C content. Below, we discuss these results in greater detail, along with the potential implications for soil C storage.
Fungal diversity and community composition had a stronger response to heating than N additions

Fungal diversity and community composition shifted in response to heating. Nitrogen additions had a small and insignificant effect on fungal community composition and a milder but significant effect on fungal diversity. Based on changes in soil respiration and C storage in the treatment plots, we expected fungal communities to be most affected by heating and heating × N followed by N additions alone, and our results generally support this prediction. Our findings are also consistent with previously described warming effects on fungal community composition at the Prospect Hill Soil Warming experiment also located at the Harvard Forest (Morrison et al., 2019; Pec et al., 2021). Greater warming versus N addition effects on fungal community composition have also been observed in a subtropical forest (Cao et al., 2020). In Cao et al. (2020) and our study, larger heating versus N addition effects may be due to the quantity and duration of added N. Because higher N additions (15 g N m⁻² yr⁻¹) shifted fungal community composition after more than 20 years of fertilization at a neighboring experiment within the Harvard Forest (Morrison et al., 2016), insignificant trends observed here are likely due to the smaller quantity and shorter duration of N additions (Moore et al., 2021). Heating was therefore a stronger factor than N additions affecting fungal community composition in our study, but the magnitude of this effect differed across fungal functional groups and with simultaneous heating and N additions. Notably, the only significant interaction between heating and N additions was observed for saprotrophic fungal community composition which shifted most in the heated × N addition compared to control plots. Interestingly, the heated × N addition plots also have the highest soil respiration rates. One possibility is that soil respiration may be linked to changes in saprotrophic fungal community composition to a greater extent than other functional groups.

There were no treatment effects on fungal biomass or the relative abundances of fungal trophic guilds. Fungal communities from all plots were dominated by mycorrhizal fungi followed by saprotrophs. Although at the same relative abundances, mycorrhizal diversity decreased with the treatment compared to control plots. In particular, community evenness declined while species richness remained unchanged. The ectomycorrhizal Russula laurocerasi and Russula subsulphurea were at more than an order of magnitude higher relative abundance in the heated and N addition plots compared to control plots, respectively, and were thus major contributors to observed declines in fungal evenness. A dominant AMF Glomus sp. that drove observed differences in community composition between the heated and control plots (Table S10) had more than 2 times greater relative abundance in the heated (45% relative abundance) versus control plots (18%; Figure S4). These observations of overdominance are consistent with other global change studies at the Harvard Forest showing that N and manganese additions select for hyperdominance by Russula vinacea (Morrison et al., 2016) and Coccinonectria ruscii (Whalen et al., 2018), respectively. Our results lend support to the idea that soil warming and N addition encourage hyperdominance by a handful of fungal species.

The functional implications of hyperdominance in mycorrhizal fungal communities remain largely unknown. One possibility is that diversity stabilizes communities to additional environmental changes (Elton, 1958; Hillebrand et al., 2008). For example, using synthetic communities, denitrification by uneven microbial communities shifted more in response to salinity stress than denitrification by even communities (Wittebolle et al., 2009). Soil fungi in the heated plots where our work was conducted were also more sensitive to the additional pressures of a nonnative, invasive plant, Alliaria petiolata (Anthony et al., 2020). Warming-induced reductions in fungal community evenness may reduce community resistance to additional environmental changes and may have myriad additional implications for forest functioning in a changing world.

Reduced fungal—fungal negative co-occurrence reflects less competition with heating

Fungal taxa exhibited reduced negative and enhanced positive co-occurrence in the heated plots. This suggests that potential species–species interactions shifted with heating, favoring coexistence over competition. Saprotrophic fungi are widely known to compete for space and nutrients, and protagonists even actively preclude colonization by antagonists when decomposing wood (Heilmann-Clausen and Boddy, 2005; Boddy and Hiscox, 2017); but these interactions are challenging to observe and quantify at the microscopic level in soil. This is why an increasing number of studies use co-occurrence analysis to identify potential species–species interactions in DNA-based surveys (Berry and Widder, 2014; Ovaskainen et al., 2017; Liu et al., 2019). The challenge with this approach is that it is typically unclear whether co-occurrence patterns are due to species interactions (Gotelli and McCabe, 2002), stochastic drift processes (Ullrich, 2004), or shared environmental preferences (Ovaskainen et al., 2017). Because we deliberately made separate co-occurrence models for the heated and unheated plots, spurious direct effects due to differences in fungal community composition, soil C contents, and SOM composition were avoided. While we cannot fully tease these scenarios apart in our study, these co-occurrence patterns establish the possibility that fungal competition was altered by heating.

