Soil fertility relates to fungal-mediated decomposition and organic matter turnover in a temperate mountain forest

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Summary

- Fungi are known to exert a significant influence over soil organic matter (SOM) turnover, however understanding of the effects of fungal community structure on SOM dynamics and its consequences for ecosystem fertility is fragmentary.
- Here we studied soil fungal guilds and SOM decomposition processes along a fertility gradient in a temperate mountain beech forest. High-throughput sequencing was used to investigate fungal communities. Carbon and nitrogen stocks, enzymatic activity and microbial respiration were measured.
- While ectomycorrhizal fungal abundance was not related to fertility, saprotrophic ascomycetes showed higher relative abundances under more fertile conditions. The activity of oxidising enzymes and respiration rates in mineral soil were related positively to fertility and saprotrophic fungi. In addition, organic layer carbon and nitrogen stocks were lower on the more fertile plots, although tree biomass and litter input were higher. Together, the results indicated a faster SOM turnover at the fertile end of the gradient.
- We suggest that there is a positive feedback mechanism between SOM turnover and fertility that is mediated by soil fungi to a significant extent. By underlining the importance of fungi for soil fertility and plant growth, these findings furthermore emphasise the dependency of carbon cycling on fungal communities below ground.

Key words: carbon cycle, ectomycorrhizal fungi, Ellenberg indicator values, enzymes, fungal guilds, Gadgil effect, plant-soil feedback, priming.

Introduction

The fertility of forest soils is a fundamental factor governing plant growth and ecosystem carbon (C) balances (Vicca et al., 2012; Binkley & Fisher, 2013). Fertility is an ecosystem characteristic describing the relative availability of below-ground resources required for plant growth (Chapin III et al., 2002), and therefore depends on soil properties such as water availability and the supply of mineral nutrients mobilised from both weathering of the geological substrate and turnover of soil organic matter (SOM) (Hansson et al., 2020). Fertile soil conditions have been associated with higher SOM decomposition rates and therefore a faster recycling of organic nutrients back to the plants (Sariyildiz & Anderson, 2003; Kyaschenko et al., 2017b, 2019). A positive feedback between fertility and SOM turnover may explain why more fertile and productive forests were shown to feature a lower soil C accumulation than less fertile and less productive forests (Vesterald & Raulund-Rasmussen, 1998; Ladegaard-Pedersen et al., 2005; Hansson et al., 2020). Recently, fungal-mediated decomposition and interactions among fungal guilds were identified to link soil fertility and SOM turnover in a boreal forest (Kyaschenko et al., 2017b). Whether this holds true for other forest ecosystems is, however, unknown.

In soils of temperate and boreal forests, saprotrophic and ectomycorrhizal fungi play a pivotal role in the transformation and decomposition of SOM (Fernandez & Kennedy, 2016; Crowther et al., 2019; Frey, 2019; Zak et al., 2019). Both fungal guilds can acquire SOM-bound resources (i.e. C and nutrients) by releasing hydrolytic and oxidative enzymes that break down organic compounds (Baldrian, 2008; Schneider et al., 2012; Kohler et al., 2015). In contrast with saprotrophic fungi, ectomycorrhizal fungi largely lack the genetic capacity to fully metabolise soil C (Baldrian, 2009; Lindahl & Tunlid, 2015; Frey, 2019; Zak et al., 2019). Instead, ectomycorrhizal fungi form symbioses with plants and receive photosynthetically assimilated C from their host in exchange for soil-derived nutrients (Read & Perez-Moreno, 2003; Van Der Heijden et al., 2015).
Suppression of saprotrophic activity by ectomycorrhizal fungi through competition for limiting nutrients (particularly nitrogen, N) has been proposed to lead to lower SOM decomposition, a mechanism referred to as the ‘Gadgil effect’ (Gadgil & Gadgil, 1971; Fernandez & Kennedy, 2016). The suppression of saprotrophic decomposition by ectomycorrhizal fungi was thereby suggested to relate to lower soil CO₂ losses, increased soil C accumulation and wider soil C:N ratios (Orwin et al., 2011; Averill & Hawkes, 2016; Sterkenburg et al., 2018; Smith & Wan, 2019). Ectomycorrhizal constraints on saprotrophic decomposition were demonstrated to be most pronounced in the litter layer and in upper soil horizons with abundant amounts of energy-rich substrate (Brzostek et al., 2015; Averill & Hawkes, 2016; Fernandez & Kennedy, 2016; Sterkenburg et al., 2018). Moreover, the occurrence of a Gadgil effect seems to be strongly determined by substrate quality; recent studies have suggested that ectomycorrhizal suppression of decomposition prevails in low quality substrates with high C:N or lignin : N ratios, respectively (Smith & Wan, 2019; Fernandez et al., 2020). These findings comply with an earlier hypothesis by Fernandez & Kennedy (2016) that ectomycorrhizal constraints on saprotrophs are more likely when soil fertility and associated N availability is low.

As a mechanism to acquire nutrients, ectomycorrhizal fungi and their host roots have also been proposed to be actively involved in SOM decomposition through the release of easily degradable carbohydrates into the rhizosphere, a process known as ‘priming’ (Phillips et al., 2012; Frey, 2019; Zak et al., 2019). Priming was suggested to stimulate saprotrophic decomposition in the litter layer as well as stronger degraded organic or mineral soil layers (Fontaine et al., 2007; Brzostek et al., 2015; Kohout et al., 2018; Fernandez et al., 2020). This mechanism is thought to increase soil CO₂ losses – by an estimated 12% in temperate forests (Bastida et al., 2019) – thereby reducing soil C stores (Huo et al., 2017; Guenet et al., 2018). The process of priming has been shown to increase with above-ground plant biomass and increased tree productivity (Hoosbeek et al., 2004; Huo et al., 2017).

