

# Tree diversity is key for promoting the diversity and abundance of forest-associated taxa in Europe

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**Decision date:** 28-Sep-2019

This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as doi: [10.1111/oik.06290].

This is the peer reviewed version of the following article: Ampoorter, E., Barbaro, L., Jactel, H., Baeten, L., Boberg, J., Carnol, M., Castagneyrol, B., Charbonnier, Y., Dawud, S.M., Deconchat, M., Smedt, P.D., Wandeler, H.D., Guyot, V., Hättenschwiler, S., Joly, F.-X., Koricheva, J., Milligan, H., Muys, B., Nguyen, D., Ratcliffe, S., Raulund-Rasmussen, K., Scherer-Lorenzen, M., van der Plas, F., Keer, J.V., Verheyen, K., Vesterdal, L. and Allan, E. (2020), Tree diversity is key for promoting the diversity and abundance of forest-associated taxa in Europe. *Oikos*, 129: 133-146, which has been published in final form at <https://doi.org/10.1111/oik.06290>. This article may be used for non-commercial purposes in accordance with Wiley Terms and Conditions for self-archiving.

## Abstract

Plant diversity is an important driver of diversity at other trophic levels, suggesting that cascading extinctions could reduce overall biodiversity. Most evidence for positive effects of plant diversity comes from grasslands. Despite the fact that forests are hotspots of biodiversity, the importance of tree diversity, in particular its relative importance compared to other management related factors, in affecting forest-associated taxa is not well known. To address this, we used data from 183 plots, located in different forest types, from Mediterranean to Boreal, and established along a climatic gradient across six European countries (FunDivEUROPE project). We tested the influence of tree diversity, tree functional composition (i.e. functional trait values), forest structure, climate and soil on the diversity and abundance/activity of nine taxa (bats, birds, spiders, microorganisms, earthworms, ungulates, foliar fungal pathogens, defoliating insects and understorey plants) and on their overall diversity and abundance/activity (multidiversity, multiabundance/activity). Tree diversity was a key driver of taxon-level and overall forest-associated biodiversity, along with tree functional composition, forest structure, climate and soil. Both tree species richness and functional diversity (variation in functional trait values) were important. The effects of tree diversity on the abundance/activity of forest-associated taxa were less consistent. Nonetheless, spiders, ungulates and foliar fungal pathogens were all more abundant/active in diverse forests. Tree functional composition and structure were also important drivers of abundance/activity: conifer stands had lower overall multidiversity (although the effect was driven by defoliating insects), while stands with potentially tall trees had lower overall multiabundance/activity. We found more synergies than trade-offs between diversity and abundance/activity of different taxa, suggesting that forest management can promote high diversity across taxa. Our results clearly show the high value of mixed forest stands for multiple forest-associated taxa and indicate that multiple dimensions of tree diversity (taxonomic and functional) are important.

**Keywords:** climate, forest-associated taxa, forest structure, soil conditions, tree functional composition , tree diversity

## Introduction

Understanding how changes in plant diversity affect the diversity of other trophic groups is of fundamental interest to understand trophic interactions. It is also crucial to predict cascading extinctions and the impact of multiple taxa on ecosystem functioning and ecosystem services (Cardinale et al. 2006, Soliveres et al. 2016). Plant diversity might have bottom-up effects on higher trophic levels and increase the diversity of consumers, such as herbivores and pathogens, by providing a greater diversity of resources for specialist consumers, more opportunities for diet mixing for generalist consumers or by increasing overall resource levels (Siemann 1998, Borer et al. 2012). Several studies have shown that diverse plant communities support a higher diversity of other taxa living in the same ecosystem (Castagneyrol and Jactel 2012, Schuldt et al. 2019). However, a lot of these studies have been carried out in grasslands (Siemann et al. 1998, Haddad et al. 2009, Scherber et al. 2010), while fewer have been conducted in mature forest stands (Sobek et al. 2009, Cavard et al. 2011, Fornoff et al. 2019). This limits our knowledge of whether switching from monocultures (the traditional focus of intensive forest management) to mixed plantings would promote high diversity and abundance of species that live in the same forest stand, i.e. forest-associated taxa. It is crucial to address this because forests provide habitats for many different organisms (Lindenmayer and Franklin 2002) and a better understanding of how tree diversity affects other groups could be used to optimise forest management for biodiversity conservation.

Tree diversity may affect the diversity and abundance of forest-associated taxa through several mechanisms. Organisms that feed on trees, such as defoliating insects and foliar fungal pathogens, are directly affected by tree species composition. Their diversity is expected to be higher in stands with more tree species (Basset et al. 2012, Castagneyrol and Jactel 2012) because diverse stands provide more and/or better quality food for consumers (Potts et al. 2003, Haddad et al. 2009). However, tree diversity may have a range of effects on the damage caused by different consumers. Tree species may have reduced damage from specialist herbivores in diverse stands (associational resistance hypothesis), however they may suffer enhanced damage from generalists (associational susceptibility hypothesis), compared to monoculture stands (Plath et al. 2012). Effects of tree diversity on forest-associated taxa may also be indirectly mediated by the impacts of the trees on environmental conditions, such as on light availability and quality (Messier et al. 1998), soil water availability (Barbier et al. 2009), soil acidity and nutrient availability (Carnol and Bazgir 2013, Augusto et al. 2015), and characteristics of the litter layer (Hobbie et al. 2006). Diverse forest stands generally have higher complexity (Brose 2003, Cavard et al. 2011), which might provide more niches and therefore increase the diversity and abundance of

understorey plants (Ampoorter et al. 2016) and earthworms (Chamagne et al. 2016). In grasslands, plant diversity has been shown to have cascading effects and to increase diversity at multiple trophic levels (Haddad et al. 2009, Scherber et al. 2010). As herbivores feed on trees and predators eat herbivores, it has been hypothesized that predator diversity also increases with increasing tree diversity (Castagneyrol and Jactel 2012). Multiple observational and experimental studies have provided support for this hypothesis (Siemann 1998, Sobek et al. 2009). Different components of tree diversity might have different effects on forest-associated taxa. For example, tree functional diversity, i.e. the variation in functional trait values, might have stronger effects than species richness, if the traits measure interspecific differences (e.g. nutritional differences) relevant for forest-associated taxa (e.g. Mason et al. 2005). Tree species evenness might also be important, because it affects the relative abundance of those traits that determine niche space for forest-associated taxa (Petchey and Gaston 2006).

Tree functional composition, i.e. the particular functional trait values found in a stand, could affect forest-associated taxa as well. For instance, the leaves of fast growing tree species have high nutrient concentrations and are likely to be more palatable for defoliating insects and foliar fungal pathogens than leaves from slow growing species (Rowe et al. 2006). Forest management may not only alter tree diversity and tree functional composition, but also the structure of forest stands. Horizontal heterogeneity refers to the spatial distribution of the trees (i.e. regular or random, clustered or dispersed), while vertical heterogeneity is related to the number of forest layers. A higher horizontal and vertical heterogeneity in forest structure increases available niche space, provides more refuges and opportunities for isolation and divergent adaptation, and can therefore enable the coexistence of more species (Cavard et al. 2011). Apart from these management-related factors, abiotic drivers, such as climate (Engelbrecht et al. 2007, Butchart et al. 2010) and soil (Rosch et al. 2002, Silvertown 2004), may also affect the diversity of the forest-associated taxa. Determining the relative importance of tree diversity, tree functional composition, forest structure, climate and soil in affecting the diversity of forest-associated taxa may help to understand the mechanisms underlying interactions between trees and forest-associated taxa.