An immediate question is therefore why fungi might compete less in response to a decade of heating. One possible explanation is a C limitation hypothesis. Because soil warming progressively reduces soil C content, fungi may have become more cooperative (i.e., sharing resources) over time with warming. Simulation studies suggest that low C availability promotes microbial cooperation through the sharing and more efficient use of decomposition products (Ebrahimi et al., 2019). Experimentally, low C content can slow microbial growth (Sawada et al., 2008; Reischke et al., 2014) and select for species that produce fewer extracellular enzymes (Malik et
al., 2018), creating the conditions for lower direct and indirect competition. Saprotrophic fungi with high competitive abilities also exhibit lower tolerance to temperature and moisture changes (Maynard et al., 2019), which supports the idea that heating might select for species with lower competitive abilities. The stress gradient hypothesis also states that facilitative versus competitive interactions should increase in frequency when physical conditions are suboptimal (Maestre et al., 2009), especially under conditions of energy and nutrient limitations, as observed for soil C and most SOM constituents in the heated versus unheated plots (Figure 6A). Thus, there is a theoretical basis for C limitations to reduce fungal competition and promote facilitative interactions consistent with our observations based on co-occurrence analysis.

**Fungal changes in decomposition genes caused by heating**

Soil warming typically increases soil respiration via accelerated decomposition (Melillo et al., 2017; Romero-Olivares et al., 2017). While this positive effect of warming can diminish over time, at the Soil Warming × Nitrogen Addition Study where this work was conducted, soil respiration remained elevated compared to control plots after 10 years of continuous heating (unpublished data). As the primary decomposers in forests, changes in fungal-mediated decomposition may drive long-term soil C losses due to heating. However, it has been technically challenging to study fungal functions since currently available approaches do not capture high resolution fungal functional genes (e.g., metagenomics that mostly capture bacterial genes) or differentiate between fungal and bacterial decomposing enzymes (e.g., potential extracellular enzyme activities). Here, we used a novel gene capture technique (see Anthony et al., 2020; Moore et al., 2021) to specifically study how changes in fungal decomposition genes shifted with heating and N additions. While we expected fungal functional gene shifts to be the greatest and similar between the heated and heated × N compared to N addition only plots, we found only partial support for this with respect to manganese peroxidase and hydrolytic enzyme encoding genes.

Heating reduced fungal manganese peroxidase gene proportions while N additions had no effect. Because manganese peroxidases are often the most common lignin-decay enzymes produced by fungi (Entwistle et al., 2018), they can have large impacts on soil C storage. Earlier work at this experiment demonstrated that heating reduces lignin-derived compounds (Pisani et al., 2015). Thus, one possibility is that lower availability of ligninolytic substrates reduced fungal production of manganese peroxidases; however, there was no correlation between lignin-derived compounds and manganese peroxidase gene proportions. It is also possible that this change is due to reductions in soil manganese availability (Whalen et al., 2018). A recent meta-analysis found that soil warming in forests increases woody plant manganese concentrations by approximately 40% (Yuan et al., 2018), which could dramatically reduce soil manganese levels by immobilizing it in woody biomass. Overall, soil fungi in the heated plots have a reduced capacity to synthesize manganese peroxidases, an indicator of lower ligninolytic decomposition potential by fungi. Interestingly, long-term warming can select for bacteria more adapted to degrading lignin (Pold et al., 2015), and thus there may be a shift in the relative importance of fungi versus bacteria in lignin decomposition under warming conditions.
Conversely, heating increased the relative abundances of fungal hydrolytic enzyme encoding genes. This is consistent with a recent meta-analysis showing that warming generally increases hydrolytic enzyme activities and that this increase in hydrolytic enzymes is strongly correlated with fungal biomass (Meng et al., 2020). These enzymes act on cellulolytic substrates which may be reduced in the heated plots at our study site. Recently, vandenEnden et al. (2021) found lower O-alkyl C compounds in heated soil at our experiment using solid-state $^{13}$C NMR spectra. This is an indicator of lower cellulolytic compound content and may be linked to the increased fungal hydrolytic enzyme gene potentials identified here. Interestingly, shifts in hydrolytic enzyme gene proportions also mirror the trajectory of soil respiration rates at this experiment, consistent with our initial expectations. In particular, both respiration and hydrolytic enzyme gene proportions are elevated in the heated plots compared to control plots. In the N addition plots, neither respiration nor hydrolytic enzyme gene proportions are different from the control plots. In the heated × N addition plots, both respiration and hydrolytic enzyme potentials are elevated to an even greater extent than in the heated plots (unpublished data). Thus, shifts in fungal hydrolytic enzyme encoding genes and changes in soil respiration may be linked across the treatment plots. While we found a small decrease in cellobiohydrolase gene proportions with N additions, there was no overall change in total hydrolytic enzyme gene proportions, suggesting that this did not constrain the liberation of cellobiose from cellulose polymers (Rani Singhamia, 2011). Regional studies over gradients of atmospheric N deposition show that hydrolytic enzyme encoding genes increase (Moore et al., 2021), but we did not verify this pattern with our experimental N addition. Our study shows that warming specifically increases fungal hydrolytic enzyme encoding gene proportions and that fungi may be key mediators of increased hydrolytic enzyme activities in heated soils.

