

1 **Title**

2 Temperate forests dominated by arbuscular or ectomycorrhizal fungi are characterized by strong shifts
3 from saprotrophic to mycorrhizal fungi with increasing soil depth

4

5 **Authors**

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16 **Abstract**

17 In temperate and boreal forests, competition for soil resources between free-living saprotrophs and
18 ectomycorrhizal (EcM) fungi has been suggested to restrict saprotrophic fungal dominance to the most
19 superficial organic soil horizons in forests dominated by EcM trees. By contrast, lower niche overlap
20 with arbuscular mycorrhizal (AM) fungi could allow fungal saprotrophs to maintain this dominance
21 into deeper soil horizons in AM-dominated forests.

22 Here we used a natural gradient of adjacent forest patches that were dominated by either AM or EcM
23 trees, or a mixture of both to determine how fungal communities characterized with high-throughput
24 amplicon sequencing change across organic and mineral soil horizons.

25 We found a general shift from saprotrophic to mycorrhizal fungal dominance with increasing soil depth
26 in all forest mycorrhizal types, especially in organic horizons. Vertical changes in soil chemistry,
27 including pH, organic matter, exchangeable cations, and extractable phosphorus, coincided with shifts
28 in fungal community composition.

29 Although fungal communities and soil chemistry differed among adjacent forest mycorrhizal types,
30 variations were stronger within a given soil profile, pointing to the importance of considering horizons
31 when characterizing soil fungal communities. Our results also suggest that in temperate forests, vertical
32 shifts from saprotrophic to mycorrhizal fungi within organic and mineral horizons occur similarly in
33 both ectomycorrhizal and arbuscular mycorrhizal forests.

34

35 Keywords: Fungal guilds; Soil physico-chemistry; Podzolic soil; Vertical segregation; *Acer*
36 *saccharum*; *Fagus grandifolia*.

37 **1. Introduction**

38 Soil fungi drive the biogeochemical cycling of carbon (C) and nutrients in terrestrial ecosystems. Free-
39 living saprotrophic fungi are major decomposers of soil organic matter, but mycorrhizal fungi also play
40 an important role [1–3]. In northern temperate forests, there are two major types of root-associated
41 fungi: arbuscular mycorrhizal (AM) and ectomycorrhizal (EcM) fungi [4, 5]. Mycorrhizal fungi acquire
42 C via plant hosts and many EcM fungi possess the enzymatic capacity to directly degrade organic
43 matter, potentially competing with free-living saprotrophs for organic nutrients such as nitrogen (N),
44 which promote soil C accumulation [6–8]. By contrast, AM fungi have limited degrading abilities and
45 therefore might compete less strongly with saprotrophic fungi for nutrients [9–11]. Such interactions
46 among saprotrophic and mycorrhizal fungi could have far-reaching implications for the C cycle,
47 especially in northern forests where a large fraction of global soil C is stored [3, 12, 13]. In particular,
48 it has been suggested that these interactions might help to explain differences in the amount and
49 vertical distributions of soil C between ectomycorrhizal- and arbuscular mycorrhizal-dominated forests
50 [7, 14, 15].

51

52 A first step towards understanding of interactions among saprotrophic and mycorrhizal fungi and their
53 functional consequences is to identify their co-occurrence patterns in soils [e.g. 16]. Different groups of
54 fungi can compete with each other for soil resources because of overlapping niches [7, 16–18]. In
55 particular, fungal types and taxa differ in their vertical distribution, especially in well-stratified soil
56 [19–21]. In EcM-dominated ecosystems such as boreal forests, strong vertical segregation of fungal
57 guilds occurs in the soil profile, where the litter layer is dominated by saprotrophic fungi and in older
58 and deeper layers are increasingly dominated by EcM fungi [21–23]. However, it remains unclear
59 whether this spatial separation reflects niche differentiation or competitive exclusion of saprotrophic
60 fungi by EcM fungi [7, 17]. Competitive interactions for nutrients among these fungal groups could
61 promote organic matter accumulation [24–26]. In AM-dominated forests, interactions and distribution
62 patterns may be different because AM fungi might not compete as strongly with saprotrophic fungi
63 than EcM fungi. However, studies of fungal vertical distribution in AM-dominated ecosystems have
64 largely focused on grasslands and crop systems [27–29] but not forests. To better understand the
65 impacts of global and land use changes on forest functioning, there is a crucial need to take different

66 mycorrhizal types fungi into consideration simultaneously [6, 7, 30], especially the AM strategy given
67 its importance in temperate forests [10].

68

69 A general hypothesis on vertical segregation among mycorrhizal types suggests that, when they co-
70 occur, EcM fungi and ericoid mycorrhizal (ErM) will dominate organic horizons while AM fungi will
71 predominantly occupy mineral horizons or soils [31, 32]. This view is supported by studies based on: i)
72 root colonization patterns in environments where mycorrhizal types co-occur [e.g. 33], ii) root patterns
73 and isotopic measurements of plants of different mycorrhizal types [e.g. 32, 34], iii) root colonization
74 patterns in "dual mycorrhizal" plants [35–37], iv) the different nutritional benefits of fungal symbionts
75 and their enzymatic capacity [31, 32] and v) global patterns of mycorrhizal distribution [31, 38].

76 However, to our knowledge this hypothesis about vertical distribution of distinct mycorrhizal types

77 (e.g. EcM and AM) across horizons has not been supported by detailed fungal community analyses.

78 For example, mycorrhizal fungal distribution does not always follow root distribution (e.g. presence of
79 AM fungi in the litter horizon [39]), and to focus on roots or rhizosphere sampling overlooks at long
80 extraradical hyphae of mycorrhizal fungi that penetrate far from root surfaces. Few studies have
81 studied vertical distribution at spatial scales that are fine (i.e. cm) and functional (i.e. by horizons). To
82 our knowledge, the vertical distribution of soil fungi in neighboring forest stands dominated by
83 different mycorrhizal types has not been reported. Therefore, it is not clear whether EcM or AM fungi
84 show similar vertical niches [32].

85

86 The difficulties associated with identifying the microorganisms directly involved in soil
87 biogeochemical cycling such as fungal saprotrophs and mycorrhizal fungi though their extraradical
88 hyphae has been a major obstacle to understand their impacts and the importance of their interactions.
89 Specific biomarkers can be used as proxy to quantify fungal biomass in soils such as phospholipid fatty
90 acid [e.g. 40], but they are common in many fungal groups and cannot discriminate between free-living
91 saprotrophic fungi and EcM fungal lineages because EcM symbiosis has arisen independently and
92 persisted numerous times in the Basidiomycetes, Ascomycetes, and Zygomycetes [41]. Also, the
93 mycelia of some fungi does not contain ergosterol [42]. With advances in high-throughput amplicon
94 sequencing [43], we are able to identify community members and their corresponding guilds [44–46].
95 Determining the taxonomic composition of fungal communities is important because different species

96 within the same fungal guild can vary in their effects on C and nutrients cycling [e.g. 47, 48]. Using
97 such sequencing methods, fungal community composition has been found to vary markedly across
98 large spatial scales, driven by broad-scale changes in climate and soil properties [49, 50]. However, the
99 mechanisms shaping distribution of fungal community and fungal groups such as free-living and root-
100 associated at small spatial scales remain comparatively little studied, and high-throughput amplicon
101 sequencing will allow to understand their potential impact on ecosystem functioning [19, 51, 52].

102

103 To determine the vertical distribution of fungal communities and guilds among temperate forests, we
104 characterized soil fungi and chemistry in adjacent forest patches dominated by trees that form AM or
105 EcM or a mixture of both strategies. Specifically, we used the natural co-occurring distribution of *Acer*
106 *saccharum* and *Fagus grandifolia* that associates exclusively with AM and EcM fungal symbionts
107 respectively [53]. These two co-occurring tree species share similar ecological strategies that they are
108 both deciduous, shade-tolerant and can dominate the canopy in adjacent forest patches in northeastern
109 North America [54, 55]. Their natural co-occurrence patterns provide an opportunity to compare
110 vertical distribution of fungal community composition in different forest mycorrhizal types, under
111 similar environmental conditions, thus minimizing variation in other important factors such as climate,
112 parent material or topography. Using this natural experimental design, we assessed how the fungal
113 community, guilds and root colonization vary across soil horizons along an AM-EcM gradient, and
114 determined to which extent this variability was linked with changes in soil chemical properties. We
115 expected the shift from saprotrophic to mycorrhizal fungi to occur deeper in AM forests compared to
116 EcM forests, and at an intermediate depth in forests containing a mixture of both strategies.

117

118 **2. Matériel and Methods**

119 *2.1 Study area*

120 The study was conducted at the University of Montréal's field station (Station de biologie des
121 Laurentides, Saint-Hippolyte, Québec, Canada). The field station is representative of temperate forests
122 of the Lower Laurentians and the Canadian Shield. The soil has a sandy loam texture derived from
123 well-drained rocky glacial till on a bedrock of Precambrian anorthosite [56, 57]. The soils are ferro-
124 humic and gleyed humo-ferric podzols with moder humus forming the forest floor [57–59]. The mean
125 annual temperature is 4.3°C and total annual precipitation is 1195 mm, with ~25% falling as snow

126 (based on 1981–2010 data, meteorological station #7037310, Saint-Hippolyte). The study area is
127 located within the sugar maple-yellow birch domain [60]. Most of the forest regrew following a major
128 fire that occurred around 1923 [61]. Mesic sites are composed mostly of a mosaic of *Acer saccharum*
129 and *Fagus grandifolia*, with *Betula alleghaniensis*, *Populus grandidentata* and *Acer rubrum* also
130 common [57]. The understory comprised various small tree species (e.g. *Acer pensylvanicum*) and
131 shrubs (e.g *Vaccinium* spp., *Viburnum* spp.).

