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COORDINATED COMMUNITY structure among trees, fungi and invertebrate groups in Amazonian rainforests

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Little is known regarding how trophic interactions shape community assembly in tropical forests. Here we assess multi-taxonomic community assembly rules using a rare standardized coordinated inventory comprising exhaustive surveys of fve highly-diverse taxonomic groups exerting key ecological functions: trees, fungi, earthworms, ants and spiders. We sampled 36 1.9-ha plots from four remote locations in French Guiana including precise soil measurements, and we tested whether species turnover was coordinated among groups across geographic and edaphic gradients. All species group pairs exhibited signifcant compositional associations that were independent from soil conditions. For some of the pairs, associations were also partly explained by soil properties, especially soil phosphorus availability. Our study provides evidence for coordinated turnover among taxonomic groups beyond simple relationships with environmental factors, thereby refning our understanding regarding the nature of interactions occurring among these ecologically important groups.

The outstanding biodiversity of tropical forests represents a major challenge for ecologists seeking to understand how species are spatially assembled in these ecosystems¹. Most studies to date have investigated assembly rules within individual species groups, especially plants^{2,3}, with less than ten percent of published articles focusing on other living groups between 2008 and 2018 (Google Scholar research using the terms "*community assembly tropical forests*" and "*determinants of community assembly tropical forests*" in separate searches; pages 1–20). At the same time, there has been considerably less investment to examine how the composition of diferent groups, notably groups of major ecological importance (e.g. primary producers, decomposers, herbivores and their predators), display coordinated turn-over across geographical and environmental gradients. Addressing this topic would shed light on the role exerted by environmental fltering and biotic interactions between communities on biodiversity organisation in the most diverse ecosystem on earth.

The majority of studies examining associations between different taxa have relied on simplified proxies such as diversity indices, ofen with conservation-oriented purposes. For instance, many works have tested whether tree diversity represents a reliable surrogate for the diversity of other taxa, mainly arthropods and vertebrates⁴⁻⁷. Nevertheless, there has only been weak and inconsistent evidence establishing signifcant diversity associations between taxa in tropical forests⁸, while studies on the subject have often been hampered by difficulties to obtain

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accurate diversity estimations in these hyper-diverse ecosystems⁹. Moreover, diversity indices provide limited information to unveil the complex assembly rules that structure communities, for which compositional information would be more appropriate¹⁰. Multi-taxon composition data are considerably lacking in ecology, especially in rainforests, because they require considerable logistical and time-consuming eforts to coordinate the work of specialists of different taxa¹¹. Additionally, we are witnessing a sharp decline of trained taxonomists able to identify specimens in many groups. This is particularly problematic in the case of arthropods, which represent most of the macroscopic diversity in the tropics⁵, as well as for other hyper-diverse and poorly known groups including fungi and non-arthropod invertebrates. The few studies reporting compositional association between taxa in tropical ecosystems have involved testing association with tree species $12-14$, which are generally well described and relatively easy to inventory¹⁵.

Even though empirical evidence is sparse, strong hypotheses can be proposed to link composition between tree communities and other groups in tropical forests. Co-evolution processes have driven trees to ofer a huge variation of food supply, anti-herbivore defences and/or housing¹⁶, and thereby to interact in multiple ways with other communities. In a recent study¹⁰, strong compositional associations have been emphasized between plant, arthropod and micro-organism communities in a diverse subtropical forest of South China. The study suggested that tree communities exert a bottom-up control on aboveground arthropod communities while, at the same time, trees are influenced by soil micro-organisms and decomposers via a top-down effect. The key role of trees in the functioning of forest ecosystems has also served as an argument to support their usefulness as a surrogate for the composition of other groups, for instance arthropod¹⁷ and fungal communities¹⁸.

Further hypotheses can be suggested to link composition among other species groups. Ants for example, are extremely diverse, both ecologically and taxonomically, and play key functional roles in forest ecosystems¹⁹. They are therefore likely to influence many other communities. Leaf-litter ants, in particular, profoundly impact soil structure through foraging and tunnelling activities, while they also represent major predators of the soil fauna20. Spiders also exhibit key functions as predators, and competitive and predator-prey interactions between ants and spiders are likely to be reflected in significant associations between both groups²¹. Other major ecosystem functions, including decomposition of organic matter, nutrient cycling and mycorrhizal associations²² are provided by fungi, which represent another highly diverse and understudied group in tropical forests. Fungal communities are expected to interface with tree communities, via feedback interactions between plants and soil²³ and pathogen-mediated negative density dependent effects^{24,25}. Other major decomposers comprise earthworm communities, which are known to be infuenced by chemical properties of the litter and soil conditions, but also to substantially modify soil properties and thereby impact other groups²⁶.

Taxonomic congruence among species groups may result not only from biotic interactions but also be explained by a similar response to environmental conditions. The assumption that environmental variation (altitudinal, topographical and edaphic) drives beta-diversity in tropical forests has been abundantly demonstrated for trees^{2,3}. Such work has built a strong foundation for conservation programs to identify areas of high biodiversity²⁷. However, there has been considerably less evidence for the influence of environment among other groups19,28. Moreover, very few attempts have been undertaken to examine to what extent coordinated taxonomic turn-overs among diferent species groups are driven by the fltering of common environmental flters. Previous studies in the Amazon have reported congruence of taxonomic composition between Melastomataceae and Pteridophytes, partly explained by Ca, Mg and K content, and soil texture²⁹. More recently, Lemes Landeiro *et al.*30 have emphasized the existence of coordinated composition turn-overs among 22 taxonomic groups that were explained by environmental variables, especially soil clay and phosphorus content.