**Fungal characteristics linked to SOM compounds and carbon storage**

The fate of soil C in a warmer world has important feedbacks to climate change. Since fungi are the dominant decomposers in forest ecosystems (Schneider et al., 2012), it was surprising that SOM compounds and C content were not strongly correlated with fungal diversity, community composition, nor that of any individual functional guild. However, fungal hydrolytic enzyme gene proportions were negatively correlated with soil organic C content and positively correlated with cutin- and suberin-derived compounds. Higher proportions of fungal hydrolytic enzyme encoding genes in the heated plots may indicate greater hydrolysis of SOM with implications for soil C storage. Thus, one possibility is that fungal functional shifts, more so than compositional changes, help to explain reduced soil C storage in chronically heated soil.

Striking above and beyond fungal functional gene changes was the positive correlation between fungal negative co-occurrence and soil C content. As discussed above, negative co-occurrence may be a reflection of fungal–fungal competition which was reduced in the heated plots. Increasing evidence suggests that specific forms of microbial competition may trade-off with other functions such as growth and the acquisition of particular resources (Maynard et al., 2017; Malik et al., 2019; Maynard et al., 2019). The production of antimicrobial toxins by fungi can constrain other metabolic functions (Čzárán et al., 2002). For example, EMF species that are highly competitive for space on tree roots (e.g., *Thelephora terrestris*) have been linked to lower hydrolytic enzyme activities, whereas less dominant species (e.g., *Suillus pungens*) are associated with higher hydrolytic enzyme activities (Moeller and Peay, 2016). Thus, the decomposition of hydrolysable SOM constituents may be constrained by competing metabolic demands associated with competition. If fungi compete less in heated soil, then they may be free to invest in other processes, such as the hydrolysis of particular SOM constituents, which can reduce soil C content. Higher hydrolytic enzyme encoding gene proportions in the heated plots lends support to this idea. A presumed exception to a competition-decomposition trade-off is indirect competition for resources liberated via oxidative decomposition.

Of course, the processes underpinning soil C loss with long-term warming are more complex than can be understood from a single sampling campaign (Melillo et al., 2017). In this study, we are likely only capturing a unique experimental period characterized by fungal functional and species coexistence shifts linked to a particular phase of soil C loss. It is also important to consider what the directionality and direct versus indirect connections may be between fungal shifts and soil carbon storage, which cannot be disentangled here. One unanswered question worth investigating in the future is whether C limitation reduces fungal competition or if soil C storage decreases when fungi compete less and decompose more. While it may be challenging to answer these questions in the field, this could be addressed experimentally. For example, using competition mesocosms involving negatively co-occurring taxa (sensu Maynard et al., 2017), it would be possible to measure decomposition in the presence or absence of competition and under manipulated C concentrations. Despite remaining uncertainties, our results suggest that fungal competition may be tightly coupled to soil C storage and sensitive to warming.

**Conclusion**

We studied how soil fungal diversity, community composition, and decomposition genes responded to 10 years of soil warming, N additions, and their combination. Our results show that soil fungal community composition shifts with warming to a greater extent than N additions, whereas both global changes reduced fungal community evenness by approximately 40%. Our results indicate that global changes can encourage hyperdominance by a handful of mycorrhizal fungal species. Fungal functional potential for the production of manganese peroxidases were reduced while hydrolytic enzyme potentials increased in the heated plots but not in the N addition plots. Warming also reduced negative fungal co-occurrence and promoted
species coexistence. Negative co-occurrence was positively correlated with soil organic C content. Through the lens of life history trade-offs, we hypothesize that warming relaxes fungal competition and promotes soil C loss via increased hydrolytic enzyme mediated decomposition. This response by fungi to warming might contribute to a decline in soil C during this phase of soil warming.

Data accessibility statement
All data and code to replicate data manipulation, statistical analyses, and figure generation are available at https://gitlab.ethz.ch/manthony/swan-fungal-responses.

Supplemental files
The supplemental files for this article can be found as follows:
- Tables S1–S10. Figures S1–S4.

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Competing interests
The authors declare no competing interests.

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Conceptualization: MAA, SDF.
Investigation: MAA, SDF.
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Writing—original draft, review, and editing: MAA, SDF, MK, JAMM, MJS.
Supervision: SDF.

References
Abarenkov, K, Nilsson, RH, Larsson, K-H, Alexander, IJ, Eberhardt, U, Erland, S, Höiland, K, Kjoller, R, Larsson, E, Pennanen, T, Sen, R, Taylor, A, Tedersoo, L, Ursing, B, Vrålstad, T, Liimatainen, K, Peinter, U, Köljalg, U. 2010. The UNITE database for molecular identification of fungi—Recent updates and future perspectives. New Phytologist 186(2): 281–285. DOI: http://dx.doi.org/10.1111/j.1469-8137.2009.03160.x.

