

This is the accepted manuscript version of the contribution published as:

Desie, E., Zuo, J., Verheyen, K., Djukic, I., Van Meerbeek, K., **Auge, H.**, Barsoum, N., Baum, C., Bruelheide, H., Eisenhauer, N., Feldhaar, H., Ferlian, O., Gravel, D., Jactel, H., Kappel Schmidt, I., Kepfer-Rojas, S., Meredieu, C., Mereu, S., Messier, C., Morillas, L., Nock, C., Paquette, A., Ponette, Q., Reich, P.B., Roales, J., Scherer-Lorezen, M., Seitz, S., **Schmidt, A.**, Stefanski, A., Trogisch, S., van Halder, I., Weih, M., Williams, L.J., Yang, B., Muys, B. (2023):

Disentangling drivers of litter decomposition in a multi-continent network of tree diversity experiments

Sci. Total Environ. **857**, Part 3 , art. 159717

The publisher's version is available at:

<http://dx.doi.org/10.1016/j.scitotenv.2022.159717>

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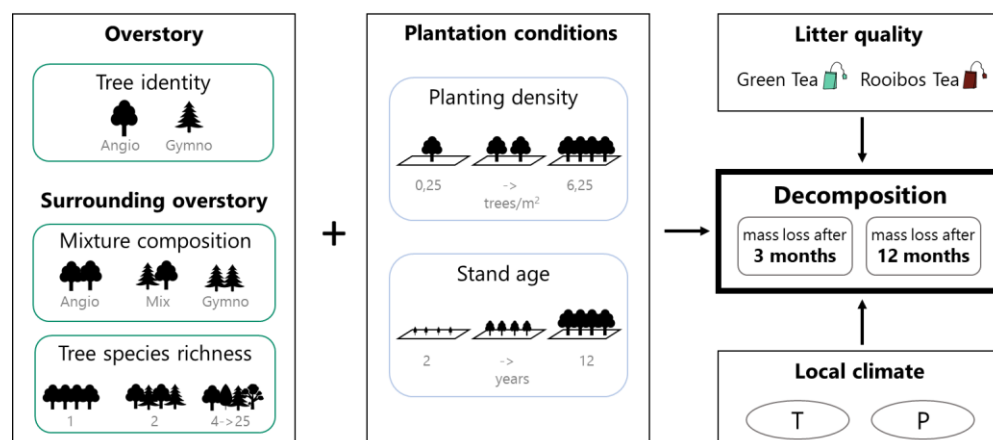
Authors' contributions

ID designed the experiment; AP, AS, BY, BM, CB, CM, CN, CAN, HB, HF, HJ, JR, KV, LM, LW, MSL, MW, NB, NE, OF, PBR, QP, StS, ST, SS collected data; ED, ID, JZ assembled the data from the sites; ED, JZ, KVM analyzed the data; ED and JZ wrote the first draft of the manuscript; BM, KV, ID and KVM provided thorough feedback on the manuscript, all coauthors commented on the advanced manuscript.

Highlights

- Tree species identity and composition affected decomposition of low-quality litter
- Young gymnosperm overstories promoted decomposition
- Tree species identity effects on decomposition depended on the age of the stand
- Tree species richness explained <1% variation in mass loss across continents
- Planting density did not impact decomposition dynamics

Graphical abstract



68 **Abstract**

69 Litter decomposition is a key ecosystem function in forests and varies in response to a range of
70 climatic, edaphic, and local stand characteristics. Disentangling the relative contribution of these
71 factors is challenging, especially along large environmental gradients. In particular, knowledge of
72 the effect of management options, such as tree planting density and species composition, on
73 litter decomposition would be highly valuable in forestry. In this study, we made use of 15 tree
74 diversity experiments spread over eight countries and three continents within the global
75 TreeDivNet network. We evaluated the effects of overstory composition (tree identity,
76 species/mixture composition and species richness), plantation conditions (density and age), and
77 climate (temperature and precipitation) on mass loss (after 3 months and 1 year) of two
78 standardized litters: high-quality green tea and low-quality rooibos tea. Across continents, we
79 found that early-stage decomposition of the low-quality rooibos tea was influenced locally by
80 overstory tree identity. Mass loss of rooibos litter was higher under young gymnosperm
81 overstories compared to angiosperm overstories, but this trend reversed with age of the
82 experiment. Tree species richness did not influence decomposition and explained almost no
83 variation in our multi-continent dataset. Hence, in the young plantations of our study, overstory
84 composition effects on decomposition were mainly driven by tree species identity on
85 decomposer communities and forest microclimates. After 12 months of incubation, mass loss of
86 the high-quality green tea litter was mainly influenced by temperature whereas the low-quality
87 rooibos tea litter decomposition showed stronger relationships with overstory composition and
88 stand age. Our findings highlight that decomposition dynamics are not only affected by climate

89 but also by management options, via litter quality of the identity of planted trees but also by
90 overstory composition and structure.