Despite some sparse evidence for cross-taxon congruence in tropical forests, there remains a great need for multi-taxon composition data combined with environmental information to better understand the factors explaining coordinated turn-over of composition among diferent species groups. Tis would also help to shed light on the nature of interactions between groups, so as to improve our predictions for the potential consequences of species loss on ecosystem functioning in tropical forests 31 .

The present study aims to (i) investigate congruence of taxonomic composition among five species groups of major ecological importance (trees, ants, spiders, fungi and earthworms), of which some remain critically understudied in the tropics, and (ii) to examine whether coordinated species-turn-over among pairs of groups is explained by soil variables. To do so, we carried out exhaustive and standardized multi-taxa inventories in 36 plots distributed across four remote areas of French Guiana representing the biogeographic and environmental gradients of this hyper-diverse tropical region. We address the following specifc questions:

- 1. (**a**) To what extent is the taxonomic composition of the fve living groups correlated to each other? **(b)** How do we interpret the signifcant pairwise associations?
- 2. (**a**) To what extent is correlation of composition between groups due to a coordinated response to environmental heterogeneity among plots? **(b)** Is the coordinated response explained by the same environmental variables among pairs of groups?

Results

Co-inertia between taxonomic groups. A Multiple Co-Inertia Analysis (MCOIA), an ordination method allowing visual representation of associations among multiple data tables, was performed to explore patterns of compositional associations among taxa. The analysis showed positive correlation of taxonomic composition between the fve species groups (Fig. 1a). A marked diference of the overall composition (all groups together) was observed between the *Mitaraka* and *Saül*-*Limonade* sites on the frst MCOIA axis (Fig. 1b). Within each of these sites, diferences were also well marked between seasonally fooded soils and the two other habitats (hilltop and slope) on the second axis of the MCOIA (see lef graph in Fig. 1b). For simplicity, Fig. 1a,b show the

Figure 1. (**a**) Correlations between the frst axes of separate PCA (performed on each species group) and the three frst axes of the MCOIA (axes 1–2 and 1–3). (**b**) Projection of plot scores on axes 1–2 and axes 1–3 of the MCOIA, emphasizing the two sites where compositional data was available for the fve groups (Mitaraka=MIT and Saül-Limonade=SL) as well as topographical habitats (hilltop, slope and seasonally fooded). Histograms represent the eigenvalues of the MCOIA axes. (c) Adjusted RV coefficients calculated for each pair of taxonomic groups (portion of the table shaded in grey), and between each taxonomic group and soil variables (frst column). Asterisks indicate whether values were signifcant (in bold) according to the MSR test: **P*<0.05; ***P* < 0.01; ****P* < 0.001. The last column represents the mean (\pm standard deviation) of RV values per group. The MCOIA performed using other combinations of species groups are presented in Figs S4 and S5.

results obtained in the two sites where data for all groups were available, but MCOIA performed with three or two sites (where more than two groups were inventoried) produced redundant results (Figs S4 and S5). Pairwise compositional associations among groups were quantified by the RV coefficient of Co-Inertia Analyses (COIA), which was tested using Moran Spectral Randomisations (MSR) to take spatial autocorrelation into account. RV values were significant ($P \le 0.05$) for each pair of groups, regardless of the soil heterogeneity (Fig. 1c). The highest RV value reached 0.746 (*P* < 0.001; MSR test) between trees and ants. Average RV values (calculated over the four values per group) ranged from 0.488 (SD = \pm 0.121) for earthworms to 0.575 for spiders (SD = \pm 0.038). All

Table 1. Effect of soil variables on the composition co-inertia for each pair of taxonomic groups. The first line corresponds to adjusted R^2 values quantifying the effect of all soil variables (ASV) on the co-inertia. Significant R^2 values ($P \le 0.05$ using a MSR test with a Sidak correction for multiple tests) are indicated in bold. The next lines represent the sum of weighted AIC_c (SWA) of each soil variable, calculated from the model averaging for multivariate generalized linear models. S: spatial structure; B=broad-scale (inter-site) spatial structures detected only; $BF = both broad- and fine-scale (intra-site) structures detected.$ ns $=$ no significant structure detected. Mean: mean SWA values (in italics) for the four pairs that were signifcantly explained by ASV.

groups also displayed signifcant co-inertia with soil variables, with trees and ants showing the highest associations ($RV = 0.623$ and 0.601, respectively).

RV values were highly correlated to Procrustes residuals sum of squares (PRSS), afer removing or not the plots on seasonally fooded forest and by using either taxonomic abundance/occurrence or presence-absence data (with *r*-Pearson values ranging from 0.62 to 0.96; Figs S6 and S7), hence showing that both methods produced consistent results. All RV and PRSS values obtained afer removing or not plots on seasonally fooded forest, and using abundance or presence-absence data, are presented in Tables S1 to S4.

Coordinated response to environmental variables for each pair of groups. Four out of the ten pairs of groups (40%) remained signifcantly explained (*P*≤0.05) by all soil variables afer correcting for multiple tests: trees-ants, trees-spiders, trees-fungi and ants-spiders (Table 1). The relative contribution of each soil variable was quantifed using the sum of weighted AICc (corrected Akaike Information Criterion) values (SWA) from a Model Averaging with Multivariate Generalized Linear Model. On average, soil phosphorus availability (P) was the variable that best explained the co-inertia among the four significant pairs (mean SWA = 0.62 ; SD = \pm 0.03; Table 1). The best following explanatory variable displayed mean SWA values ranging from 0.49 for the percentage of sand to 0.54 and 0.55 for Ca and Mg, respectively. P was the best explanatory variable for each of the four signifcant pairs (from 0.59 for the trees-fungi to 0.66 for the trees-spiders pair), along with Ca (0.58) and Mg (0.60) for the trees-fungi pair.