132

133 2.2 Selection of forest plots

134 Plots were selected based on the dominance of different mycorrhizal tree types: AM-dominated stands
135 (>80% relative basal area by AM trees; mainly *Acer saccharum*) and EcM-dominated stands (generally
136 >80% relative basal area by EcM trees except one plot at 63%; mainly *Fagus grandifolia*), and mixed
137 stands (approximately equal basal area of AM and EcM trees, mainly *A. saccharum* maple and *F.*
138 *grandifolia*). Tree basal area was based on all trees ≥ 5 cm diameter at breast height (DBH) within a
139 plot. Plots were 20 m \times 20 m in size. We selected five blocks, each containing one plot of each
140 corresponding to one of the three mycorrhizal types (i.e. EcM, AM, mixed), for a total of 15 plots (Fig.
141 S1). Plots were selected as to minimize variation in environmental conditions (i.e. altitude, slope,
142 aspect, total basal area; Table S1) among plots within a block, and to be as close as possible from each
143 other (<400 m). For each plot, precise geographic coordinates, altitude, topographic location, slope and
144 orientation were measured (Table S1).

145

146 2.3 Soil sampling

147 Soil sampling was conducted in July and August 2015. In each plot, 10 samples were taken along two
148 oriented north-south transects (five samples per transect). Samples were collected to 20 cm depth using
149 PVC cores (7.5 cm in diameter). Samples were kept in coolers with ice and transported to the
150 laboratory to be processed within 96 hours of sampling. The PVC cores were split open to measure
151 horizon thickness then separated by: litter (L), where original structures are easily distinguishable,
152 fragmented (F), where there had been partial decomposition where structures were difficult to
153 recognize, and humus (H), comprised of highly decomposed organic matter, where original structures
154 are indistinguishable (see Fig. S2). The mineral horizons were Ae, as characterized by
155 leaching/eluviation of clay, Fe, Al or organic matter; and B, as characterized by illuviation/enrichment

156 in organic matter [62]. The 10 samples per plot were pooled by horizon. One sub-sample per horizon
157 per plot was immediately frozen for subsequent DNA extraction. Fine roots (<2 mm in diameter) were
158 set aside for mycorrhizal colonization analyses and a sub-sample of soil was air-dried for chemical
159 analyses.

160

161 *2.4 Soil analysis*

162 Air-dried soils were analyzed for pH, total carbon (C), total nitrogen (N), total phosphorus (P), organic
163 P, inorganic P and labile P. The pH was determined in 10 mM CaCl₂ in a 1:2 soil to solution ratio with
164 a glass electrode. Total C and N were determined simultaneously by automated combustion and gas
165 chromatography with thermal conductivity detection on a Flash EA112 analyzer (CE Elantech, New
166 Jersey, USA). After NaOH-EDTA extraction, inorganic P in the extraction material was determined by
167 molybdate colorimetry at 880 nm with a 1-cm path length. Total P in the NaOH-EDTA extracts was
168 determined by molybdate colorimetry at 880 nm with a 1-cm path length, following acid-persulfate
169 digestion at 80 °C overnight in sealed glass tubes. Organic P was calculated as the difference between
170 NaOH-EDTA total P and NaOH-EDTA P_i. Labile (plant-available) P was determined by Bray-1
171 extraction, with phosphate detected using automated molybdate colorimetry on a Lachat Quikchem
172 8500 (Hach Ltd, Loveland, CO). Exchangeable cations were determined by extraction in 0.1 M BaCl₂
173 (2 hours, 1:30 soil to solution ratio) and detection by inductively-coupled plasma optical-emission
174 spectrometry (ICP-OES) with an Optima 7300 DV (Perkin-Elmer Ltd, Shelton, CT, USA). Total
175 exchangeable bases (TEB) was calculated as the sum of the charge equivalents of Ca, K, Mg and Na.
176 Effective cation exchange capacity (ECEC) was calculated as the sum of the charge equivalents of Al,
177 Ca, Fe, K, Mg, Mn and Na. Base saturation was determined as TEB / ECEC ×100.

178

179 *2.4 Root colonization by fungi*

180 Fungal colonization was determined on fine roots (<2 mm diameter) of F, H, Ae and B horizons (no
181 roots in the L). Roots were cleared in 10% w/v KOH, then stained in an ink and vinegar solution for 5
182 min at 90 °C [63–65]. Roots were then rinsed in slightly acidified tap water for 30–40 min to remove
183 excess ink, after which they were placed in a 50% (v/v) lacto-glycerol solution for storage until
184 colonization could be evaluated. The gridline intersection method was performed under
185 stereomicroscope to quantify the length of roots colonized by AM and EcM fungi [63, 66]. Due to

186 magnification limits, some structures of ericoid mycorrhizal fungi might have been included in the AM
187 colonization percentage.

188

189 *2.5 Fungal community characterization*

190 The fungal community was characterized by amplicon sequencing. Soil DNA was extracted using the
191 PowerSoil DNA Isolation Kit (#12888-100 - Mo-Bio Laboratories Inc., Carlsbad, USA) following the
192 instructions of the manufacturer. Around 100 mg of soil for organic horizons (L, F and H), and 200 mg
193 for mineral horizons (Ae and B) were used for the extraction.

194 Soil amplification of the Internal Transcribed Spacer of the ribosomal RNA was performed by Genome
195 Québec (Montréal, Canada) with the ITS3_KYO2 and ITS4 primer pair [67]. This pair of primer limits
196 coverage bias toward Ascomycetes or Basidiomycetes and is also known to amplify Glomeromycetes
197 [e.g. 68]. The final reaction mix contained 0.02 U μl^{-1} Taq Roche HiFi polymerase, 1X Buffer 10X
198 with 18 mM MgCl₂, 5% DMSO, 0.2mM of each dNTP and 0.5 μM of each primer and DNA sample
199 diluted at 1/100. Thermal cycling was done in an Eppendorf Mastercycler Gradient (Eppendorf,
200 Hamburg, Germany) with the following cycling conditions: 2 min initial denaturation at 94 °C; 40
201 cycles of 30 s denaturation at 94 °C, 30 s annealing at 55 °C and 30 s elongation at 72 °C; and a 7 min
202 final elongation at 72 °C. The PCR products were loaded on 1% agarose gels with 1× sodium borate
203 buffer run at 220 V, and visualized after ethidium bromide staining (1 $\mu\text{g ml}^{-1}$).

204 Soil amplicon sequencing was performed by using the MiSeq Illumina technology by Genome Québec
205 (Montréal, Canada). The final concentration of the reaction mix contained 0.025 U μl^{-1} Taq Roche
206 HiFi polymerase, 1X Buffer 10X, 1.8mM of MgCl₂, 5% DMSO. Sequencing was done in an MiSeq
207 Illumina with the following conditions: 10 min initial denaturation at 95 °C; 15 cycles of 15 s
208 denaturation at 95 °C, 30 s annealing at 60 °C and 1 min elongation at 72 °C; and a 3 min final
209 elongation.

210

211 *2.6 Bioinformatics*

212 The fungal community was determined by filtering, denoising and assigning taxonomy to paired
213 amplicons using a customized script
214 (https://github.com/alexiscarter/Fungal_com_SBL/tree/master/dada2) adapted from the DADA2
215 pipeline [69]. In brief, using the *filterAndTrim* function, reads were truncated at 280 bp and discarded if

216 they had more than three expected errors or a quality score lower than six. Then, amplicon sequence
217 variants (ASV) were inferred for each sample with the *dada* function. Forward and reverse reads were
218 merged using the *mergePairs* function with a minimum overlap of 12 bp. Potentially chimeric
219 sequences were identified by the pooled method of the *removeBimeras* function. The amplicon
220 sequence variant approach was used instead of the classical operational taxonomic as proposed by
221 Callahan *et al.* [70] and others [71]. This method does not use a particular threshold for classifying
222 sequences into operational taxonomic units, as no threshold appears to be universally applicable for
223 fungi [72]. Instead, it used the divisive amplicon denoising algorithm aimed at finding ASV that refer
224 back to original biological sequences [69, 73]. The taxonomy of the ASV was assigned with the
225 UNITE database, version 7.2 [74]. ASV that belong to the same species were grouped together. The
226 functional information for ASV was obtained from the online FUNGuild database [44].
227

228 *2.7 Statistical analyses*

229 To describe the fungal community and assess the effects of environmental parameters we used
230 ordination approaches and multivariate analyses of variance. The community matrix was composed of
231 the number of sequences per ASV of 75 soil samples from five soil horizons in each of 15 plots (one
232 sample of L horizon in an EcM plot was excluded due to poor amplification). Due to some inherent
233 limitations of the approach, either biological (e.g., varying number of DNA copies per organism) or
234 technical (varying sequencing depth, extraction and amplification biases among samples), the number
235 of sequence reads is not a direct measure of taxa abundance in the environment, but comparisons
236 among samples remain useful as they can be considered semi-quantitative [75, 76]. Explanatory
237 variables for each sample were classified into three groups: (i) soil chemistry, (ii) soil horizon (L, F, H,
238 Ae or B), and (iii) forest type (AM, EcM or mixed).
239 Differences in soil properties, root colonization, guild abundance and richness among horizons and
240 forest type were tested using linear mixed-effect models; block was treated as random factor in these
241 analyses. Model assumptions were assessed by visual inspections of residuals. Comparison were
242 determined using post-hoc Tukey tests were used to determine significant differences.
243 In β -diversity analyses, we used the Bray-Curtis dissimilarity index for the community structure and its
244 binary version, the Sørensen index for the community composition [77]. These asymmetrical

245 coefficients do not consider double zeroes and can therefore be used with raw abundances or counts
246 [77].