91 **Keywords:** biodiversity, biogeochemical cycle, carbon turnover, decomposition, forest, mass
92 loss, Tea Bag Initiative, tree communities, tree species richness, TreeDivNet

1. Introduction

Leaf litter decomposition is a central function in forest ecosystems that significantly affects global terrestrial carbon and nutrient cycling (Hättenschwiler, 2005; Gessner et al., 2010; Handa et al., 2014; Bradford et al., 2016). Litter decomposition rate mainly depends on the net effect of litter quality, (micro)climatic conditions, soil properties and the composition of the decomposer community (Coûteaux et al., 1995). Accordingly, the decomposition process is very sensitive to a changing climate while simultaneously forming an important feedback to the global carbon budget, resulting in potential mitigation or amplification of climate change (Aerts, 1997; Davidson and Janssens, 2006; García-Palacios et al., 2016). Several studies have already targeted the impact of macroclimate on litter decomposition dynamics across large spatial scales (Djukic et al., 2018) and along gradients of global change drivers such as atmospheric nitrogen (N) deposition (Kwon et al., 2021). However, the impact on litter decomposition of forest silvicultural management decisions such as overstory tree species composition and planting density, relative to litter quality and climate, has rarely been explored on a multi-continental scale (Joly et al., 2017).

Anthropogenic influences on tree composition and diversity in forest ecosystems include direct effects of silvicultural management decisions (planting and thinning) as well as indirect effects of environmental changes. Although it is obvious that the surrounding tree community considerably affects decomposition, its specific role in decomposition processes at broader scales is still poorly understood (Scherer-Lorenzen et al., 2007; Prescott and Vesterdal, 2013). Furthermore, our knowledge of the impact of silvicultural management decisions such as planting density or thinning regimes on decomposition remains incomplete (Lado-monserrat et al., 2015; Bueis et

al., 2018). A better understanding of tree community and silvicultural management decisions on forest nutrient and carbon cycles would allow us to design mixed forest plantations that are not only more resilient to climate change (Messier et al., 2022) but also better at mitigating climate change (Silva Pedro et al., 2015).

Multiple factors related to overstory tree composition influence litter decomposition (Joly et al., 2017). First and foremost, tree identity strongly affects decomposition by influencing litterfall mass (Prescott, 2002), litter quality (Cornwell et al., 2008; Vivanco and Austin, 2008; Zuo et al., 2018; Hoeber et al., 2020), microclimate (Gottschall et al., 2019), and even soil properties (Reich et al., 2005; Desie et al., 2019) and decomposer communities (Hobbie et al., 2006; Zhang et al., 2020; Peng et al., 2022). Given the large biogeographic area encompassed by this study, we define species identity according to the two major lineages of trees, i.e. angiosperm species or gymnosperm species, assuming that they impact microclimate, water availability and nutrient cycling in different ways (Augusto et al., 2015) and harbor different decomposer communities which dominate different stages of decomposition (Zhang et al., 2020). Typically, gymnosperm species have higher LAI and rainwater interception resulting in drier soil surfaces (Aranda et al., 2012) and tend to acidify soils due to their low litter quality more than angiosperm species (Finzi et al., 1998; De Schrijver et al., 2012).

Second, the species composition of the tree community can influence decomposition through complementary nutrient use (Tilman et al., 2014; Lin et al., 2021) and rhizosphere processes (Binkley and Giardina, 1998; Wardle et al., 2004). To date, evidence for the influence of tree species diversity on decomposition remains ambiguous, with studies reporting inconsistent

(Naeem et al., 1994; Wardle and Nicholson, 1996), mere additive (Scherer-Lorenzen et al., 2007), synergistic (Handa et al., 2014; Maxwell et al., 2020; Strukelj et al., 2021), and antagonistic effects (Blair et al., 1990; Wardle et al., 1997; Seidelmann et al., 2016; Joly et al., 2017). These context-dependent results are probably a consequence of the multitude of pathways through which tree species identity, composition, and diversity could affect decomposition (Jewell et al., 2017), which are simultaneously affected by stand characteristics, management, and climate (Lin et al., 2021). For example, planting density, another important management decision besides tree species selection, could change decomposition by affecting the forest microclimate, soil nutrient-availability, and the total quantity of litter that is produced (Bueis et al., 2018). Furthermore, the driving factors of decomposition can change with a) litter type (Bradford et al., 2016) as high-quality litter is more controlled by abiotic (edaphic and climatic) factors whereas low-quality litter with more structurally complex C substrates is more affected by biotic (overstory) factors (Fanin et al., 2020); b) the age of the stand: as the opening of the forest canopy due to stem exclusion at late successional stages can reduce humidity and thus slow down litter decomposition (Trogisch et al., 2016); and c) the decomposition process itself (Berg & McClaugherty, 2020) where there is a shift in control from biotic to abiotic factors with ongoing litter decay (García-Palacios et al., 2016). Hence, it remains a challenge to control for multiple factors of influence in observational studies, particularly due to confounding factors like tree composition and climate affecting decomposition. Consequently, the relative contribution of these factors and their context-dependencies are even less studied.