Anthony, MA, Frey, S, Stinson, K. 2017. Fungal community homogenization, shift in dominant trophic guild, and appearance of novel taxa with biotic invasion. Ecosphere 8(9): e01951.

Anthony, MA, Stinson, KA, Moore, JAM, Frey, SD. 2020. Plant invasion impacts on fungal community structure and function depend on soil warming and nitrogen enrichment. Oecologia 194(4): 659–672. DOI: http://dx.doi.org/10.1007/s00442-020-04797-4.

Anthony, MA, Stinson, KA, Trautwig, A, Coates-Connor, E, Frey, S. 2019. Fungal communities do not recover after removing invasive Alliaria petiolata (garlic mustard). Biological Invasions 21(10): 3085–3099.

Averill, C, Hawkes, CV. 2016. Ectomycorrhizal fungi slow soil carbon cycling. Ecology Letters 19(8): 937–947.

Banerjee, S, Kirkby, CA, Schmutter, D, Bissett, A, Kirkkegaard, JA, Richardson, AE. 2016. Network analysis reveals functional redundancy and keystone taxa amongst bacterial and fungal communities during organic matter decomposition in an arable soil. Soil Biology and Biochemistry 97: 188–198. DOI: http://dx.doi.org/10.1016/j.soilbio.2016.03.017.

Bengtsson-Palme, J, Ryberg, M, Hartmann, M, Branco, S, Wang, Z, Godhe, A, De Wit, P, Sánchez-García, M, Ebersberger, I, de Sousa, F. 2013. Improved software detection and extraction of ITS1 and ITS 2 from ribosomal ITS sequences of fungi and other eukaryotes for analysis of environmental sequencing data. Methods in Ecology and Evolution 4(10): 914–919.

Berry, D, Widder, S. 2014. Deciphering microbial interactions and detecting keystone species with co-occurrence networks. Frontiers in Microbiology 5. DOI: http://dx.doi.org/10.3389/fmicb.2014.00219.

Blanchet, FG, Cazelles, K, Gravel, D. 2020. Co-occurrence is not evidence of ecological interactions. Ecology Letters 23(7): 1050–1063. DOI: http://dx.doi.org/10.1111/ele.13525.

Boddy, L, Hiscox, J. 2017. Fungal ecology: Principles and mechanisms of colonization and competition by saprotrophic fungi, in The fungal Kingdom. John Wiley & Sons, Ltd: 293–308. DOI: http://dx.doi.org/10.1128/9781555819583.ch13.

Bolger, AM, Lohse, M, Usadel, B. 2014. Trimmomatic: A flexible trimmer for Illumina sequence data. Bioinformatics 30(15): 2114–2120.

Boose, E. 2021. Fisher Meteorological Station at Harvard Forest since 2001. Harvard Forest Data Archive: HF001 (v.26). Environmental Data Initiative. Available at https://doi.org/10.6073/pasta/69e92642b512897032446cefe795fbb8.
Cao, J, Lin, T-C, Yang, Z, Zheng, Y, Xie, L, Xiong, D, Yang, Y. 2020. Warming exerts a stronger effect than nitrogen addition on the soil arbuscular mycorrhizal fungal community in a young subtropical Cunninghamia lanceolata plantation. Geoderma 367: 114273. DOI: http://dx.doi.org/10.1016/j.geoderma.2020.114273.

Caporaso, JG, Kuczynski, J, Stombaugh, J, Bittinger, K, Bushman, FD, Costello, EK, Fierer, N, Penna, AG, Goodrich, J, Gordon, JI. 2010. QIIME allows analysis of high-throughput community sequencing data. Nature Methods 7(5): 335.

Clemmensen, KE, Finlay, RD, Dahlberg, A, Stenlid, J, Wardle, DA, Lindahl, BD. 2015. Carbon sequestration is related to mycorrhizal fungal community shifts during long-term succession in boreal forests. New Phytologist 205(4): 1525–1536.

Contosta, AR, Frey, SD, Cooper, AB. 2011. Seasonal dynamics of soil respiration and N mineralization in chronically warmed and fertilized soils. Ecosphere 2(3): 1–21.

Czárán, TL, Hoekstra, RF, Pagie, L. 2002. Chemical warfare between microbes promotes biodiversity. PNAS 99(2): 786–790.

DeAngelis, KM, Pold, G, Topcuoğlu, BD, van Diepen, LT, Varney, RM, Blanchard, JL, Melillo, J, Frey, SD. 2015. Long-term forest soil warming alters microbial communities in temperate forest soils. Frontiers in Microbiology 6: 104.

Ebrahimi, A, Schwartzman, J, Cordero, OX. 2019. Cooperation and spatial self-organization determine rate and efficiency of particulate organic matter degradation in marine bacteria. PNAS 116(46): 23309–23316. DOI: http://dx.doi.org/10.1073/pnas.1908512116.

Edgar, RC. 2010. Search and clustering orders of magnitude faster than BLAST. Bioinformatics 26(19): 2460–2461.