Previous studies conducted in forests have found support for both the Gadgil effect and priming, primarily by building on results derived from manipulation experiments that reduced C flow to roots, such as root trenching, tree girdling, or clear cutting (Brzostek et al., 2015; Averill & Hawkes, 2016; Kohout et al., 2018; Sterkenburg et al., 2018; Fernandez et al., 2020). Yet, the proposed mechanisms of fungal-mediated decomposition and interguild fungal interactions have rarely been used to explore soil fertility–SOM feedback (Sterkenburg et al., 2015; Kyaschenko et al., 2017b). Our mechanistic understanding of how fungi affect SOM dynamics at an ecosystem level is therefore still limited (Fernandez & Kennedy, 2016; Frey, 2019; Zak et al., 2019).

Here, we studied a natural fertility gradient in a mountain forest of European beech that is typical for the temperate montane region of Central Europe. Less fertile plots at the site are characterised by a low stand productivity, topsoils with a high organic matter accumulation, and stony subsoils, while more fertile plots featured a higher productivity, topsoils with lower organic matter accumulation and loamy subsoils (Fig. 1a). The general aim of this study was to evaluate the interrelation between fertility, SOM turnover and soil fungi in forest decomposition dynamics. Based on earlier findings (Ladegaard-Pedersen et al., 2005; Sterkenburg et al., 2015; Kyaschenko et al., 2017b, 2019) we hypothesised that: (1) SOM decomposition is positively related to fertility, resulting in faster SOM turnover under fertile conditions with higher litter input; and (2) fertility coincides with a shift in the soil fungal community composition that consequently influences SOM decomposition dynamics (Fig. 1b).

Specifically, we discuss whether fungal interguild competition for limiting nutrients (i.e. a Gadgil effect) and/or stimulation of saprotrophic activity (i.e. priming) may be potential mechanisms underlying SOM decomposition dynamics along the fertility gradient.

The fertility gradient was quantified by analysing stand, organic layer and mineral soil properties (e.g. litter input, C and N stocks). Soil was further analysed for potential enzymatic activities, CO₂ losses from microbial respiration, fungal community composition and interrelations between these variables. Enzymatic activities, microbial respiration and mean residence time of the organic layer were used as proxies for SOM decomposition. The assessment of fertility was based on vascular plant species and associated Ellenberg indicator values (Ellenberg et al., 1992).

Materials and Methods

Site description

The study was conducted in the Reichraminger Hintergebirge, a mountain range located in the Austrian Calcareous Alps (47°49′08″N, 14°23′34″E). The forest stand is dominated by European beech (Fagus sylvatica L.) with a stand age of c. 146 yr in 2015. The site is southeast exposed with a slope inclination of 35° at an elevation of 1000 to 1100 m above sea level (asl) (Supporting Information Fig. S1). The parent bedrock is limestone, the dominant soil types are Rendzic Leptosol and Chromic Cambisol. Average annual air temperature and precipitation are 7.8°C and 1645 mm, respectively (Kobler et al., 2015). Sixteen plots (c. 25 m × 25 m) were selected along a natural fertility gradient (extent c. 700 m; Fig. S1), characterised by a change in topography, soil type and depth, and hydrology. The fertility gradient followed the contour line, with less fertile plots featuring a convex topography, shallower, stony soils and more fertile plots holding deeper, loamy soils and a concave topography (Fig. 1a).

Stand inventory, vegetation survey, microclimatic measurements and litter fall collection

In May 2015, diameter at breast height and height of trees were measured at each plot; allometric functions were applied to calculate above-ground woody biomass of trees (Wutzler et al., 2008). For each plot, the identity and dominance of plant species in the herbal layer was determined at four subplots; species dominance was visually estimated and expressed in percentage soil cover. Subplots were defined as triangular areas between three beech
trees each. Plant species were assigned Ellenberg indicator values (Table S1) and cover-weighted mean indicator values were calculated per plot (Ellenberg et al., 1992). Measurements of soil temperature (5 cm depth) and moisture (0–7 cm depth) were conducted six times between May and August 2015 using a handheld thermometer and a TDR moisture meter (Spectrum Technologies, Aurora, IL, USA); plot-wise means of the six measurement campaigns were used for further analyses. In total, 20 litter traps (d: 52 cm, h: 70 cm) were installed along the gradient in September 2015. Traps were installed in close vicinity to the established plots (Fig. S1). Until October 2016, litter traps were emptied regularly and collected litter was dried (105°C) and weighed (±0.01 g); litter fall was subsequently summed up for each trap and is given in g m⁻² yr⁻¹. Depending on the location along the gradient, litter traps were grouped into three levels of fertility for further analyses (low, medium, high fertility; Fig. S1).

**Organic layer and mineral soil sampling and processing**

In June 2015, organic layer samples were taken at 12 selected plots along the gradient (Fig. S1, dotted line rectangles). Litter (OLF) and humus horizons (O H, if present) were sampled separately with a large corer (d: 19 cm). Four replicates were taken per plot. Samples were pooled per plot and horizon, and roots and stones were removed. Total mass of litter and humus samples was determined by weighing (±0.01 g); litter fall was subsequently summed up for each trap and is given in g m⁻² yr⁻¹. Depending on the location along the gradient, litter traps were grouped into three levels of fertility for further analyses (low, medium, high fertility; Fig. S1).

In August 2015, mineral soil samples were taken at each plot along the gradient (Fig. S1, solid line rectangles). Samples were taken from the mineral topsoil (0–10 cm, c. 1 l soil volume) using a shovel. Four replicates were taken per plot. Due to the high stone content (up to 80% of soil volume), sampling deeper than c. 10 cm was hardly feasible at many locations. Topsoil was, however, the largest pool for organic C and N (see Methods S1; Table S2). Soil samples were immediately sieved (2 mm) in the field. For fungal community analysis, 0.5 g of homogenised mineral soil was weighed into 1.5 ml LifeGuard Soil Preservation Solution (MO BIO, Carlsbad, CA, USA). Mineral soil for enzyme analyses was frozen at −20°C on the same day, and soil for other analyses was kept at 4°C until further processing.