The combination of different tree diversity experiments across biomes (TreeDivNet, Verheyen et al., 2016; Djukic et al., 2018; Paquette et al., 2018) provides a unique opportunity to investigate

effects of tree species identity, composition and diversity on litter decomposition, and their relative importance to climate related variables. This coordinated multi-site experiment across different continents (Fraser et al., 2013) has made use of the standardized Tea Bag method (Keuskamp et al., 2013) to evaluate decomposition processes without having litter quality confounded with biome and local environment, or with diversity in litter composition (of the litterbag), or with overstory diversity effects (Lin et al., 2021). We incubated standardized litter bags for three and 12 months under 29 focal tree species, encompassing both high-quality angiosperm litter and low-quality gymnosperm litter, occurring in 90 different compositions (monocultures and different combinations of angiosperm and gymnosperm species) with tree species richness ranging from one to 24 species (most experiments have a species richness gradient between one and four species) grown on 15 different experimental sites (Figure S1). We used green tea and rooibos tea which are representative for fast (high-quality) and slow (low-quality) decomposing leaf litter (Didion et al., 2016). Our main objective was to evaluate how the multiple factors related to the overstory composition (tree identity, species/mixture composition, and species richness), plantation conditions (age and planting density) and local climate (temperature and precipitation) affect mass loss of high and low-quality litter at two different stages of decay (after 3 and 12 months) (Figure 1). We hypothesized the following:

- 1) Overstory tree species composition impacts decomposition directly via litter quality with angiosperm and/or more diverse communities promoting faster decomposition rates in comparison to the recalcitrant gymnosperm overstories. Moreover there is an indirect effect of overstory tree species composition via the type of micro-environment found on the forest floor: dense and evergreen canopies of gymnosperms reduce water availability at the forest

floor through higher rainwater interception and evaporation, despite dense canopies better preserving a humid forest microclimate, thus impeding decomposition.

2) The effect of overstory composition on decomposition becomes stronger with time (i.e., the age of the stand) as differences among microbial communities and microclimate have had more time to accumulate.

3) Planting density impacts decomposition through changes in the micro-climate, with higher densities leading to less favorable conditions for decomposition due to higher rainwater interception.