Elton, CS. 1958. The ecology of invasions by plants and animals. London, UK: Methuen.

Entwistle, EM, Romanowicz, KJ, Argiroof, WA, Freedman, ZB, Morris, JJ, Zak, DR. 2018. Anthropogenic N deposition alters the composition of expressed class II fungal peroxidases, in Elliot, MA ed. Applied and environmental microbiology 84(9): e02816-17. DOI: http://dx.doi.org/10.1128/AEM.02816-17.

Fernandez, CW, Nguyen, NH, Stefanski, A, Han, Y, Hobbie, SE, Montgomery, RA, Reich, PB, Kennedy, PG. 2017. Ectomycorrhizal fungal response to warming is linked to poor host performance at the boreal-temperate ecotone. Global Change Biology 23(4): 1598–1609. DOI: http://dx.doi.org/10.1111/gcb.13510.

Fox, J, Friendly, GG, Graves, S, Heiberger, R, Monette, G, Nilsson, H, Riple, B, Weisberg, S, Fox, MJ, Suggests, M. 2007. The car package. Vienna, Austria: R Foundation for Statistical Computing.

Franklin, O, Nåsholm, T, Högborg, P, Högborg, MN. 2014. Forests trapped in nitrogen limitation—An ecological market perspective on ectomycorrhizal symbiosis. New Phytologist 203(2): 657–666. DOI: http://dx.doi.org/10.1111/nph.12840.

Frey, SD. 2019. Mycorrhizal fungi as mediators of soil organic matter dynamics. Annual Review of Ecology, Evolution, and Systematics 50: 237–259.

Gibbons, SM, Lekberg, Y, Mummey, DL, Sangwan, N, Ramsey, PW, Gilbert, JA. 2017. Invasive plants rapidly reshape soil properties in a grassland ecosystem. mSystems 2(2). DOI: http://dx.doi.org/10.1128/mSystems.00178-16.

Gotelli, NJ, McCabe, DJ. 2002. Species co-occurrence: A meta-analysis of J. M. Diamond’s assembly rules model. Ecology 83(8): 2091–2096. DOI: http://dx.doi.org/10.2307/372040.

Griffith, DM, Veech, JA, Marsh, CJ. 2016. Cooccur: Probabilistic species co-occurrence analysis in R. Journal of Statistical Software 69(2): 1–17.

Heilmann-Clausen, J, Boddy, L. 2005. Inhibition and stimulation effects in communities of wood decay fungi: Exudates from colonized wood influence growth by other species. Microbial Ecology 49(3): 399–406. DOI: http://dx.doi.org/10.1007/s00248-004-0240-2.

Heinsonso, J, Sun, H, Santalahli, M, Bäcklund, K, Hari, P, Pumpenan, J. 2015. Evidences on the ability of mycorrhizal genus piloderma to use organic nitrogen and deliver it to scots pine. PLos ONE 10(7): e0131561. DOI: http://dx.doi.org/10.1371/journal.pone.0131561.

Hillebrand, H, Bennett, DM, Cadotte, MW. 2008. Consequences of dominance: A review of evenness effects on local and regional ecosystem processes. Ecology 89(6): 1510–1520. DOI: http://dx.doi.org/10.1890/07-1053.1.

Ihrmark, K, Bödeker, I, Cruz-Martinez, K, Friberg, H, Kubartova, A, Schenck, J, Strid, Y, Stenlid, J, Brandström-Durling, M, Clemmensen, KE. 2012. New primers to amplify the fungal ITS2 region—evaluation by 454-sequencing of artificial and natural communities. FEMS Microbiology Ecology 82(3): 666–677.

Lee, J, Lee, S, Young, JPW. 2008. Improved PCR primers for the detection and identification of arbuscular mycorrhizal fungi. FEMS Microbiology Ecology 65(2): 339–349.

Lekberg, Y, Gibbons, SM, Rosendahl, S, Ramsey, PW. 2013. Severe plant invasions can increase mycorrhizal fungal abundance and diversity. ISME Journal 7(7): 1424–1433. DOI: http://dx.doi.org/10.1038/ismej.2013.41.

Lenth, R. 2018. Package “lsmeans.” The American Statistician 34(4): 216–221.

Li, H. 2013. Aligning sequence reads, clone sequences and assembly contigs with BWA-MEM. arXiv preprint arXiv:13033997. Available at https://arxiv.org/abs/1303.3997.

Lilleskov, E, Hobbie, EA, Horton, T. 2011. Conservation of ectomycorrhizal fungi: Exploring the linkages between functional and taxonomic responses to
anthropogenic N deposition. *Fungal Ecology* 4(2): 174–183.

Lindahl, BD, Kyschenko, J, Varenius, K, Clemmensen, KE, Dahlberg, A, Karlton, E, Stendahl, J. 2021. A group of ectomycorrhizal fungi restricts organic matter accumulation in boreal forest. *Ecology Letters* 24(7): 1341–1351. DOI: http://dx.doi.org/10.1111/ele.13746.