To determine mineral soil bulk density, stone content and root biomass, three small soil pits (c. 15 cm × 15 cm surface area) were dug down to 10 cm mineral soil depth per plot. After the organic layer was removed, the mineral soil was sampled; the sampled soil volume was estimated by refilling the pit with quartz sand and measuring the volume sand used. In the laboratory, samples were weighed; roots and stones were picked per hand and rinsed clean. Fine roots (d<2 mm), coarse roots (d>2 mm), stones, and subsamples (c. 10 g) of sieved (2 mm) soil were dried (105°C, 24 h) and weighed (±0.0001 g). Root biomass volume (g m⁻²) and stone content (vol%) were determined using dry weight and specific densities. Fine soil dry mass (g m⁻²) and fine soil bulk density (g cm⁻³) was subsequently calculated for each plot. Coarse roots were not used for statistical analysis.

**Organic layer and mineral soil analyses**

Total C and N concentration (%) of organic layer and mineral soil samples was analysed on a 300 mg subsample using a TruSpec CHN analyser (Leco, St Joseph, MI, USA); subsamples were dried (105°C, 24 h) and ground before analysis (Pulverisette...
Carbon and N stocks of the organic layer were subsequently calculated by multiplying the total dry mass of the horizon by C and N concentrations. Inorganic C concentration of mineral soil subsamples was determined using the Scheibler method (ÖNORM L 1084, 1999). Organic soil C concentration of mineral soil layers was calculated as the difference in inorganic and total C concentrations. Mineral soil organic C and N stocks (0–10 cm; g m⁻²) were calculated from respective organic C and N concentrations multiplied by dry weight of mineral soil.

The mean residence time of the organic layer was calculated per plot (Berger et al., 2009). For that, the organic layer dry mass was divided by the average annual litter fall rate of the closest litter traps (Fig. S1). This approach assumes root litter input into the organic layer to be negligible.

Soil CO₂ efflux from microbial respiration was measured on fresh mineral soil (equivalent to c. 25 g oven-dried soil) within a few days after sampling. For respiration measurements, soil was sieved to 2 mm and filled in 200 cm³ steel cylinders at field bulk density (Reichstein et al., 2000; Schindlbacher et al., 2015). After c. 3 d of equilibration at 4°C, cylinders were placed into 21 plastic containers connected to an infrared gas analysser unit (SBA-4; PP Systems International, Amesbury, MA, USA). In brief, microbial respiration of each sample was determined as ΔCO₂ within closed containers for 6 min. Microbial respiration was determined at a standardised temperature of 15°C. Respiration rates are expressed in mg C g⁻¹ C d⁻¹, and in mg C m⁻² d⁻¹ using total dry mass of mineral soil for conversion. Details on the measurement system and protocol can be found elsewhere (Mayer et al., 2017a,b).

Potential activities of hydrolytic soil enzymes were measured fluorometrically according to Marx et al. (2001) and German et al. (2011). Briefly, 0.5 g of mineral soil was suspended in 50 ml of a 100 mM Tris buffer, pH 7.5, and homogenised for 1 min in a sonication bath (48 kHz, 50 W). Aliquots of 200 µl were pipetted under constant stirring (on a magnetic plate) into black 96-well microplates, with four technical replicates for each sample. Optimal substrate concentrations and incubation times for leucine aminopeptidase (1 mM), N-acetyl-β-d-glucosaminidase (1 mM), β-galactosidase (0.5 mM), acid phosphatase (2 mM), β-xylosidase (1 mM) and cellobiohydrolase (0.3 mM) were evaluated in advance to avoid potential substrate inhibition. Next, 50 µl of substrate (dissolved in deionised water) were added to each well and the plate shaken horizontally for 30 s for mixing. The microplates were incubated at 20°C in the dark for 120 min (acid phosphatase) or 180 min (for all other enzymes). Fluorescence was measured using a multiplate reader with an excitation of 365 nm and an emission of 450 nm, at 20 and 100 flashes (EnSpire; Perkin Elmer, Waltham, MA, USA). Standard curves were prepared in buffer solution using four standard solutions with concentrations between 10 µM and 250 µM, for substrates based on 4-methylumbelliferone (all enzymes except leucine aminopeptidase) and two standard curves with concentrations of 20 µM and 50 µM, for substrates based on 7-amino-4-methylcoumarin (leucine aminopeptidase). Corresponding sets of standard curves were prepared in soil slurry to account for quenching. To measure potential phenol oxidase activity, 3,4-dihydroxy-1-phenylalanine (DOPA) was used as substrate. Here, 900 µl of soil suspension (or 900 µl of buffer solution for blank wells) were mixed with an equivalent amount of a 10 mM DOPA solution (prepared in 100 mM TRIS buffer), shaken horizontally for 10 min and centrifuged at c. 1500 g force for 5 min. Then, 250 µl of this suspension was transferred into a clear 96-well plate with three-fold repetition. Absorption was measured immediately (time 0), and after c. 6 h of incubation in the dark (20°C), at 450 nm using a multiplate reader (as above). Potential phenol oxidase activity was calculated as the difference between absorption at time 0 and after incubation. Potential activities of hydrolytic and oxidative enzymes are expressed in mmol or mol g⁻¹ C h⁻¹ and in mmol or mol m⁻² h⁻¹ using total dry mass of mineral soil for conversion.

