1	Arbuscular mycorrhizal fungal community composition is altered by long-
2	term litter removal but not litter addition in a lowland tropical forest
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40	SUMMARY
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42	• Tropical forest productivity is sustained by the cycling of nutrients through
43	decomposing organic matter. Arbuscular mycorrhizal (AM) fungi play a key role in
44	the nutrition of tropical trees, yet there has been little experimental investigation into
45	the role of AM fungi in nutrient cycling via decomposing organic material in tropical
46	forests.
47	
48	• We evaluated the responses of AM fungi in a long-term leaf litter addition and removal
49	experiment in a tropical forest in Panama. We described AM fungal communities
50	using 454-pyrosequencing, quantified the proportion of root length colonised by AM
51	fungi using microscopy, and estimated AM fungal biomass using a lipid biomarker.
52	
53	• AM fungal community composition was altered by litter removal but not litter addition.
54	Root colonisation was substantially greater in the superficial organic layer compared
55	to the mineral soil. Overall colonisation was lower in the litter removal treatment,
56	which lacked an organic layer. There was no effect of litter manipulation on the
57	concentration of the AM fungal lipid biomarker in the mineral soil.
58	
59	• We hypothesise that reductions in organic matter brought about by litter removal may
60	lead to AM fungi obtaining nutrients from recalcitrant organic or mineral sources in
61	the soil, besides increasing fungal competition for progressively limited resources.
62	
63	
64	INTRODUCTION
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66	The productivity of most tropical forests is sustained by symbiotic associations between
67	plants and arbuscular mycorrhizal (AM) fungi (Read, 1991; Alexander & Lee, 2005). AM
68	fungi play crucial roles in nutrient cycling and are also major vectors of carbon (C) in the
69	global C cycle (Johnson et al., 2013). AM fungi obtain up to 20-30% of total plant

photosynthates (Drigo *et al.*, 2010) and may enhance the decomposition of organic matter,
releasing substantial quantities of CO<sub>2</sub> to the atmosphere through their respiration
(Nottingham *et al.*, 2010).

73

Tropical forest growth currently constitutes the largest terrestrial sink for anthropogenic CO<sub>2</sub> (Oren *et al.*, 2001) and thus makes a substantial contribution to the regulation of the global climate system (Field *et al.*, 1998). Anticipating future effects of anthropogenic change on tropical forests demands a clearer understanding of how nutrient availability limits forest productivity, and the roles of AM fungi in complex scenarios of nutrient limitation and colimitation. Nonetheless, AM fungi are under-investigated in tropical systems in general, and tropical forests in particular (Alexander & Selosse, 2009).

81

It is widely hypothesised that the symbiotic function of AM fungi is determined by the 82 relative availability of C, nitrogen (N), and phosphorus (P; Johnson, 2010; Johnson et al., 83 2013). This is based on evidence which shows that fertilisation with N and P can reduce AM 84 fungal colonisation of roots (Johnson et al., 2003), and that the relative amounts of N and P 85 determine mycorrhizal symbiotic function (Johnson, 2010). In some cases this may cause 86 87 AM fungi to behave less mutualistically (Johnson, 1993); where neither N or P is limited, the only limitation to fungal growth is the supply of plant C, meaning that fungal C demand can 88 increase to the point where plant growth is depressed (Johnson, 2010). 89

90

Much current understanding concerning the function of AM fungal symbioses comes from 91 studies that explore how variation in nutrient availability affects AM fungal characteristics 92 (eg. Treseder, 2004; Wurzburger & Wright, 2015). Amongst these, nutrient addition 93 experiments are one of the most widely used approaches, particularly in field settings 94 95 (Treseder, 2004). Nutrient addition is hypothesised to affect AM fungi either directly, by alleviating fungal nutrient limitation and thereby stimulating fungal growth (Treseder & 96 Allen, 2002), or indirectly, by causing plants to reduce investment of carbohydrate in their 97 AM fungal partners (Mosse & Phillips, 1971; Johnson, 2010). 98

99

100 Besides altering AM fungal biomass, nutrient addition may affect AM fungal community

101 composition and diversity. Changes in community composition and diversity are likely to

arise from differences in the functional properties of AM fungal taxa and their ability to

103 compete with other fungi (AM or saprobe) for key resources (Hart & Reader, 2002; Maherali

& Klironomos, 2007; Powell et al., 2009). For instance, different AM fungal taxa can vary in 104 the translocation of P (Ravnskov & Jakobsen, 1995) or N (Veresoglou et al., 2012) to plant 105 partners, carbon storage and demand (Pearson & Jakobsen, 1993), relative allocation to intra-106 and extra-radical biomass (Hart & Reader, 2002), and growth and life-history strategy (Hart 107 & Reader, 2002; Maherali & Klironomos, 2007; Powell et al., 2009). Furthermore, plant-AM 108 fungal combinations perform differently in alternative settings, with wide range of symbiotic 109 outcomes (Klironomos, 2003; Powell et al., 2009). Consequently, the advantage of AM 110 fungal associations will vary according to the prevailing conditions and the ecological niche 111 112 of the fungal partner. Evaluation of community parameters thus provide important information to supplement the aggregate metrics of root colonisation and concentration of the 113 AM fungal biomarker lipid (a proxy for AM fungal biomass), which cannot distinguish 114 between members of the AM fungal community. 115

116

In addition, AM fungal species that share a common evolutionary history may also share 117 traits and ecological functions (Maherali & Klironomos, 2007; Powell et al., 2009), and 118 community data can thus be used to infer the ecological processes structuring AM fungal 119 communities. Phylogenetically over-dispersed communities (communities consisting of taxa 120 121 that are less related to each other than expected by chance) are hypothesised to be structured by competition, preventing closely related and functionally similar taxa (those sharing a 122 common niche) from co-occuring. By contrast, phylogenetically under-dispersed (or 123 clustered) communities are hypothesised to be structured by habitat filters; features of the 124 environment that permit only the co-occurence of species with specific traits or ecological 125 tolerances, and which can cause taxa with similar traits to respond in similar ways to 126 environmental pressures (Webb et al., 2002; Maherali & Klironomos, 2007). 127

128

The great majority of nutrient addition studies apply inorganic fertilisers (eg. see Treseder, 129 2004). These studies are useful in highlighting the roles of individual nutrients and simulating 130 the effects of inorganic nutrient deposition. However, fertilisation treatments are artificial and 131 do not mimic pathways of nutrient cycling under natural conditions (Sayer & Banin, 2016). 132 Furthermore, the regulation of plant-AM fungal relations is strongly dependent on the relative 133 availability of different nutrients (Treseder & Allen, 2002; Johnson, 2010), whereas the 134 addition of large quantities of one or more inorganic nutrients (e.g. N, P, K) strongly distorts 135 stoichiometric relationships, and largely neglects the role of organic matter in nutrient cycling 136 (Sayer & Banin, 2016). 137

138

Under natural conditions, nutrient cycling in forests occurs largely through litterfall, root 139 death, root exudates, decomposition, and the growth and death of microorganisms (Attiwill & 140 Adams, 1993; Leff et al., 2012). It is via these processes that the regulatory processes 141 governing plant-AM fungal exchange have evolved. Indeed, over large latitudinal gradients 142 there is a strong relationship between leaf litter quality, the organic matter resulting from its 143 degradation, and the predominant mycorrhizal type in a given bioregion (Read, 1991). 144 Nonetheless, there have been few experimental investigations into the effects of leaf litter 145 146 amendments on AM fungi in highly diverse tropical forests.

147

Although multiple lines of evidence suggest a key role for AM fungi in cycling nutrients via 148 organic sources, the majority of studies investigating the effects of organic amendments on 149 AM fungi have been conducted in experimental microcosms, and most have examined 150 changes in biomass rather than community parameters (Hodge, 2014). These experiments 151 demonstrate that AM fungal hyphae preferentially proliferate in organic substrates in 152 experimental microcosms (Hodge & Fitter, 2010), are able to capture N from organic 153 substrates (Leigh et al., 2009), and can enhance the decomposition of organic material 154 155 (Hodge, 2014). The few existing field studies show that organic matter additions in agricultural systems tend to increase AM fungal colonisation of plant roots and hyphal 156 abundance in soils (Gryndler et al., 2005; Gosling et al., 2010). Furthermore, AM fungal 157 hyphae can grow into decomposing leaf litter on tropical forest floors (Herrera et al., 1978; 158 Posada et al., 2012; Camenzind & Rillig, 2013). Together, these studies strongly suggest that 159 AM fungal hyphae are important in recycling nutrients from leaf litter. This is likely due to 160 tightly coupled interactions between AM fungi and saprophytic fungi and bacteria (Herman 161 et al., 2012) given that AM fungi have not been shown to possess saprophytic capabilities 162 (Hodge, 2014). 163

164

We investigated AM fungal responses to altered organic matter inputs in a lowland tropical forest in Panama using an existing long-term litter manipulation experiment in which nine years of litter removal and addition treatments have altered fine root biomass (Sayer *et al.*, 2006a), litter production, foliar and litter nutrient concentrations, and soil nutrient pools (Vincent *et al.*, 2010; Sayer & Tanner, 2010b). This platform provided a unique opportunity to evaluate the responses of AM fungal communities to changes in organic matter inputs in a well-studied lowland tropical forest setting. 172

We hypothesised that: i) litter addition would increase net AM fungal abundance, given the 173 well-documented stimulatory effects of organic matter additions on AM fungal growth, ii) 174 litter removal would also increase net AM fungal abundance, given that plants may increase 175 investment in AM fungi when nutrient availability is reduced (Johnson, 2010), iii) that the 176 addition or removal of organic matter would result in changes in the AM fungal community 177 composition, and iv) that litter manipulation would alter the ecological processes structuring 178 AM fungal communities, and that this would be reflected in changes in the degree of 179 180 relatedness (or phylogenetic structure), of AM fungal communities.

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## 183 MATERIALS AND METHODS

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# 185 Site description and experimental design

186The Gigante Litter Manipulation Experiment (GLMP) is located on the Gigante Peninsula

187 (9°06' N, 79°54' W) within the Barro Colorado Nature Monument (BCNM) in Panama,

188 Central America. Nearby Barro Colorado Island (BCI; c. 5 km from the study site) has a

mean annual rainfall of 2600 mm, with a strong dry season between January and April and a

<sup>190</sup> mean annual temperature of 27  $^{\circ}$ C (Leigh, 1999). Tree species composition and canopy

height are characteristic of mature (>200 year old) secondary forest (Wright *et al.*, 2011) and

the soils are classed as moderately acidic Oxisols (Dieter et al., 2010; Turner & Wright,

- 193 2013), with low concentrations of available P and moderate concentrations of base cations
- 194 (Turner et al., 2013). The GLMP consists of fifteen 45 m  $\times$  45 m plots; starting in 2003, leaf
- litter from five plots was raked up once a month (litter removal treatment; L-), immediately

added to five plots where it was distributed as evenly as possible (litter addition treatment;

L+), and five plots were left undisturbed as controls (C; see Sayer & Tanner 2010 for details).