247 To visualize differences in fungal community composition and abundance among samples, we used
248 non-metric multidimensional scaling (NMDS). To test for differences between samples across horizons
249 and forest types, we used permutational multivariate analysis of variance (PERMANOVA). *P*-values
250 for pairwise tests were adjusted using the Benjamini-Hochberg method [78]. Because the
251 PERMANOVA method is sensitive to differences in multivariate dispersions among groups, the
252 homogeneity of dispersion was tested to assess differences and tested for significance by permutations
253 [79].

254 Distance-based redundancy analysis (RDA) was used to quantify the extent to which changes in fungal
255 community structure were related to soil chemistry, horizon and forest type [77]. Soil chemistry data
256 were standardized and linear dependencies were explored using variance inflation factors and avoided
257 if >10 [80]. To test how much variance was independently explained by the explanatory matrices,
258 variation partitioning was performed using partial RDA [pRDA, 81]. In RDA and pRDA, coefficients
259 of determination were adjusted (i.e. adjusted- R^2 values) to take into account the number of explanatory
260 variables in the model [82, 83].

261 Analyses were performed and visualized using the R software [84] with the following main packages:
262 *dada2* [69], *dplyr* [85], *emmeans* [86], *ggplot2* [87], *ggpubr* [88], *nlme* [89], *phyloseq* [90] and *vegan*
263 [91]. Code for bioinformatical and statistical analyses are available at:
264 <https://doi.org/10.5281/zenodo.3631982>. Sequence and chemistry data can be accessed at
265 <https://doi.org/10.5281/zenodo.3631861>.

266

267 **3. Results**

268 *3.1 Soil chemistry variation across horizons and forest types*

269 All soil chemical properties varied significantly across horizons (Fig. 1), and these differences were
270 consistent across forest types (soil horizon \times forest type interaction, $P > 0.05$; except for pH where $P =$
271 0.026). The pH of the L horizon declined from pH ~4 (in 0.01 M CaCl₂) to ~3.25 in the H horizon, but
272 this decline was not as pronounced for AM forests than for EcM or mixed forests (Fig. 1a). The pH
273 then increased from the H to the B horizon in all forests. Effective cation exchange capacity and base
274 saturation declined with increasing depth (Figs. 1b-c), except for ECEC in the Ae horizon. Organic C

275 generally declined with depth, but AM forests tended to have lower organic C concentration in the H
276 horizons than EcM or mixed forests (Fig. 1d). By contrast, total N increased from the L to the Ae
277 horizon and then declined in the B horizon (Fig. 1e). As a result, the C:N ratio decreased with
278 increasing depth from the L to the Ae horizon (Fig. 1f). Inorganic and organic P increased in deeper
279 horizons while labile (Bray) P decreased (Figs. 1g-i).

280 Forest types differed significantly in their pH, C:N ratio, NaOH-EDTA total P, NaOH-EDTA organic
281 and inorganic P concentrations ($P < 0.05$). AM-dominated forest plots tended to have higher pH, total
282 P, inorganic P and organic P but lower C:N ratio compared to EcM-dominated forest plots.

283

284 *3.2 Root colonization by mycorrhizal fungi*

285 Colonization of fine roots by AM and EcM fungi was significantly different among mycorrhizal type
286 ($P < 0.0001$, Fig. 2) but only differ across horizons in the EcM-dominated forest ($P = 0.007$). Fine
287 roots in AM forest were more strongly colonized by AM fungi than those from mixed and EcM forests
288 ($P < 0.05$, Fig. 2a). By contrast, fine roots in EcM forests were more strongly colonized by EcM fungi
289 compared to those from AM forests ($P < 0.05$, Fig. 2b). Root colonization by EcM fungi tended to
290 decrease with soil depth in EcM forest down to ~20% in the B horizon (Fig. 2b). In mixed and AM
291 forests, EcM colonization was highest in the H or Ae horizons but always lower than 30%.

292

293 *3.3 Overall fungal community*

294 We found 781 fungal taxa (at the species level or below) from a total of 2521 ASV detected using
295 high-throughput amplicon sequencing across all horizons and plots. Fungal ASV richness tended to
296 decrease with soil depth regardless of the forest type (Fig. S3). The highest fungal ASV richness was
297 found in L horizons of the AM forests.

298

299 *3.4 Fungal guilds*

300 Saprotrrophic and symbiotrophic (EcM, AM and ErM) guilds showed distinct vertical distributions
301 among horizons and across forest types (Fig. 2c-f). Saprotrrophic fungal taxa dominated the upper
302 horizons (especially L and F; Fig. 2c), and mycorrhizal fungi were almost absent in the L horizon
303 (Figs. 2d-f). Fungal taxa assigned to the saprotrophic guild were slightly more abundant in the organic
304 horizons of the AM and mixed forests compared to EcM forest (Fig. 2c). Abundance of saprotrophic

305 fungi were significantly different among forest types ($P < 0.031$) but differences were not significant
306 across horizons of different forest types (soil horizon \times forest type, $P = 0.325$). In deeper horizons,
307 sequences attributed to mycorrhizal fungi were more abundant (Figs. 2d-f). Sequences of AM (i.e.
308 Glomeromycetes) fungi were much more abundant in the AM forest (Fig. 2d), and the opposite was
309 true for EcM fungi (Fig. 2f). Both AM and EcM taxa were well represented in the mixed forests (Figs.
310 2d-e). Sequences of ericoid mycorrhizal (ErM) fungi were less abundant in AM forest except for the F
311 horizon where their abundance was high in all forests (Fig. 2f). Richness patterns of fungal guilds
312 tended to follow abundance data (Fig. S4). Saprotrophic fungi had the higher number of taxa followed
313 by EcM, ErM and AM fungi. Saprotrophic fungal richness was highest in the upper horizons and
314 decreased with depth. There was a higher richness of EcM fungi in EcM and mixed forests and very
315 few EcM taxa in the L horizon.

316

317 *3.5 Fungal community structure*

318 Soil horizons had the strongest influence over fungal community structure (includes abundance data) in
319 the three forest types, as shown by the NMDS ordination (Fig. 3). The composition (based on presence-
320 absence data) of the fungal community showed similar patterns (Fig. S5), suggesting that results
321 primarily reflected changes in ASV composition rather than relative abundance. Differences in
322 multivariate dispersions with Bray-Curtis and Sørensen measures were not significant among forest
323 types ($P > 0.05$) but were significant among horizons ($P < 0.05$), with the L horizon showing the
324 lowest multivariate dispersions. In other words, fungal communities from the L horizons were more
325 similar to each other than fungal communities from the other horizons. Fungal community composition
326 and abundance significantly differed among all horizons but also among forest types ($P < 0.001$, Table
327 S2). However, the differences among horizons did not depend on forest type and vice-versa (soil
328 horizon \times forest type interaction not significant; Table S2). Pairwise comparisons revealed that fungal
329 community composition and abundance in AM and EcM forests significantly differed from each other,
330 but not from mixed forests (Fig. 3).

331

332 *3.6 Edaphic drivers of fungal community structure*

333 Variation in soil chemistry explained a large fraction of the total variation in fungal community
334 structure (adjusted- $R^2 = 23.3\%$, $P = 0.001$, see Table S3 for results of the constrained ordinations). In

335 the L horizons, fungal communities were associated with higher pH, ECEC, labile L and C:N ratio
336 (Fig. 4). Fungal communities in mineral horizons (Ae and B) were associated with high organic and
337 inorganic P but low labile P (Fig. 4). Between L and mineral horizons, fungal communities were
338 associated with low pH (H horizon) and high labile P (F horizon).

339 Forest mycorrhizal type explained a lower but still significant amount of variation (adjusted- $R^2 = 2.7\%$,
340 $P = 0.006$). There was a clear difference in the fungal community structure of AM and EcM forests,
341 whereas the mixed forests were intermediate or more similar to EcM forest (Fig. 5).

342 Abiotic and biotic variables together explained ~35% ($P = 0.001$) of the total variation in the fungal
343 community structure. Variation in fungal community structure depended on horizons and forest
344 mycorrhizal types, and was also influenced by soil chemistry (Fig. 6). Within forest types, fungal
345 communities were not significantly different among blocks. Horizon, forest type and soil chemistry
346 still explained a significant fraction of the variation in the fungal community structure when
347 considering the effects of the other variables (Table S3). Most of the explained variation was shared
348 between soil chemistry and horizon (Fig. 6). However, forest type still had a unique and significant
349 impact on the variation of the fungal community. A small fraction of variation was shared between soil
350 chemistry and forest type (Fig. 6).