Soil fungal community analysis

For DNA isolation from mineral soil samples, half of the suspension in LifeGuard Soil Preservation Solution (see above) was transferred to the wells of a Bead Plate from the PowerSoil-htp 96 Well Soil DNA Isolation Kit (Mo Bio, Carlsbad, CA, USA). After centrifugation and removal of the supernatant, the combined vacuum and centrifugation protocol of the manufacturer was followed. Cell lysis was carried out in a FastPrep-96 bead beater (MP Biomedicals, Santa Ana, CA, USA) twice at 6 m s⁻¹ for 45 s with a 1 min break before the second lysis. To increase recovery of DNA from soil (Feinstein et al., 2009), new Bead Solution and Solution C1 were added to the soil pellet after the first extraction and the full extraction was repeated. Library preparation and Illumina MiSeq sequencing of fungal amplicons was conducted as described in Keiblinger et al. (2018). In brief, the fungal ITS2 region was amplified with the primer pair of ITS3Mix_NeXTf and ITS4Mix_NeXTr. Forward and reverse primers were equimolar mixes of modified versions of original primers published by White et al. (1990) as suggested by Teder-soo et al. (2015). Nextera XT adapters were attached to the 5’-end of the fungal-specific primers for subsequent indexing and high-throughput sequencing. Illumina MiSeq PE250 sequencing was performed at the NGS Unit of the Vienna Biodiversity Core Facility GmbH (Vienna, Austria). Quantification of total fungal DNA was carried out using qPCR with FungiQuant primers targeting the SSU region (Liu et al., 2012) and following the protocol described in Unterwurzacher et al. (2018) with a modified assay volume of 10 µl. The qPCR standard was prepared by mixing equal amounts of genomic DNAs from Penicillium canescens NG_p02, Trichoderma harzianum NG_p29, and Triticarium sp. gab0401. Total fungal DNA is expressed in µg DNA m⁻² using total dry mass of mineral soil for conversion. Primers for sequencing and qPCR targeted different regions (ITS2 and SSU, respectively) in the rDNA gene cluster due to different requirements for specificity and sequence variability. Both regions are present in the same copy number per genome.

Sequence data analysis followed the steps outlined in Unterwurzacher et al. (2018) and Gorfer et al. (2021). USEARCH scripts were used for chimera detection and filtering of
underrepresented sequences (<10 reads in the full dataset). VSEARCH (Rognes et al., 2016) was used for clustering and counting sequences per cluster given a 97% sequence similarity, which compensated for an artificial inflation of operational taxonomic units (OTU) numbers. Taxonomic affiliation of OTUs was carried out with the U TAX script against the UNITE database (Kõljalg et al., 2013), while manual editing of the data increased phylogenetic accuracy (Hofstetter et al., 2019). When no accurate classification at the genus level was possible, the closest taxonomic level, to which a clear affiliation was possible, was used instead. Nonfungal sequences were excluded from further analyses. Fungal OTUs were affiliated to ecological lifestyles/guilds (Deltedesco et al., 2020; Gorfer et al., 2021); the lifestyles/guilds are: ectomycorrhizal fungi, other symbiotic fungi (e.g. species with unspecific mycorrhizal lifestyle or forming arbuscular mycorrhizas), saprotrophic ascomycetes, saprotrophic basidiomycetes, other saprotrophic fungi (e.g. Mortierella, Rhizophysiales, Mucor), pathogenic fungi, and those of unknown lifestyle (Table S3). For statistical community analysis (see below), fungal OTUs were taxonomically grouped at genus level or closest higher taxonomic level (e.g. family). Ratios between relative abundances of ectomycorrhizal fungi and saprotrophic fungal guilds were calculated for each plot.

Statistical analysis

All variables were averaged per plot before analyses. To assess soil fertility continuously, vascular plant’s Ellenberg indicator values for nutrients, soil reaction (a proxy for soil pH), and moisture were used. In brief, the indicator variables were analysed by means of principal component analysis (PCA; Fig. S2). The scores of the first PCA axis, integrating the availability of soil resources to plants, were used to represent a ‘fertility index’. See Methods S2 for a detailed description of the calculated fertility index.

Variables were related to each other using linear regression models. Models were extended by a variogram correlation structure when residuals were spatially autocorrelated (Zuur et al., 2009). Differences in annual litter fall among fertility levels (Fig. S1) was tested by means of analysis of variance (ANOVA); to meet the criteria of ANOVA, data were log transformed before analysis.

Detrended correspondence analysis (DCA) was used to determine patterns among the fungal community in mineral soil (Palij & Shankar, 2016). To study the effect of fertility on the fungal community, the fertility index was correlated to the DCA scores. To explore the role of fungi in SOM decomposition in greater detail, potential soil enzyme activities, microbial respiration, and mineral soil C : N ratios were additionally correlated to DCA scores. Canonical correspondence analysis (CCA) was used to investigate how much of the total variation was explained by the individual variables. The significances of the variables were tested by means of Monte Carlo permutation tests (n = 999). DCA and CCA were based on relative abundance of 352 taxonomic groups that occurred on ≥3 plots.

Level of significance was set at P < 0.05. Statistical analysis and plotting was conducted in R (R Core Team, 2017) using packages NLME (Pinheiro et al., 2014) and VEGAN (Oksanen et al., 2016).

Results

Characteristics of the fertility gradient

The investigated fertility gradient was characterised by plant species such as Carex alba, Helleborus niger, and Daphne laureola at less fertile plots, and Mercurialis perennis and Allium ursinum at more fertile plots (Table S1). A particularly high abundance of Allium ursinum indicates a high nutrient and moisture availability. The fertility index (from this point forwards referred to as ‘fertility’) was strongly related to vascular plant’s Ellenberg indicator values for soil nutrients and moisture (Fig. S2). Above-ground woody biomass of trees and fine root biomass were positively correlated with fertility (Fig. 2a,b), being c. 2-fold and 1.5-fold higher at the more fertile plots of the gradient when compared with the less fertile plots, respectively. Similarly, litter fall increased with increasing fertility levels (Fig. 2c; F2,17 = 3.687, P = 0.047).