#### 199 Sampling

In May 2012, after nine years of treatments, we sampled at six points in the inner 30 m x 30

201 m of each of the 15 experimental plots (a total of 30 samples per treatment); we selected

sampling points at random using random number sheets to delineate point coordinates, with

- the provision that all points were separated by at least 3 m. At each sampling point, we
- collected the litter (Oi) and fermentation (Oe) horizons from a 78.5 cm<sup>2</sup> area, using a knife to
- 205 cut around the edge of a metal disk (C and L+ treatments only; the L- treatment lacked an

organic horizon), and two cores from the mineral soil (0-10 cm depth) using a 5-cm diameter 206 corer (all treatments). To prevent cross-contamination, we wiped down and flame-sterilised 207 all equipment in between samples, handled all samples with fresh latex gloves, and double-208 bagged samples in sealed Ziploc<sup>TM</sup> bags. All samples were stored at 4°C and processed 209 within 36 hours of returning from the field. Root samples were obtained from one of the two 210 cores per sampling point by washing away soil and organic matter under a continuous stream 211 of filtered water over a sieve with a mesh size of  $500 \,\mu$ m. We retained fine roots (< 1 mm in 212 diameter) for further analysis, drying a subsample over silica gel for DNA extraction, and 213 storing a second subsample in 70% ethanol for microscopic analysis. The remaining soil 214 cores were sieved to remove stones and roots, composited to make one sample per plot, and 215 thoroughly homogenised. 20 g subsamples for lipid analysis were frozen at -80°C for 12 h, 216 lyophilised, and stored dry at -80°C until further processing. 217

218

Prior to lipid and nutrient extractions, an equal mass of each sample was pooled to make one
composite sample per plot (a total of 15 samples). Prior to DNA extraction, the six root
samples per plot were individually pulverised in a homogeniser (TissueLyser II, Qiagen), and
an equal mass of each sample was pooled to make one composite sample per plot (a total of
15 samples). Microscopic analysis of root samples was performed on individual samples
(total = 6 samples per plot, 90 samples in total).

225

#### 226 AM fungal abundance

We used the percentage of root length colonised as a measure of intra-radical AM fungal 227 abundance (McGonigle et al., 1990). We soaked and rinsed the root samples with distilled 228 water to remove the ethanol. Roots were then cleared by autoclaving in 5% KOH for 5-60 229 minutes; bleached in solution of ammonia in 3% H<sub>2</sub>O<sub>2</sub> for 15-60; acidified in 2% HCl for 30 230 minutes; and stained with 0.05% trypan blue (in a 1:1:1 solution of distilled water, glycerol 231 and lactic acid) for 20 minutes at 60°C. The optimum clearing and bleaching time varied 232 depending on the thickness and pigmentation of the roots. We quantified AM fungal 233 colonisation by hyphae, vesicles and arbuscules using a compound light microscope at  $200 \times$ 234 magnification, according to the method of McGonigle et al. 1990 with at least 100 235 intersections per sample, and one sample per core. AM fungal colonisation was expressed as 236 the percentage fine root length colonised by AM fungal hyphae, vesicles or arbuscules. 237

- 239 We used the neutral lipid fatty acid (NLFA)  $16:1\omega 5$  as a biomarker for extra-radical AM
- <sup>240</sup> fungal biomass. We performed lipid extraction and analysis according to Frostegård et al.
- 241 (1993) with modifications (Nilsson *et al.*, 2007). Briefly, lipids extracted from 4 g lyophilised
- soil per plot were fractionated into neutral lipids, glycolipids, and polar lipids on silica
- columns by successive elution with chloroform, acetone and methanol. Methyl
- nonadecanoate (FAME 19:0) was added as an internal standard, and neutral and polar
- fractions were converted to fatty acid methyl esters (FAMEs) prior to analysis on a gas
- chromatograph with a flame ionisation detector and a 50 m HP5 capillary column (Hewlett
- Packard, Wilmington, DE, USA). The mean NLFA to PLFA ratio across all samples was 1.3,
- suggesting that NLFA 16:1ω5 is an effective AM fungal biomarker in these soils (Olsson,
  1999).
- 250

#### 251 Soil chemistry

- Measurement of inorganic N (NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup>), resin-extractable P, organic P, and pH was 252 performed as described in Turner et al. (2013). Analysis of total N and C was performed on 253 air-dried soils by automated combustion and gas chromatography on a Thermo Flash EA1112 254 analyzer (CE Elantech, New Jersey, USA). Organic P was extracted in a mixture of 0.25 M 255 NaOH and 0.05 M EDTA, and analysed as described by Turner et al. 2008. Exchangeable 256 cations were extracted in 0.1 M BaCl<sub>2</sub>, with detection by ICP-OES (Hendershot et al. 2008), 257 and effective base saturation (EBS) was calculated by dividing the cmol of positive charge 258 per kg dry soil of exchangeable bases (Ca + K + Mg + Na) by that of the total cations (Al + 259
- 260 261

#### 262 **DNA extraction and sequencing**

Ca + Fe + K + Mg + Mn + Na; Hendershot et al. 2008).

We extracted DNA from 50 mg of pulverised root using MoBio PowerPlant DNA isolation
kits according to the manufacturer's instructions (MoBio Laboratories Inc., Carlsbad, CA,
USA).

266

We amplified the partial small subunit (SSU) region of 18S ribosomal DNA (*c*. 550 bp) with the universal eukaryotic primer NS31 (Simon *et al.*, 1992) and the AM fungal-specific primer AM1, which amplifies the major families of the Glomeromycota (Helgason *et al.*, 1998). We chose this primer set because it is widely represented in sequence databases, and because we wanted to facilitate comparisons with previous work using these primers. In addition, these primers have been demonstrated to have extremely low PCR bias against artificially

- assembled community templates (Cotton *et al.*, 2014). Prior to amplification, the primers
- were modified by the addition of the 454 pyrosequencing adaptors A and B, in addition to a
- 10 bp multiplex identifier (MID) on the forward primer (NS31). We conducted duplicate
- polymerase chain reactions (PCRs) in 25 µl sample volume using Phire hot start II DNA
- polymerase (Life Technologies LTD, Paisley, UK). Conditions were: 98°C for 1 minute; 32
- cycles of  $98^{\circ}$ C for 10 s and  $72^{\circ}$ C for 15 s; and a final extension phase of  $72^{\circ}$ C for 2 minutes.
- 279
- 280 We gel-purified the PCR products using MinElute PCR purification kits (Qiagen Ltd, West
- Sussex, UK) and pooled the samples in equimolar concentrations, evaluating the
- concentration of DNA in the cleaned PCR products using Quant-iT PicoGreen dsDNA Assay
- 283 Kit (Invitrogen, Life Technologies LTD, Paisley, UK). Amplicon libraries were distributed
- on PicoTiter Plates and sequenced on an FLX Titanium system using Lib-L shotgun
- chemistry (Roche, Basel, Switzerland). No sequences were detected in the blanks included as
- negative controls at each of the extraction, PCR, gel purification, and quantification steps.
- 287

## 288 Bioinformatic analysis

- All bioinformatic analyses were performed using the software mothur (Schloss *et al.*, 2009) 289 290 unless otherwise stated. Sequence filtering was performed with the sff.multiple quality filtering protocol. Reads were removed from the dataset if they did not contain the 10 bp 291 292 MID, had > 1 error in the barcode sequence, > 2 errors in the forward primer, or were shorter than 200 bp in length. After quality filtering and removal of barcode and primer sequences, 293 clustering was performed using the algorithm Clustering 16S rRNA for Operational 294 Taxonomic Unit (OTU) Prediction (CROP), an unsupervised Bayesian clustering method that 295 forms clusters based on the organisation of sequences without setting a hard similarity cutoff 296 (Hao *et al.*, 2011). To provide finer taxonomic resolution, we set the *i* and *u* parameters to 2% 297 cluster difference rather than the conventional 3% because the SSU region has relatively low 298 variation (Öpik et al., 2013; Davison et al., 2015). The centre sequence from each cluster was 299 used as a representative sequence in subsequent analyses. 300
- 301

302 Sequence alignment was performed with the software MAFFT v7.149b (Katoh *et al.*, 2002)

- 303 using the L-INS-i algorithm (iterative refinement using local pairwise alignment) and the
- alignment from Krüger et al. (2012) as a backbone. Alignments were improved with
- 305 MUSCLE (Edgar, 2004) using the –refine option. Trees were built using RAxML v. 8.0

- 306 (Stamatakis, 2014) with GTR GAMMA implementation, and bootstrap values based on 1000
   307 runs.
- 308
- 309 We used the Basic Local Alignment Search Tool (BLAST, Altschul et al., 1990; minimum e-
- value  $10^{-30}$ ) on one representative sequence from each cluster iteratively against three
- databases in the following order of preference: i) sequences from Krüger et al. (2012); ii) all
- 312 virtual taxa (VT) from the MaarjAM AM fungal sequence database
- 313 (www.maarjam.botany.ut.ee); and iii) all 18S Glomeromycotan sequences from SILVA
- database. Non-Glomeromycotan clusters were removed when the highest blast match did not
- correspond to an AM fungal sequence in any of the three datasets.
- 316
- 317 Clusters were named based on matches to database entries at > 97% covering a minimum of
- <sup>318</sup> 80% of the query sequence. We used the generic names from Krüger et al. (2012), and VT
- numbers from the MaarjAM database. Where clusters did not match a VT at > 97% we
- assigned a name based on the highest VT match and phylogeny (eg. Glomus\_OTU1). We
- fused clusters based on matches to database sequences > 97% and the tree topology obtained
- from RaXML. Clusters that occurred in < 2 samples, and with < 5 reads total were removed
- from the dataset. Raw sequence data were deposited in the International Nucleotide Sequence
- 324 Database Sequence Read Archive (accession no. SRP076949).
- 325

#### 326 Statistical analysis

- All statistical analyses were conducted in R version 3.1.2 (R Development Core Team, 2014).
- 329 Multivariate analysis of AM fungal communities
- 330 We accounted for variation in the number of sequences between samples by using a variance
- stabilising (VS) transformation of the OTU table, implemented with the DESeq2 package
- (Love *et al.*, 2014), according to McMurdie and Holmes (2014). This approach avoids the
- need for rarefying, which can result in data that misrepresent the original community
- 334 (McMurdie & Holmes, 2014). All subsequent analysis was performed on the VS transformed
- OTU table, using the copy number of DNA sequences as a measure of relative abundance of each OTU.
- 337

338 To examine the effect of litter manipulation on AM fungal community composition, we used

- 339 multivariate generalised linear models (M-GLMs) with negative binomial error structures
- using the mvabund package (Wang *et al.*, 2012). M-GLMs provide a more robust way to

analyse multivariate community data than do distance-based approaches such as

342 PERMANOVA (Warton *et al.*, 2015). We ascertained the degree to which individual OTUs

were affected by litter manipulation using DESeq2 (Anders & Huber, 2010), which estimates

the effect size (as logarithmic fold change) and reports *P*-values adjusted for multiple

- 345 comparisons.
- 346

To visualise differences in AM fungal communities across litter manipulation treatments we used non-metric multidimensional scaling (NMDS) ordination, using the metaMDS function in the vegan package (Anderson 2001, Oksanen et al. 2010). Ordination was based on Bray-Curtis dissimilarity calculated from square-root transformed abundances. The range of data values was large, and a square root transformation was applied to improve the quality of the ordination by reducing the weighting of the most abundant OTUs (Legendre & Legendre, 2012; Oksanen *et al.*, 2010).