351

352 **4. Discussion**

353 In this study, we determined vertical shifts in soil fungal community composition across soil horizons
354 and forest mycorrhizal types (AM, EcM, and mixed AM/EcM) and compared how saprotrophic fungal
355 dominance extends to deeper horizons in AM vs. EcM forests. Although there was a tendency for
356 lower abundance of saprotrophic fungi in organic F and H horizons in EcM forests than in AM or
357 mixed forests, all three forest types showed a similar saprotrophic-to-mycorrhizal shift in fungal
358 composition with increasing soil depth. This shift in fungal dominance was most pronounced in
359 organic horizons. Moreover, we found that changes in fungal community composition were largely
360 driven by differences in soil chemistry, which were far stronger across horizons (i.e. depth) within a
361 single forest than across forest mycorrhizal types for the same horizon. Our results highlight the
362 importance of considering soil vertical structure and associated changes in chemistry when
363 characterizing soil fungal communities. They also suggest that, at least in northern forests, AM fungi

364 are not being restricted where inorganic nutrients predominate and might have more similar edaphic
365 vertical niches with EcM fungi than what has been suggested in the literature [31, 32, 35].

366

367 Fungal communities were strongly stratified with depth along the soil profile, being most distinct in the
368 L horizon (composed of recently-fallen leaves). Litter of the EcM, AM and mixed forests had high
369 fungal richness and distinct fungal communities that were dominated by saprotrophic fungi. This has
370 also been observed in forests of tropical, temperate and boreal biomes dominated by EcM trees [19, 21,
371 22, 92, 93]. Dominance by saprotrophic fungi in the most superficial litter layer has also been observed
372 in other AM-dominated ecosystems [29, 94], as we have found in this northern temperate forest. Our
373 results therefore provide further evidence of this general pattern whereby the L horizon possesses a
374 distinct fungal community dominated by fungal saprotrophs, compared to deeper horizons in which
375 mycorrhizal fungi are more abundant.

376

377 As suggested by Bahram et al. [51], studies that have reported weak vertical segregation of fungal
378 communities have often excluded the most superficial L horizon from their analyses [16, e.g. 49]. The
379 L horizon of the EcM, AM and mixed forests tended to have higher C:N ratio, pH, concentration of
380 cations and labile P than deeper horizons. While this pattern seems generalizable for pH [e.g. 21, 93], it
381 remains uncertain or unexplored for the other chemical variables. Our results suggest that the L horizon
382 which is characterized by the presence of organic matter in which the original structures can be visually
383 distinguished [62] should be considered separately in future studies of fungal community composition,
384 given its chemical, microbial and functional distinctiveness.

385

386 From the F to the B horizon, fungal communities showed strong turnover across soil horizons, with
387 distinct fungal communities in each horizon. The fungal composition, abundance and guilds tended to
388 progressively change among horizons in the soil profile but these changes were less pronounced than
389 with the L. This was also observed in other study systems [21, 93, 95]. There are reports of evenly
390 distributed guilds among the organic and mineral horizons [e.g. 16], but vertical segregation of fungi
391 and especially root-associated fungi is often strongly impacted by determinant factors such as soil
392 chemistry and host plants [19, 20, 51]. In our study, there was major variation in the vertical
393 distribution of soil fungi that was largely driven by soil chemical characteristics, with these changes

394 being observed in all three forest mycorrhizal types. Our results further support those of other studies
395 that have found the vertical variability of mycorrhizal and saprotrophic fungal communities across
396 different soil horizons to be much larger than horizontal or temporal variability [51, 96]. Studies that
397 focus on ecosystem topsoil processes in terrestrial environments should consider the strong physical,
398 chemical and biological heterogeneity that occurs within the first few centimeters, by sampling distinct
399 soil horizons separately.

400

401 We showed that underground fungal community structure varied significantly between neighboring
402 forest dominated by AM or EcM trees. As expected, AM forests showed higher abundance of AM
403 fungi, whereas EcM forests showed higher abundance of EcM fungi. Direct observation of fungal
404 colonization in roots confirmed these patterns. Forests with a mix of both strategies supported
405 intermediate communities between the two extremes of the gradient, as reported in a study focusing on
406 ecosystem processes [e.g. 97]. It is worth noting that fungal saprotrophs tended to be more abundant in
407 organic horizons of mixed and AM forests compared to EcM forests. Together with higher pH and
408 lower organic C in these AM forests, this result might indicate a tendency toward a more “inorganic
409 nutrient economy” compared to the studied forests dominated by EcM fungi. The latter would
410 represent a more “organic nutrient economy”, associated with a slower turnover of plant-derived C due
411 to lower abundance of free-living saprotrophs [10]. These small differences observed at local scale may
412 be responsible for observed patterns found at the ecosystem scale [14]. It has been found elsewhere that
413 forests dominated by different species of broadleaf trees of the same mycorrhizal strategy can also
414 show differences in fungal community structure [98]. However, in our study, fungal composition,
415 abundance and guilds tended to differ between EcM and AM forests. Such a distinction has previously
416 been reported in a study comparing very distinctive EcM forests of broadleaf trees vs. conifers [99], the
417 effect of mycorrhizal type was relatively small but nonetheless present, and could also be linked to
418 differences in nutrient availability.

419

420 Our study design provides a useful system for exploring the relative importance of mycorrhizal type on
421 soil biogeochemical cycling. The soil profile in these northern temperate forests have low vertical
422 mixing, resulting in podzols with high stratification, as commonly encountered in boreal soils. Soil
423 horizons were easily identifiable mainly through their color and such sampling may allow for better

424 association between DNA sequences and soil chemistry as well as more valuable comparison across
425 sites [100]. Variation in important factors such as parent material, topography and regional climate
426 were minimized but other factors (e.g. productivity, soil texture) could still co-vary with mycorrhizal
427 dominance at the plot scale. Importantly, this study system allowed us to study different mycorrhizal
428 types within the same site [7, 30, 51] and across a gradient of mycorrhizal dominance [15]. The
429 observed differences in soil chemistry among forests could be linked with dominant mycorrhizal
430 strategies. Higher saprotrophic fungal diversity has been observed in the upper soil layers of AM-
431 dominated tropical forests compared to EcM forests [101]. Our study provides further evidence, in a
432 temperate system, host plants are an important factor controlling mycorrhizal community composition
433 [51, 102]. To some extent, this was expected given that AM and EcM fungi are obligate symbionts
434 with their host plants [32]. As such, considering tree mycorrhizal strategies and their interactions with
435 saprotrophs may help to better predict carbon storage at small and global scale [8].

436

437 Our use of high-throughput amplicon sequencing approach allowed us to assess the distribution of the
438 soil fungal community and to discriminate among AM, EcM and saprotrophic fungi. However, result
439 from high-throughput sequencing approaches need to be interpreted with caution because of
440 unavoidable biases at different levels [43, 103]. For example, how to adequately normalize for taxa
441 abundance among samples remains unresolved [104, 105]. Furthermore, although we acknowledge that
442 soil and root compartments might host different fungal communities [e.g. 106], but sampling bulk soil
443 allows to capture the potential free-living saprotrophs as well as root-associated fungi and their
444 extraradical hyphae. Finally, our choice of the primers might have resulted in an under-representation
445 of some fungal groups such as Glomeromycetes, but comparisons in taxa abundance between samples
446 remain relevant [76]. Using specific primers targeting Glomeromycetes [107, 108], and plants using
447 DNA from the root tissue [68, 109] would certainly allow to further understand the importance of these
448 underground interactions and the vertical segregation among root and fungi of different mycorrhizal
449 types.

450

451 Our results show that fungal communities in horizons vertically separated by a few centimeters are
452 very different from each other in terms of composition and abundance. This contributes to high fungal
453 and functional diversity in the topsoil. Moreover, our work suggests that the forest mycorrhizal type

454 influences the overall and saprotrophic fungal community, advancing our current understanding of the
455 potential impacts of mycorrhizal strategies on the distribution of key organisms for ecosystem
456 functioning such as C and nutrient cycling [10]. We also reported for the first time that broad patterns
457 of vertical fungal distribution across the upper five horizons in AM-dominated northern forest are
458 comparable to neighboring EcM-dominated or mixed forests. This result challenges the traditional
459 view that AM fungi have a more restricted niche toward mineral soils compared to EcM fungi due to
460 their incapability to directly decompose organic matter [31]. Our study suggests that the ecological and
461 functional roles of AM fungi in organic horizons of temperate forests, including recently deposited
462 litter, deserves more attention [39].

463

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476

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478

479 **Availability of data and material**

480 Sequence and chemistry data can be accessed at: <https://doi.org/10.5281/zenodo.3631861>.
481 Custom code for bioinformatical and statistical analyses are available at:
482 <https://doi.org/10.5281/zenodo.3631982>.

483

484 **Authors' contributions**

485 EL and AC conceived the ideas and designed methodology; AC, BT, SJ and MB collected the data; AC
486 analyzed the data; AC and EL interpreted the results; AC led the writing of the manuscript. All authors
487 contributed critically to the drafts and gave final approval for publication.