P was signifcantly spatially structured at both fne (within-site) and broad (among-site) spatial scales, as were the overall soil heterogeneity (all soil variables together), OM and K. N only displayed broad-scale spatial structures. No signifcant structure was detected for soil texture, Ca, Mg, Na, Al and Mn.

Discussion

We observed signifcant compositional associations between fve species groups that provide major ecological functions in tropical forests. For four out of the ten pairs of groups, the co-inertia between the two groups was signifcantly explained by the overall soil heterogeneity, with P availability being the most infuential variable.

All pairs of taxonomic groups showed coordinated turn-over of composition. All pairs of taxonomic groups displayed signifcant co-inertia, refecting signifcant coordinated compositional turn-overs across plots, regardless of the influence of soil conditions (Fig. 1c). Two out of the three RV values < 0.5 were obtained with earthworms (the trees-earthworms and ants-earthworms pairs). These relatively low values may have partly resulted from lower statistical power since earthworms' data were available in two sites only. However, this group was strongly associated with spiders and fungi and thus additional data on earthworms are needed to verify its association with other groups. The relatively low RV value (0.385) observed for the fungi-ants pair may be explained by the absence of ant-associated fungi in our sampling. Indeed, we targeted Basidiomycota and visible fungi, while Chaetothyriales and most ant-associated fungi are leaf-endophytes belonging to Ascomycota. The RV value was nonetheless signifcant, which may suggest the existence of other interactions between both groups for which we lack hypotheses.

The highest co-inertia ($RV = 0.746$) was found between trees and ants, while all groups were significantly associated to trees. This result is in agreement with previous studies that emphasized the influence of trees, as key drivers of resource (energy, nutrients) fuxes, on the composition of other communities following bottom-up effects^{4,10}. It also supports the idea that tree species composition may serve as a useful surrogate for other communities and thereby contribute to improving current estimations of local and regional tropical diversity which remain greatly inaccurate³². The key role of trees in structuring tropical forest ecosystems has often been a foundation for conservation management programs and to help policy makers to choose protected areas and to improve models of biodiversity dynamics in response to climate and land use change scenarios 33 .

The strong association between trees and ants may possibly results from the quality of the litter fall that influences the habitat structure as well as the community of ant preys^{34,35}. The significant association obtained between ants and all other groups supports previous evidence for the major ecological functions that they provide in tropical forests^{19,20,36,37}. Ant species being well-described in the Amazon region and relatively easy to inventory, our results reveal their utility as another potential proxy in future biodiversity assessment surveys³⁸.

RV values were, on average, lower among pairs including trees (mean $RV = 0.549$) than among pairs including spiders (mean RV = 0.575 ; Fig. 1c). The latter results may depict a network of interactions that is more complex than proposed by most bottom-up and top-down models¹⁰, with spiders interacting more directly with the other studied groups than do trees. For example, the signifcant association observed between spiders and ants $(RV = 0.518)$ may reflect the strong intra-guild competition occurring between the two groups, which both comprise diverse generalist predators²¹. The strong taxonomic congruence between trees and spiders (RV = 0.598) may be explained by the same reasons as we proposed for the tree-ant association (see above). Interactions between spiders and fungi and between spiders and earthworms have never been reported to date and we lack hypotheses to interpret the signifcant associations observed for these pairs of groups.

The highly significant association between trees and fungi $\text{RV}=0.454$) supports previous evidence highlighting the importance and the variety of interactions occurring between these two groups in the Amazon³⁹. For example, diferent fungi guilds (arbuscular mycorrhizal and ectomycorrhizal fungi, as well as saprophytic fungi) have been shown to exert a top-down control on tree species composition in a tropical forest of south-eastern China9 . Interestingly, changes in fungi communities where also highly correlated with changes in earthworm communities ($RV = 0.588$). As saprophytic fungi and especially wood-decaying fungi were the most abundant, we can hypothesize that fungi and earthworms share similar niches in tropical forests. Moreover, fungi have already been demonstrated to represent an important food source for earthworms, and diferent earthworm species can have preferences for distinct fungal taxa⁴⁰.

It is worth pointing out that compositional associations were observed despite uneven taxonomic resolution among groups (species for ants and trees, genera for fungi and spiders, OTU's for earthworms). This makes sense when considering previous evidence reporting that genera represent good ecological surrogates for both spiders⁴¹ and fungi⁴². In the case of earthworms, current species delimitations have been suggested to be more precise when considering the molecular classifcations we employed to determine OTUs, due to substantial cryptic morphological variation among putatively distinct taxa⁴³.

Coordinated responses to edaphic heterogeneity among pairs of groups. Four out of the ten pairs of groups signifcantly reacted in a coordinated way with the overall edaphic variation (Table 1). Te absence of signal for the four pairs of groups that included earthworms might result from low statistical power due to the fewer number of sites (two) in which they were inventoried. It may also be due to a limited efect of the soil stoichiometry variation on these decomposers compared to the role exerted by the carbon quality of the soil⁴⁴.