354

Soil physical characteristics were standardised to zero mean and unit variance, and fit to the NMDS ordinations (function envfit from the vegan package) with significance ascertained using 9999 permutations. Individual values of exchangeable cations were collapsed into the metric of effective base saturation (EBS). Organic phosphorus correlated closely with resinextractable phosphorus ( $r^2 > 0.7$ ) and was omitted, since resin-extractable phosphorus better approximates the plant-available phosphorus fraction (Condit *et al.*, 2013).

361

## 362 *Community phylogenetic structure*

We asked whether litter manipulation altered the degree of relatedness between taxa in AM 363 fungal communities. We used two indices of community phylogenetic structure: Net 364 Relatedness Index (NRI) and Nearest Taxa Index (NTI; (Webb, 2000). Positive values of 365 these metrics indicate that taxa in a community are on average more closely related to each 366 other than to members of the regional taxon pool (phylogenetically clustered), and negative 367 values indicate that taxa in a community are less closely related (phylogenetically over-368 dispersed). NRI is sensitive to tree-wide phylogenetic patterns, and NTI is sensitive to 369 phylogenetic community patterns close to the tips of the phylogeny. Observed values of these 370 metrics were compared to 10,000 null communities generated using the 'independentswap' 371

algorithm, which maintains column and row totals and accounts for differences in community
richness and taxon prevalence (Gotelli, 2000). Statistical significance of phylogenetic
structure was ascertained using a two-tailed *t*-test. Community phylogenetic analysis was

performed using the picante package (Kembel & Ackerly, 2010).

376

Univariate analysis of AM fungal abundance and diversity, and soil physical characteristics $We analysed the effects of litter manipulation on the concentrations of NLFA 16:1<math>\omega$ 5 in the soil, AM fungal colonisation of plant roots, AM fungal OTU richness and predominance, and metrics of phylogenetic community structure (NRI and NTI) using linear models having confirmed that all variables met the assumptions. Where the main effect of litter manipulation was significant, we performed Dunnett's *post-hoc* analysis to compare each treatment with the controls.

384

To ascertain whether AM fungal colonisation of roots was greater in the mineral soil or organic layer we built linear mixed effects models (using the lme4 package; Bates et al. 2014). Models included 'layer' and 'treatment' as fixed effects, and 'plot' as a random effect. The significance of fixed effects was assessed by comparing nested models using parametric bootstrapping with 10000 simulations, using the PBmodcomp function from the pbkrtest package (Halekoh & Højsgaard, 2014). Results are reported as significant at  $\alpha < 0.05$ .

391

## 392 **RESULTS**

393

## 394 Soil chemistry

Soil nutrients were lower in litter removal compared to litter addition treatments for inorganic N; resin and organic P, pH, and extractable Ca, Mg, and Mn (K was not significantly lower).
Compared to the controls, the soils in the L- plots had lower concentrations of inorganic N, resin and organic P, Ca, and Mg, whereas soils in the L+ plots had higher concentrations of resin P, and Ca (Figure 1, Table S1). A full discussion of the effects of litter manipulation on soil chemistry is provided in Sayer & Tanner (2010) and Sayer *et al.* (2012).

401

#### 402 AM fungal abundance

There was no significant effect of litter manipulation on the proportion of root length
colonised by any AM fungal structure in the mineral soil (total colonization, hyphae, vesicles
or arbuscules), although for each of the structures there was a trend towards higher root

- colonisation in both litter removal and litter addition treatments compared to the controls 406 (Figure 2; total colonisation:  $F_{2,12} = 1.7$ , P = 0.23; hyphae:  $F_{2,12} = 1.4$ , P = 0.29; vesicles: 407  $F_{2,12} = 2.5$ , P = 0.13; arbuscules:  $F_{2,12} = 1.3$ , P = 0.31). In the control and litter addition plots, 408 the proportion of root length colonised by all AM fungal structures was substantially greater 409 in the superficial organic layer than in the mineral soil (significant 'layer' term; hyphae: 410 likelihood-ratio test (LRT) = 50.0 P < 0.001; vesicles: LRT = 19.6, P < 0.001; arbuscules: 411 LRT = 28.6, P < 0.001; all structures: LRT = 51.6, P < 0.001; Figure 2). Because root 412 colonization was highest in the superficial organic layer, the overall abundance of AM fungi 413 414 was lower in the litter removal treatment, which lacked this layer.
- 415

There was no effect of litter manipulation on AM fungal biomass in the mineral soil

(concentration of NLFA 16:1 $\omega$ 5; Figure S1), nor was AM fungal biomass correlated with any of the measured soil variables (soil pH:  $F_{1,13} < 0.001$ , P = 0.98; effective base saturation:  $F_{1,13} = 0.01$ , P = 0.92, resin-extractable phosphorus:  $F_{1,13} = 0.12$ , P = 0.74; and inorganic nitrogen:  $F_{1,13} = 0.54$ , P = 0.48).

421

#### 422 AM fungal community composition and structure

Four AM fungal families were represented in the sequencing dataset (Acaulosporaceae, 423 Archaeosporaceae, Gigasporaceae, Glomeraceae; Figure 3), indicating reasonable taxonomic 424 coverage of the Glomeromycota (based on the classification of Redecker et al. 2013). No 425 members of the Diversisporaceae, Paraglomeraceae, Geosiphonaceae, Ambisporaceae, 426 Claroideosporaceae or Pacisporaceae were detected. Rarefaction curves for each sample 427 indicated that sequencing intensity was sufficiently high to detect the majority of OTUs. 428 Rarefaction curves pooled by experimental treatment approached asymptotes, indicating that 429 sampling effort was sufficient to capture the range of AM fungal taxa across the sites (Figure 430 S2). A total of 10,197 sequences were retained after quality control, clustered into 72 OTUs, 431 and 95.9% of all sequences matched Glomeromycota in the databases. Fifty-six OTUs 432 remained after blasting, filtering, merging, and trimming (exclusion of OTUs with a total of 5 433 or less reads), representing a total of 8825 sequences. Each sample (1 per plot) contained a 434 mean of 18 OTUs (range: 11-24), and the mean number of sequences per sample was 588 435 (range: 237-1225; Table S2). A phylogenetic tree is provided in Figure S3. 436 437 Overall AM fungal community composition was altered by litter removal but was not 438

- 439 significantly affected by litter addition (Multivariate GLM: Wald  $_{2,12} = 11.5$ , P < 0.003;
- 13

treatment contrast for litter removal: Wald = 9.2, P < 0.003 and for litter addition: Wald = 440 5.9, P = 0.24; Figure 4). There were no significant differences among treatments when the 441 analysis was repeated at levels of genus and family (multivariate GLM; genus: Wald  $_{2,12}$  = 442 3.9, P = 0.24; family: Wald <sub>2.12</sub> = 2.1, P = 0.66; Figure 3). In the analysis of individual OTUs 443 (using the DESeq2 package), litter removal significantly (P < 0.05) reduced the relative 444 abundance of four OTUs and increased the relative abundance of three OTUs (P < 0.05; 445 Figure 5). By contrast, litter addition significantly increased the relative abundance of two 446 OTUs (Figure 5; Table S3). All of the significantly affected OTUs were in the family 447 Glomeraceae apart from a single OTU in the Acaulosporaceae, which had lower relative 448 abundance in the litter removal treatment. Neither litter treatment altered the total number of 449 AM fungal OTUs (richness; ANOVA:  $F_{2,12} = 0.15$ , P = 0.86), nor the proportional 450 abundance of the dominant AM fungal taxon (predominance; ANOVA:  $F_{2,12} = 0.37$ , P =451 0.69; Figure S4). Of the variables fitted to the NMDS ordination, soil pH, effective base 452 saturation (EBS), resin-extractable P, and inorganic N concentrations were significantly 453 correlated with AM fungal community composition (Figure 4). 454

455

#### 456 AM fungal community assembly

Litter manipulation moderately altered the degree of relatedness between taxa in AM fungal 457 communities, as summarised by the Net Relatedness Index (NRI). There was a greater 458 likelihood of detecting closely related taxa in litter addition plots than litter removal plots 459 (ANOVA:  $F_{2.12} = 4.02$ , P = 0.05; Figure 6), although neither treatment differed significantly 460 from controls. Furthermore, whereas neither treatment showed significant phylogenetic 461 structure of AM fungal communities relative to null model distributions, the NRI was >0 in 462 the litter addition treatment and <0 in the litter removal treatment, indicating a trend towards 463 phylogenetic under-dispersion (taxa more related to each other than expected by chance) in 464 the litter addition treatment (Figure 6) and phylogenetic over-dispersion (taxa less related to 465 each other than expected by chance) in the litter removal treatment. However, when we used 466 the Nearest Taxa Index (NTI), which is sensitive to patterns in relatedness close to the tips of 467 the phylogeny, AM fungal communities were neither significantly structured relative to null 468 distributions, nor affected by litter manipulation (ANOVA:  $F_{2,12} = 0.25$ , P = 0.79). 469

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#### 472 **DISCUSSION**

Litter removal altered AM fungal community composition (Figure 4), indicating that inputs 474 of organic matter are important in structuring AM fungal communities. Together with 475 substantially greater AM fungal root colonisation in the superficial organic layer than the 476 mineral soil (70% versus 30% respectively; Figure 2e-h), our findings suggest that AM fungi 477 obtain a substantial part of their nutrition from decomposing organic matter in this lowland 478 tropical forest. We observed a trend towards increased AM fungal colonisation of roots 479 growing in the mineral soil both in litter addition and litter removal treatments relative to 480 controls (Figure 2), providing some support for our hypotheses that plants may increase 481 482 investment in AM fungal associations in both litter addition and litter removal treatments. 483