Fertility was negatively correlated with organic layer C and N stocks (Fig. 3a,b). No effect of fertility was observed on litter layer C : N ratios (Fig. 3c). Due to the absence O3 horizon, at some plots C : N ratios were not determined. Fertility was negatively correlated with mineral soil C and N concentrations (Fig. S3). Mineral soil C and N stocks were similar across the gradient (Fig. 3d,e). Fertility was negatively correlated with both mineral soil C : N ratios (Fig. 3f) and soil pH (Fig. S3); C : N ratios ranged between c. 15 and 16 at more fertile and less fertile plots, respectively.

Soil volumetric water content at 0–7 cm mineral soil depth was similar across the fertility gradient, and soil temperature at 5 cm depth was negatively correlated with fertility (Fig. S3).

Fertility effects on decomposition processes and fungal guilds

Fertility was correlated with our proxies for SOM decomposition in both the organic layer, and the mineral soil. The mean residence time of the organic layer was negatively correlated with fertility and varied from c. 13 yr at less fertile plots to c. 5 yr at more fertile plots, respectively (Fig. 4a). A fertility effect on SOM decomposition in the mineral soil was identified by a positive correlation with the microbial respiration rates and the potential activities of phenol oxidase and leucine aminopeptidase (Fig. 4b,c,h). No correlation was found between fertility and the five other extracellular soil enzymes (Fig. 4).

Fertility effects on proxies for SOM decomposition in mineral soil were also given indirectly by correlations with above-ground woody biomass of trees. Above-ground woody biomass of trees was positively correlated with microbial respiration and the
Fig. 2 Relationship between the fertility index and fertility levels of a beech dominated mountain forest and (a) above-ground woody biomass of trees, (b) fine root biomass, and (c) annual leaf litter fall, respectively. The fertility index is based on the first axis of a principal component analysis of Ellenberg indicator values for vascular plants at sampling plots (Supporting Information Fig. S2). Given are test statistics of linear regression models ($n = 16$). Solid and dashed lines show fitted models and 95% confidence intervals, respectively.

Fig. 3 Relationship between the fertility index of a beech dominated mountain forest and carbon (C) and nitrogen (N) stocks, and C : N ratios of the organic layer (a–c) and the mineral soil (0–10 cm) (d–f), respectively. The fertility index is based on the first axis of a principal component analysis of Ellenberg indicator values for vascular plants at sampling plots (Supporting Information Fig. S2). Given are test statistics of linear regression models ($n = 16$). Solid and dashed lines show fitted models and 95% confidence intervals, respectively.
potential activities of all tested extracellular enzymes except cellobiohydrolase (Table 1).

Fertility was positively correlated with the relative abundance of saprotrophic ascomycetes while other fungal guilds were unaffected by fertility (Fig. 5). Fertility was positively correlated with total fungal DNA (Fig. 5g) and negatively correlated with the ratio of relative abundances of ectomycorrhizal fungi to saprotrophic ascomycetes (Fig. 5h). Ratios between ectomycorrhizal fungi and other saprotrophic groups were not related to fertility (Fig. 5i).

Fig. 4 Relationship between the fertility index of a beech dominated mountain forest and (a) mean residence time (MRT) of the organic layer, (b) microbial respiration and the potential activity of (c) phenol oxidase, (d) $\beta$-glucosidase, (e) cellobiohydrolase, (f) $\beta$-xylosidase, (g) N-acetyl-glucosaminidase, (h) leucine aminopeptidase, and (i) acid phosphatase in the mineral soil (0–10 cm). The fertility index is based on the first axis of a principal component analysis of Ellenberg indicator values for vascular plants at sampling plots (Supporting Information Fig. S2). Given are test statistics of linear regression models ($n = 16$). Solid and dashed lines show fitted models and 95% confidence intervals, respectively.
Table 1 Relationship between above-ground woody biomass of trees (m$^{-2}$ ha$^{-1}$) and microbial respiration, and potential soil enzyme activities, determined for the mineral soil (0–10 cm) of a fertility gradient in a beech dominated mountain forest.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Unit</th>
<th>Slope</th>
<th>P-value</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microbial respiration</td>
<td>g C m$^{-2}$ d$^{-1}$</td>
<td>(+)</td>
<td>0.008</td>
<td>0.41</td>
</tr>
<tr>
<td>Phenol oxidase</td>
<td>mol m$^{-2}$ h$^{-1}$</td>
<td>(+)</td>
<td>0.006</td>
<td>0.43</td>
</tr>
<tr>
<td>β-Glucosidase</td>
<td>mmol m$^{-2}$ h$^{-1}$</td>
<td>(+)</td>
<td>0.035</td>
<td>0.28</td>
</tr>
<tr>
<td>Cellulohydrolase</td>
<td>mmol m$^{-2}$ h$^{-1}$</td>
<td>(+)</td>
<td>0.136</td>
<td>0.15</td>
</tr>
<tr>
<td>β-Xylosidase</td>
<td>mmol m$^{-2}$ h$^{-1}$</td>
<td>(+)</td>
<td>0.037</td>
<td>0.28</td>
</tr>
<tr>
<td>Leucine aminopeptidase</td>
<td>mmol m$^{-2}$ h$^{-1}$</td>
<td>(+)</td>
<td>0.001</td>
<td>0.56</td>
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<tr>
<td>N-Acetylg glucosaminidase</td>
<td>mmol m$^{-2}$ h$^{-1}$</td>
<td>(+)</td>
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<td>0.54</td>
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<tr>
<td>Acid phosphatase</td>
<td>mmol m$^{-2}$ h$^{-1}$</td>
<td>(+)</td>
<td>0.005</td>
<td>0.44</td>
</tr>
</tbody>
</table>

Test statistics of linear regression models are given; directions of slope coefficients are indicated ($n = 16$).