Previous research mainly focused on the impact of one potential driver on forest biodiversity. However, to develop better forest management plans and to understand the importance of tree diversity as a driver of associated diversity, we need to determine the relative importance of tree diversity alongside other management-related and abiotic drivers in affecting forest biodiversity. Moreover, these relations have predominantly been explored for individual taxa (Zhang et al. 2016) while taxa are likely to respond differently to changes in tree diversity, tree functional composition, forest structure, climate and soil. For

example, vertical stratification predominantly influences taxa that occur in several forest layers, like birds, as a more complex forest structure would result in more abundant and diverse prey resources, allowing higher levels of predator functional diversity (Barbaro et al. 2014). Tree diversity could have stronger effects on lower trophic levels because these are more directly associated with the trees, in accordance with the effect of plant diversity in grasslands (Scherber et al. 2010). In addition, diversity and abundance of a specific taxon may be driven by different factors (Boulangeat et al. 2012) making it important to compare responses of diversity with those of abundance. To determine the overall responses of diversity and abundance, multidiversity or multiabundance indices can be used (Allan et al. 2014) and these may enable us to identify general responses across a wide range of organism groups.

In this study, part of the FunDivEUROPE project (Baeten et al. 2013), we examined multiple drivers of the diversity and abundance of various forest-associated taxa. The study plots were located in mature European forests of six contrasting types that differ in tree species composition and that are spread along a large climatic gradient. We aimed to quantify the relative effects of tree diversity, tree functional composition, forest structure, climate and soil on the diversity and abundance (or activity, see methods) of nine key forest-associated taxa (bats, birds, spiders, micro-organisms, earthworms, ungulates, pathogens, defoliating insects and understorey plants). These taxa represent different trophic levels above and belowground, they are important for conservation or nature tourism and/or are key drivers of ecosystem functioning and ecosystem services providers. We examined each taxon separately but to quantify overall responses, we also used multidiversity and multiabundance/activity indices. In addition, we investigated how the diversity and abundance/activity of the different forest-associated taxa were related to each other.

## Material and methods

### *Forest plots*

The Exploratory Platform of the FunDivEUROPE project includes six regions representing the main forest types in Europe: boreal forest (Finland), hemiboreal forest and nemoral coniferous and mixed broadleaved-coniferous forest (Poland), beech forest (Germany), mountainous beech forest (Romania), thermophilous deciduous forest (Italy) and Mediterranean mixed forest (Spain) (Barbati et al. 2007). In each region, three to five target tree species, which are regionally common and/or of silvicultural importance, were selected. The total tree species pool consisted of the coniferous species *Abies alba*, *Picea abies*, *Pinus nigra* and *Pinus sylvestris*, and the broadleaved species *Acer pseudoplatanus*, *Betula*

*pendula/pubescens*, *Carpinus betulus*, *Castanea sativa*, *Fagus sylvatica*, *Fraxinus excelsior*, *Ostrya carpinifolia*, *Quercus robur/petraea*, *Quercus cerris*, *Quercus faginea* and *Quercus ilex*. Based on these regional tree species pools, 28 to 43 plots of 30 m × 30 m were chosen per region, in mature forests, to cover a gradient in species richness from monoculture stands to mixtures containing all species from the regional pool (i.e. 3 to 5 species mixtures). Plots were surrounded by a buffer zone 10 m wide, which had a similar forest composition and structure, to avoid edge effects. For each level of species richness, plots with different tree species compositions were included. This resulted in 209 plots in total over the six regions. More information on region-specific tree species pools, tree species combinations and number of plots can be found in Supporting Information Appendix S1. Intraregional covariation between environmental variables (bedrock, soil, topography) and tree species richness was avoided, so that past silvicultural management and/or random species assembly were the key drivers of tree diversity. Some admixture of non-target species was unavoidable but we ensured that the target species contributed: 1) >90% of the number of individuals, 2) >90% of total basal area, and 3) were always represented by more than two individuals per plot. We therefore focused on target species richness in our analyses, but we also tested for the effect of admixed tree species richness. All forests where these plots are located are considered ancient forests, i.e. they have been continuously forested at least since the oldest available land-use maps. They have also been (at least historically) managed for forestry and are currently in the mid to late stem exclusion, understorey reinitiation or old-growth development stage (i.e. excluding very young stands). For more information on the regions, tree species combinations, plot selection criteria and plot-level information, see Baeten et al. (2013) and Jucker et al. (2014).

#### *Forest-associated taxa*

Within all plots, in all regions, data on community composition and abundance/activity were collected for bats, birds, arboreal spiders in the shrub layer, microorganisms, earthworms, ungulates (no data on community composition), foliar fungal pathogens, defoliating insects and understorey plants. Different ways of quantifying diversity and abundance/activity were used for the different taxa. We could not obtain abundance data for all groups and used the activity of defoliating insects (herbivory), pathogens (infection) and ungulates (browsing damage) as proxies. Information on the data collection can be found in Supporting Information Appendix S2 and raw diversity and abundance/activity data can be seen in Fig. 3 and Fig. 4. We visited all forest plots once to ensure that we sampled as many plots as possible because we wanted to maximise sample size and spatial coverage to increase statistical power, and to sample all plots within the same region within a short time period to limit seasonal variation. Sampling data in many

plots in several countries is challenging and it would simply not have been possible to sample each plot more intensively. Sampling periods varied across the year between regions to account for differences in climatic conditions and leaf phenology. Sampling efforts were relatively small for taxa like bats and birds but analysis of sample completeness for these taxa indicated that we sampled an adequate proportion of the European species pool (see Barbaro et al. 2019 for a fuller analysis of sample completeness for birds and bats). Species pools of forest-associated taxa differed considerably between regions. An overview of the variation in species numbers between regions is shown for each of our 8 organism groups in the Supporting Information Appendix S3. In general, our goal was not to quantify absolute diversities of the different taxonomic groups (and hence to sample the complete communities in all plots) or to understand the response of each taxonomic group in detail. In contrast, this study focussed on relative differences in diversity and abundance of forest-associated taxa and on drawing general conclusions on how the forest-associated taxa respond to different drivers.

In order to test for the overall response of diversity and abundance/activity of forest-associated taxa, we calculated “multidiversity” and “multiabundance/activity” indices following Allan et al. (2014). These measures quantify the average proportional diversity and abundance/activity respectively across taxonomic groups. For each taxon, diversity and abundance/activity values were scaled to the maximum value observed in any plot (however, scaling to maxima per region did not change the results, see Table S9.1), so that all taxa were weighted equally, i.e. species rich taxa like understorey plants are weighted the same as species poor taxa like bats. Then, scaled values were averaged per plot over the different taxa.