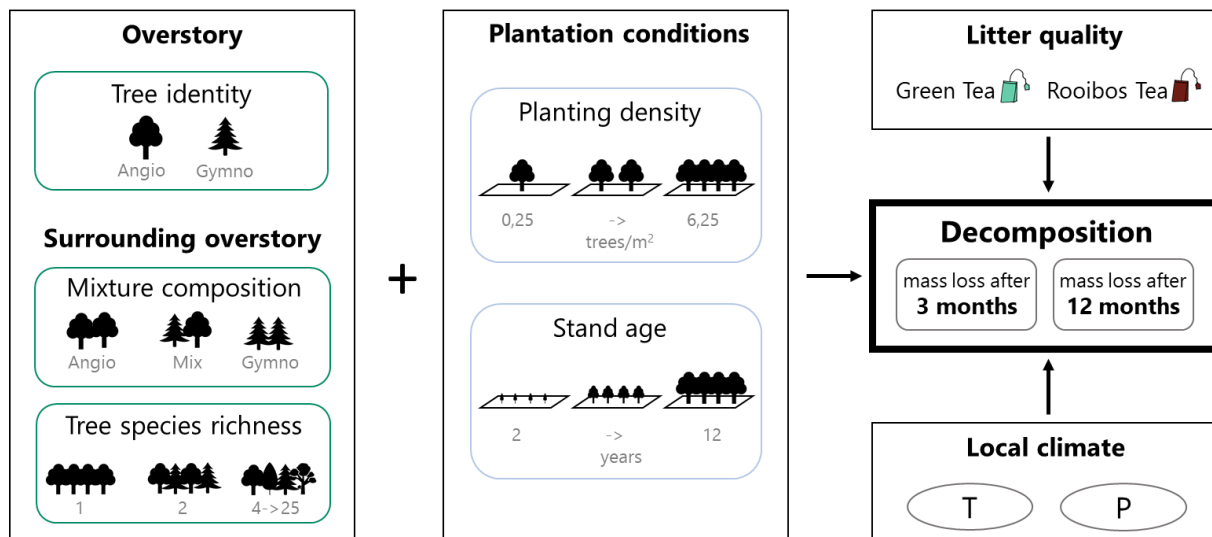


Figure 1: Conceptual model showing the possible factors affecting litter decomposition in the tree diversity experiments. Angio = angiosperms, Gymno = gymnosperms, Mix = mixture of angiosperm and gymnosperm species; T = temperature; P = precipitation.

2. Materials and Methods

2.1 Study sites

The experiment was carried out at 15 sites belonging to the global Tree Diversity Network (TreeDivNet network, <http://www.treedivnet.ugent.be/>) (Verheyen et al., 2016; Paquette et al., 2018) (Table 1, Figure S1), designed to test tree diversity effects on ecosystem functions worldwide. The studied sites are distributed over boreal, temperate, Mediterranean, and subtropical biomes. Planting densities of the experiments range between 0.25 and 6.25 trees.m⁻². At the time of the litterbag experiment (2016), the forests were still rather young with tree age ranging between 1 and 12 years. Local climate data for the year 2016 (extracted from TerraClimate (Abatzoglou et al., 2018)) varied from 6.9°C to 19.4°C for mean annual temperature (MAT) (12.5°C to 24.1°C for the summer of 2016) and mean annual precipitation (MAP) 591 mm to 1903 mm (28 mm to 615 mm for summer) (Table 1).

204 **Table 1.** Basic characteristics of the experimental sites and number of tea bags used in this study.

				Location		Climate		Local weather during experiment				SR gradient ^e	Tree planting year	Tree density (tree m ⁻²)	No. bags 3M	No. bags 12M	
No.	Country	Name of Experiment	Site	Latitude	Longitude	Biome	MAT ^a	MAP ^b	MPT ^c 3M	MPT 12M	CPP ^d 3M						CPP 12M
A	Belgium	FORBIO	Gedinne	49.99	4.98	Temperate	10.40	670	15.45	10.81	322.20	897.30	1, 2, 4	2010	0.44	62	52
B	Belgium	FORBIO	Hechtel- Eksel	51.16	5.31	Temperate	8.60	1030	16.99	11.43	278.20	631.80	1, 2, 4	2012	0.44	69	69
C	Belgium	FORBIO	Zedelgem	51.15	3.12	Temperate	10.10	708	16.73	11.90	265.00	688.30	1, 2, 4	2009	0.44	71	71
D	Canada	IDENT	Auclair	48.23	-69.10	Boreal	2.30	1015	12.52	4.88	615.90	1400.90	1, 2, 6	2010	6.25	126	NA ^g
E	Canada	IDENT	Montreal	45.86	-73.93	Temperate	6.20	976	20.78	8.64	236.90	1166.40	1, 2, 4	2009	4.00	120	120
F	China	BEF-China	Xingangshan	29.12	117.91	Warm- temperate, humid	17.10	1777	24.10	19.41	534.60	1903.30	1, 8, 24	2009	0.60	94	101
G	Germany	BIOTREE	Kaltenborn	50.78	10.22	Temperate	7.80	650	15.83	10.22	266.00	885.90	1, 2, 4	2004	0.25/0.44	48	47
H	Germany	ECOLINK- Salix	Rostock	54.06	12.08	Temperate	8.50	590	17.92	10.17	194.20	810.20	1, 2	2014	1.56	36	NA ^g
I	Germany	Kreinitz	Zeithain	51.23	13.15	Temperate	8.40	575	18.04	11.24	212.60	607.30	1, 3, 6	2005	1.25	68	72
J	Germany	MyDiv	Bad Lauchstädt	51.39	11.88	Temperate	9.00	492	19.32	11.17	164.70	538.40	1, 2, 4	2015	1.00	72	71
K	France	ORPHEE	Pierroton	44.74	-0.80	Temperate	12.75	876	17.55	14.63	136.00	705.30	1, 2, 3	2008	0.25	141	141
L	Italy	IDENT	Macomer	40.24	8.70	Mediterranean	13.80	866	22.33	15.74	28.80	867.00	1, 2, 6	2014	4.00	108	108
M	Sweden	ECOLINK- Salix	Uppsala	60.44	18.08	Temperate	5.60	470	13.79	7.93	241.80	591.30	1,2	2014	1.56	26	31
N	UK	Climate- match	Kent	53.40	-0.30	Temperate	9.30	763	15.88	11.41	116.40	681.2	1, 4	2011	0.25	62	NA ^g
O	USA	IDENT	Cloquet	46.68	-92.52	Temperate	2.60	717	16.48	6.99	545.90	1113.50	1, 2, 6	2010	6.25	123	118