Lindahl, BD, Tunlid, A. 2015. Ectomycorrhizal fungi—Potential organic matter decomposers, yet not saprotrophs. *New Phytologist* 205(4): 1443–1447. DOI: http://dx.doi.org/10.1111/nph.13201.

Liu, F, Li, Z, Wang, X, Xue, C, Tang, Q, Li, RW. 2019. Microbial co-occurrence patterns and keystone species in the gut microbial community of mice in response to stress and chondroitin Sulfate disaccharide. *International Journal of Molecular Sciences* 20(9): 2130. DOI: http://dx.doi.org/10.3390/ijms20092130.

Maestre, TM, Callaway, RM, Valladares, F, Lortie, CJ. 2015. Ectomycorrhiza fungus—Li, F, Liu, F, Lin, K, Liu, F, Li, Z, Wang, X, Xue, C, Tang, Q, Li, RW. 2019. Microbial co-occurrence patterns and keystone species in the gut microbial community of mice in response to stress and chondroitin Sulfate disaccharide. *International Journal of Molecular Sciences* 20(9): 2130. DOI: http://dx.doi.org/10.3390/ijms20092130.

Maestre, TM, Callaway, RM, Valladares, F, Lortie, CJ. 2015. Ectomycorrhiza fungus—Li, F, Liu, F, Lin, K, Liu, F, Li, Z, Wang, X, Xue, C, Tang, Q, Li, RW. 2019. Microbial co-occurrence patterns and keystone species in the gut microbial community of mice in response to stress and chondroitin Sulfate disaccharide. *International Journal of Molecular Sciences* 20(9): 2130. DOI: http://dx.doi.org/10.3390/ijms20092130.

Malik, AA, Puissant, J, Buckeridge, KM, Goodall, T, Jehmlich, N, Chowdhury, S, Gweon, HS, Peyton, JM, Mason, KE, van Agtmaal, M, Blaud, A, Clark, I, Whitaker, J, Pywell, R, Oksanen, J, Blanchet, FG, Duan, L, Dunson, D, Roslin, T, Abreu, G. 2018. Package “vegan.” *Community Ecology Package, Version 2.2.2019.01.025.

Maynard, DS, Bradford, MA, Covey, KR, Lindner, D, Glaeser, J, talbert, DA, Tinker, PJ, Walker, DM, Crowther, TW. 2019. Competitive trait expressed in fungal trait expression across broad spatial scales. *Nature Microbiology* 4(5): 846–853. DOI: http://dx.doi.org/10.1038/s41556-019-0361-5.

Maynard, DS, Crowther, TW, Bradford, MA. 2017. Competitive network determines the direction of the diversity–function relationship. *PNAS* 114(33): 11464–11469. DOI: http://dx.doi.org/10.1073/pnas.1712211114.

Melillo, JM, Frey, SD, DeAngelis, KM, Werner, WJ, Bernard, MJ, Bowles, FP, Pold, G, Knorr, MA, Grandy, AS. 2017. Long-term pattern and magnitude of soil carbon feedback to the climate system in a warming world. *Science* 358(6359): 101–105. DOI: http://dx.doi.org/10.1126/science.aan2874.

Meng, C, Tian, D, Zeng, H, Li, Z, Chen, HY, Niu, S. 2020. Global meta-analysis on the responses of soil extra-cellular enzyme activities to warming. *Science of the Total Environment* 705(135992). DOI: https://doi.org/10.1016/j.scitotenv.2019.135992.

Moeller, HV, Peay, KG. 2016. Competition-function trade-offs in ectomycorrhizal fungi. *PeerJ* 4: e2270. DOI: http://dx.doi.org/10.7717/peerj.2270.

Moore, JAM, Anthony, MA, Pec, GJ, Trocha, LK, Trzebny, A, Geyer, KM, van Diepen, LTA, Frey, SD. 2021. Fungal community structure and function shifts with atmospheric nitrogen deposition. *Global Change Biology* 27(7): 1349–1364. DOI: http://dx.doi.org/10.1111/gcb.15444.

Morrison, EW, Frey, SD, Sadowsky, JJ, van Diepen, LT, Thomas, WK, Pringle, A. 2016. Chronic nitrogen additions fundamentally restructure the soil fungal community in a temperate forest. *Fungal Ecology* 23: 48–57.

Morrison, EW, Pringle, A, van Diepen, LTA, Grandy, AS, Melillo, JM, Frey, SD. 2019. Warming alters fungal communities and litter chemistry with implications for soil carbon stocks. *Soil Biology and Biochemistry* 132: 120–130. DOI: http://dx.doi.org/10.1016/j.soilbio.2019.02.005.