#### *Drivers of biodiversity*

We tested several components of tree diversity, tree functional composition, forest structure, climate and soil as potential drivers of the diversity and abundance/activity of forest-associated taxa. *Tree diversity* was measured as target tree species richness (SR), admixed tree SR, target species evenness and target species functional diversity. Functional diversity was calculated as abundance weighted functional dispersion, which measures the distance between tree species in multivariate trait space (Laliberté and Legendre 2010; using the function dbFD in the FD package in R; Laliberté 2014). Higher values indicate greater functional dissimilarity. Functional traits were measured, for each tree species in each region, on freshly fallen leaf litter. Relative abundances of the tree species were calculated from basal area. We selected traits related to palatability for defoliating insects and foliar fungal pathogens and to decomposability: specific leaf area (SLA), leaf pH and the proportion of nitrogen (N), lignin,



hemicellulose, condensed tannins and soluble phenolics in the leaf. Maximum tree height and mean tree lifespan were also included because they refer to the structure and stability of the tree layer (information on the data collection in Supporting Information Appendix S2).

*Tree functional composition* was determined in several ways. We fitted the community weighted mean (CWM; Ricotta and Moretti 2011) of leaf N in all models, as this is a key trait for defoliating insects and foliar fungal pathogens and is therefore expected to have direct or indirect effects on many taxa. The CWM is calculated by multiplying the trait value of each tree species by its relative abundance (basal area) and then summing across all tree species in the plot. In addition to CWM N, we included measures related to tree functional composition that aimed to represent the two major axes of plant strategy variation, which are (i) a resource acquisition axis, differentiating fast growing acquisitive species from slow growing conservative species (leaf % N is also related to this axis but we fitted it separately as it could directly affect nutrient levels for consumers and decomposers), and (ii) a tree height axis, differentiating small and large plants (Diaz et al. 2016). We explored different ways of representing these two axes in our models (Supporting Information Appendix S2). The best models had the resource axis represented as the proportion of conifers in the plots and the size axis by maximum tree height.

*Forest structure* was measured in each plot as (i) mean basal area of the target tree species, (ii) horizontal heterogeneity, i.e. the coefficient of variation of the tree-to-tree distances, and (iii) vertical heterogeneity. To calculate vertical heterogeneity, we first used measures of the cover of the tree layer (diameter > 7.5cm), shrub layer (height > 1.3m and diameter < 7.5cm) and understorey layer (height < 1.3m) which were taken in three quadrats per plot (Supporting Information Appendix S2). The cover of the tree layer was estimated for each quadrat while looking upwards and it measures the crown cover of all trees with stems > 7.5cm diameter. We first averaged the cover of each of layer to produce one average cover per plot and then calculated vertical heterogeneity as the Shannon diversity of these three layers. Maximum heterogeneity occurs when tree, shrub and understorey cover values are the same.

Information on *climate* (annual mean temperature, annual precipitation; WorldClim 2015) and *soil* conditions (soil pH and C/N) in each plot was also included (soil sampling methodology in Supporting Information Appendix S2).

#### *Data analysis*

All analyses were performed in R 3.1.1 (R Core Team 2014). From the 209 plots, we only used those plots for which diversity (ungulates excluded) and abundance/activity values of all nine taxa were



available (183 plots). More information on which plots were included can be found in the Supporting Information Appendix S1. All explanatory and response variables were scaled to a mean of 0 and a standard deviation of 1, to allow us to compare effect sizes between explanatory and response variables. Abundance/activity values were first log transformed. All correlations between explanatory variables were  $\leq 0.6$ , except for tree species richness and functional diversity (Supporting Information Appendix S4). We fitted both variables in the models as we were interested in both effects. However, we explored the sensitivity of our results to this correlation by fitting models with only tree species richness or only functional diversity (Supporting Information Appendix S5-6). Variance inflation factors (VIFs) were also calculated to check for multicollinearity. We used the *car* package in R to calculate VIFs for all explanatory variables and found that all VIFs were  $< 4.5$  (Supporting Information Appendix S7). VIFs  $< 10$  indicate that collinearity is not an issue for the analysis (Dormann et al. 2013). All of the effects that we present are partial effects, after variables have been corrected for each other (Schiegg 2010). As the variables are also scaled, these partial coefficients provide a measure of the relative importance of each variable, similar to a variance decomposition analysis. Next, we calculated correlations between the diversity of all taxa, between log-transformed abundance/activity of all taxa, and between diversity and log-transformed abundance/activity for each taxon (Supporting Information Appendix S8).

We used linear mixed effects models to determine the relative effects of all explanatory variables on the response variables. All models included random effects for region and for target tree species composition to account for the non-independence between plots within the same region and replicate tree species compositions, respectively. Models were fitted using *lmer* in the *lme4* package (Bates et al. 2013). Model structure, in R code, was:

$$y = \text{Target SR} + \text{Evenness} + \text{Admixed SR} + \text{Functional diversity} + \text{Proportion conifers} + \text{CWM height} + \text{CWM N} + \text{Basal area} + \text{Horizontal heterogeneity} + \text{Vertical heterogeneity} + \text{Temperature} + \text{Precipitation} + \text{Soil pH} + \text{Soil CN} + (1|\text{Region}) + (1|\text{Tree species composition})$$

Parameter estimation was done using restricted maximum likelihood in full models. For each full model, we calculated the 95 % confidence intervals around each parameter by bootstrapping, with 10 000 iterations, using the *confint* function in the *lme4* package. Means and confidence intervals are reported in Supporting Information Appendix S9. For each model, we calculated the proportion of the deviance explained by the fixed factors only (marginal  $R^2$ ) or by the random and fixed effects (conditional  $R^2$ ), according to Nakagawa and Schielzeth (2013) (piecewiseSEM package; Lefcheck 2016).

## Results

Explanatory variables explained 37-65% of the variation in diversity and 36-53% of the variation in abundance/activity for spiders, defoliating insects and understorey plants but only 4-23% of the variation in diversity and 10-16% of the variation in abundance/activity for bats, birds, microorganisms, earthworms, ungulates and foliar fungal pathogens (Fig. 1, Fig. 2). Most taxa were affected by several parameters, only microorganism diversity was not affected by any variables tested. The forest parameters explained 44% of the variation in overall multidiversity and 35% of the variation in overall multiabundance/activity. Conditional  $R^2$  values were clearly higher than marginal  $R^2$  values, suggesting that the random effects of region and target tree species composition also accounted for a lot of variation in the data. This means that our fixed forest parameters did not capture all of the variation between regions and forest stands with different tree species combinations. However, as we were not interested in the effects of these terms, and only wanted to account for the non-independence between plots within the same region or with the same tree species combination, we do not further discuss the variation accounted for by the random effects.