#### 484 Litter removal

Litter removal may have altered AM fungal community composition by reducing N-485 availability, either via direct fungal N-limitation, or by altering plant N status, leading to 486 changes in plant allocation to AM fungi. Litter removal reduced the amount and availability 487 of soil inorganic N, and crucially, reduced N concentrations in leaf litter after five years 488 (Sayer & Tanner, 2010b; Sayer et al., 2012), suggesting that N- availability to plants had 489 decreased. N concentrations of AM fungal hyphae are substantially higher than that of plant 490 491 tissues (Hodge et al., 2010) and comparison of the C:N ratios of plant and fungal tissues indicate that severe N-limitation may be more likely to suppress fungal growth than plant 492 growth (Kaye & Hart, 1997; Johnson, 2010). Given that different AM fungal taxa are known 493 to vary in growth strategy and biomass allocation, and vary in the translocation of N to plant 494 partners (Veresoglou et al., 2012), litter removal may have selected for low-N AM fungal 495 specialists. Alternatively, litter removal may have altered AM fungal community composition 496 by increasing AM fungal competition (both with other AM fungi and saprobes) for a more 497 limited resource. 498

499

An alternative possibility is that changes in AM fungal community composition in the litter 500 removal plots reflect niche separation arising from a shift in AM fungal P-acquisition 501 strategies. The availability of P is thought to limit many biological processes in lowland 502 tropical forests (Vitousek & Sanford, 1986), and is a limiting nutrient in these forests (Wright 503 et al., 2011; Turner & Wright, 2013). As a large proportion of the P required for plant growth 504 is cycled through leaf litter (Sayer & Tanner, 2010b), we would expect the litter removal 505 treatment to affect plant P status. However, there was no reduction in leaf litter P in litter 506 removal plots, nor a reduction in litterfall or plant productivity in the first 6 years of litter 507

manipulation (Sayer & Tanner, 2010b; Sayer *et al.*, 2012), indicating that trees in the litter
removal plots were able to access sufficient P from alternative sources to maintain
productivity and foliar P concentrations.

511

At least some of the additional P available to plants in the litter removal treatment was 512 probably acquired from stable organic P pools in the mineral soil. Organic P in forests occurs 513 in fresh organic matter (such as leaf litter), microbial biomass, and non-biomass stable 514 organic phosphorus (Vincent et al., 2010). Under normal conditions, P is rapidly released 515 516 from leaf litter via leaching (Schreeg et al., 2013) or mineralisation (Richardson & Simpson, 2011) before being taken up directly by plants by mycorrhizal fungi (Herrera et al., 1978). 517 This results in 'direct' nutrient cycling by which nutrient losses through leaching might be 518 minimised (Went & Stark, 1968). After three years of litter removal, the stable organic P pool 519 in the upper 2 cm of the mineral soil was reduced by 23%, while the overall inorganic P pool 520 remained unchanged (Vincent et al., 2010). Given that our study took place after nine years 521 of litter removal, and the depletion of the stable organic P pool had conceivably continued, it 522 is probable that additional P could also have been mobilised from recalcitrant mineral P 523 stocks in the soil. 524

525

The role of AM fungi in P acquisition is well-known (Smith & Read, 2008), and is likely that 526 a shift in plants' primary source of P from decomposing litter to stable organic P and stocks 527 of mineral P would involve a change in the primary function of plants' AM fungal 528 associations. Although limited, there is evidence that different AM fungal species differ in 529 their ability to acquire (Cavagnaro et al., 2005) and transport P to plant hosts (Munkvold et 530 al., 2004), and that AM fungal taxa may benefit plants to different degrees based on the type 531 of soil P available (eg. mineral versus organic; Reynolds et al., 2005). Consequently, it is 532 possible that the taxa with increased relative abundance in litter removal plots were mineral P 533 specialists, and those with decreased relative abundance were litter specialists (Figure 5). 534 Nonetheless, it is striking that the dominant taxon and the relative abundances of most taxa in 535 the litter removal treatment remained unchanged. Given the probable shift in plants' primary 536 P source in the litter removal treatment, this would suggest that most of the AM fungal taxa 537 observed at this site are readily able to adapt to the changed conditions. This is interesting in 538 the light of studies of ectomycorrhizal fungi, which document wide differences in the ability 539 of different taxa to mobilise and acquire P from different sources (Plassard et al., 2011). 540

Other factors besides changes in nutrient availability could explain the shift in community 542 composition observed in the litter removal treatment. Organic amendments such as leaf litter 543 can affect a number of other soil properties besides nutrient availability, such as habitat space 544 available for decomposers (Sayer, 2006). It is thus possible that AM fungal communities 545 were affected by changes in the non-AM microbial community or soil fauna, which can 546 impact AM fungal growth and function (Johnson et al., 2005; Sayer et al., 2006b; Gryndler et 547 al., 2008; Hodge, 2014), and which play a key role in AM fungal uptake of nutrients from 548 leaf litter given the lack of documented saprophytic effects of AM fungi (Hodge, 2014). 549 550 Previous studies at this site show no major changes in either temperature or soil water content among treatments (Sayer & Tanner, 2010a), and it is thus unlikely that these factors are 551 responsible for the observed effects. 552

553

Soil pH was correlated with the NMDS ordinations of AM fungal community shifts, and may have been responsible for some of the observed shifts in community composition. However, studies documenting the effects of pH on AM fungi have largely reported a reduction in root colonisation and extra-radical hyphal biomass with decreasing pH (Wang *et al.*, 1993; Clark, 1997; van Aarle *et al.*, 2002) as well as reduced AM fungal  $\beta$  diversity (Dumbrell *et al.*, 2009), none of which were observed in this study.

560

Regardless of the mechanism underlying the shifts in AM fungal community composition, 561 the trend towards more phylogenetically over-dispersed (less closely related) AM fungal 562 communities in the litter removal plots relative to the litter addition plots (Figure 6) may 563 reflect increasing competition between AM fungal taxa following litter removal. This is 564 because more closely related AM fungal taxa tend to share functional traits (Maherali & 565 Klironomos, 2007; Powell et al., 2009), a phenomenon known as phylogenetic trait 566 conservatism (Webb et al., 2002). Consequently, phylogenetically over-dispersed 567 communities are thought to be structured more by competition than by habitat filtering, 568 which reduces the likelihood that closely related and functionally similar taxa will co-occur 569 (Webb *et al.*, 2002). 570

571

## 572 Litter addition

AM fungal colonisation of roots was substantially higher in the organic horizons than the

- 574 mineral soil in the control and litter addition treatments (70% versus 30% respectively;
- 575 Figure 2e-h). This finding agrees with a sizeable body of evidence which shows that the

addition of organic material may increase AM fungal colonisation of plant roots (Gryndler et 576 al., 2005; 2008; Gosling et al., 2010), and AM fungal sporulation (Gosling et al., 2010). 577 Indeed, AM fungal hyphae proliferate in organic substrates (Hodge & Fitter, 2010), and grow 578 into decomposing leaf litter in tropical forests (Herrera et al., 1978; Posada et al., 2012; 579 Camenzind & Rillig, 2013). Together with the finding that fine roots proliferated into the 580 organic horizons in the litter addition treatment (Sayer et al., 2006a), our results suggest that 581 AM fungi may represent important pathways for plant uptake of nutrients from sites of 582 organic matter decomposition in this tropical forest. However, given that AM fungi lack 583 584 substantial saprophytic capability (Hodge, 2014), it is unlikely that AM fungi themselves are actively involved in litter decomposition, but rather are able to efficiently acquire nutrients as 585 they are released from decomposing organic matter by the action of saprobes. 586

587

Given much greater root colonisation by AM fungi in the organic horizons of the litter 588 addition and control plots relative to the mineral soil, it is surprising that we observed no 589 significant increase in root colonisation in the mineral soil of litter addition treatments 590 relative to controls (Figure 2a-d), where organic matter content is elevated relative to controls 591 (Tanner et al., 2016). It is possible that plant investment in AM fungi in litter addition plots is 592 593 lower, due to the increases in soil fertility and tree nutrient status (indicated by marginal increases in litterfall and foliar N and P; Figure 1, Table 2; (Sayer & Tanner, 2010b; Sayer et 594 al., 2012). This interpretation follows from the functional equilibrium hypothesis, by which 595 plants allocate resources to the structures that are the most helpful in acquiring the most 596 limiting nutrients (Johnson, 2010), and by which plants should reduce investment in AM 597 fungal associations when soil fertility increases because the carbon costs outweigh the 598 nutritional benefits (Mosse & Phillips, 1971; Johnson, 2010). Reduced plant investment in 599 AM fungi would counter the stimulatory effects of organic matter on AM fungal 600 colonisation. 601

602

#### 603 Limitations of this study

We did not measure NLFA in the superficial organic layer, or below 10 cm so we were not able to determine if total AM biomass was affected by litter treatment. In addition, we did not characterise AM fungal communities from roots sampled from the superficial organic layer due to technical constraints. As such, we are unable to address the extra-radical responsiveness of AM fungi to increased inputs of organic matter, and directly address the selection of litter-specific AM fungal communities. Vertical stratification of ectomycorrhizal 610 communities has been described in boreal forest podzols (Rosling *et al.*, 2003), and increased

AM fungal colonisation of roots in the superficial organic layer could be hypothesised to

<sup>612</sup> reflect shifts in the structure and composition of AM fungal communities. This warrants

further investigation. Finally, we made no direct measure of nutrient transfer, and our

- discussion of how leaf litter manipulation altered AM fungal function is thus necessarily
- 615 speculative.
- 616

## 617 Potential sequencing bias

AM fungal communities were strongly dominated by taxa in the Glomeraceae (Figure 3),
which was due in part to our choice of marker region because the SSU is biased towards
Glomeraceae (Kohout *et al.*, 2014) and may underestimate diversity in some Diversisporales
(Davison *et al.*, 2015). Indeed, a previous study in the Barro Colorado Nature Monument
(BCNM) using Sanger sequencing and the same AM1/NS31 primer set similarly found a
strong dominance of AM fungal species in the Glomeraceae (Husband *et al.*, 2002).

Furthermore, a compilation of globally sampled AM fungal sequences obtained from the

amplification of a similar SSU region (with the primers AML1/NS31) described a similar

pattern: 79% of OTUs were from the order Glomerales (compared to 84% in this study), and

15% were from the Diversisporales (compared to 14% in this study; (Öpik *et al.*, 2013). By

contrast, a study in a montane forest in Ecuador using the ribosomal large subunit (LSU)

region found their dataset dominated by the Diversisporales (Camenzind *et al.*, 2014).