Mucha, J, Peay, KG, Smith, DP, Reich, PB, Stefański, A, Hobbie, SE. 2018. Effect of simulated climate warming on the ectomycorrhizal fungal community of boreal and temperate host species growing near their shared ecotonal range limits. *Microbial Ecology* 75(2): 348–363. DOI: http://dx.doi.org/10.1007/s00248-017-1044-5.

National Institute of Standards and Technology. 2008. Mass spectral library (NIST/EPA/NIH). Available at https://chemdata.nist.gov/dokuwiki/doku.php?id=chemdata:downloads:start.

Nguyen, NH, Song, Z, Bates, ST, Branco, S, Tedersoo, L, Menke, J, Schilling, JS, Kennedy, PG. 2016. FUN-Guild: An open annotation tool for parsing fungal community datasets by ecological guild. *Fungal Ecology* 20: 241–248.

Oksanen, J, Blanchet, FG, Kindt, R, Legendre, P, Minchin, PR, O’hara, R, Simpson, GL, Solymos, P, Stevens, MHH, Wagner, H. 2013. Package “vegan.” *Community Ecology Package, Version 2.2.2019.01.025.

Ópik, M, Vanatoa, A, Vanatoa, E, Moora, M, Davidson, J, Kalwij, JM, Reier, Ú, Zobel, M. 2010. The online database MaarJAM reveals global and ecosystemic distribution patterns in arbuscular mycorrhizal fungi (Glomeromycota). *New Phytologist* 188(1): 223–241. DOI: http://dx.doi.org/10.1111/j.1469-8137.2010.03334.x.

Ovaskainen, O, Tikhonov, G, Norberg, A, Guillaume Blanchet, F, Duan, L, Dunson, D, Roslin, T, Abrego, N. 2017. How to make more out of community data? A conceptual framework and its implementation as models and software. *Ecology Letters* 20(5): 561–576. DOI: http://dx.doi.org/10.1111/ele.12757.

Pan, Y, Birdsey, RA, Fang, J, Houghton, R, Kauppi, PE, Kurz, WA, Phillips, OL, Shvidenko, A, Lewis, SL, Canadell, JG, Ciais, P, Jackson, RB, Pacala, SW, McGuire, A, Piao S, Rautiainen, A, Sitch, S, Hayes, D. 2011. A large and persistent carbon sink...
in the world’s forests. *Science* **333**(6045): 988–993. DOI: http://dx.doi.org/10.1126/science.1201609.

**Peay, KG, Kennedy, PG, Talbot, JM.** 2016. Dimensions of biodiversity in the Earth mycobiome. *Nature Reviews Microbiology* **14**(7): 434–447. DOI: http://dx.doi.org/10.1038/nrmicro.2016.59.

**Pec, GJ, van Diepen, ITA, Knorr, M, Grandy, AS, Melillo, JM, DeAngelis, KM, Blanchard, JL, Frey, SD.** 2021. Fungal community response to long-term soil warming with potential implications for soil carbon dynamics. *Ecosphere* **12**(5): e03460. DOI: http://dx.doi.org/10.1002/ecs2.3460.

**Pisani, O, Frey, SD, Simpson, AJ, Simpson, MJ.** 2015. Soil warming and nitrogen deposition alter soil organic matter composition at the molecular-level. *Biogeochemistry* **123**(3): 391–409.

**Pold, G, Melillo, JM, DeAngelis, KM.** 2015. Two decades of warming increases diversity of a potentially lignolytic bacterial community. *Frontiers Microbiology* **6**(480). DOI: http://dx.doi.org/10.3389/fmicb.2015.00480.

**R Development Core Team.** 2019. *R: A language and environment for statistical computing*. Vienna, Austria: R Foundation for Statistical Computing.

**Rani Singhania, R.** 2011. Chapter 8—Production of cellulolytic enzymes for the hydrolysis of lignocellulosic biomass, in Pandey, A, Larroche, C, Ricke, SC, Dus, C-G, Gnansounou, E eds., *Biotechnology of Biomass, in Pandey, A, Larroche, C, Ricke, SC, Duszynski, H, Asi, E, Atkinson, B, Benham, S, Carroll, C, Cool, N, De Vos, B.* 2018. Environment and host as large-scale controls of ectomycorrhizal fungi. *Nature* **558**(7707): 243–248. DOI: http://dx.doi.org/10.1038/s41586-018-0189-9.

**Read, D, Perez-Moreno, J.** 2003. Mycorrhizas and nutrient cycling in ecosystems—A journey towards relevance? *New Phytologist* **157**(3): 475–492. DOI: http://dx.doi.org/10.1046/j.1469-8137.2003.00704.x.

**Reischke, S, Rousk, J, Báath, E.** 2014. The effects of glucose loading rates on bacterial and fungal growth in soil. *Soil Biology and Biochemistry* **70**: 88–95. DOI: http://dx.doi.org/10.1016/j.soilbio.2013.12.011.