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206 ^a MAT = mean annual temperature in °C from Djukic et al. (2018); ^b MAP = mean annual precipitation in mm from Djukic et al. (2018); ^c MPT = Mean
207 Period Temperature in °C with Period referring to the experimental incubation period; ^d CPP = Cumulative Period Precipitation in mm; 3M = 3
208 months of incubation; 12M = 12 months of incubation; ^e SR gradient = species richness gradient; ^f No. of bags = number of installed tea bags, ^g NA
209 = 12M not included in the experiment.

2.2 Decomposition experiment

We used two varieties of Lipton tea bags as our standard litter: fast decomposing green tea and slowly decomposing rooibos tea (Keuskamp et al., 2013). Before field installation, the bags were dried at 70°C for 48 h, and the initial mass was recorded. During the summer of 2016, tea bags of each tea type were installed in the topsoil (0-5 cm below the surface) underneath different focal tree species (comprising a total of 29 different tree species) in plots with different tree species compositions along a gradient of different species richness (including 1, 2, 3, 4, 6, 8, and 24 species, depending on the site). In total, 2227 bags were incubated and recovered after 3 months (1226 bags) and after 12 months (1001 bags) of incubation (Figure 1, Table 1). In mixtures, tea bags were placed under specific focal trees at the base of the stem. The list of tree species is provided in Table S1 (Supporting information). Total weight of the tea bags was recorded by weighing the filled tea bags with the string and label, and an averaged weight for the empty bags with string and label (0.248 g) was subtracted from this value to estimate the amount of tea before incubation. After the incubation period, tea bags were carefully collected, dried at 70°C for 48h, and the remaining weight of the tea was recorded, assuming that the bag itself did not lose any mass. For a more detailed description of the method, please refer to Djukic et al. (2018) where more information on the methodology is provided.

2.3 Data analysis

Decomposition was modeled as a function of various drivers using linear mixed models including site and plot as random intercepts (plot nested in site) using the package lme4 (Bates et al., 2015). Fixed effects were: the identity of the focal tree (i.e., the tree under which the litterbag was

installed); whether it was an angiosperm or gymnosperm; mixture composition of the plot in which the litterbag was installed (pure angiosperms, pure gymnosperms, or mixture of angiosperms and gymnosperms); tree species richness of the plot; planting age of the experiment; planting density; mean period temperature (MPT) and cumulative period precipitation (CPP) during the experimental period. As not all tea bags were incubated for exactly three months (mean=103 days, SD=21) or twelve months (mean=364 days, SD=26), we included length of the incubation period in the statistical analysis to account for this variation. All models were executed on centered variables. The different levels of the fixed effects (e.g., tree identity varying within plot and site vs climate and plantation conditions only varying between sites) were accounted for by the hierarchical structures of the linear mixed effects models (Zuur et al., 2009). The response variables of the mixed models were litter mass loss per litter type (green and rooibos). Because the impact of tree identity (here defined as the difference between angiosperm and gymnosperm overstories) can change over time due to differences in early growing rates (Zhang et al., 2022), the interaction between focal tree identity and planting age was included in the models. Likewise, the interaction between MPT and CPP was included in all models. Type I anova tests were executed on all models (Table 2). Alternative models including aridity indices or excluding BEF-China (with the outlying species richness level 24) are provided in the appendix (Table S4). The normality and homogeneity of residuals of models were checked by plotting the fitted values versus the standardized residuals, and by graphically evaluating the histograms of the standardized residuals. The partial effects of these regressions were plotted separately for each variable (Figures 3-6) accounting for the other variables in the models using the effects package (Fox et al., 2022).

253 Variance partitioning analysis was executed using the partR2 package (Stoffel et al., 2021).
254 Variables were grouped to reduce complexity: temperature (T) and precipitation (P) are 'climate';
255 identity of the target tree and type of mixture of the surrounding trees are 'tree composition',
256 and age and density are 'plantation conditions'. We determined both the unique effects of single
257 predictors and the effects shared by each pair of predictors. The proportion of variance explained
258 by the fixed effects and the random effects was calculated by comparing the marginal and
259 conditional R^2 of the mixed model (sensu Nakagawa and Schielzeth, 2013) calculated using
260 r.squaredGLMM using the MuMIn package (Barton, 2022). All statistical analyses were
261 performed using R version 4.0.5 (R Core Team, 2018).