Relationship between soil fungi, microbial respiration and enzyme activities

The relative abundance of saprotrophic ascomycetes was positively related to microbial respiration rates and phenol oxidase activity (Tables 2, S5). Saprotrophic basidiomycetes were negatively related to microbial respiration and phenol oxidase activity (Tables S4, S5). Other fungal guilds were neither related to microbial respiration nor to enzyme activities on an area-based unit (Table S4), however a negative correlation between these variables and the guild of other symbiotic fungi was found when units were expressed per g soil C (Table S5). Total fungal DNA was positively related to microbial respiration and all measured enzymes (Table 2).

Fungal community composition

The fungal community in the mineral soil was dominated by saprotrophic ascomycetes from the taxonomic groups of Hyaloscyphaceae, Tetracladium and Trichoderma, pathogenic fungi from Neonectria, Dactylonectria and Ilyonectria, ectomycorrhizal fungi from Inocybe, and Sebacina and other symbiotic fungi from the unspecific mycorrhizal group of Sebacinaeae, respectively (Fig. 6; Table S3). Other saprotrophic fungi were dominated by Mortierella. Under more fertile conditions, saprotrophic ascomycetes from the genus Tetracladium, and pathogenic fungi from the genus Dactylonectria were more abundant. Under less fertile conditions, saprotrophic basidiomycetes from the genera Geminibasidium and Saitozyma were more abundant (Fig. 6).

The first DCA axis identified a correlation between the fungal community composition and fertility and associated changes in potential enzyme activity and microbial respiration (Fig. 6); the second DCA axis revealed a correlation with mineral soil C : N ratios. Individual CCA analysis revealed fertility to explain 13.2% of the variance among the fungal community composition (total inertia was 1.026). Phenol oxidase, cellulohydrolase, microbial respiration and soil C : N ratios were significant and explained 12.1, 9.1, 9.3 and 10.6% of the variance, respectively; other enzymes were not significant and are therefore not displayed. A stepwise backward variable selection (starting with all

Fig. 5 Relationship between fertility index of a beech dominated mountain forest and the relative abundance of (a) ectomycorrhizal fungi, (b) other symbiotic fungi, (c) saprotrophic ascomycetes, (d) saprotrophic basidiomycetes, (e) other saprotrophic fungi (e.g. moulds), (f) pathogenic fungi, (g) the total amount of fungal DNA, and (h) the ratio between relative abundance of ectomycorrhizal fungi (EMF) to saprotrophic (SAP) ascomycetes in the mineral soil (0–10 cm). The fertility index is based on the first axis of a principal component analysis of Ellenberg indicator values for vascular plants at sampling plots (Supporting Information Fig. S2). Given are test statistics of linear regression models ($n = 16$). Solid and dashed lines show fitted models and 95% confidence intervals, respectively.
variables) revealed that fertility and soil C:N ratios together explained 21.5% of the variance.

Discussion

Exploring feedback mechanisms between fertility and SOM turnover is critical to a better understanding of C and nutrient cycling in forest ecosystems. Here, we showed that proxies for SOM decomposition related to a shift in the soil fungal community composition along a fertility gradient in a temperate mountain forest of European beech.

Our first hypothesis – SOM decomposition will be positively related to fertility (Fig. 1b) – was strongly supported by our results: more fertile plots featured a shorter residence time of the

Table 2 Relationship between relative abundance of saprotrophic ascomycetes (%) and total fungal DNA in soil (µg m⁻²) and microbial respiration, and potential enzyme activities, determined for the mineral soil (0–10 cm) of a fertility gradient in a mountain beech forest.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Unit</th>
<th>Saprotrophic ascomycetes</th>
<th>Total fungal DNA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Slope</td>
<td>P-value</td>
</tr>
<tr>
<td>Microbial respiration</td>
<td>g C m⁻² d⁻¹</td>
<td>(+)</td>
<td>0.041</td>
</tr>
<tr>
<td>Phenol oxidase</td>
<td>mol m⁻² h⁻¹</td>
<td>(+)</td>
<td>0.002</td>
</tr>
<tr>
<td>β-Glucosidase</td>
<td>mmol m⁻² h⁻¹</td>
<td>(+)</td>
<td>0.052</td>
</tr>
<tr>
<td>Cellulbiohydrolase</td>
<td>mmol m⁻² h⁻¹</td>
<td>(+)</td>
<td>0.064</td>
</tr>
<tr>
<td>β-Xylosidase</td>
<td>mmol m⁻² h⁻¹</td>
<td>(+)</td>
<td>0.054</td>
</tr>
<tr>
<td>Leucine aminopeptidase</td>
<td>mmol m⁻² h⁻¹</td>
<td>(+)</td>
<td>0.124</td>
</tr>
<tr>
<td>N-Acetyl-glucosaminidase</td>
<td>mmol m⁻² h⁻¹</td>
<td>(+)</td>
<td>0.458</td>
</tr>
<tr>
<td>Acid phosphatase</td>
<td>mmol m⁻² h⁻¹</td>
<td>(+)</td>
<td>0.266</td>
</tr>
</tbody>
</table>

Given are test statistics of linear regression models; directions of slope coefficients are indicated (n = 16). Information on correlations with other fungal guilds is given in Supporting Information Table S4.