P was the soil factor that best explain the coordinated response among the four pairs signifcantly explained by the overall soil heterogeneity (mean SWA = 0.62). The significant effect of P on coordinated taxonomic turn-overs has rarely been emphasized before, but a previous study³⁰ demonstrated that the availability of this element, along with the percentage of clay, explained the congruence between lianas of the Bignoniaceae family and other living groups in the Brazilian Amazon. We may assume that the joint infuence of P and other soil properties result from bottom-up efects exerted by trees on the composition of other groups: it has been well supported that tree species composition varies with soil P availability⁴⁵ and other soil variables³, thereby influencing other groups depending at least partially on tree communities for resource use. Soil texture was not among the variables that best explained compositional congruence among pairs of groups, although the infuence of this property on tree community composition⁴⁶ but also on ant community composition³⁴ has been well established in tropical forests. This may be due to the low soil textural variation among plots (the coefficient of variation ranged from 0.46 to 0.74 for the three soil texture variables, while it reaches, on average, 1.18 among all the other variables). In a second position, we found that compositional congruence among groups was also well explained by Mg and Ca (mean SWA over the four pairs=0.55), for which the role has, to our knowledge, never been demonstrated for explaining coordinated beta-diversity patterns in tropical forests. Nevertheless, there has been previous evidence for its influence on composition within individual groups, especially in rainforest tree² and fungi²⁸ communities.

Questions raised and research perspectives. In light of our results and the previous works that have highlighted compositional associations among other groups than the ones studied here¹⁰⁻¹⁴, it is cogent to consider to what extent taxonomic congruence occurs among other communities in the ecosystem, such as vertebrates, herbaceous and epiphytic plants, and other micro-organisms. For instance, we could hypothesise that birds and mammals display coordinated beta-diversity with most of the taxa investigated in our study, e.g. via predation on invertebrates and plant use (herbivory, zoogamy, zoochory). However, the disparities of dispersal capacities between large and small animals would jeopardize the chance of detecting compositional associations at the 1.9ha scale of our plots47. Sampling designs should therefore take multiple spatial scales into account so that congruent patterns between animal groups displaying contrasted spatial structures are analysed at appropriate spatial scales. One means to accomplish this will be to integrate the recent progress in assessing biodiversity using metabarcoding methods48, which represent a promising approach for investigating compositional association between macro and micro-organism communities. We hope that our work will encourage further similar studies

Table 2. Characteristics and taxonomic diversity of each living group in all study sites combined and within each site. *nd*: no data available. ENT_2 = Effective Number of Taxa expected for a random sampling (with replacement) of 2 individuals, following Dauby & Hardy (2011). Bold values represent the highest taxonomic richness and $ENT₂$ among sites for each living group.

in tropical forests but also extend our approach to other ecosystems, in order to better understand the importance of coordinated beta-diversity and the underlying mechanisms driving these patterns in natural communities.

Conclusion

We demonstrate the existence of coordinated beta-diversity patterns among five species groups of major ecological importance in tropical forests, a rarely investigated subject in ecology. The coordinated species turn-overs partly and signifcantly resulted from the common efect of soil variables, especially soil phosphorus availability, but were also partly unrelated to the heterogeneity of the measured soil variables. Our results shed new light in our understanding of species assembly rules among groups of which some remain critically understudied in the tropics. They also confirm previous studies reporting the usefulness of trees but also ants as proxies for assessing the composition of other species groups, which will improve the design and impact of conservation programs using tree and arthropod inventories in tropical forests. Our study paves the way for deeper investigations of biotic interactions in the tropics and in other ecosystems, and may bring new hypotheses to understand biodiversity patterns and even diversifcation processes.

Materials and Methods

Study area and inventory sites. Between 2012 and 2017, multi-taxonomic inventories were carried out in four sites of mature lowland and lower mountain tropical moist forests across French Guiana (Fig. S1). French Guiana is located in the northern part of South America, between latitudes 2° and 6°N, and between longitude 51° and 55°W. Soils of the region are heavily weathered and highly depleted in soil nutrients. The relief is extremely eroded and generally fat, with elevation rarely exceeding 200 m, except in two mountain ranges with peaks beyond 800m. Mean annual rainfall in the four inventory sites ranges between 1500 and 3000mm and is distributed seasonally throughout the year⁴⁹. The wet season stretches from December to July, and is usually interrupted in February or March by a short dry period, while the dry season occurs from August to November with monthly rainfall rarely exceeding 100mm. Mean temperature oscillates around 25 °C with low seasonal variation49.

The four inventory sites comprised six to 12 plots of 1.9 ha each (see next section for details), spaced by at least 500 m to each other, and located on habitats presenting contrasted topographical features⁵⁰: hilltop, slope and seasonally flooded forest (Table 2 and Appendix S1 for details). The most northerly site, *Trinité* (4°24′36″N, 53°24′47″W), comprised six plots, four on hilltops and two on seasonally fooded forest. In the second site, *Saül-Limonade* (3°33′36″N, 53°12′W), 12 plots were inventoried, four on each topographical habitat. In the third site, *Itoupé* (3°1′12″N, 53°6′W), nine plots were disposed along an altitudinal gradient, at 400, 600 and 800 m (three plots per elevation level). In the fourth and most southerly site, *Mitaraka* (2°13′12″N, 54°27′36″W), nine plots were inventoried, three on each topographical habitat. *Saül-Limonade*, *Itoupé*, *Mitaraka* are part of the National Amazonian Park of French Guiana (PAG, www.pag.fr). *Trinité* is a natural reserve part of the Network of Natural Reserve of French Guiana ([www.guyane-parcregional.fr\)](http://www.guyane-parcregional.fr).