3 Results

3.1 Litter quality and climate effects on mass loss over time

Across experimental sites and biomes, green tea decomposed significantly faster ($65\% \pm 9\%$ after 3 months and $72\% \pm 10\%$ after 12 months, mean \pm SE) compared to rooibos tea ($20\% \pm 12\%$ after 3 months and $35\% \pm 12\%$ after 12 months, mean \pm SE) (Figure S3-S5). The increase in mass loss rate between 3 and 12 months was higher for rooibos tea, as indicated by the significant interaction effect between time and litter type (Figure S2, Table S2). Sites were located over a large climatic gradient, with mean temperature during the experiment ranging from 4.8°C at Auclair, Canada, to 19.4°C at Xingangshan, China, and mean annual precipitation from 591 mm at Bad Lauchstädt, Germany, to 1903 mm at Xingangshan, China (Table 1). Highest mass loss was found in Xingangshan and lowest in Montreal, Rostock and Auclair (Figure S3 and S4). Generally, mass loss increased from boreal < Mediterranean < temperate < warm temperate humid biomes (Figure S5). Mass loss was, however, not significantly influenced by MPT or CPP, irrespective of tea type (Table 2).

Table 2. ANOVA output of linear mixed models testing the effect of tree species composition, planting, and climate related variables on mass loss of green and rooibos tea after 3 months and 12 months, respectively. The interaction between age and focal tree identity (angiosperm or gymnosperm) and between mean period temperature (MPT) and cumulative period precipitation (CPP) were included in all models. Site and plot were included as random factors with plot nested in site. Models were executed on centered variables.

Fixed effect	Mass loss of green tea after 3 months						Mass loss of rooibos tea after 3 months					
	Sum Sq	Mean Sq	Num DF	Den DF	F value	P	Sum Sq	Mean Sq	Num DF	Den DF	F value	P
Incubation period length	0.024	0.024	1	503.29	7.83	0.005	0.003	0.003	1	487.30	0.97	0.32
Focal tree identity	0.002	0.002	1	174.13	0.93	0.33	0.019	0.019	1	78.21	4.87	0.03
Age	0.000	0.000	1	9.30	0.02	0.88	0.008	0.008	1	9.30	2.05	0.18
Tree composition	0.000	0.000	2	187.72	0.10	0.89	0.004	0.002	2	94.07	0.60	0.54
Species richness	0.001	0.001	1	84.57	0.43	0.50	0.001	0.001	1	45.32	0.27	0.59
Planting density	0.000	0.000	1	8.93	0.10	0.75	0.001	0.001	1	9.04	0.25	0.62
MPT	0.000	0.000	1	9.12	0.14	0.71	0.001	0.001	1	9.23	0.45	0.51
CPP	0.000	0.000	1	9.14	0.20	0.65	0.001	0.001	1	9.20	0.30	0.59
Focal tree identity: Age	0.020	0.020	1	205.25	6.50	0.01	0.079	0.079	1	126.59	19.63	<0.001
MPT: CPP	0.001	0.001	1	9.01	0.46	0.51	0.002	0.002	1	9.10	0.66	0.43

Fixed effect	Mass loss green tea after 12 months						Mass loss rooibos tea after 12 months					
	Sum Sq	Mean Sq	Num DF	Den DF	F value	P	Sum Sq	Mean Sq	Num DF	Den DF	F value	P
Incubation period	0.037	0.037	1	213.86	7.12	0.008	0.170	0.170	1	139.54	26.85	<0.001
Focal tree identity	0.000	0.000	1	155.04	0.05	0.81	0.008	0.008	1	130.22	1.42	0.23
Age	0.003	0.003	1	6.50	0.66	0.44	0.024	0.024	1	6.05	3.81	0.09
Tree composition	0.011	0.005	2	168.62	1.06	0.34	0.045	0.022	2	146.08	3.61	0.02
Species richness	0.017	0.017	1	40.57	3.27	0.07	0.002	0.002	1	108.45	0.34	0.55
Planting density	0.001	0.001	1	6.26	0.29	0.60	0.009	0.009	1	5.78	1.54	0.26
MPT	0.061	0.061	1	6.08	11.71	0.01	0.002	0.002	1	5.68	0.31	0.59
CPP	0.013	0.013	1	6.31	2.56	0.15	0.000	0.000	1	6.02	0.14	0.71
Focal tree identity: Age	0.000	0.000	1	163.90	0.02	0.87	0.008	0.008	1	134.38	1.38	0.24
MPT: CPP	0.007	0.007	1	6.83	1.48	0.26	0.006	0.006	1	6.76	1.00	0.35