Tree diversity increased the diversity and abundance/activity of many forest-associated taxa. Target species richness (SR) increased the diversity and abundance/activity of spiders and foliar fungal pathogens, the diversity of defoliating insects and the activity of ungulates. However, note that the effects on spider abundance may have been driven by more intense sampling in diverse stands (no effect on mean abundance per tree species, see Supporting Information Appendix S2). Target SR reduced the diversity and abundance of birds. Tree functional diversity tended to increase the diversity of all taxa, especially birds. Overall, both target SR and tree functional diversity significantly increased multidiversity, showing that they have generally positive effects on the diversity of forest-associated taxa. Due to the strong correlation between tree SR and tree functional diversity (Supporting Information Appendix S4), we repeated all analyses without tree functional diversity. In these models tree SR had a neutral effect on bird diversity while other effects were unchanged (Supporting Information Appendix S5). However, when bird models were fitted without tree SR, the effect of functional diversity remained (data not shown), showing that functional diversity effects are robust to any correlation with species richness. Multiabundance/activity did not significantly increase with tree species richness and functional diversity. Tree species evenness reduced spider diversity and increased bat activity. In contrast, admixed species richness had no effects on any taxa.

Tree functional composition was also important for several forest-associated taxa. The diversity and abundance/activity of foliar fungal pathogens and defoliating insects was lower in stands with a high proportion of conifers, while spider diversity was higher in conifer stands. Overall multidiversity was also lower in stands dominated by conifers, but this result should be treated with caution as it is mainly driven by the defoliating insects (no significant effect on multidiversity without defoliators:  $\beta=0.02\pm0.3$ , slope  $\pm$  95% confidence intervals) and their diversity is difficult to assess accurately in coniferous stands (Guyot et al. 2016). Forest stands with a higher CWM height (thus a higher proportion of potentially tall trees) had a less diverse and less abundant earthworm community and a lower understorey cover but a higher abundance of microorganisms. In general, forest stands with a higher CWM height had a significantly lower multiabundance/activity (although this effect was not significant if multiabundance was standardised per region, Table S9.1). Forest stands dominated by trees with nitrogen-rich leaves had a higher diversity of spiders and a higher understorey cover but a lower diversity of foliar fungal pathogens.

Forest structure seemed to be less important for the diversity and abundance/activity of forest-associated taxa than tree diversity and tree functional composition. Plots with a higher mean basal area contained a less diverse spider and understorey plant community. Higher vertical heterogeneity was associated with lower bat diversity and activity but a higher diversity of defoliating insects and higher activity of ungulates. Understorey plant cover was used in the calculation of vertical heterogeneity. We therefore included vertical heterogeneity in the understorey plant abundance model to account for it but do not discuss its effect. Overall, vertical heterogeneity increased multiabundance/activity, also when understorey plants were excluded from the calculation ( $\beta=0.13\pm0.12$ ; slope  $\pm$  95% confidence intervals).

The climate variables (temperature, precipitation) and soil conditions (soil pH, soil C/N) had rather minor effects on the forest-associated taxa. Multiabundance/activity was lower in warmer areas (only if standardised across regions, Table S9.1) and although the results were not significant, all taxa, except microorganisms, had lower diversity at higher precipitation levels. Higher diversity and activity of foliar fungal pathogens and lower activity of defoliating insects were found in soils with higher pH. The activity of defoliating insects was lower in soils with higher soil C/N.

The diversities of the different forest-associated taxa were often related. In general, there were more positive than negative correlations between diversities (60% were positive). The average correlation was 0.02 and thus also slightly positive (Fig. 3). We found strong correlations between multidiversity and the diversities of defoliating insects, foliar fungal pathogens and earthworms. Diversities of pathogens and

defoliating insects were strongly positively correlated, while diversities of understorey plants, bats and birds were all negatively correlated with diversities of microorganisms and pathogens. The proportion of positive correlations was higher between the abundance/activity of the taxa (68%) where the average correlation was 0.04 (Fig. 4). Multiabundance/activity was strongly correlated with the abundance/activity of ungulates, earthworms, spiders and bats. Abundance/activity of ungulates, earthworms, spiders and bats were negatively correlated with the abundance of microorganisms.

Diversity and abundance/activity were correlated for defoliating insects, pathogens, earthworms, spiders, birds and bats (Supporting Information Appendix S8), which means that effects on diversity may be driven by changes in abundance. For this reason, we refitted the same linear mixed effect models with the residuals of diversity (or multidiversity), after correcting for abundance/activity (or multiabundance/diversity). This indicated that all of the effects on multidiversity and most of the effects on diversity of individual taxa remained significant and were thus not driven by abundance (Fig. 5). Some effects on diversity disappeared: the effect of Target SR on spiders, the effect of CWM height on earthworms, the effects of vertical heterogeneity on bats and the effect of soil pH on foliar fungal pathogens. As abundance is also affected in these cases, effects on diversity are driven by abundance. A few effects on diversity disappeared even though abundance did not respond significantly: effects of CWM N on spiders and foliar fungal pathogens, effects of vertical heterogeneity on defoliating insects and in these cases effects seem jointly mediated by abundance and diversity changes. There were also new effects on diversity that appeared, which indicate that abundance variation was obscuring the effects of certain variables on diversity: a positive effect of CWM height on foliar fungal pathogens, a positive effect of temperature on foliar fungal pathogens and multidiversity, negative effects of precipitation on birds but a positive effect on spiders, and a positive effect of soil C/N on defoliating insects.

## Discussion

Our analysis showed that tree diversity and functional composition were the most important drivers of the diversity and abundance/activity of forest-associated taxa. Tree diversity measures were the only ones that had consistent effects on overall multidiversity. Tree functional composition tended to have more contrasting effects on individual groups and therefore fewer overall effects on diversity. However, a high proportion of conifers reduced multidiversity (although this effect is driven by the defoliating insects that are hard to assess in conifer stands). The overall abundance of associated taxa was driven more by tree functional composition (CWM height) and forest structure (vertical heterogeneity). Abiotic factors were

less important but affected some groups. In general, the diversity and abundance/activity of spiders, defoliating insects and understorey plants were better explained by key forest features in comparison with bats, birds, microorganisms, earthworms, ungulates and foliar fungal pathogens. This would partially support the idea that lower trophic levels (producers, primary consumers) are more directly affected by stand characteristics and that effects on higher trophic levels are more indirect. However, not all groups responded in the expected fashion (spiders, foliar pathogens), which shows that it is challenging to predict the responses of individual groups and this underlines the importance of multitaxa approaches to understand the drivers of forest biodiversity.

Tree diversity was a key driver of forest-associated multidiversity and both target tree species richness and tree functional diversity were important. Target tree species richness was particularly important in increasing the diversity of foliar fungal pathogens and defoliating insects. These taxa may be species-specific and may therefore respond more strongly to target tree species richness than tree functional diversity (Gilbert et al. 2012). Although a high diversity of pathogens and insects is likely to be beneficial from a conservation perspective, it might not always be welcomed by forest managers (see extended discussion in Supporting Information Appendix S2). Tree functional diversity increased overall multidiversity, showing it tended to have positive effects on average. Functional diversity was always positively associated with the diversity of the separate taxa but only significantly increased bird diversity individually. Forest stands with trees that vary in their resource use strategies are likely to provide a greater diversity of resources for consumers and higher heterogeneity in soil resources and microclimate (Keller et al. 2013). This should provide more niches and therefore allow a larger number of forest-associated species to coexist (Levine and HilleRisLambers 2009). The negative effects of target species richness on bird diversity (and abundance) are surprising but are driven by the inclusion of functional diversity in the models. This means that bird diversity was lower in stands dominated by many functionally similar tree species and that a high functional diversity of trees is crucial to maintain high bird diversity. These effects on birds are likely to be indirect and may arise if low tree functional diversity results in a low diversity and activity of defoliating insects. This could have negative cascading effects on bird diversity and abundance, if bird species differ in their feeding preferences (Zhang et al. 2013, Charbonnier et al. 2016). The importance of both tree species richness and tree functional diversity show that multiple dimensions of tree diversity are important for promoting overall forest biodiversity.