Fig. 6 Patterns of variation in soil fungal community in the mineral soil (0–10 cm) across a fertility gradient in a beech dominated mountain forest as explained by detrended correspondence analysis (DCA). The DCA is based on 16 pooled sampling plots and 352 fungal taxonomic groups present at ≥3 sampling plots. Taxonomic grouping occurred at genus level or closest taxonomic level. Colour-coded fungal lifestyles/guilds were assigned; symbol size gives an indication for relative abundance. To increase readability, not all groups are labelled. Vectors are significant mineral soil variables related to fungal community patterns (POX, phenol oxidase; LEU, leucine aminopeptidase). Fertility is an index based on the first axis of a principal component analysis of Ellenberg’s indicator values for vascular plants at sampling plots (Supporting Information Fig. S2). Details on fungal taxonomic groups, relative abundances and associated lifestyles/guilds are given in Table S3.
organic layer and higher rates of microbial respiration, potential phenol oxidase and leucine aminopeptidase activities in the mineral soil (Fig. 4). The potential activity of most enzymes was also positively related to above-ground woody biomass of trees, indicating an indirect fertility effect via productivity (Table 1). This pattern was mirrored in lower organic layer C and N stocks (Fig. 3a,b) and lower mineral soil C and N concentrations at more fertile plots (Fig. S3). As stone content and fine soil bulk density showed opposing trends along the fertility gradient (Fig. S3), C and N stocks of the mineral soil (0–10 cm) were, however, similar across the gradient (Fig. 3d,e). Nevertheless, as tree biomass stocks and annual litter fall increased along with fertility (Fig. 2), the results supported a hypothesised, faster SOM turnover under more fertile conditions (Fig. 1b). Such a positive relationship between fertility and SOM decomposition is in line with previous studies from temperate and boreal forests (Ladegaard-Pedersen et al., 2005; Kyaschenko et al., 2017b, 2019). In addition to a better water supply and an inherently better mineral nutrient supply (e.g. potassium, magnesium) from the loamy subsoils (Figs 1a, S2), a faster SOM turnover and a related nutrient mobilisation point towards a positive feedback on ecosystem fertility, expressed in a higher above-ground woody biomass of trees and a modified composition of the herbal layer. We acknowledge that we could not confidently assess fertility effects on total soil C storage, as subsoils were only analysed at selected locations (Table S2). However, it is reasonable to speculate that loamy subsoils at more fertile plots store considerable amounts of C, which may additionally be stabilised by aggregate formation or association with mineral surfaces (Schmidt et al., 2011; Lehmann & Kleber, 2015).

Our second hypothesis – fertility coincides with a shift in the soil fungal community composition that consequently influences SOM decomposition (Fig. 1b) – was also supported by our results: fungal community structure and total fungal DNA (as proxy for fungal biomass) changed along the gradient, with saprotrophic ascomycetes being relatively more abundant at more fertile plots (Figs 5, 6). Moreover, the results indicate a strong control of total fungal DNA and saprotrophic ascomycetes on microbial respiration and potential phenol oxidase activity (Table 2), suggesting ascomycetes to be a prominent regulator of SOM decomposition in the studied mountain forest. Although saprotrophic ascomycetes are known to be less efficient decomposers than basidiomycetes, they can produce hydrolysing and oxidising enzymes that degrade SOM (Baldrain, 2006; Grinhut et al., 2007; Eichlerová et al., 2015; Kohlert et al., 2015). In particular fungi from the groups of Tetracladium and Hyaloscyphaceae (higher abundance at more fertile plots; Fig. 6) were reported to have considerable decay capacities with respect to organic substrates (Abdel-Raheem, 1997; Boberg et al., 2011; Anderson & Marvanová, 2020). By contrast, previous gradient studies in boreal pine and spruce forests reported a lower ascomycetes abundance and a higher basidiomycetes abundance under more fertile conditions; basidiomycetes were therefore suggested to be the principal regulator of SOM decomposition in these forest ecosystems (Sterkenburg et al., 2015; Kyaschenko et al., 2017b). In the mountain beech forest of this study, the abundance of saprotrophic basidiomycetes was, except for a few taxa (e.g. Geminibasidium), largely independent of fertility (Figs 5d, 6).

Ratios between relative abundance of ectomycorrhizal fungi and saprotrophic ascomycetes were higher under less fertile conditions (Fig. 5b), suggesting a greater competition pressure of ectomycorrhizal fungi on this saprotrophic guild. A negative correlation between fertility and mineral soil C:N ratios (Fig. 3) might also point towards a higher ectomycorrhizal N foraging in mineral soils at the less fertile end of the gradient (Averill et al., 2014; Fernandez & Kennedy, 2016; Smith & Wan, 2019). Particularly Tomentella and Clavulina were related to wider mineral soil C:N ratios (Fig. 6). In a recent study, fungi from the genus Tomentella were suggested to be strongly involved in the suppression of saprotrophic decomposition thereby being key for driving the ‘Gadgil effect’ (Fernandez et al., 2020). However, neither microbial respiration nor potential phenol oxidase activity were related to relative ectomycorrhizal abundance or its ratios to saprotrophic fungi (Tables S4, S5). An ectomycorrhizal suppression of SOM decomposition under less fertile conditions could therefore not clearly be identified. This contradicts previous indications for a ‘Gadgil effect’ along a boreal forest gradient (Kyaschenko et al., 2017b). Slow SOM turnover rates in boreal coniferous forest can result in the build-up of thick organic layers and the establishment of perennial fungal communities with a high potential for interguild competition (Clemmensen et al., 2015; Kyaschenko et al., 2017a). Temperate forests have a generally faster SOM turnover compared with boreal forests (Wang et al., 2018). Moreover, ectomycorrhizal constraints on decomposition were shown to be present only when substrate quality is low (Smith & Wan, 2019; Fernandez et al., 2020). It is therefore likely that the studied temperate beech forest features less (intense) competitive interaction between fungal guilds, potentially related to a more ephemeral fungal community and/or a higher substrate quality compared with boreal coniferous forests. However, we cannot rule out that saprotrophic decomposition was suppressed by ectomycorrhizal fungi also at the studied forest site.