630

631 We used the number of DNA sequences as a measure of relative abundance of OTUs.

Although sequence abundance may reflect biases introduced through PCR and sequencing

- 633 protocols, the NS31-AM1 primer set exhibited very low levels of PCR bias when used to
- amplify artificial community templates of known composition (Cotton *et al.*, 2014). This is
- possibly because of the consistent length (c. 1.5% variation) and GC content (c. 3% variation)

of the amplified region across different AM fungal taxa (Helgason *et al.*, 1999), as variation

in amplicon length and GC content are known to cause biases in PCR reactions (Ihrmark et

*al.*, 2012), and may cause biases in the 454 sequencing process as well (Kauserud *et al.*,

639 2011). In any case, in a comparative analysis of our dataset using both quantitative and

<sup>640</sup> presence-absence approaches led to identical conclusions (Figure S5).

641

## 642 CONCLUDING REMARKS

Our findings show that the presence of decomposing leaf litter is important both in 644 structuring AM fungal communities, and in determining the extent of root colonisation by 645 AM fungi. Alterations in AM fungal community composition in response to litter removal 646 may be due to a range of factors including the reduction of key nutrients supplied by 647 decomposing leaf litter, notably N and P, changes in the action of saprobes, and changes in 648 water availability and pH. We hypothesise that a reduction in the quantity of decomposing 649 fresh organic matter brought about by litter removal may lead to AM fungi obtaining scarce 650 nutrients such as P from recalcitrant organic or mineral sources in the soil. Our hypothesis 651 652 helps to explain how trees were able to maintain their P-status despite the chronic removal of a major P input in this lowland tropical forest, and merits further investigation. 653

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658

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## 672 AUTHOR CONTRIBUTIONS

673

MS and SM designed the study. EVJT and EJS established the leaf litter manipulation

experiment. MS and DR performed the lab and field work. PAO and HW supported the lipid

analysis. BT conducted the nutrient analysis. NR conducted the bioinformatic analysis. MS

677 conducted the statistical analysis and wrote the manuscript with input from all authors.

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# 679 **REFERENCES**

- 680
- Alexander I, Lee S. 2005. Mycorrhizas and ecosystem processes in tropical rain forest:
   implications for diversity. Burslem D, Pinard M, Hartley S eds. Biotic Interactions in the
   Tropics. Cambridge: Biotic Interactions in the Tropics, 165–203.
- Alexander I, Selosse M-A. 2009. Mycorrhizas in tropical forests: a neglected research imperative. *New Phytologist* 182: 14–16.
- Altschul SF, Gish W, Miller W, Myers EW. 1990. Basic local alignment search tool.
   *Journal of Molecular Biology* 215: 403-410.
- Anders S, Huber W. 2010. Differential expression analysis for sequence count data.
   *Genome biology* 11: R106.
- 690 Attiwill PM, Adams MA. 1993. Nutrient Cycling in Forests. *New Phytologist* 124: 561–582.
- 691 **Camenzind T, Rillig MC. 2013**. Extraradical arbuscular mycorrhizal fungal hyphae in an 692 organic tropical montane forest soil. *Soil Biology and Biochemistry* **64**: 96–102.
- 693 Camenzind T, Hempel S, Homeier J, Horn S, Velescu A, Wilcke W, Rillig MC. 2014.
- Nitrogen and phosphorus additions impact arbuscular mycorrhizal abundance and molecular diversity in a tropical montane forest. *Global Change Biology* **20**: 3646–3659.
- Cavagnaro TR, Smith FA, Smith SE, Jakobsen I. 2005. Functional diversity in arbuscular
   mycorrhizas: exploitation of soil patches with different phosphate enrichment differs among
   fungal species. *Plant, Cell and Environment* 28: 642-650.
- Condit R, Engelbrecht BMJ, Pino D, Perez R, Turner BL. 2013. Species distributions in
   response to individual soil nutrients and seasonal drought across a community of tropical
   trees. *Proceedings of the National Academy of Sciences* 110: 5064–5068.
- Cotton TEA, Dumbrell AJ, Helgason T. 2014. What Goes in Must Come out: Testing for
   Biases in Molecular Analysis of Arbuscular Mycorrhizal Fungal Communities (R Balestrini,
   Ed.). *PLoS ONE* 9: e109234–7.
- 705 Davison J, Moora M, Öpik M, Adholeya A, Ainsaar L, Bâ A, Burla S, Diedhiou AG,
- Hiiesalu I, Jairus T, *et al.* 2015. Global assessment of arbuscular mycorrhizal fungus
   diversity reveals very low endemism. *Science* 349: 970–973.
- Dieter D, Elsenbeer H, Turner BL. 2010. Phosphorus fractionation in lowland tropical
   rainforest soils in central Panama. *Catena* 82: 118–125.
- 710 Drigo B, Pijl AS, Duyts H, Kielak AM, Gamper HA, Houtekamer MJ, Boschker HTS,
- Bodelier PLE, Whiteley AS, van Veen JA, et al. 2010. Shifting carbon flow from roots into
- associated microbial communities in response to elevated atmospheric CO2. *Proceedings of*
- *the National Academy of Sciences* **107**: 10938–10942.
- 714 Dumbrell AJ, Nelson M, Helgason T, Dytham C, Fitter AH. 2009. Relative roles of niche

- and neutral processes in structuring a soil microbial community. *The ISME Journal* 4: 337–345.
- Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and high
   throughput. *Nucleic Acids Research* 32: 1792–1797.
- 719 Field C, Behrenfeld M, Randerson J, Falkowski P. 1998. Primary production of the
- biosphere: integrating terrestrial and oceanic components. *Science* **281**: 237–240.
- 721 Frostegård Å, Tunlid A, Bååth E. 1993. Phospholipid Fatty Acid Composition, Biomass,
- and Activity of Microbial Communities from Two Soil Types Experimentally Exposed to
- 723 Different Heavy Metals. *Applied and Environmental Microbiology* **59**: 3605–3617.
- 724 Gosling P, Ozaki A, Jones J, Turner M, Rayns F, Bending GD. 2010. Organic
- management of tilled agricultural soils results in a rapid increase in colonisation potential and
   spore populations of arbuscular mycorrhizal fungi. *Agriculture, Ecosystems & Environment* **139**: 273–279.
- Gotelli NJ. 2000. Null model analysis of species co-occurrence patterns. *Ecology* 81: 2606 2621.
- 730 Gryndler M, Hršelová H, Cajthaml T, Havránková M, Řezáčová V, Gryndlerová H,
- 731
   Larsen J. 2008. Influence of soil organic matter decomposition on arbuscular mycorrhizal

   733
   10
   255
   266
- fungi in terms of asymbiotic hyphal growth and root colonization. *Mycorrhiza* **19**: 255–266.
- 733 Gryndler M, Larsen J, Hršelová H, Řezáčová V, Gryndlerová H, Kubát J. 2005. Organic
- and mineral fertilization, respectively, increase and decrease the development of external
   mycelium of arbuscular mycorrhizal fungi in a long-term field experiment. *Mycorrhiza* 16:
   159–166.
- Hao X, Jiang R, Chen T. 2011. Clustering 16S rRNA for OTU prediction: a method of
   unsupervised Bayesian clustering. *Bioinformatics* 27: 611–618.
- Hart MM, Reader RJ. 2002. Taxonomic Basis for Variation in the Colonization Strategy of
   Arbuscular Mycorrhizal Fungi. *New Phytologist* 153: 335–344.
- Helgason T, Daniell TJ, Husband R, Fitter AH, Young JPW. 1998. Ploughing up the
  wood-wide web? 394: 431–431.
- Helgason T, Fitter AH, Young JPW. 1999. Molecular diversity of arbuscular mycorrhizal
   fungi colonising Hyacinthoides non-scripta (bluebell) in a seminatural woodland. *Molecular Ecology* 8: 659–666.
- Herrera R, Merida T, Stark N, Jordan CF. 1978. Direct phosphorus transfer from leaf
   litter to roots. *Naturwissenschaften* 65: 208-209.
- Hendershot WH, Lalande H, Duquette M 2008. Ion exchange and exchangeable cations.
   Carter MR, Gregorich E eds. *Soil Sampling and Methods of Analysis*. Boca Raton, FL:
- Canadian Society of Soil Science and CRC Press, 173-178.
- 751 Hodge A. 2014. Interactions between arbuscular mycorrhizal fungi and organic material
- substrates. *Advances in applied microbiology* **89**: 47–99.

- 753 Hodge A, Fitter AH. 2010. Substantial nitrogen acquisition by arbuscular mycorrhizal fungi
- from organic material has implications for N cycling. *Proceedings of the National Academy*
- 755 *of Sciences* **107**: 13754–13759.
- Hodge A, Helgason T, Fitter AH. 2010. Nutritional ecology of arbuscular mycorrhizal
   fungi. *Fungal Ecology* 3: 267–273.
- 758 Husband R, Herre EA, Turner SL, Gallery R, Young JPW. 2002. Molecular diversity of
- arbuscular mycorrhizal fungi and patterns of host association over time and space in a
- tropical forest. *Molecular Ecology* **11**: 2669–2678.
- 761 Ihrmark K, Bödeker ITM, Cruz-Martinez K, Friberg H, Kubartova A, Schenck J, Strid
- 762 Y, Stenlid J, Brandström Durling M, Clemmensen KE, et al. 2012. New primers to
- amplify the fungal ITS2 region evaluation by 454-sequencing of artificial and natural
- communities. *FEMS Microbiology Ecology* **82**: 666–677.
- Johnson D, Krsek M, Wellington EMH, Stott AW, Cole L, Bardgett RD, Read DJ,
- Leake JR. 2005. Soil invertebrates disrupt carbon flow through fungal networks. *Science* 309: 1047.
- Johnson N. 1993. Can fertilization of soil select less mutualistic mycorrhizae? *Ecological Applications* 3: 749–757.
- Johnson NC. 2010. Resource stoichiometry elucidates the structure and function of arbuscular mycorrhizas across scales. *New Phytologist* 185: 631–647.
- Johnson NC, Angelard C, Sanders IR, Kiers ET. 2013. Predicting community and ecosystem outcomes of mycorrhizal responses to global change. *Ecology letters* 16: 140–153.
- Johnson NC, Rowland D, Corkidi L, Egerton-Warburton L. 2003. Nitrogen enrichment alters mycorrhizal allocation at five mesic to semiarid grasslands. *Ecology* 84: 1895-1908.
- Katoh K, Misawa K, Kuma K-I, Miyata T. 2002. MAFFT: a novel method for rapid
  multiple sequence alignment based on fast Fourier transform. *Nucleic Acids Research* 30:
  3059–3066.
- Kauserud H, Kumar S, Brysting AK, Nordén J, Carlsen T. 2011. High consistency
   between replicate 454 pyrosequencing analyses of ectomycorrhizal plant root samples.
- 781 *Mycorrhiza* **22**: 309–315.
- Kaye JP, Hart SC. 1997. Competition for nitrogen between plants and soil microorganisms.
   *Trends in ecology & evolution* 12: 139–143.
- 784 Kembel SW, Cowan PD, Helmus MR, Cornwell WK, Morlon H, Ackerly DD, Blomberg
- SP, Webb CO. 2010. Picante: R tools for integrating phylogenies and ecology.
   *Bioinformatics* 26: 1463–1464.
- 786 787
- Klironomos JN. 2003. Variation in plant response to native and exotic arbuscular
   mycorrhizal fungi. *Ecology* 84: 2292–2301.
- 790 Kohout P, Sudová R, Janoušková M, Čtvrtlíková M, Hejda M, Pánková H, Slavíková R,
- 791 Štajerová K, Vosátka M, Sýkorová Z. 2014. Comparison of commonly used primer sets for