Tree species evenness and admixed species richness had few effects on forest-associated taxa. This is probably due to the plot selection. Forest stands were only included if they had an evenness of at least

60% of the maximum possible, which resulted in a rather small variation between the plots. In addition, admixed species never exceeded 10% of the total basal area and therefore are unlikely to have had large effects on any of the forest-associated taxa (Baeten et al. 2013). In general, although plots were selected to differ in tree diversity, whilst minimising confounding effects of environmental variation or spatial position, in an observation study such as ours we cannot rule out the possibility that tree diversity effects were caused by unmeasured variables that are correlated with tree diversity. Experimental manipulations, e.g. biodiversity experiments, in which tree diversity is varied in a controlled fashion, are necessary to fully test the causality of the effects found here. However, the currently available tree diversity experiments are still relatively young and the approach used here remains the only option to evaluate the effects of tree diversity in mature forest stands.

The abundance/activity of forest-associated taxa also responded to tree diversity although none of the diversity dimensions tested here had a significant effect on multiabundance/activity. Target tree species richness increased the activity of foliar fungal pathogens. Trees may experience associational susceptibility to foliar fungal pathogens infestation in forest stands with a higher target tree species richness (Barbosa et al. 2009, Nguyen et al. 2016), as a result of more suitable microclimates for fungi (Jules et al. 2014), lower fitness of the trees due to competition (Pollastrini et al. 2013) or spillovers of generalist foliar fungal pathogen species between trees (Parker et al. 2015). The positive effect of tree species richness on spider abundance may be due to cascading effects, where tree diversity affects predators via a change in prey diversity/abundance (Scherber et al. 2010, Castagneyrol and Jactel 2012). However, we found little evidence for correlations between herbivores and predators and effects of tree diversity on spider abundance need to be treated with caution (see Supplementary Information Appendix S2). In addition, tree species richness increased the activity of ungulates. The optimal foraging theory states that these browsers should consume more in areas of high forage quality to maximize their instantaneous rate of intake (Stephens and Krebs 1986). Foraging in tree species rich stands offers the benefit of diet mixing, allowing greater and complementary intake of nutrients, as well as buffering against the accumulation of a particular source of digestion-inhibiting or toxic secondary metabolites (Wang et al. 2010, Milligan and Koricheva 2013). In general, tree diversity had stronger effects on the diversity rather than on the abundance/activity of forest-associated taxa and effects of tree diversity were not mediated by changes in abundance (in contrast with some of the effects of tree functional composition). Tree diversity effects on the diversity of forest-associated taxa are therefore likely to be mediated more by changes in resource heterogeneity than resource quantity (Borer et al. 2012). These

differing effects of tree diversity on species richness and abundance/activity show the importance of considering both aspects to understand effects of plant diversity on forest-associated taxa.

Tree functional composition also had a clear impact on several forest-associated taxa. Forest stands dominated by conifers had a lower diversity and activity of foliar fungal pathogens and defoliating insects. Conifers typically have a denser canopy (and cooler, wetter microclimate) and produce litter with a lower pH and nutrient concentrations and a higher lignin content than broadleaved species (e.g. Sardans et al. 2011, Augusto et al. 2015). This likely has a direct negative effect on organisms feeding on leaves (e.g. Brändle and Brandl 2001). Moreover, it is well known in European forests that the number of insect herbivores species, and specifically defoliators like Lepidoptera, is significantly lower on conifer than on broadleaved species (Kennedy and Southwood 1984, Brändle and Brandl 2001). However, it is also more challenging to assess herbivory in conifer stands, which may have led to an underestimation of herbivore activity in conifer forests (Guyot et al. 2016). In addition, the level of leaf area loss differs among the observed insect guilds, which means that our measure of insect herbivore activity might be dominated by certain feeding guilds, such as leaf chewers. In contrast to the primary consumers, spider diversity was higher in conifer-dominated plots. Conifers probably provide a more diverse spectrum of microhabitats than deciduous trees (Korenko et al. 2011) and their dense canopies lower the searching efficiency of birds (Whelan 2001). Multidiversity was lower in conifer stands, although this was driven by the defoliating insects.

In addition to the dominance of conifers, tree size and nutrient traits were also important in affecting the forest-associated taxa. Forest stands dominated by potentially taller trees (high CWM height) had a lower abundance overall and lower understorey cover and earthworm abundance in particular. Martens et al. (2000) found that light availability on the forest floor decreased as tree height increased. As light is the major limiting factor for forest understorey cover (Jennings et al. 1999, Barbier et al. 2008), higher CWM height probably reduced understorey productivity by reducing light levels. The negative effect of CWM height on earthworm abundance may be an indirect effect mediated by soil quality, as CWM height was negatively correlated with soil pH and low soil pH is known to reduce earthworm abundance (Reich et al. 2005). However, we did not find direct effects of the soil variables on earthworm abundance. For other groups, variation in temperature and humidity might be more important (Barkman 1992). Forest stands with taller trees are likely to be moister and to have a more stable temperature, which might provide an ideal habitat for foliar fungal pathogens and microorganisms, whose diversity and abundance respectively were higher in stands with potentially taller trees. Tree nutrient levels (CWM N) affected the diversity of



spiders and pathogens and these effects were partially mediated by biomass. Effects on spiders are likely indirect and mediated by an increase in prey. The effects on pathogens are surprising but could indicate that trees with more nitrogen rich leaves also produce nitrogen-based defences against pathogens. Tree functional composition is therefore a key driver of forest-associated taxa and in addition to well-known effects of conifers, other tree traits seem to be important in determining the abundance and diversity of several groups.

Effects of forest structure were less important than the effects of tree diversity and tree functional composition. However, plots with a higher basal area had a less diverse understorey and spider community. A higher basal area means denser and darker stands, which reduces the understorey (Barbier et al. 2008) and therefore potentially the habitat for spiders. Bat activity was reduced by vertical heterogeneity, probably due to the reduction of foraging space in heterogeneous stands. Most bat species avoid dense forest layers because it makes flying and locating prey more difficult (Jung et al. 2012). Ungulate activity was higher in forest stands with a higher vertical heterogeneity. This probably results from the way vertical heterogeneity was calculated. A higher understorey cover resulted in a higher value of vertical heterogeneity. As ungulates only browse in the lower forest layers, a higher understorey cover may also increase ungulate activity. Vertical heterogeneity increased multiabundance/activity as well because most individual taxa tended to increase in abundance/activity in more structurally heterogeneous plots. Hence, promoting structurally heterogeneous forests, and ensuring an abundant understorey, generally increases the abundance/activity of forest-associated taxa, but there are trade-offs because it reduces the activity of bats.