Priming might be a process underlying higher SOM decomposition under more fertile conditions. Rhizodeposition can account for up to a third of the photosynthetically fixed C and depends on stand productivity and root biomass (Jones et al., 2009; Phillips et al., 2011; Liese et al., 2018). Moreover, priming is positively affected by above-ground biomass and tree productivity (Hoosbeek et al., 2004; Huo et al., 2017). While we did not measure rhizodeposition, it is reasonable to speculate that the greater fine root biomass at more fertile plots is indicative of more labile C being released to the rhizosphere, thereby potentially stimulating saprotrophic decomposition. This assumption is backed up by earlier findings, where tree girdling, root trenching and clearcutting in temperate oak-beech, oak, and spruce forests decreased litter decomposition and soil enzymatic activity, indicating that rhizosphere C input primes SOM decomposition (Brzostek et al., 2015; Kohout et al., 2018; Fernandez et al., 2020). In addition to priming by the input of labile C exudates from roots and ectomycorrhizas, decomposition might also be stimulated by the input of dead fine roots and fungal mycelium (Brabcová et al., 2016; Fernandez & Kennedy, 2016; Frey, 2019).
A positive relationship between fertility and SOM decomposition may also be linked to other abiotic and biotic factors. Soil temperature and moisture, for example, are known to strongly control microbial activity (Moyano et al., 2012; Mayer et al., 2017b). Here, fertile conditions were associated with greater soil water availability, as assessed by Ellenberg indicator values (Fig. S2); measured volumetric water content in the topsoil was, however, constant along the fertility gradient (Fig. S3). Soil temperature was even slightly lower at more fertile plots (Fig. S3), potentially due to greater shading by canopies. This would suggest microclimatic differences in the upper soil horizons to be of minor importance for SOM decomposition along the fertility gradient. An enhanced water supply to trees, however, might again result in a positive feedback on SOM decomposition via increased productivity and a related priming effect (Hoosbeek et al., 2004; Huo et al., 2017). Other microbial and faunal groups (e.g. bacteria, earthworms) and potential interactions with soil fungi may also be of great relevance for SOM decomposition (Baldrain, 2017). Particularly earthworms could play a crucial role in litter decay at fertile plots with loamy subsoils (Barthod et al., 2020). Moreover, high quality litter from geophytes (e.g. Allium ursinum; Table S1) (Jandl et al., 1997; Prescott, 2010) and higher N mineralisation rates at fertile plots could be important factors contributing to a positive fertility feedback (Norris et al., 2013).

Taken together, our results indicated a strong interdependency between soil fertility, SOM decomposition and soil fungal community composition in the studied mountain forest ecosystem. The results furthermore show a fertility–SOM decomposition feedback mechanism, which is mediated by soil fungi to a significant extent. We did not find a clear indication that ectomycorrhizal fungi constrained saprotrophs along the gradient, but we suggest that SOM decomposition at more fertile plots was stimulated by rhizosphere priming. Moreover, we suggest that a higher SOM decomposition and associated nutrient mobilisation under fertile conditions result in a positive feedback on tree growth. As our conclusions are based on correlation analyses, manipulation experiments (e.g. tree girdling) and/or additional in-depth measurements (e.g. on rhizodeposition, interaction of fungi with other soil organisms) are needed to further improve our mechanistic understanding of the important yet complex influence of soil fungal interactions on the carbon and nutrient cycle.

Acknowledgements

This study was partially funded by the projects ‘C-Alp’ and ‘C-Alp II’ (Austrian Academy of Sciences, ÖAW – Research initiative ‘Earth System Sciences (ESS)’), granted to DLG, and by the Austrian Science Fund (FWF), project number P29087-B25, granted to TWB. MM has received further support from an Erwin Schrödinger Fellowship from the FWF (project number J-4369). The research site was provided by the Austrian Federal Forests (ÖBi AG); we particularly thank Monika Kanzian and Georg Lettner. We thank Petra Berger, Linda Fleck, Viktoria Gapp, Daniel Gröticke, Marcel Hirsch, Hannah Hochsattel, Johannes Kobler, Frauke Neumann, Maria von Rochow, Helmut Schume, and Tiina Strand for help with field work and/or laboratory analysis, and Elmar Pröll, Kalkalpen National Park for logistical assistance. Finally, we acknowledge the support of Melitta/Toppits®. Detailed comments by three reviewers and the editor helped to improve an earlier version of the manuscript.

Author contributions

MM, BR, BM, HS, KK, TWB and DLG designed the study. MM, BR, BM, HS, MG and TWB conducted field work and laboratory analyses. CR performed enzymatic analysis. MG, CT and HB performed fungal community analyses. MM wrote the first draft of the manuscript and all authors contributed to later versions of the manuscript and to data interpretation.

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Data availability

All data used for the analyses of this study are publicly accessible. Sequencing and associated data have been deposited at NCBI BioProject PRJNA521677, BioSamples SAMN12582230–SAMN12582341 and GenBank accession numbers MK626959–MK627467.

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

Fig. S1 Layout of the 16 plots along the fertility gradient.

Fig. S2 Principal component analysis (PCA) analysing Ellenberg indicator values of vascular plants of the ground vegetation layer.

Fig. S3 Relationship between fertility index and mineral soil organic C concentration, N concentration, inorganic C concentration, pH, bulk density, stone content, moisture and temperature, respectively.

Fig. S4 Relationship between fertility index and selected fungal guild ratios.

Methods S1 Determination of total mineral soil organic carbon and nitrogen stocks.

Methods S2 Calculation of integrative fertility index based on Ellenberg indicator values.

Table S1 List of vascular plant species in the ground vegetation layer and their Ellenberg indicator values.

Table S2 Total mineral soil organic C and N stocks.

Table S3 List of fungal taxonomic groups including abundance and lifestyle/guild.
Table S4 Relationship between relative abundance of selected soil fungal guilds, guild ratios, and microbial respiration, and potential enzyme activities in mineral soil.

Table S5 Relationship between relative abundance of soil fungal guilds and microbial respiration, and potential enzyme activities per g soil C.

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