- valuating arbuscular mycorrhizal fungal communities: Is there a universal solution? *Soil*
- 793 Biology and Biochemistry **68**: 482–493.
- <sup>794</sup> Leff JW, Wieder WR, Taylor PG, Townsend AR, Nemergut DR, Grandy AS, Cleveland
- 795 **CC. 2012.** Experimental litterfall manipulation drives large and rapid changes in soil carbon 796 cycling in a wet tropical forest. *Global Change Biology* **18**: 2969–2979.
- 797 Legendre P, Legendre L. 2012. *Numerical Ecology*. Elsevier Science.
- 798 Leigh EG Jr. 1999. *Tropical Forest Ecology*. Oxford, UK: Oxford University Press.
- Leigh J, Hodge A, Fitter AH. 2009. Arbuscular mycorrhizal fungi can transfer substantial
   amounts of nitrogen to their host plant from organic material. *New Phytologist* 181: 199–207.
- Love MI, Huber W, Anders S. 2014. Moderated estimation of fold change and dispersion
   for RNA-seq data with DESeq2. *Genome biology* 15: 550.
- Maherali H, Klironomos JN. 2007. Influence of Phylogeny on Fungal Community
   Assembly and Ecosystem Functioning. *Science* 316: 1746–1748.
- 805 McGonigle, Miller, Evans, Fairchild. 1990. A New Method which Gives an Objective
- 806 Measure of Colonization of Roots by Vesicular-Arbuscular Mycorrhizal Fungi. *New*
- 807 *Phytologist* **115**: 495–501.
- 808 **McMurdie PJ, Holmes S. 2014**. Waste not, want not: why rarefying microbiome data is 809 inadmissible. *PLoS computational biology* **10**: e1003531.
- 810 Mosse B, Phillips JM. 1971. The Influence of Phosphate and Other Nutrients on the
- Development of Vesicular-arbuscular Mycorrhiza in Culture. *Journal of General Microbiology* 69: 157–166.
- 813 **Munkvold L, Kjøller R, Vestberg M, Rosendahl S, Jakobsen I. 2004**. High functional 814 diversity within species of arbuscular mycorrhizal fungi. *New Phytologist* **164**: 357–364.
- Nilsson LO, Bååth E, Falkengren-Grerup U, Wallander H. 2007. Growth of
   ectomycorrhizal mycelia and composition of soil microbial communities in oak forest soils
- along a nitrogen deposition gradient. *Oecologia* **153**: 375–384.
- Nottingham AT, Turner BL, Winter K, Van Der Heijden MGA, Tanner EVJ. 2010.
- Arbuscular mycorrhizal mycelial respiration in a moist tropical forest. *New Phytologist* 186:
  957–967.
- Oksanen J, Blanchet FG, Kindt R, Legendre P. 2010. Multivariate analysis of ecological
   *communities in R: vegan tutorial. R package version 1.17-0.* http://vegan.r-forge.r-project.org
- Olsson PLA. 1999. Signature fatty acids provide tools for determination of the distribution and interactions of mycorrhizal fungi in soil. *FEMS Microbiology Ecology* 29: 303–310.
- 825 Oren R, Ellsworth DS, Johnsen KH, Phillips N, Ewers BE, Maier C, Schäfer KV,
- McCarthy H, Hendrey G, McNulty SG, et al. 2001. Soil fertility limits carbon
- sequestration by forest ecosystems in a CO2-enriched atmosphere. *Nature* **411**: 469–472.

- <sup>828</sup> Öpik M, Zobel M, Cantero JJ, Davison J, Facelli JM, Hiiesalu I, Jairus T, Kalwij JM,
- **Koorem K, Leal ME**, *et al.* **2013**. Global sampling of plant roots expands the described molecular diversity of arbuscular mycorrhizal fungi. *Mycorrhiza* **23**: 411–430.
- Pearson JN, Jakobsen I. 1993. Symbiotic Exchange of Carbon and Phosphorus between
  Cucumber and Three Arbuscular Mycorrhizal Fungi. *New Phytologist* 124: 481–488.
- 833 **Posada RH, Madrinan S, Rivera EL**. 2012. Relationships between the litter colonization by
- saprotrophic and arbuscular mycorrhizal fungi with depth in a tropical forest. *Fungal biology* **116**: 747–755.
- 836 Powell JR, Parrent JL, Hart MM, Klironomos JN, Rillig MC, Maherali H. 2009.
- Phylogenetic trait conservatism and the evolution of functional trade-offs in arbuscular
  mycorrhizal fungi. *Proceedings of the Royal Society B: Biological Sciences* 276: 4237–4245.
- **Read DJ. 1991.** Mycorrhizas in ecosystems. *Experientia* **47**: 376-391.
- Reynolds HL, Vogelsang KM, Hartley AE, Bever JD, Schultz PA. 2005. Variable
- responses of old-field perennials to arbuscular mycorrhizal fungi and phosphorus source.
- 842 *Oecologia* **147**: 348–358.
- Richardson AE, Simpson RJ. 2011. Soil Microorganisms Mediating Phosphorus
   Availability Update on Microbial Phosphorus. *Plant physiology* 156: 989–996.
- Rosling A, Landeweert R, Lindahl BD, Larsson KH, Kuyper TW, Taylor AFS, Finlay
   RD. 2003. Vertical distribution of ectomycorrhizal fungal taxa in a podzol soil profile. *New*
- **KD**. 2003. Vertical distribution of ectomycorrnizal fungal taxa in a podzol soil profile. *New Phytologist* 159: 775–783.
- Sayer EJ. 2006. Using experimental manipulation to assess the roles of leaf litter in the
   functioning of forest ecosystems. *Biological Reviews* 81: 1–31.
- 850 Sayer EJ, Banin LF. 2016. Tree Nutrient Status and Nutrient Cycling in Tropical Forest—
- Lessons from Fertilization Experiments. In: Goldstein G, Santiago L, eds. *Tropical Tree*
- 852 *Physiology*. Springer International Publishing, 275-297.
- Sayer EJ, Tanner EVJ. 2010a. A new approach to trenching experiments for measuring
   root–rhizosphere respiration in a lowland tropical forest. *Soil Biology and Biochemistry* 42:
   347–352.
- Sayer EJ, Tanner EVJ. 2010b. Experimental investigation of the importance of litterfall in
   lowland semi-evergreen tropical forest nutrient cycling. *Journal of Ecology* 98: 1052–1062.
- Sayer EJ, Tanner EVJ, Cheesman AW. 2006a. Increased Litterfall Changes Fine Root
   Distribution in a Moist Tropical Forest. *Plant and Soil* 281: 5–13.
- 860 Sayer EJ, Tanner EVJ, LACEY A. 2006b. Effects of litter manipulation on early-stage
- decomposition and meso-arthropod abundance in a tropical moist forest. *Forest Ecology and Management* **229**: 285–293.
- 863 Sayer EJ, Wright SJ, Tanner EVJ, Yavitt JB, Harms KE, Powers JS, Kaspari M,
- 64 Garcia MN, Turner BL. 2012. Variable Responses of Lowland Tropical Forest Nutrient

- 865 Status to Fertilization and Litter Manipulation. *Ecosystems* **15**: 387-400.
- 866 Schloss PD, Westcott SL, Ryabin T, Hall JR, Hartmann M, Hollister EB, Lesniewski
- **RA**, Oakley BB, Parks DH, Robinson CJ, et al. 2009. Introducing mothur: Open-Source,
- Platform-Independent, Community-Supported Software for Describing and Comparing
   Microbial Communities. *Applied and Environmental Microbiology* **75**: 7537–7541.
- 870 **Schreeg LA, Mack MC, Turner BL**. **2013**. Nutrient-specific solubility patterns of leaf litter 871 across 41 lowland tropical woody species. *Ecology* **94**: 94–105.
- 872 Simon L, Lalonde M, Bruns TD. 1992. Specific amplification of 18S fungal ribosomal
- genes from vesicular-arbuscular endomycorrhizal fungi colonizing roots. *Applied and Environmental Microbiology* 58: 291–295.
- 875 Smith SE, Read DJ. 2008. *Mycorrhizal Symbiosis*. Cambridge, UK: Academic Press.
- Stamatakis A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis of
   large phylogenies. *Bioinformatics* 30: 1312–1313.
- Tanner EVJ, Sheldrake MWA, Turner BL. 2016. Changes in soil carbon and nutrients
- following six years of litter removal and addition in a tropical semi-evergreen rain forest.
   *Biogeosciences Discussions*: doi:10.5194/bg-2016-229.
- 881
- Treseder K. 2004. A meta-analysis of mycorrhizal responses to nitrogen, phosphorus, and
   atmospheric CO2 in field studies. *New Phytologist* 164: 347–355.
- Treseder KK, Allen MF. 2002. Direct nitrogen and phosphorus limitation of arbuscular
   mycorrhizal fungi: a model and field test. *New Phytologist* 155: 507–515.
- Turner BL, Wright SJ. 2013. The response of microbial biomass and hydrolytic enzymes to
   a decade of nitrogen, phosphorus, and potassium addition in a lowland tropical rain forest.
   *Biogeochemistry* 117: 115–130.
- Veresoglou SD, Chen B, Rillig MC. 2012. Arbuscular mycorrhiza and soil nitrogen cycling.
   Soil Biology and Biochemistry 46: 53-62.
- Vincent AG, Turner BL, Tanner EVJ. 2010. Soil organic phosphorus dynamics following
   perturbation of litter cycling in a tropical moist forest. *European Journal of Soil Science* 61:
   48–57.
- Vitousek PM, Sanford RL. 1986. Nutrient Cycling in Moist Tropical Forest. *Annual Review* of Ecology and Systematics 17: 137–167.
- Wang Y, Naumann U, Wright ST, Warton DI. 2012. mvabund- an Rpackage for modelbased analysis of multivariate abundance data. *Methods in Ecology and Evolution* 3: 471–
  474.
- Warton DI, Foster SD, De'ath G, Stoklosa J. 2015. Model-based thinking for community
   ecology. *Plant Ecology* 261: 669-682.
- Webb C. 2000. Exploring the Phylogenetic Structure of Ecological Communities: An
  Example for Rain Forest Trees. *The American Naturalist* 156: 145–155.