488

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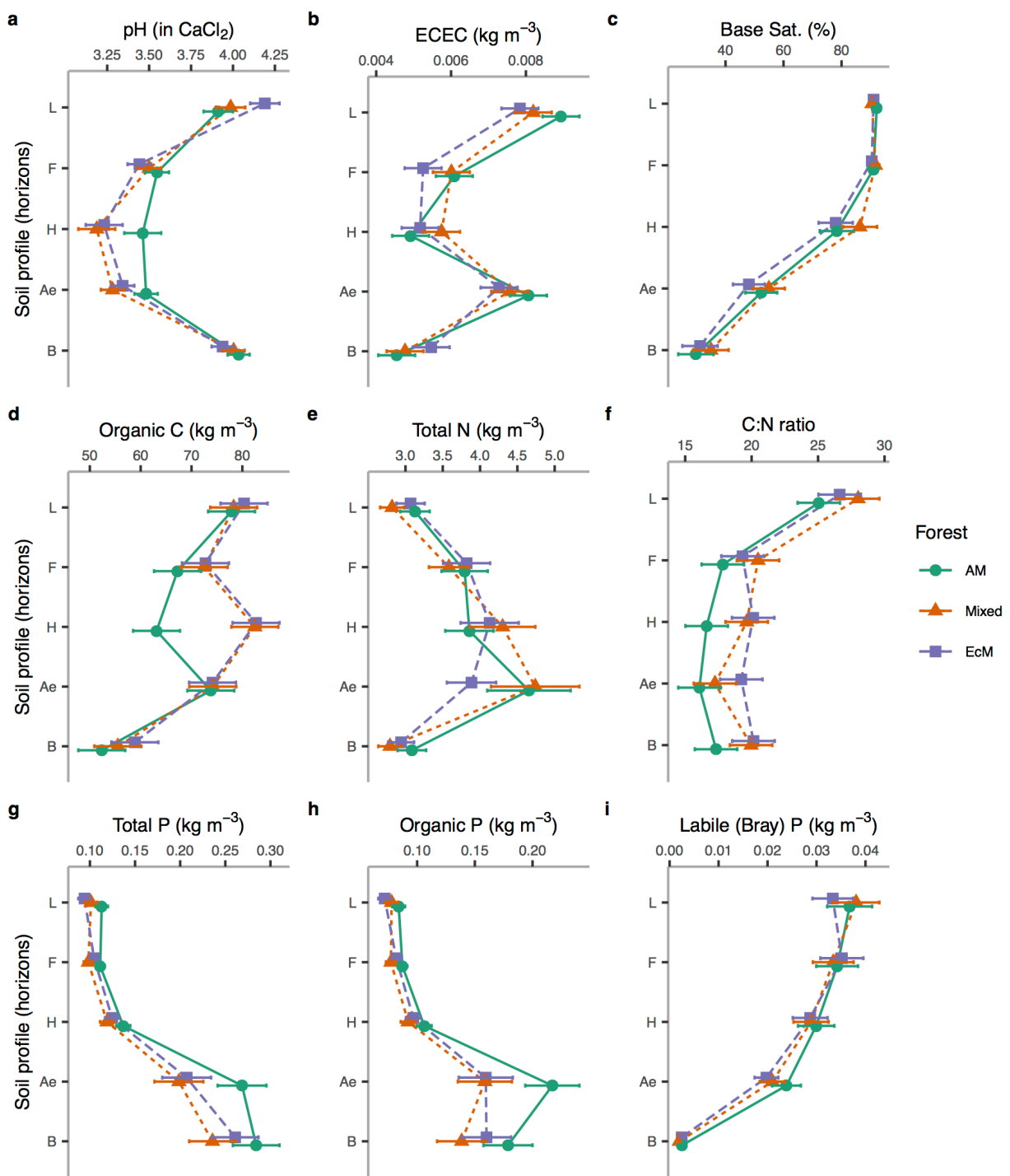
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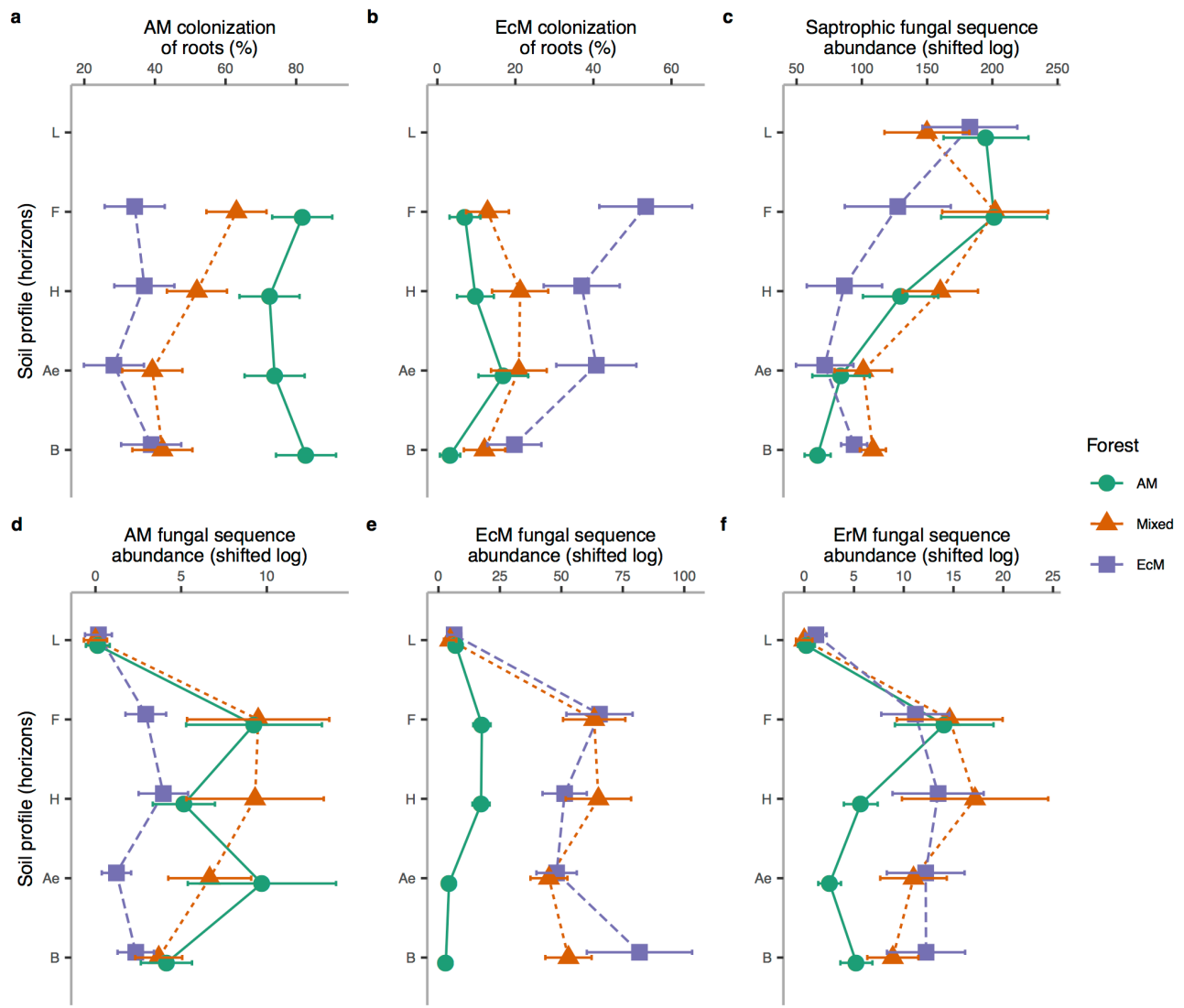
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783 **FIGURES**