Abiotic drivers were also less important but had effects on some groups. Higher temperatures were associated with lower abundance overall but with a higher diversity of foliar fungal pathogens and a higher multidiversity, which is likely to be because Mediterranean forests had a higher diversity but lower abundance of several groups, such as plants and spiders. Increasing precipitation increased spider diversity, which agrees with results of Robertson and Aviles (2019) who found that rainfall positively affected spiders by increasing vegetation cover. The lower activity of foliar fungal pathogens and higher activity of defoliating insects that were found in more acid soils, could be due to changes in leaf chemistry (Izuta et al. 2001), which affected the palatability of the leaves for foliar fungal pathogens and defoliating insects. It is likely that considering a wider range of soil variables would have shown greater effects on soil microorganisms and earthworms, two groups that were not well explained by our models. In general, abiotic factors often had contrasting effects on diversity and abundance, e.g. precipitation for

spiders and soil CN for defoliating insects, which often led to larger effects of abiotic factors on diversity after correcting for abundance.

In general, we found more synergies than trade-offs between the diversities of different taxa, even though effects of explanatory variables were not always significant. This might be driven by functional links, i.e. high diversity at lower trophic levels increases the diversity of organisms that feed on them (Haddad et al. 2009, Scherber et al. 2010), or by shared responses to forest or environmental drivers, which is the most likely explanation for the correlation between pathogen and defoliator diversity. The synergies mean that changes in forest management could benefit multiple taxa. However, some negative correlations did occur, consistent with findings that the species richness of above- and belowground taxa may respond to different drivers (Gossner et al. 2016). These trade-offs might indicate some top-down control in addition to the bottom-up effects of tree diversity, and indicate that it is not possible to maximise the diversity of all taxa at the local scale of a forest stand. Instead, a variety of different stands might be necessary to maintain high diversity at the forest scale (van der Plas et al. 2016b).

Our analysis of a unique dataset, from a network of mature forest plots distributed across Europe, shows that high tree diversity is most important for promoting a high overall diversity of forest-associated taxa. This supports previous results from grasslands and suggests that bottom-up effects of plant diversity are a major driver of diversity at other trophic levels. Since the key driver of differences in tree diversity between our plots was silvicultural management, this study suggests that well-designed forest management can support a high biodiversity of many taxa. In general, forest managers should consider promoting mixed stands, and more specifically, favouring functionally dissimilar tree species. Tree functional composition, and in particular the presence of conifers, had contrasting effects on diversity of different taxa, so management that promotes variation in conifer abundance, or other aspects of functional composition, between stands is likely to support the highest biodiversity. Lowering basal area could also have beneficial effects on forest-associated biodiversity. In general, biodiversity at the landscape-scale could be optimized by creating a mosaic of forest stands differing in tree species composition, as trade-offs exist between diversities of some forest-associated taxa.

Overall, a change in tree diversity can influence multiple forest-related ecosystem functions and services (Scherer-Lorenzen 2014, van der Plas et al. 2016a). For example, previous research indicated that increasing tree diversity increases tree productivity (Paquette and Messier 2011), soil carbon storage (Ruiz-Benito et al. 2014), berry and game production (e.g. Gamfeldt et al. 2013). Our results clearly

showed that mixing tree species is essential for maintaining a generally high biodiversity in forests. Forest managers should therefore also consider increasing tree species richness and functional diversity as an effective way to provide the multitaxa and multiguild biodiversity that is needed to support the resilient provisioning of multiple ecosystem services under global change.

#### **Data accessibility statement**

Data are available from the Dryad Digital Repository: <<http://dx.doi.org/10.5061/dryad.xxxxx>> (Ampoorter et al. 2019).

**Acknowledgements** – We want to thank the site managers O. Bouriaud, H. Bruelheide, F. Bussotti, L. Finér, B. Jaroszewicz and F. Valladares and their field teams for installing the FunDivEUROPE Exploratory Platform and for performing basic measurements. We also thank S. Müller for organisational support for the field surveys and M. Liebergesell for collating the tree trait data on mean tree life span and maximum tree height.

**Funding** – This study is performed within the framework of the FunDivEUROPE project and has received funding from the European Union Seventh Framework Programme (FP7/2007-2013) under grant agreement no. 265171. EA was supported via an ERC Consolidator Grant awarded to KV (PASTFORWARD project, Grant no 614839).

**Author contributions** – AmE, BarL, JH, BaeL, CY, DWH, KJ, MH, SLM, vdPF, VK and AIE conceived the ideas and designed methodology; AmE, BarL, JH, BJ, CM, CB, CY, DSM, DM, DSP, DWH, GV, HS, JFX, KJ, MH, MB, ND, RS, RRK, VKJ and VL collected the data; AIE led the data analysis; AmE led the writing of the manuscript. All authors contributed critically to the drafts and gave final approval for publication.

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## Figure Legends

Figure 1. Drivers of diversity of forest-associated taxa. Relative effects of the explanatory variables on each of the taxon-level diversities and on multidiversity. Colours show the relative size and direction of the effects of each explanatory variable, in units of standard deviations. Blue colours represent a positive effect and red colours a negative effect. An asterisk indicates that bootstrapped confidence intervals around the parameter did not include 0 and that the effect was significant. Marginal and conditional  $R^2$  are indicated for each model. Brackets for the effect of proportion of coniferous species on multidiversity indicate that this effect may be partially spurious, see text (SR: species richness; funct.: functional; propn.: proportion; CWM: community weighted mean; horiz: horizontal; vert: vertical; Marg: marginal; Cond: conditional).