Plot confguration and taxonomic inventories. Plots were designed using a modifcation of Gentry plots^{51,52}. The protocol consists of establishing ten parallel transects of 10×50 m emanating perpendicular to a 200 m central line (every 20 m along this line), successively oriented in alternate directions. In this sampling design, fve taxonomic groups were inventoried over diferent surfaces: tree species (1.9ha), leaf-litter ants (hereafer "ants" for simplicity, 0.12 ha), spiders (0.12 ha), fungi (1 ha) and earthworms (1 ha). Voucher and tissue

samples were collected for all taxa sampled and used to inform the taxonomic hypotheses for species lists in each site. These samples are stored in local laboratories in Kourou and are available for consultation upon request from the authors, with duplicates deposited in regional taxonomic museums (and herbaria for trees). The sampling intensity difered across groups (see details in Appendix S1), with composition data available in the four sites for soil variables, trees, ants and fungi, three sites for spiders and two sites for earthworms. The protocol used to inventory each taxonomic group in the feld and identify taxa is detailed in Appendix S1. We identifed among all sites: 1054 tree species, 171 fungi genera, 445 ant species, 81 spider genera and 65 OTU's of earthworms. Table 2 summarizes the overall taxonomic diversity among sites and within each site, represented by the species richness and the effective number of species expected from a random sampling of 2 individuals. The latter statistic was used to provide a diversity measure that gives relatively more weight than species richness to the most abundant taxa⁵³.

Soil and topographical data. Ten bulked soil cores were collected at 0-10, 10-20 and 20-30 cm depth in each plot, at each of the ten crossing points between parallel transects and the main central line (Appendix S1). They were combined into a composite 500 g sample. The latter was dried at 25 °C to reach a constant mass, then sieved to 2 mm and shipped within 3 months for physical and chemical analyses at CIRAD (France). Physical and chemical soil properties were then measured using standard soil analysis protocols⁵⁴. Physical properties corresponded to soil texture (percentage of clay, silt and sand). Chemical variables corresponded to the percentage of soil organic matter (OM), the available content of phosphorus, N and six bioavailable cations (Ca, Mg, K, Na, Al and Mn). After eliminating outlier values⁵⁵, the soil heterogeneity among plots was decomposed using a Principal Component Analysis (PCA) on the correlation matrix between normalised (Box-Cox transformation) centred-reduced soil variables values (Figs S2 and S3). The PCA showed that most plots located on seasonally flooded habitat displayed soil conditions that were very different from the two other habitats (Fig. S2). These habitats diferences were therefore emphasized in the ordination analyses described in the following section.

Data analysis. *Testing Associations between data tables*. In order to quantify and test the compositional associations among pairs of taxa (question 1a), we frst used a *Multiple Co-Inertia Analysis* (MCOIA)56 to have a visual overview of the correlations between taxonomic groups, and to detect overall compositional (all groups confounded) structures in our data. Abundance and occurrence data were Hellinger-transformed⁵⁷ prior to all analyses to reduce the weight of overly abundant taxa in each group.

We then performed "classical" *Co-Inertia Analysis* (COIA)⁵⁸ to quantify pairwise associations of composition between data tables. The *RV coefficient*⁵⁹ was calculated (at the plot level) to quantify the co-inertia of taxonomic composition between groups and the co-inertia between the composition of each group and soil variables. RV values were adjusted⁶⁰ to avoid biases due to different numbers of columns between the two compared composition matrices. The RV coefficient varies between 0 and 1, a value of 1 indicating that compared matrices are perfectly correlated.

To test RV values while taking spatial autocorrelation into account, we compared the observed coefficients with 999 null values obtained using *Moran Spectral Randomizations* (MSR)⁶¹. MSR is a flexible method providing a way to generate artifcial variables displaying spatial structures that accurately mimic the original structures. To do so, the method first consists in fitting *Moran Eigenvector's Maps* (MEM)⁶² to univariate or multivariate data, in order to detect spatial patterns at multiple spatial scales. MEM correspond to eigenvectors modelling multi-scale spatial structures in any univariate or multivariate data⁶². Then, based on which plots are connected in the "best" spatial weighting matrix (i.e., the matrix that produce the set of MEM that best fit the data⁶³), the MSR method uses a conditional simulation procedure that preserves the original structure of the data⁶¹. The spatial connections between plots in the spatial weighting matrix were modelled using graph-based confgurations (*Gabriel's* graph and *minimum spanning tree*) which have been demonstrated to be well-suited for modelling spatial structures with nested sampling designs like ours (e.g. distant sites containing six to 12 plots) 63 . A more detailed description of the method used to obtain MEM and perform the MSR test of RV values is described in Appendix S2.

RV coefficients were considered significant when no more than 5% of the null RV values were equal to or higher than the observed value. A signifcant RV between two groups indicates a signifcant coordinated turn-over of composition. We took multiple test efects into account in all of our tests by using a Šidák correction64.

It is worth noting that conceptually, COIA is very close to the *Procrustes Analysis* which usually leads to redundant results65,66. We thus performed Procrustes analyses (calculation of the Procrustes residual sum of square to quantify association between data tables⁶⁷) as a complementary approach to further verify consistency in our results.