3.2 Tree identity, composition, and diversity effects

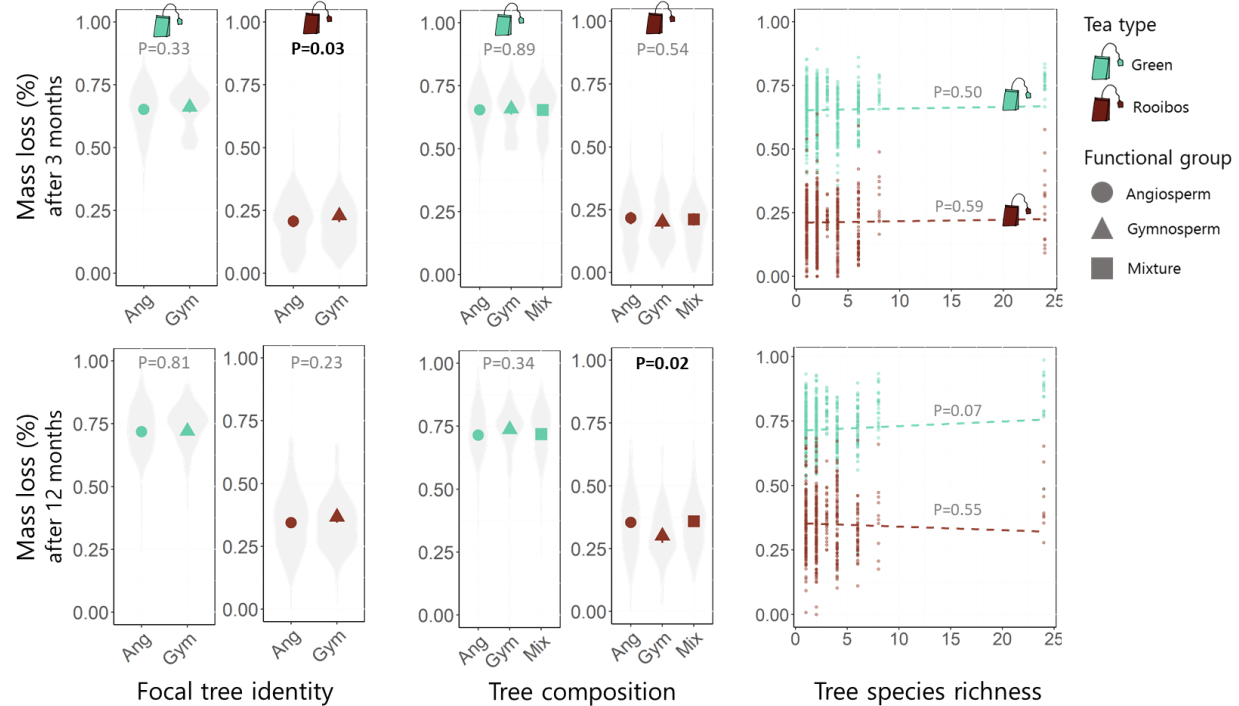


Figure 2. Mass loss (%) as a function of focal tree identity (left), surrounding tree functional composition (middle) and tree species richness (right) for the two litter types. Relations were tested using linear mixed models (Table 2) accounting for other overstory related variables, climate, plantation conditions, plot and site. Significant relations are indicated by a full line whereas non-significant (NS) relations are indicated by a dotted line. The shaded parts indicate the standard error interval. Observations are indicated for litter types, green tea (green) and rooibos tea (red) and per incubation periods, 3 month (top) and 12 months (bottom).

Mass loss of rooibos tea after 3 months of incubation was significantly higher under gymnosperm compared to angiosperm overstories (Figure 2). The functional group of the focal tree under

which tea bags were placed did not influence mass loss after 12 months of incubation (Figure 2 and S2). After 12 months of incubation, tree composition significantly affected mass loss of rooibos tea with gymnosperm overstories resulting in lower mass loss compared to mixed overstories (Figure 2). This is after accounting for tree identity, where gymnosperm focal trees promote mass loss (although not significant $P=0.23$) (Figure 2, Table 2). Species richness did not significantly impact mass loss, irrespective of substrate type or stage of decay (Figure 2). The marginally significant increase of green tea mass loss with species richness after 12 months is based on the highest species richness level of BEF-China (Table S4).

3.3 Plantation conditions effects

Planting density did not impact mass loss. Mass loss of rooibos tea after 12 months decreased marginally significantly ($P=0.09$) with the age of the experiment whereas decomposition of green tea was not affected by age or planting density (Figure 3). We did, however, find a significant interaction effect of age and focal tree species identity on mass loss after 3 months incubation ($P=0.01$ for green tea and $P<0.001$ for rooibos tea): for gymnosperm focal trees we observed more negative relationships between mass loss and stand age (Figure 4).