- Webb CO, Ackerly DD, McPeek MA, Donoghue MJ. 2002. Phylogenies and Community
   Ecology. Annual Review of Ecology and Systematics 33: 475–505.
- 905 Went FW, Stark N. 1968. Mycorrhiza. *BioScience* 18: 1035–1039.
- 906 Wright SJ, Yavitt JB, Wurzburger N, Turner BL, Tanner EVJ, Sayer EJ, Santiago LS,
- Kaspari M, Hedin LO, Harms KE, *et al.* 2011. Potassium, phosphorus, or nitrogen limit
  root allocation, tree growth, or litter production in a lowland tropical forest. *Ecology* 92:
  1616–1625.
- 910 **Wurzburger N, Wright SJ. 2015**. Fine root responses to fertilization reveal multiple 911 nutrient limitation in a lowland tropical forest. *Ecology* **96**: 2137–2146.
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# 914 Supporting Information

- Figure S1. Effect of litter manipulation on the levels of NLFA 16:105 in the top 10 cm of
   forest soil.
- <sup>917</sup> Figure S2. Rarefaction curves pooled by experimental treatment and for each sample.
- Figure S3. Maximum-likelihood phylogenetic tree of all operational taxonomic units (OTUs)
   detected in this study.
- Figure S4. Effect of litter manipulation on AM fungal OTU richness (total number of OTUs
- in a sample; a) and predominance (the proportional abundance of the dominant AM fungaltaxon; b).
- Figure S5. Comparison of AM fungal communities described by the quantitative Bray-Curtis metric of dissimilarity (a), the Jaccard presence-absence based metric of dissimilarity (b), and correlation between the two (c).
- Table S1. Response of soil physical characteristics to nine years of litter removal and addition in a tropical forest.
- Table S2. AM fungal OTUs altered by nine years of leaf litter addition and removal.
- Table S3. Number of sequences per sample after blasting, filtering, merging and trimming.
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## 938 FIGURE LEGENDS

Figure 1. Effects of litter manipulation on soil physical characteristics. Values are means  $\pm$ Fisher's Least Significant Difference. Grey shaded regions represent control treatments. Litter treatments are significantly different from controls at  $\alpha < 0.05$  (n = 5) where error bars do not overlap the grey shaded regions. Standard normal deviates are plotted to facilitate visual comparison of effect size. Al, Mn and N (inorganic) were log transformed prior to analysis due to heteroscedasicity. N (inorg.) = inorganic N; P (res.) = resin extractable P; P (tot.) = total P; TEB = total exchangeable bases; EBS = effective base saturation; L- = litter

946 removal treatment; L+ = litter addition treatment.

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Figure 2. Percent root length colonised by AM fungi (total colonisation, colonisation by 948 hyphae, colonisation by vesicles and colonisation by arbuscules). Left-hand panels (a-d) 949 show the effect of litter manipulation on AM fungal colonisation of roots in the mineral soil. 950 Right-hand panels (e-h) compare colonisation in roots between the mineral soil ('soil') and 951 superficial organic horizon ('organic') across control and litter addition treatments. L- is litter 952 removal, C is control, and L+ is litter addition. In left hand panels (a-d) values are means  $\pm$ 953 Fisher's Least Significant Difference, and non-overlapping error bars indicate significance at 954  $\alpha < 0.05$  (*n* = 5). In right-hand panels, values are means  $\pm 95\%$  confidence intervals obtained 955 by parametric bootstrapping with 10000 simulations. 956

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Figure 3. Mean proportional abundance of AM fungal genera (a) and families (b) in mixed root samples across litter manipulation treatments (n = 5); L- is litter removal, C is control, and L+ is litter addition.

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Figure 4. NMDS ordination plot showing changes in AM fungal community composition in long-term litter removal plots (circles), but not litter addition plots (triangles), compared to controls (squares) in a lowland tropical forest. Site scores are shown and ellipses describe

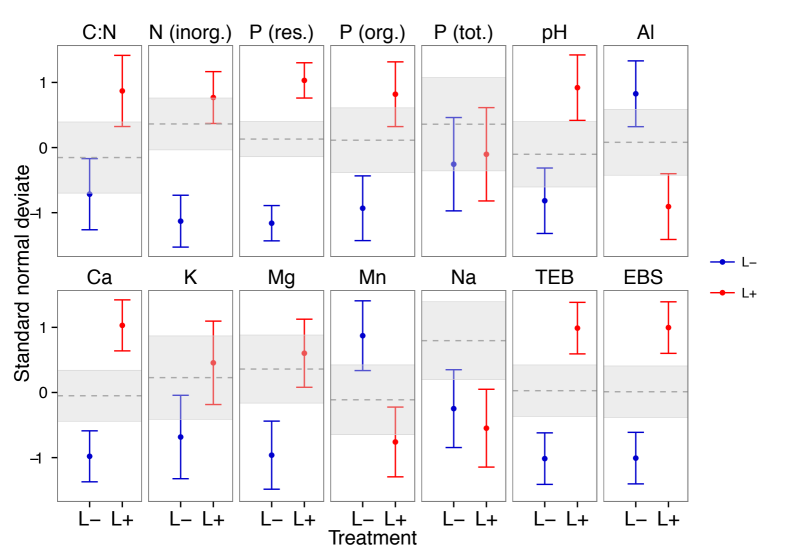
- 965 95% confidence areas. Arrows indicate the direction and degree of significant correlations
- between NMDS axes and soil physical characteristics (n = 5). EBS = effective base
- 967 saturation; P (resin) = resin extractable phosphate; N (inorg.) = inorganic N; L- = litter
- removal, C = control, and L+ = litter addition. Axes are scaled to half-change (HC) units, by
- which one HC unit describes a halving of community similarity.

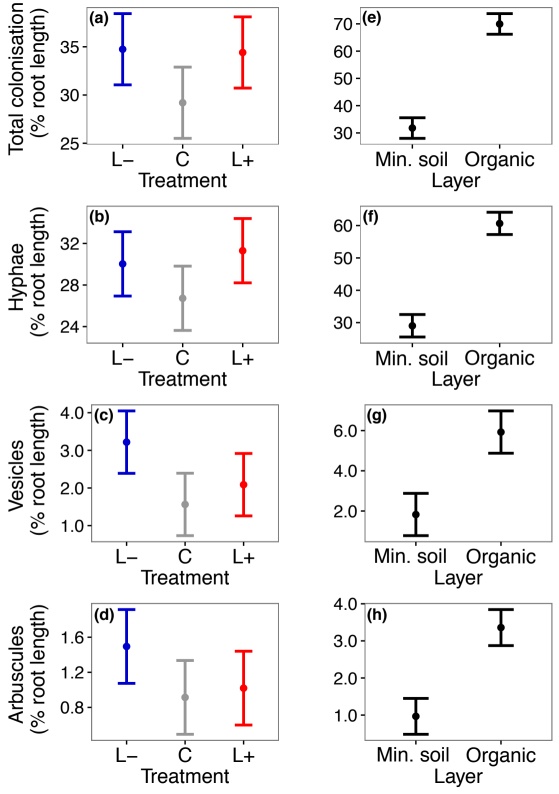
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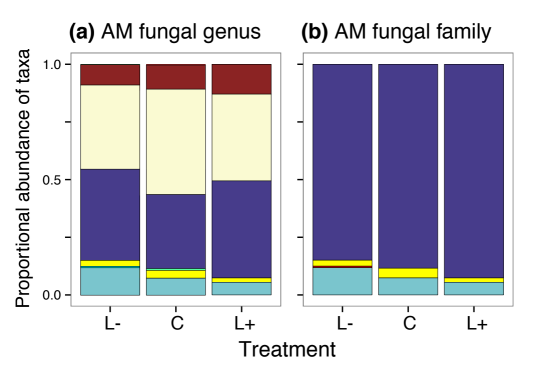
Figure 5. Effect of litter addition (red) and removal (blue) on the relative abundance of 971 972 individual AM fungal operational taxonomic units (OTUs). Significantly altered (P < 0.05) OTUs are shown based on both adjusted and unadjusted P values. The names of OTUs that 973 974 are significantly affected by litter manipulation are emboldened. x- axis indicates the effect size as log2 fold change, and error bars show standard errors. OTUs are arranged in order of 975 decreasing rank abundance (more highly ranked OTUs are those that are more prevalent 976 across all samples in the dataset). Significance was ascertained based on negative binomial 977 Wald tests using standard maximum likelihood estimates for generalised linear models, as 978 implemented in the DESeq2 package. 979

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Figure 6. Litter manipulation moderately altered the degree of relatedness between taxa in AM fungal communities when described using the metric of Net Relatedness Index (NRI). Higher numeric values correspond to more closely related AM fungal communities. Values are means  $\pm$  Fisher's Least Significant Difference: non-overlapping error bars indicate significance at  $\alpha < 0.05$  (n = 5). Dotted lines indicate significance threshold of  $\alpha = 0.05$ derived from comparison with 10000 null communities generated using the 'independentswap' algorithm. L- is litter removal, C is control, and L+ is litter addition.