**Rillig, MC, Ryo, M, Lehmann, A, Aguilar-Trigueros, CA, Buchert, S, Wulf, A, Iwasaki, A, Roy, J, Yang, G.** 2019. The role of multiple global change factors in driving soil functions and microbial biodiversity. *Science* **366**(6467): 886–890. DOI: http://dx.doi.org/10.1126/science.aay2832.

**Rineau, F, Court, P-E.** 2011. Secreted enzymatic activities of ectomycorrhizal fungi as a case study of functional diversity and functional redundancy. *Annals of Forest Science* **68**(1): 69–80. DOI: http://dx.doi.org/10.1007/s13595-010-0008-4.

**Romero-Olivares, AL, Allison, SD, Treseder, KK.** 2017. Soil microbes and their response to experimental warming over time: A meta-analysis of field studies. *Soil Biology and Biochemistry* **107**: 32–40. DOI: http://dx.doi.org/10.1016/j.soilbio.2016.12.026.

**Sawada, K, Funakawa, S, Kosaki, T.** 2008. Soil microorganisms have a threshold concentration of glucose to increase the ratio of respiration to assimilation. *Soil Science and Plant Nutrition* **54**(2): 216–223. DOI: http://dx.doi.org/10.1111/j.1747-0765.2007.00235.x.

**Schneider, T, Keibligner, KM, Schmid, E, Sterflinger-Gleixner, K, Ellersdorfer, G, Roschitzki, B, Richter, A, Eberl, L, Zechmeister-Boltenstern, S, Riedel, K.** 2012. Who is who in litter decomposition? Metaproteomics reveals major microbial players and their biogeochemical functions. *The ISME Journal* **6**(9): 1749–1762. DOI: http://dx.doi.org/10.1038/ismej.2012.11.

**Simon, L, Lalonde, M, Bruns, T.** 1992. Specific amplification of 18 S fungal ribosomal genes from vesicular-arbuscular endomycorrhizal fungi colonizing roots. *Applied and Environmental Microbiology* **58**(1): 291–295.

**Treseder, KK, Marusenko, Y, Romero-Olivares, AL, Maltz, MR.** 2016. Experimental warming alters potential function of the fungal community in boreal forest. *Global Change Biology* **22**(10): 3395–3404. DOI: http://dx.doi.org/10.1111/gcb.13238.

**Ulrich, W.** 2004. Species co-occurrences and neutral models: Reassessing J. M. Diamond’s assembly rules. *Oikos* **107**(3): 603–609. DOI: http://dx.doi.org/10.1111/j.0030-1299.2004.12981.x.

**Van der Linde, S, Suz, LM, Orme, CDL, Cox, F, Andreae, H, Asi, E, Atkinson, B, Benham, S, Carroll, C, Cool, N, De Vos, B.** 2018. Environment and host as large-scale controls of ectomycorrhizal fungi. *Nature* **558**(7709): 243–248. DOI: http://dx.doi.org/10.1038/s41586-018-0189-9.

**vandenEngen, L, Anthony, MA, Frey, SD, Simpson, M.** 2021. Biogeochemical evolution of soil organic matter composition after a decade of warming and nitrogen addition. *Biogeochemistry*. DOI: http://dx.doi.org/10.1007/s10533-021-00837-0.

**Veech, JA.** 2013. A probabilistic model for analysing species co-occurrence. *Global Ecology and Biogeography* **22**(2): 252–260.

**Whalen, ED, Smith, RG, Grandy, AS, Frey, SD.** 2018. Manganese limitation as a mechanism for reduced decomposition in soils under atmospheric nitrogen deposition. *Soil Biology and Biochemistry* **127**: 252–263. DOI: http://dx.doi.org/10.1016/j.soilbio.2018.09.025.

**Wheeler, J, Frey, S, Stinson, K.** 2017. Tree seedling responses to multiple environmental stresses: Interactive effects of soil warming, nitrogen fertilization, and plant invasion. *Forest Ecology and Management* **403**: 44–51.

**White, TJ, Bruns, T, Lee, S, Taylor, J.** 1990. Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. *PCR Protocols: A Guide to Methods and Applications* **18**(1): 315–322.

**Wittebolle, L, Marzorati, M, Clement, L, Balloi, A, Dafonchio, D, Heylen, K, De Vos, P, Verstraete, W, Boon, N.** 2009. Initial community evenness favours functionality under selective stress. *Nature* **458**(7238): 623–626. DOI: http://dx.doi.org/10.1038/nature07840.
Yuan, Y, Ge, L, Yang, H, Ren, W. 2018. A meta-analysis of experimental warming effects on woody plant growth and photosynthesis in forests. *Journal of Forest Research* **29**(3): 727–733. DOI: http://dx.doi.org/10.1007/s11676-017-0499-z.

Zhou, Z, Wang, C, Luo, Y. 2020. Meta-analysis of the impacts of global change factors on soil microbial diversity and functionality. *Nature Communications* **11**(1): 3072. DOI: http://dx.doi.org/10.1038/s41467-020-16881-7.