Testing coordinated response to soil variables for each pair of groups. In order to test the infuence of environmental heterogeneity in explaining coordinated species turn-overs among pairs of taxa (questions 2a and 2b), we used the soil data and the co-inertia axes obtained from a COIA performed on each pair of taxonomic groups. We then computed the $R²$ value (adjusted to account for the number of explanatory variables) of a Redundancy Analysis (RDA) of the co-inertia axes on the set of all soil variables. Tis value was tested by comparing it with 999 null values obtained using the MSR method described above to account for spatial autocorrelation in the residuals of the RDA model. Tests were considered signifcant if less than fve percent of the null values were higher than the observed one. The relative effect of each individual variable was quantified using the sum of weighted AICc (corrected Akaike Information Criterion) values (hereafer, SWA) from a *Model Averaging with Multivariate Generalized Linear Model*⁶⁸. The R² test and SWA values allowed testing whether the co-inertia between two groups was explained by soil heterogeneity, and if so, which soil variable(s) best explained it. Finally, MEM were also used to test at which spatial scale(s) the overall soil heterogeneity and each individual soil variable explained the co-inertia among pairs of groups (see Appendix S2 for details). Indeed, MEM eigenvalues are directly correlated to Moran's *I* and therefore to the spatial scales at which variables are structured.

All analyses were performed in the R statistical environment⁶⁹, using packages *car*, *ade4*, *adespatial*, *usdm*, *cocorresp*, *MatrixCorrelation*, *vegan*, *ape* and *mglmn* (see references of these packages in Appendix S2). Te R script as well as the soil and composition data tables used to perform our analyses are provided in Appendices S3 to S9.

Data Availability

The authors confirm that, should the manuscript be accepted, the data supporting the results will be archived in an appropriate public repository such as Dryad or Figshare and the data DOI will be included at the end of the article.