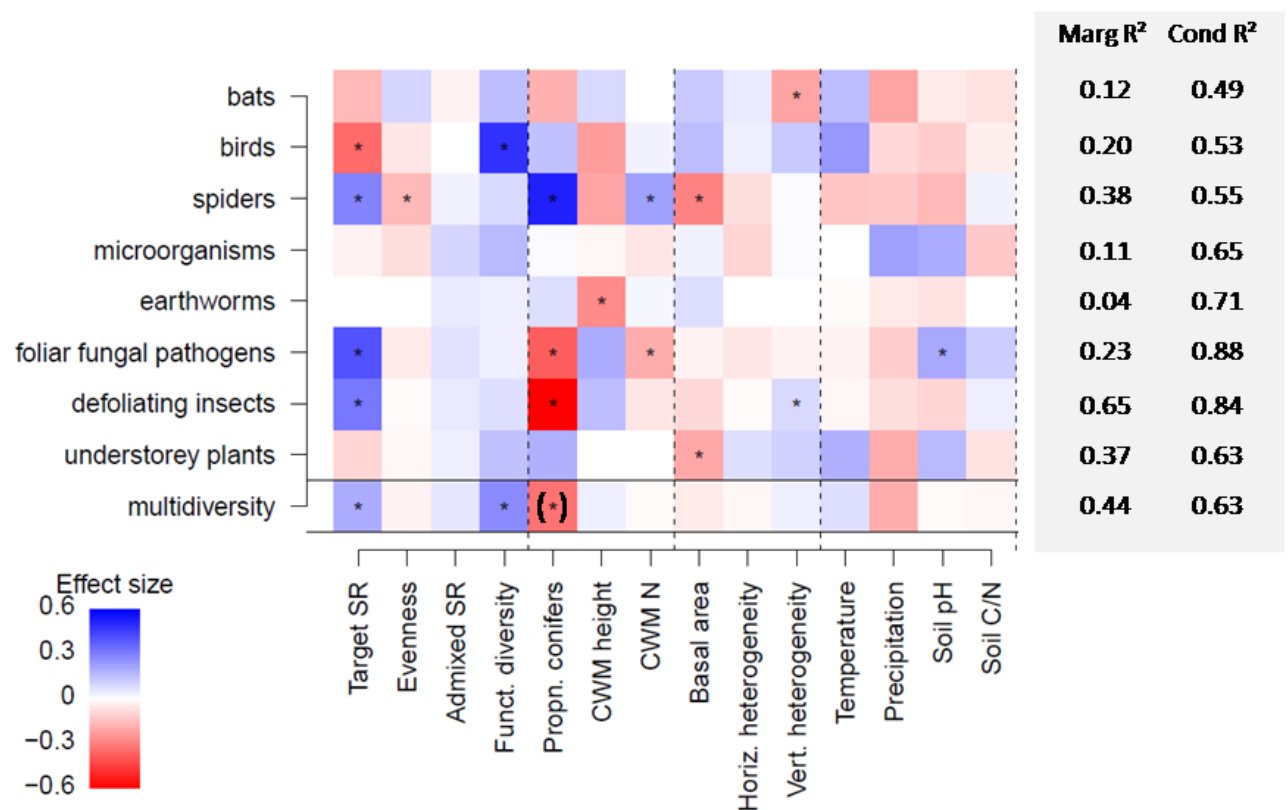


Figure 2. Drivers of abundance/activity of forest-associated taxa. Relative effects of the explanatory variables on the taxon-level abundance/activity and on multiabundance/activity. For details, see legend of Fig. 1. The effect of vertical heterogeneity on understorey plant abundance is not shown (black box) as this effect is spurious.

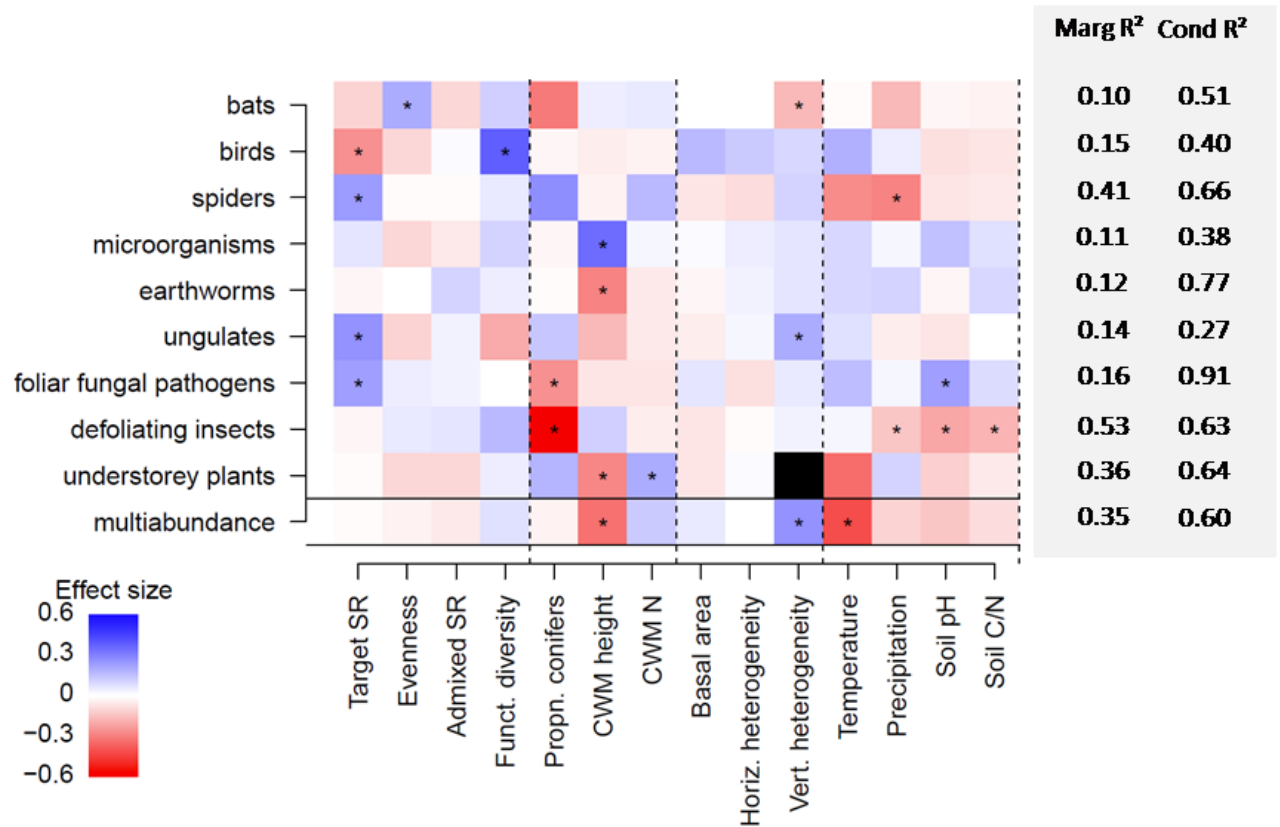


Figure 3. Correlations between diversities of all taxa. Numbers in the upper triangle represent the Pearson correlation coefficients ('\*\*\*':  $p \leq 0.001$ ; '\*\*':  $p \leq 0.01$ ; '\*':  $p \leq 0.05$ ). Figures in the lower triangle and on the diagonal represent scatter plots and distribution plots, respectively.