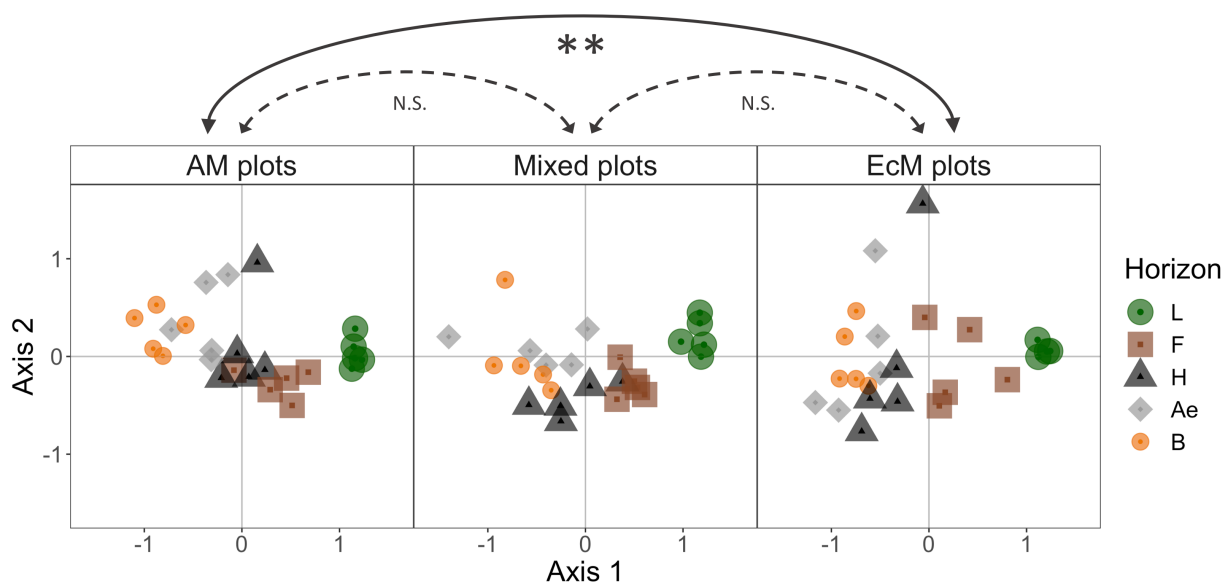
784

785 **Fig. 1** Soil physico-chemical characteristics from organic-to-mineral horizons (L, F, H, Ae, B) in each
 786 mycorrhizal forest type (AM, arbuscular mycorrhizal; EcM, ectomycorrhizal; Mixed, mixture of AM
 787 and EcM): (a) pH (in CaCl_2), (b) effective cation exchange capacity, (c) base saturation, (d) organic
 788 carbon, (e) total nitrogen, (f) carbon over nitrogen ratio, (g) total phosphorus, (h) organic phosphorus
 789 and (h) labile (Bray) phosphorus. All data are means ± 1 SE ($n = 5$)



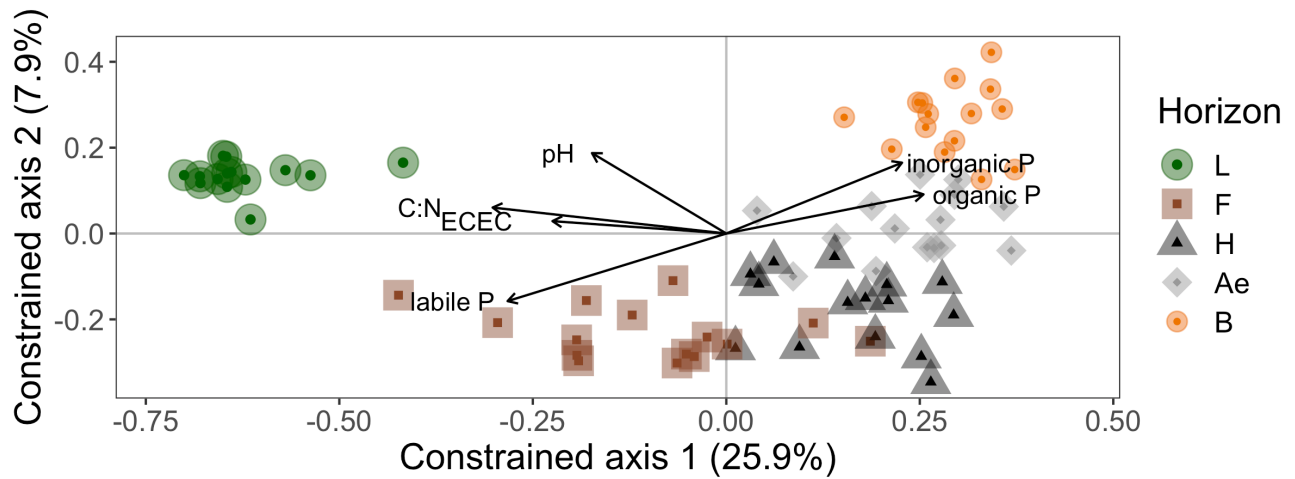
790

791 **Fig. 2** Soil profiles from organic-to-mineral horizons (L, F, H, Ae, B) on each mycorrhizal forest type
 792 (AM, arbuscular mycorrhizal; EcM, ectomycorrhizal; Mixed, mixture of AM and EcM) showing
 793 variations in: root colonized by (a) AM fungi, (b) EcM fungi, and abundances (on shifted log data) of
 794 sequences belonging to (c) saprotrophic fungi, (d) AM fungi, (e) EcM fungi, (d) ericoid mycorrhizal
 795 (ErM) fungi. Upper organic horizon (L) had no roots so colonization was set to zero. All data are
 796 means \pm 1 SE ($n = 5$, except $n = 4$ for the L horizon in EcM forest)



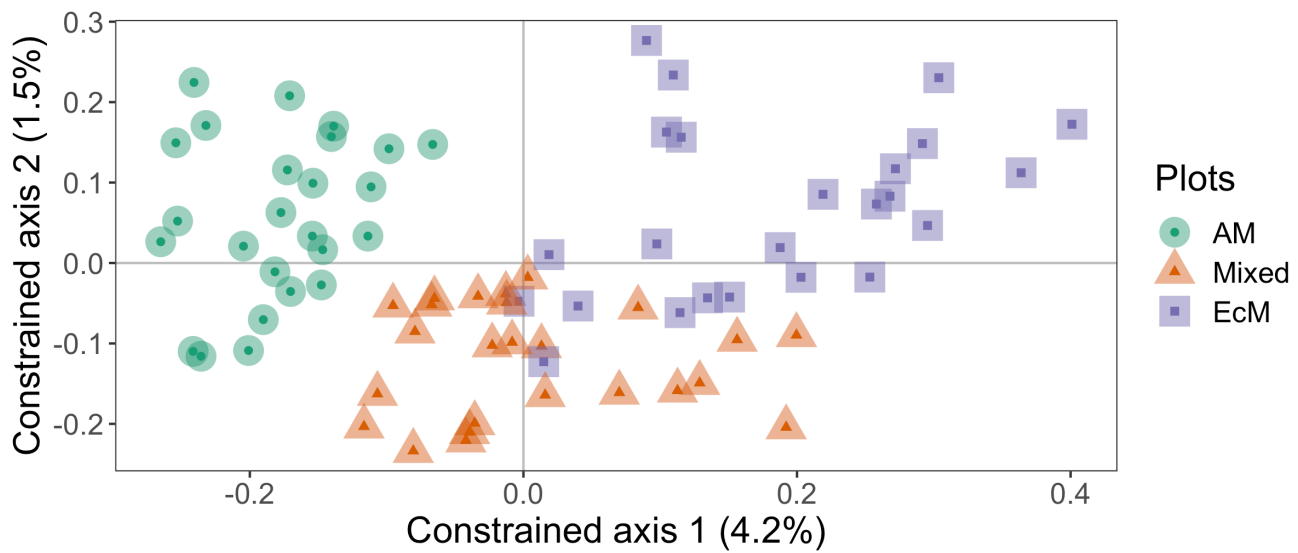
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798 **Fig. 3** Ordination of the fungal community composition (Bray-Curtis dissimilarities) plotted in the
 799 different forest types using a non-metric multidimensional scaling with two dimensions and a stress of
 800 0.17. ** indicates difference in fungal community structure between arbuscular mycorrhizal (AM) and
 801 ectomycorrhizal (EcM) plots (P -value ≤ 0.01), N.S. indicates non-significant differences (see Table S2
 802 for details)



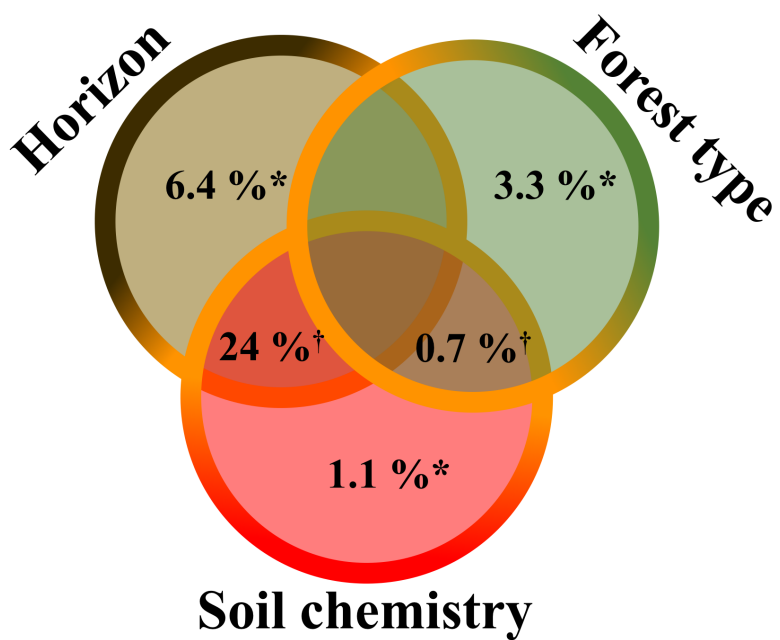
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804 **Fig. 4** Constrained ordination of the overall fungal community by soil chemistry variables using a
 805 distance-based redundancy analysis with Bray-Curtis dissimilarities. Horizons are shown in different
 806 shape and colors. The two first constrained axes explaining most variation are drawn. Adjusted- R^2 =
 807 23.3 %, P -value = 0.001