References

- 1. Kraf, N. J. B., Valencia, R. & Ackerly, D. D. Functional traits and niche-based tree community assembly in an Amazonian forest. *Science* **322**, 580–582 (2008).
- 2. Condit, R., Engelbrecht, B. M. J., Pino, D., Perez, R. & Turner, B. L. Species distributions in response to individual soil nutrients and seasonal drought across a community of tropical trees. *Proc. Natl Acad. Sci. USA* **110**, 5064–5068 (2013).
- 3. Vleminckx, J. *et al.* The influence of spatially structured soil properties on tree community assemblages at a landscape scale in the tropical forests of southern Cameroon. *J. Ecol.* **105**(2), 354–366 (2017).
- 4. Scherber, C. *et al*. Bottom-up efects of plant diversity on multitrophic interactions in a biodiversity experiment. *Nature* **468**, 553–556 (2010).
- 5. Basset, Y. *et al*. Arthropod diversity in a tropical forest. *Science* **338**, 1481–1484 (2012).
- 6. Schuldt, A. *et al*. Multitrophic diversity in a biodiverse forest is highly nonlinear across spatial scales. *Nat. Commun.* **6**, 10169, <https://doi.org/10.1038/ncomms10169>(2015).
- 7. Vasconcelos, H. L. *et al*. Congruent spatial patterns of ant and tree diversity in Neotropical savannas. *Biodivers. Conserv.* **28**, 1075–1089 (2019).
- 8. Filete de Morais, G., Guilherme dos Santos Ribas, L., Carlo Gonçalves Ortega, J., Heino, J. & Mauricio Bini, L. Biological surrogates: A word of caution. *Ecol. Indic.* **88**, 214–218 (2018).
- 9. Pimm, S. L. *et al*. Te biodiversity of species and their rates of extinction, distribution, and protection. *Science* **344**(6187), 1246752 (2014)
- 10. Schuldt, A. *et al*. Belowground top-down and aboveground bottom-up efects structure multitrophic community relationships in a biodiverse forest. *Sci. Rep.* **7**, 4222, <https://doi.org/10.1038/s41598-017-04619-3>(2017).
- 11. Didham, R. K., Edwards, O. R., Leather, S. R. & Basset, Y. Arthropod diversity and the future of all-taxa inventories. *Insect Conserv. Diver.* **6**, 1–4 (2013).
- 12. Bagchi, R. *et al*. Pathogens and insect herbivores drive rainforest plant diversity and composition. *Nature* **506**, 85–88 (2014).
- 13. Duan, M. *et al*. Disentangling efects of abiotic factors and biotic interactions on cross-taxon congruence in species turnover patterns of plants, moths and beetles. *Sci. Rep.* **6**, 23511,<https://doi.org/10.1038/srep23511> (2016).
- 14. Zhang, K. *et al*. Plant diversity accurately predicts insect diversity in two tropical landscapes. *Mol. Ecol.* **25**, 4407–4419 (2016).
- 15. Santi, E. *et al*. Simple to sample: Vascular plants as surrogate group in a nature reserve. *J. Nat. Conserv.* **18**(1), 2–11 (2010).
- 16. Fine, P. V. A. *et al*. Insect herbivores, chemical innovation and the evolution of habitat specialization in Amazonian trees. *Ecology* **94**, 1764–1775 (2013).
- 17. Schafers, A. P., Raemakers, I. P., Sykora, K. V. & Ter Braak, C. J. F. Arthropod assemblages are best predicted by plant species composition. *Ecology* **89**, 782–794 (2008).
- 18. Schmitt, J. P. *et al*. Assessment of tree species richness as a surrogate for macrofungal species richness. *Biol. Conserv.* **121**, 99–110 (2004).
- 19. Lawes, M. J., Moore, A. M., Andersen, A. N., Preece, N. D. & Franklin, D. C. Ants as ecological indicators of rainforest restoration: community convergence and the development of an Ant Forest Indicator Index in the Australian wet tropics. *Ecol. Evol.* **7**, 8442–8455 (2017).
- 20. Jouquet, P., Dauber, J., Lagerlöf, J., Lavelle, P. & Lepage, M. Soil invertebrates as ecosystem engineers: Intended and accidental eVects on soil and feedback loops. Agriculture, ecosystems and environment. *Appl. Soil Ecol.* **32**, 153–164 (2006).
- 21. Schuldt, A. & Staab, M. Tree species richness strengthens relationships between ants and the functional composition of spider assemblages in a highly diverse forest. *Biotropica* **47**, 339–346 (2015).
- 22. Smith, S. E. & Read, D. J. *Mycorrhizal Symbiosis*. Academic Press, London, UK (2008).
- 23. Philippot, L., Raaijmakers, J. M., Lemanceau, P. & van der Putten, W. H. Going back to the roots: the microbial ecology of the rhizosphere. *Nat. Rev. Microbiol.* **11**, 789–799 (2013).
- 24. Janzen, D. H. Herbivores and the number of tree species in tropical forests. *Am. Nat.* **104**, 501–528 (1970).
- 25. Connell, J. H. *On the Role of Natural Enemies in Preventing Competitive Exclusion in Some Marine Animals and in Rain Forest Trees*. In: Den Boer, P. J. and Gradwell, G. R., Eds, Dynamics of Populations, Centre for Agricultural Publishing and Documentation, Wageningen, The Netherlands (1971).
- 26. Reich, P. B. *et al*. Linking litter calcium, earthworms and soil properties: a common garden test with 14 tree species. *Ecol. Lett.* **8**, 811–818 (2005).
- 27. Neves, D. M. *et al.* Dissecting a biodiversity hotspot: The importance of environmentally marginal habitats in the Atlantic Forest Domain of South America. *Divers. Distrib.* **00**, 1–12 (2017).
- 28. Schappe, T. et al. The role of soil chemistry and plant neighbourhoods in structuring fungal communities in three Panamanian rainforests. *J. Ecol.* **105**, 569–579 (2017).
- 29. Ruokolainen, K., Tuomisto, H., Macía, M. J., Higgins, M. A. & Yli-Halla, M. Are foristic and edaphic patterns in Amazonian rain forests congruent for trees, pteridophytes and Melastomataceae? *J. Trop. Ecol.* **23**, 13–25 (2007).
- 30. Lemes Landeiro, V. *et al*. How far can we go in simplifying biomonitoring assessments? An integrated analysis of taxonomic surrogacy, taxonomic sufficiency and numerical resolution in a megadiverse region. *Ecol. Indic.* **23**, 366-373 (2012).
- 31. Decaëns, T. *et al*. Biodiversity loss along a gradient of deforestation in Amazonian landscapes. *Conserv. Biol.* **32**(6), 1380–1391 (2018).
- 32. Slik, F. J. W. *et al*. An estimate of the number of tropical tree species. *P. Natl. Acad. Sci. USA* **12**(33), E4628–E4629 (2015).
- 33. Socolar, J. B., Gilroy, J. J., Kunin, W. E. & Edwards, D. P. How Should Beta-Diversity Inform Biodiversity Conservation? *Trends Ecol. Evol.* **31**(1), 67–80 (2016).
- 34. Ríos‐Casanova, L., Valiente‐Banuet, A. & Rico‐Gray, V. Ant diversity and its relationship with vegetation and soil factors in an alluvial fan of the Tehuacán Valley, Mexico. *Acta Oecol.* **29**, 316–323 (2006).
- 35. Donoso, D. A., Johnston, M. K. & Kaspari, M. Trees as templates for tropical litter arthropod diversity. *Oecologia* **164**, 201–211 (2010)