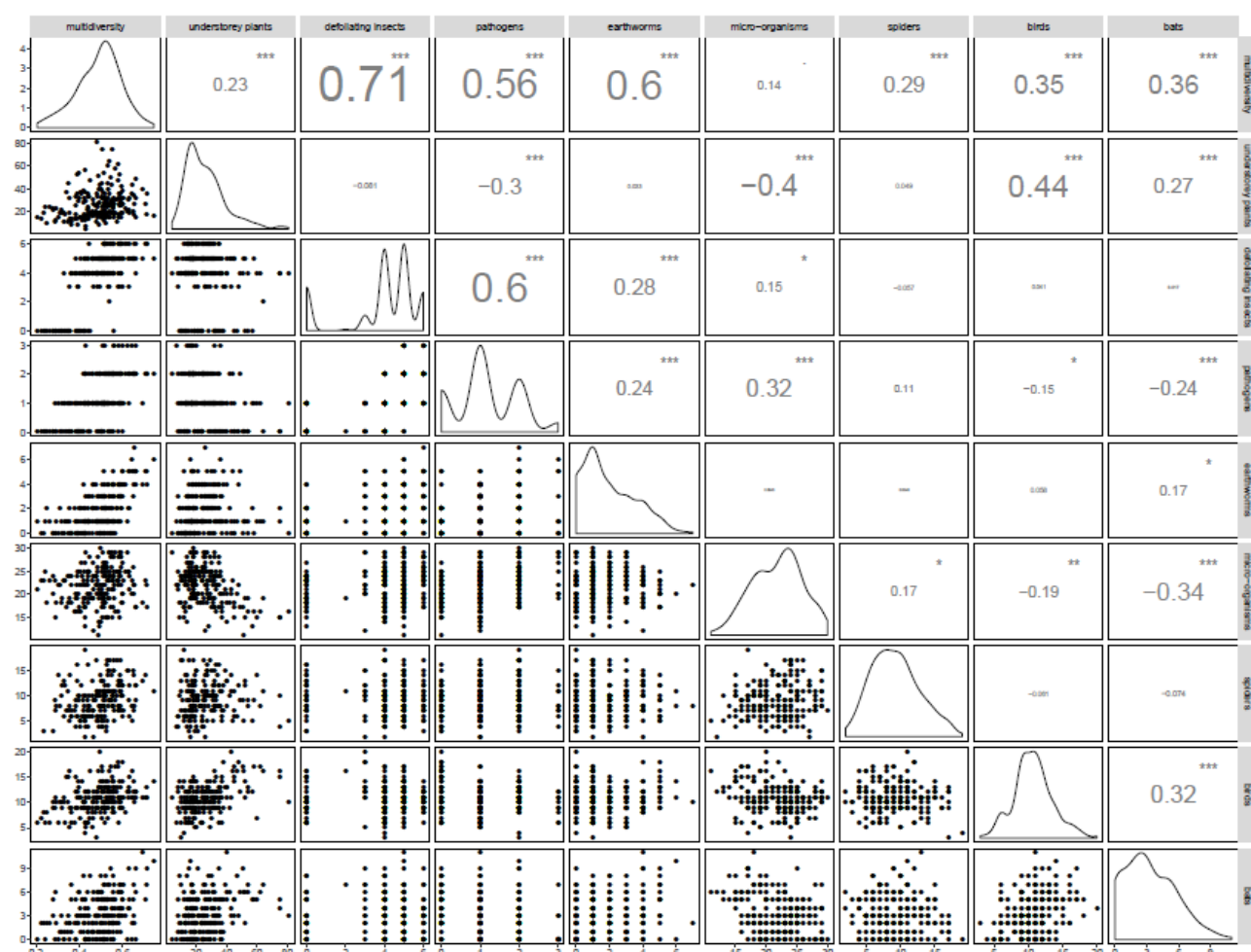


Figure 4. Correlations between log-transformed abundance/activity of all taxa. Numbers in the upper triangle represent the Pearson correlation coefficients (‘\*\*\*’:  $p \leq 0.001$ ; ‘\*\*’:  $p \leq 0.01$ ; ‘\*’:  $p \leq 0.05$ ). Figures in the lower triangle and on the diagonal represent scatter plots and distribution plots, respectively.

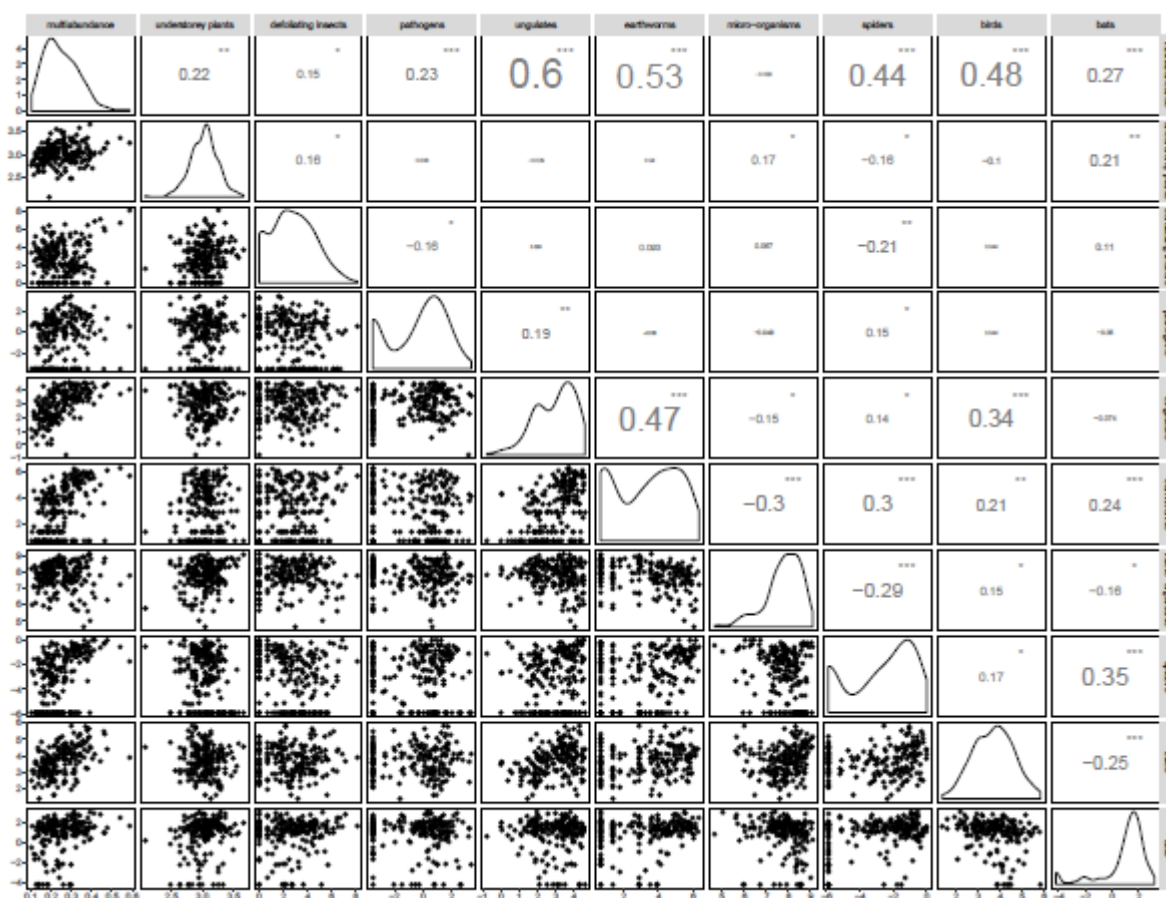




Figure 5. Drivers of diversity residuals of forest-associated taxa (after correcting for abundance/activity). Relative effects of the explanatory variables on residuals of each of the taxon-level diversities and on residuals of multidiversity, after diversity and multidiversity were corrected for abundance/activity and multiabundance/activity, respectively. Colours show the relative size and direction of the effects of each explanatory variable, in units of standard deviations. Blue colours represent a positive effect and red colours a negative effect. An asterisk indicates that bootstrapped confidence intervals around the parameter did not include 0 and that the effect was significant (SR: species richness; funct.: functional; propn.: proportion; CWM: community weighted mean; horiz: horizontal; vert: vertical).