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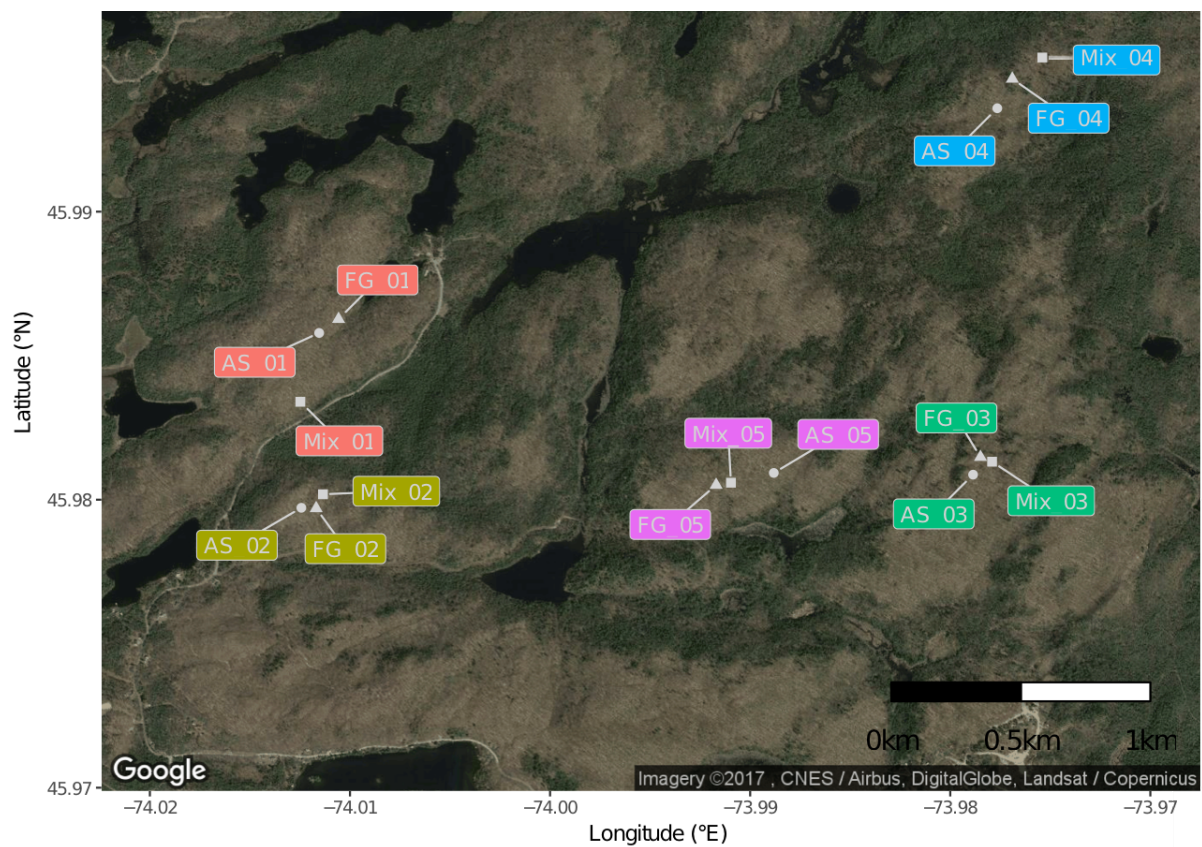
809 **Fig. 5** Constrained ordination of the fungal community structure depending on the forest mycorrhizal
 810 type (AM, arbuscular mycorrhizal; EcM, ectomycorrhizal; Mixed, mixture of AM and EcM) using a
 811 distance-based redundancy analysis with Bray-Curtis dissimilarities. Forest type are shown in different
 812 shape and colors. The two constrained axes are shown. Adjusted- $R^2 = 2.7\%$, P -value = 0.006



813

814 **Fig. 6** Venn diagram displaying the amount of variation (i.e. adjusted- R^2) of the fungal community
 815 explained by horizon, soil chemistry and forest mycorrhizal type or a combination of them. Values
 816 <0.1 % are not shown. Ellipses are not drawn to scale. Only variables with significant redundancy
 817 analysis (RDA) results were tested for partial-RDA and included in this diagram. Overall adjusted- R^2 =
 818 34.8 %, * indicates P -value < 0.05 and † indicates non-testable portion. For more details see Table S3

819 **Supplementary Material**



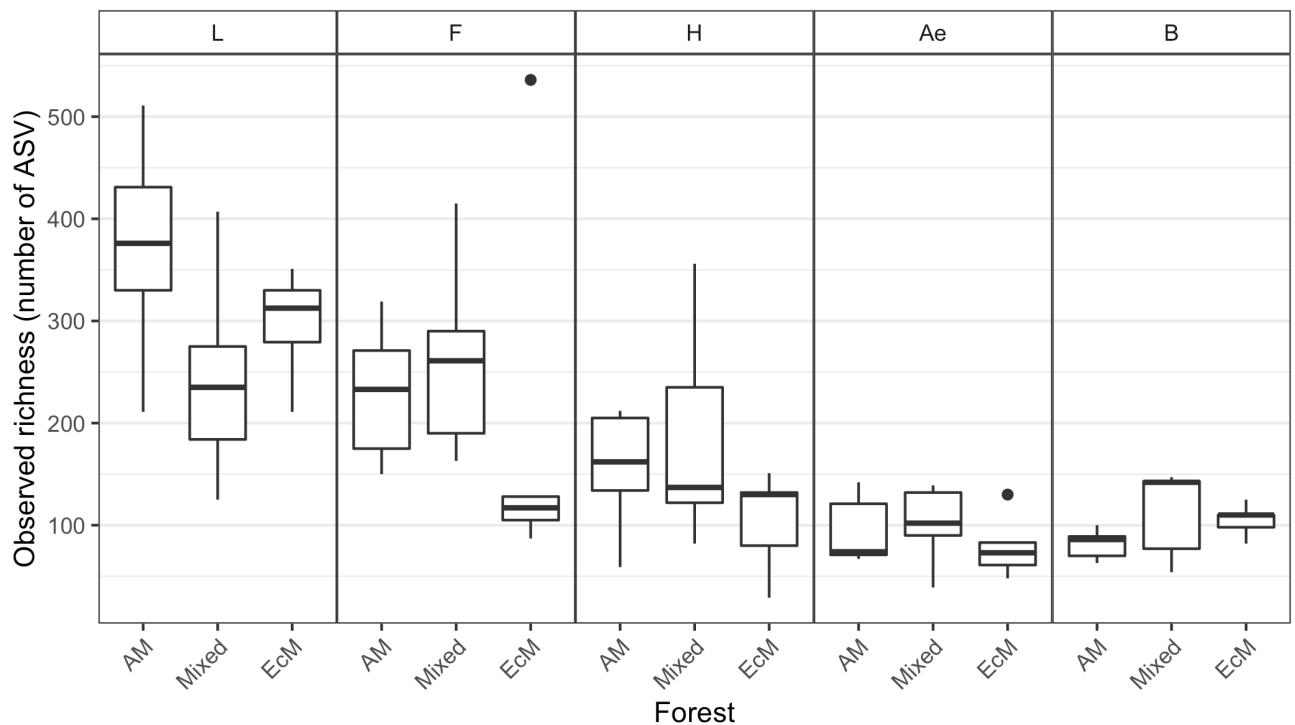
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821 **Figure S1.** Map showing the 15 plots grouped in five blocks (different colors) at the University of
822 Montréal's field station (Québec, Montréal). The characteristics of each plots are listed in Table S1.

823

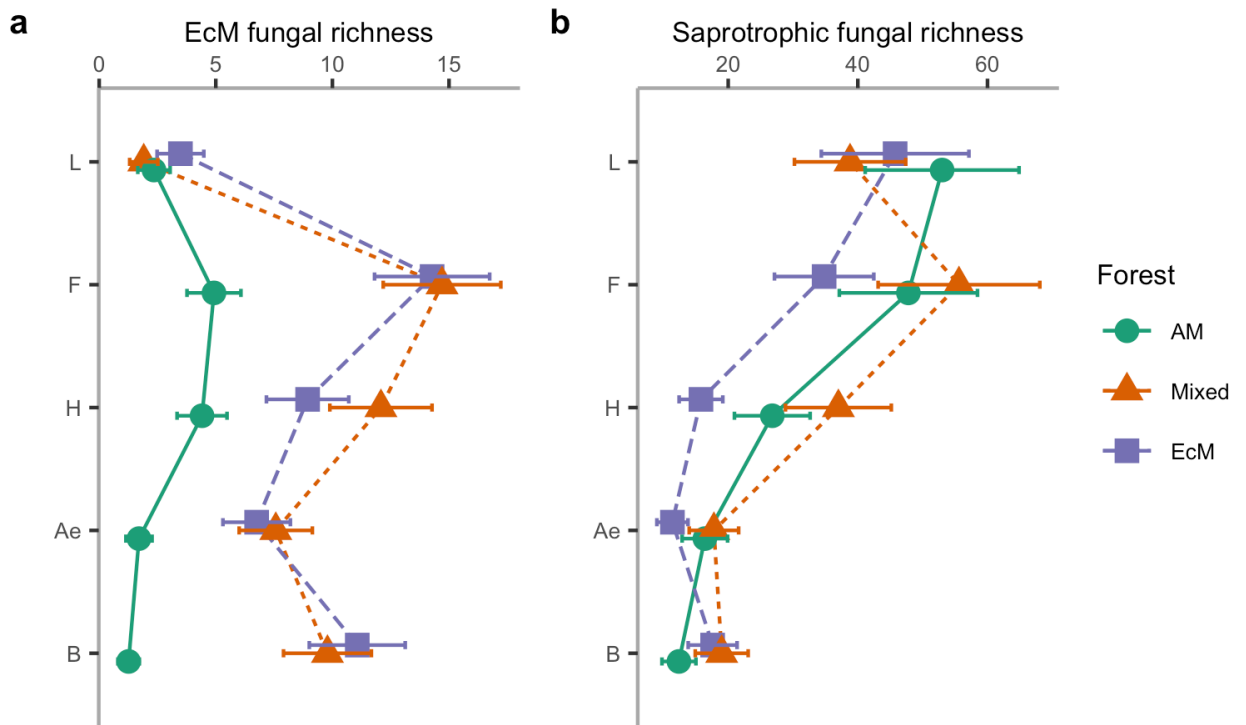


824 **Figure S2.** Picture of a soil core of approximately 25 cm deep, sampled with a rectangular auger,
825 representing a typical profile in the studied sites. The five horizons can easily be distinguished with entire
826 leaves at the top (L), then partially decomposed materials (F) and black humus (H), followed by grey Ae
827 and brown B.