- 36. Folgarait, P. J. Ant biodiversity and its relationship to ecosystem functioning: A review. *Biodivers. Conserv.* **7**, 1221–1244 (1998).
- 37. Del Toro, I., Ribbons, R. R. & Pelini, S. L. The little things that run the world revisited: A review of ant-mediated ecosystem services and disservices (Hymenoptera: Formicidae). *Myrmecol. News* **17**, 133–146 (2012).
- 38. Andersen, A. N. & Majer, J. D. Ants show the way Down Under: Invertebrates as bioindicators in land management. *Front. Ecol. Environ.* **2**, 291–298 (2004).
- 39. Peay, K. G., Baraloto, C. & Fine, P. V. A. Strong coupling of plant and fungal community structure across western Amazonian rainforests. *ISME J.* **7**(9), 1852–1861 (2013).
- 40. Bonkowski, M., Grifths, B. S. & Ritz, K. Food preferences of earthworms for soil fungi. *Pedobiologia* **44**, 666–676 (2000).
- 41. Cardoso, P., Pekar, S., Jocqué, R. & Coddington, J. A. Global Patterns of Guild Composition and Functional Diversity of Spiders. *PLoS ONE* **6**(6), e21710, <https://doi.org/10.1371/journal.pone.0021710> (2011).
- Tedersoo, L. & Smith, M. E. Lineages of ectomycorrhizal fungi revisited: Foraging strategies and novel lineages revealed by sequences from belowground. *Fungal Biol. Rev.* **27**, 83–99 (2013).
- 43. Decaëns, T. *et al*. DNA barcoding reveals diversity patterns of earthworm communities in remote tropical forests of French Guiana. *Soil Biol. Biochem.* **92**, 171–183 (2016).
- 44. Hättenschwiler, S. & Bracht Jørgensen, H. Carbon quality rather than stoichiometry controls litter decomposition in a tropical rain forest. *J. Ecol.* **98**, 754–763 (2010).
- 45. Quesada, C. A. *et al*. Basin-wide variations in Amazon forest structure and function are mediated by both soils and climate. *Biogeosciences* **9**, 2203–2246 (2012).
- 46. Vleminckx, J. *et al*. Impact of fne-scale edaphic heterogeneity on tree species assembly in a central African rainforest. *Journal of Vegetation Science* **26**(1), 134–144 (2015).
- 47. Seidler, T. G. & Plotkin, J. B. Seed dispersal and spatial pattern in tropical trees. *PloS Biol.* **4**, 2132–2137 (2006).
- 48. Barsoum, N., Bruce, C., Forster, J., Ji, Y.-Q. & Yu, D. W. Te devil is in the detail: Metabarcoding of arthropods provides a sensitive measure of biodiversity response to forest stand composition compared with surrogate measures of biodiversity. *Ecol. Indic.* **101**, 313–323 (2019).
- 49. Gourlet-Fleury, S., Guehl, J. M. & Laroussinie, O. Ecology and management of a neotropical rainforest: lessons drawn from Paracou, a long-term experimental research site in French Guiana (eds Gourlet-Fleury, S., Guehl, J.-M. & Laroussinie, O.). 326p (2004).
- 50. Ferry, B., Morneau, F. O., Bontemps, J.-D., Blanc, L. & Freycon, V. Higher treefall rates on slopes and waterlogged soils result in lower stand biomass and productivity in a tropical rain forest. *J. Ecol.* **98**, 106–116 (2010).
- 51. Phillips, O. L. *et al*. Efcient plot-based foristic assessment of tropical forests. *J. Trop. Ecol.* **19**, 629–645 (2003).
- 52. Baraloto, C. *et al*. Rapid simultaneous estimation of aboveground biomass and tree diversity across Neotropical forests: A comparison of feld inventory methods. *Biotropica* **45**(3), 288–298 (2013).
- 53. Dauby, G. & Hardy, O. J. Sampled-based estimation of diversity sensu stricto by transforming Hurlbert diversities into efective number of species. *Ecography* **34**, 001–012 (2011).
- 54. Pansu, M. & Gautheyrou, J. Handbook of Soil Analysis: Mineralogical, Organic and Inorganic Methods. Springer, Berlin, Germany (2006).
- 55. Zuur, A. F., Ieno, E. N. & Elphick, C. S. A protocol for data exploration to avoid common statistical problems. *Methods Ecol. Evol.* **1**, 3–14 (2010).
- 56. Bady, P., Dolédec, S., Dumont, B. & Fruget, J.-F. Multiple co-inertia analysis: a tool for assessing synchrony in the temporal variability of aquatic communities. *Cr. Biol.* **327**(1), 29–36 (2004).
- 57. Legendre, P. & Gallagher, E. D. Ecologically meaningful transformations for ordination of species data. *Oecologia* **129**, 271–280 (2001)
- 58. Dolédec, S. & Chessel, D. Co-inertia analysis: an alternative method for studying species–environment relationships. *Freshwater Biol.* **31**, 277–294 (1994).
- 59. Robert, P. & Escoufer, Y. "A Unifying tool for linear multivariate statistical methods: the RV-coefcient". *Appl. Statist.* **25**(3), 257–265 (1976).
- 60. Mayer, C. D., lorent, J. & Horgan, G. W. Exploratory analysis of multiple omics datasets using the adjusted RV coefcient. *Stat. Appl. Genet. Mol. Biol.* **10**, article 14 (2011).
- 61. Wagner, H. H. & Dray, S. Generating spatially constrained null models for irregularly spaced data using Moran spectral randomization methods. *Methods Ecol. Evol.* **6**, 1169–1178 (2015).
- 62. Dray, S., Legendre, P. & Peres-Neto, P. R. Spatial modelling: a comprehensive framework for principal coordinate analysis of neighbour matrices (PCNM). *Ecol. Model.* **196**, 483–493 (2006).
- 63. Bauman, D., Drouet, T., Fortin, M. J. & Dray, S. Optimizing the choice of a spatial weighting matrix in eigenvector-based methods. *Ecology* **99**(10), 2159–2166 (2018).
- 64. Šidák, Z. K. "Rectangular Confdence Regions for the Means of Multivariate Normal Distributions". *J. Am. Stat. Assoc.* **62**(318), 626–633 (1967).
- 65. Dray, S., Chessel, D. & Tioulouse, J. Co-inertia analysis and the linking of ecological tables. *Ecology* **84**(11), 3078–3089 (2003).
- 66. Legendre, P. & Legendre, L. *Numerical Ecology*, 3rd English edn. Elsevier Science BV, Amsterdam (2012).
- 67. Jackson, D. A. PROTEST: a PROcrustean Randomization TEST of community environment concordance. *Ecoscience* **2**, 297–303 (1995).
- 68. Nakamura, A. *et al.* The role of human disturbance in island biogeography of arthropods and plants: an information theoretic approach. *J. Biogeogr.* **42**(8), 1406–1417 (2015).
- 69. R Development Core Team R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria, <http://www.R-project.org> (2018).

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Author Contributions

C.B. was principal investigator for the project. J.V. led the data analysis and the writing of the manuscript, with the help of C.B., H.S., T.D., M.F. and J.O. The collection and identification of abundance/species occurrence data was coordinated by C.B., J.E., G.J., A.D. and P.P. for trees, H.S. and M.R. for fungi, J.O. and M.F. for ants, V.V. for spiders and T.D. and E.L. for earthworms. All authors contributed to the proofreading of the manuscript.

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