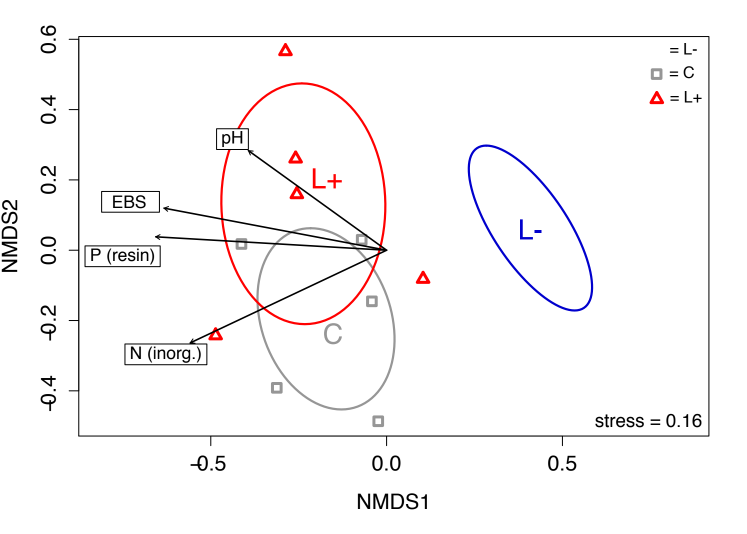


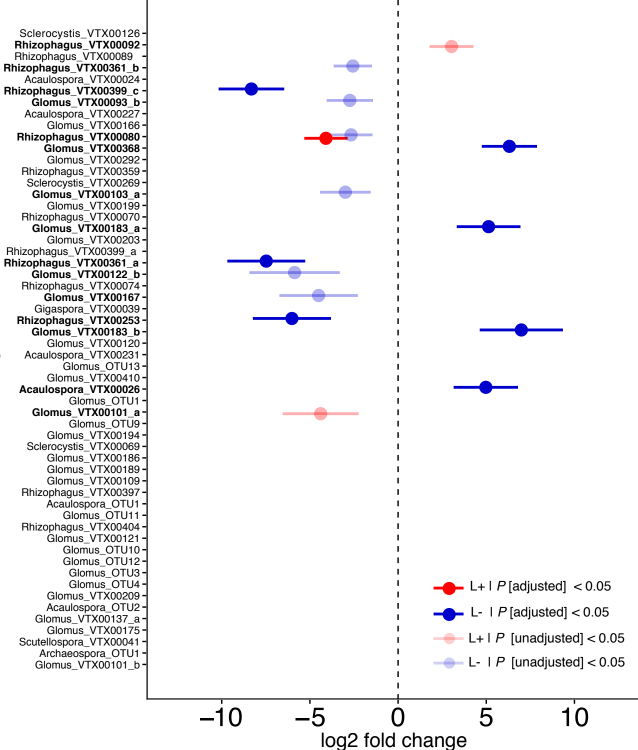
# AM fungal genus

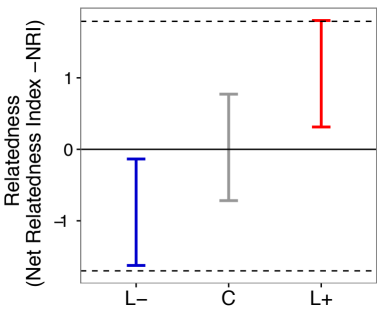
Acaulospora Archaeospora Gigaspora Glomus Rhizophagus Sclerocystis Scutellospora

# AM fungal family

Acaulosporaceae Archaeosporaceae Gigasporaceae Glomeraceae







# Arbuscular mycorrhizal fungal community composition is altered by longterm litter removal but not litter addition in a lowland tropical forest

Supporting Information: Figures S1-S5 and Tables S1-S3

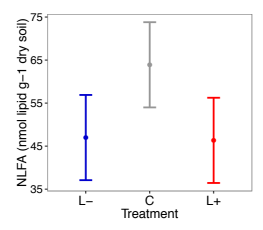


Figure S1. Effect of litter manipulation on the levels of NLFA 16:1 $\omega$ 5 in the top 10 cm of forest soil. Values are means ± Fisher's Least Significant Difference: non-overlapping error bars indicate significance at P < 0.05. L- is litter removal, C is control, and L+ is litter addition.

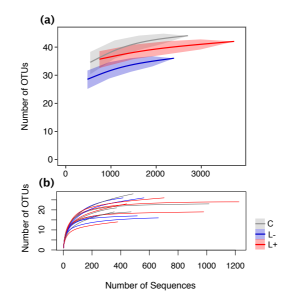


Figure S2. Rarefaction curves pooled by experimental treatment (a) approached asymptotes, indicating that sampling effort was sufficient to capture the range of AM fungal taxa across the sites. Rarefaction curves for each sample (b) indicated that sequencing intensity was sufficiently high to detect the majority of OTUs. C is control, L- is litter removal, and L+ is litter addition. Shaded bands show 95% confidence regions calculated from the standard error of the estimate using the function specaccum in the R package vegan.

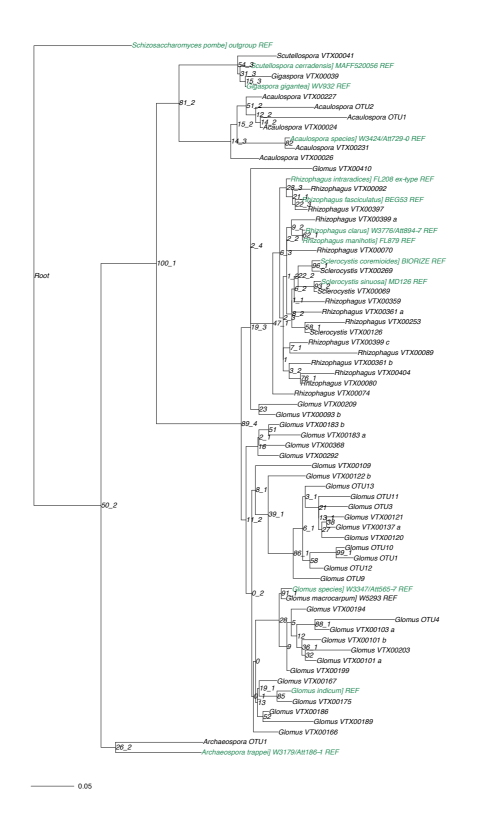


Figure S3. Maximum-likelihood phylogenetic tree of all operational taxonomic units (OTUs) detected in this study. The scale bar equals the number of substitutions per site. A subset of reference sequences from Kruger *et al.* (2012) are displayed in green text.

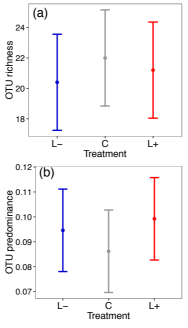


Figure S4. Effect of litter manipulation on AM fungal OTU richness (total number of OTUs in a sample; a) and predominance (the proportional abundance of the dominant AM fungal taxon; b). Values are means  $\pm$  Fisher's Least Significant Difference: non-overlapping error bars indicate significance at P < 0.05. L- is litter removal, C is control, and L+ is litter addition.

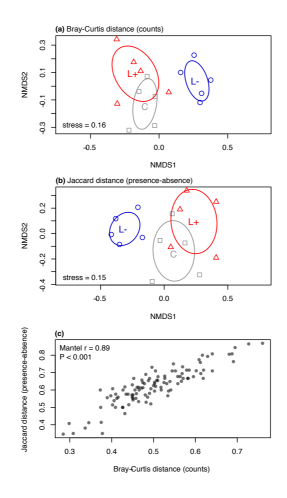


Figure S5. Comparison of AM fungal communities described by the quantitative Bray-Curtis metric of dissimilarity (a), the Jaccard presence-absence based metric of dissimilarity (b), and correlation between the two (c). (a) and (b) are two-dimensional NMDS plots with ellipses describing 95% confidence areas around the sample scores.

Variable	n = 5)		SE			
	Litter removal		Control	Litter addition		
Al	38.0		12.0	2.6		1.7
Ca	740	*	1400	2200	*	180
К	53.0		75.0	80.0		9.7
Mg	270	*	390	410		31
Mn	72.0		46.0	34.0		1.2
Na	9.8		13	8.7	*	1.4
P (resin)	6.0	*	17.0	25.0	*	1.5
P (total)	370		390	370		20
P (organic)	76		93	100		5
C:N	9.9		10.0	11.0		0.2
N (inorganic)	2.1	*	4.9	6.1		1.2
TEB <sup>1</sup>	6.1	*	10.0	14.0	*	1.1
EBS <sup>2</sup>	7.0	*	11.0	15.0	*	1.0
рН	5.3		5.5	5.8		0.1

Table S1. Response of soil physical characteristics to nine years of litter removal and addition in a tropical forest

#### Notes

Variables significantly affected by litter addition are asterisked

All nutrients are expressed as mg kg<sup>-1</sup> of dry soil

TEB is expressed as cmol kg<sup>-1</sup> dry soil

EBS is a unitless fraction

SE = standard error

<sup>1</sup> Total Exchangeable Bases

<sup>2</sup> Effective Base Saturation

AM fungal OTU	log2 Fold Change	SE <sup>1</sup>	P value		Rank abundance	Treatment <sup>2</sup>	Direction of change
Acaulospora_VTX00026	-4.98	1.83	0.03	*	32	L-	-
Glomus_VTX00183_a	-5.13	1.81	0.03	*	18	L-	-
Glomus_VTX00183_b	-6.99	2.36	0.02	*	27	L-	-
Glomus_VTX00368	-6.31	1.57	0.00	*	11	L-	-
Rhizophagus_VTX00253	6.02	2.22	0.03	*	26	L-	+
Rhizophagus_VTX00361_a	7.47	2.21	0.01	*	21	L-	+
Rhizophagus_VTX00399_c	8.32	1.86	0.00	*	6	L-	+
Rhizophagus_VTX00080	4.09	1.23	0.04	*	10	L+	+
Glomus_VTX00093_b	2.74	1.32	0.04		7	L-	+
Glomus_VTX00103_a	2.99	1.44	0.04		15	L-	+
Glomus_VTX00122_b	5.87	2.56	0.02		22	L-	+
Glomus_VTX00167	4.51	2.22	0.04		24	L-	+
Rhizophagus_VTX00080	2.67	1.22	0.03		10	L-	+
Rhizophagus_VTX00361_b	2.57	1.09	0.02		4	L-	+
Glomus_VTX00101_a	4.39	2.16	0.04		34	L+	+
Rhizophagus_VTX00092	-3.03	1.25	0.02		2	L+	-

Table S2. AM fungal OTUs altered by nine years of leaf litter addition and removal, as ascertained using the DESeq2 package.

Asterisks (\*) denotes P values corrected for multiple comparisons

<sup>1</sup> SE = standard error

<sup>2</sup> L- is litter removal and L + is litter addition

trimming (exclusion of OTUs with a total of 5 or less reads)					
Plot	Treatment	Number of sequences			
1	С	1016			
11	С	372			
15	С	473			
5	С	355			
7	С	487			
12	L-	516			
13	L-	417			
4	L-	237			
6	L-	663			
8	L-	562			
10	L+	441			
14	L+	379			
2	L+	979			
3	L+	1225			
9	L+	703			
		total = 8825			

Table S3. Number of sequences per sample after blasting, filtering, merging, and trimming (exclusion of OTUs with a total of 5 or less reads)

# References

Krüger M, Krüger C, Walker C, Stockinger H, Schüßler A. 2012. Phylogenetic reference data for systematics and phylotaxonomy of arbuscular mycorrhizal fungi from phylum to species level. *New Phytologist* 193: 970–984.