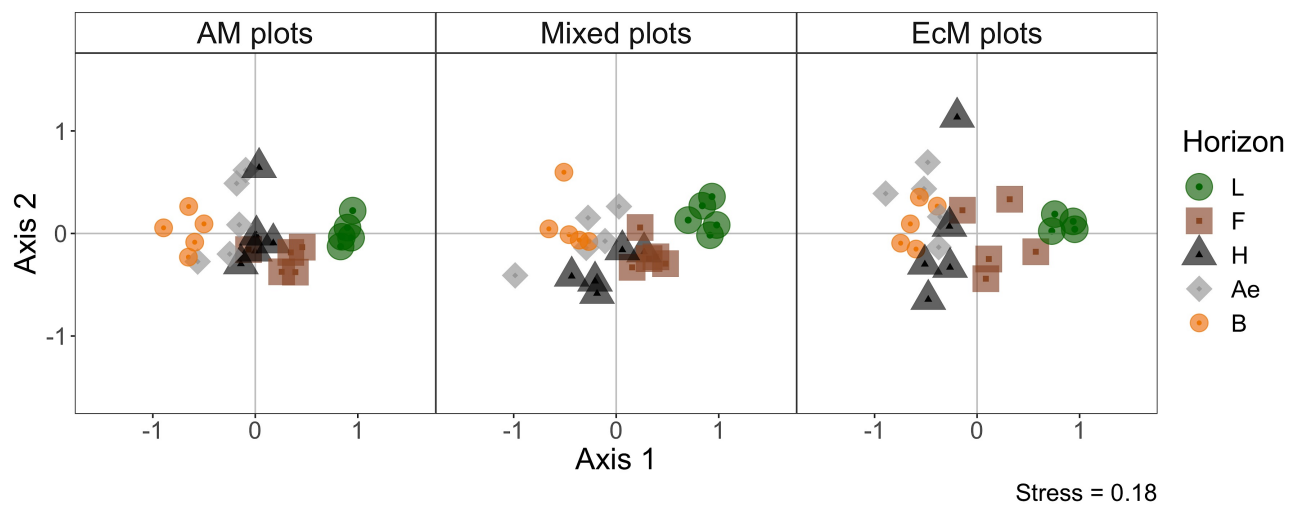
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829 **Figure S3.** Boxplots illustrating differences in the number of amplicon sequence variant (ASV) by
 830 horizon (L to B, from left to right). Singletons and doubletons were excluded. Bold horizontal lines
 831 represent median values; box margins 25th and 75th percentile; vertical lines represent largest and lowest
 832 value within 1.5 times interquartile range above 75th and below 25th percentile respectively; dots
 833 represent outliers that fall outside of that range.



834

835 **Figure S4.** Soil profiles from organic-to-mineral horizons (L, F, H, Ae, B) on each mycorrhizal forest
 836 types (AM, arbuscular mycorrhizal; EcM, ectomycorrhizal; Mixed, mixture of AM and EcM) showing
 837 variations in richness of (a) EcM fungi and (b) saprotrophic fungi. All data are means \pm 1 SE ($n = 5$,
 838 except $n = 4$ for the L horizon in EcM forest). Note: Due to low value, richness of AM and EcM guilds
 839 were not modeled.



840

841 **Figure S5.** Ordination of the fungal community composition (Sørensen distances)
 842 of the different forest type on two axes using a non-metric multidimensional scaling with two dimensions and a stress of 0.18.
 843 To visually assess the impact of Sørensen distance, the scale is kept identical to the one of Fig. 1.

844 **Table S1.** Characteristics of each plot under study. Plots in the same block were selected to have
 845 homogenous environmental conditions.

Plot ID	Block	Altitude	Slope	Aspect	Total basal area (m ² ha ⁻¹)	EcM basal area (% of total)	tree AM (%) of total)	tree basal area (% of total)	Dominant canopy species	Mycorrhizal dominance
AS_01	1	403	10	95	23.2	7.5	92.5	AS	AM	
FG_01	1	381	20	103	37.4	81.6	18.4	FG	EcM	
Mix_01	1	383	18	160	36.1	53.4	46.6	AS and FG	Mixed	
AS_02	2	398	13	140	30.9	8.4	91.6	AS	AM	
FG_02	2	391	9	105	40.4	95.8	4.2	FG	EcM	
Mix_02	2	381	10	110	41.9	57.4	42.6	AS and FG	Mixed	
AS_03	3	374	9	140	33.3	8.4	91.6	AS	AM	
FG_03	3	388	0	0	29.9	79.7	20.3	FG	EcM	
Mix_03	3	396	0	0	37.0	43.9	56.1	AS and FG	Mixed	
AS_04	4	376	16	220	38.9	8.3	91.7	AS	AM	
FG_04	4	395	14	120	39.1	62.8	37.2	FG	EcM	
Mix_04	4	375	18	180	27.8	55.5	44.5	AS and FG	Mixed	
AS_05	5	366	15	140	40.2	6.4	93.6	AS	AM	
FG_05	5	365	9	150	36.3	89.7	10.3	FG	EcM	
Mix_05	5	366	20	190	30.8	56.1	43.9	AS and FG	Mixed	

846 Acronyms: AS = *Acer saccharum*, FG = *Fagus grandifolia*, AM = arbuscular mycorrhiza, EcM =
 847 ectomycorrhiza.

848 **Table S2.** Multivariate analyses of differences in structure (Bray-Curtis dissimilarities) and composition
 849 (Sørensen distances) among different types of fungal communities. Analyzed using permutational
 850 multivariate analysis of variance (PERMANOVA). *P*-values were determined using 9999 permutations.
 851 * *P* ≤ 0.05; ** *P* ≤ 0.01, *** *P* ≤ 0.001. Df stands for degree of freedom.

Fungal Community	Dissimilarity measure	<i>P</i> -Value	Df	Fungal Community	<i>P</i> -Value
Among forest types	Structure	0.00014***	2	AM vs Mixed	0.0935
				AM vs EcM	0.0039**
				Mixed vs EcM	0.0935
	Composition	0.00074***	2	AM vs Mixed	0.118
				AM vs EcM	0.026*
				Mixed vs EcM	0.089
Among horizons	Structure	0.00001***	4	L vs F	0.00001***
				F vs H	0.00001***
				H vs Ae	0.0023**
				Ae vs B	0.00007***
				L vs F	0.00001***
	Composition	0.00001***	4	F vs H	0.0001***
				H vs Ae	0.0118*
				Ae vs B	0.0022**
				-	-
				-	-
Forest × Horizon	Structure	0.11718	8	-	-
	Composition	0.15424	8	-	-

852 Note: Only PERMANOVA results with *P*-Value ≤ 0.05 were considered for multiple comparisons. L vs
 853 H, L vs Ae, L vs B, F vs Ae, F vs B, H vs B not included but *P*-values < 0.0001.

854 **Table S3.** Partition of variation of the fungal community due to soil chemistry (Chemistry), experimental
 855 blocking design (Block), soil layers (Horizon) and forest mycorrhizal type (Forest) using distance-based
 856 redundancy analysis (RDA) and partial-RDA analyses. *P*-values were determined using 999
 857 permutations. ** *P*-values ≤ 0.01 ; *** *P*-values ≤ 0.001 . In the formula Y is the fungal community
 858 matrix, X is the explained matrix and Z is the conditional matrix which is partialed out. Only significant
 859 RDA results were tested for partial-RDA.

Model	Formula (Y~X Z)	adjusted-	
		R ² (%)	<i>P</i> -value
Overall RDA	Y~ Chemistry + Block + Horizon + Forest	34.8	0.001***
	Y ~ Chemistry	23.3	0.001***
Single RDA	Y~ Block	0.3	0.372
	Y~ Horizon	27.8	0.001***
	Y~ Forest	2.7	0.006**
partial-RDA	Y~ Chemistry Block + Horizon + Forest	1.1	0.036*
	Y~ Horizon Chemistry + Block + Forest	6.4	0.001***
	Y~ Forest Chemistry + Block + Horizon	3.3	0.001***

860