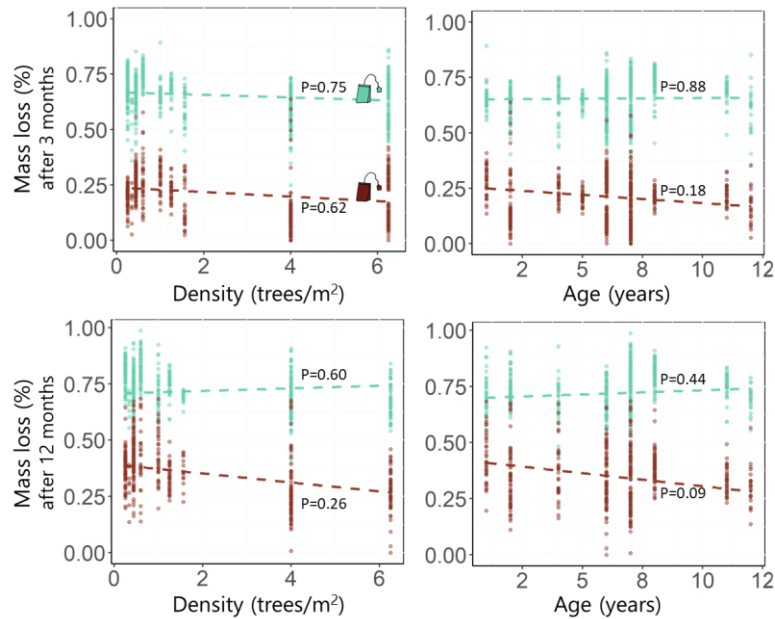


Figure 3. Mass loss (%) as a function of planting density (left) and age since the plantation (right) for two litter types after 3 months (top) and 12 months (bottom) of incubation. Relationships were tested using linear mixed models (Table 2) accounting for overstory effects, local climate, and random effects. There were no significant relationships ($\alpha < 0.05$) between the variables shown in these figures. Observations are given for litter types: green tea (green) and rooibos tea (red).

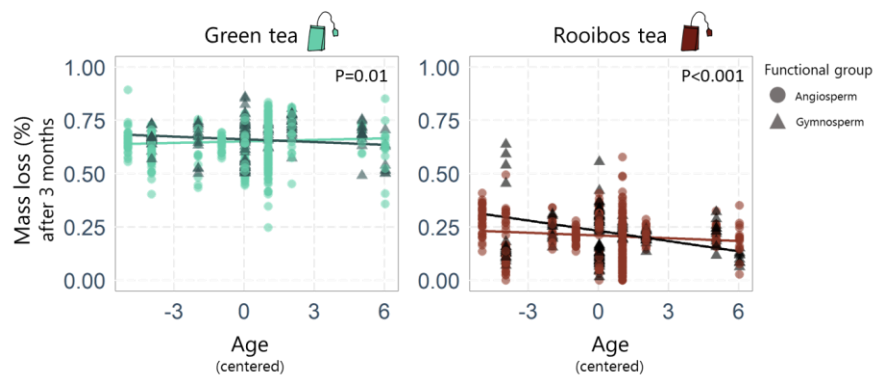


Figure 4. Mass loss (%) as a function of age (centered variable) for green tea (left) and rooibos tea (right) for overstory functional group (pale circles: angiosperms and dark triangles: gymnosperms). Relationships were tested using linear mixed models (Table 2) accounting for overstory, climate and random effects. The significance level of the interaction is indicated in top left corners.

3.4 Variance partitioning

Litter type explained most of the variation (80%) in mass loss after three months in our study with minimal contributions of climate (temperature and precipitation combined, 0.9%) and plantation conditions (age and density combined, 0.1%). Overstory composition (identity or

composition of the mixture) did not explain any variation (~0 %). After 12 months of incubation, the proportion of variation explained by climate variables increased (to 2 %). When analyzed for tea type separately, random effects (site and plot) explained most of the variation in green tea mass loss (58 %) after 3 months, with very limited variance explained by climate (1 %), plantation conditions (0.4 %), overstory species composition (identity and composition combined, 0.1%) and species richness (0.1 %). Variance in rooibos mass loss after 3 months was explained by climate (3 %), plantation conditions (3 %) and species composition (0.3%), whereas species richness explained almost no variation (~0 %). After 12 months of incubation, climate explained more of the variance in green tea mass loss (23 %). For rooibos tea, mass loss after 12 months was increasingly explained by species composition and by incubation period length and its interaction with plantation conditions (14%) (Figure 5).

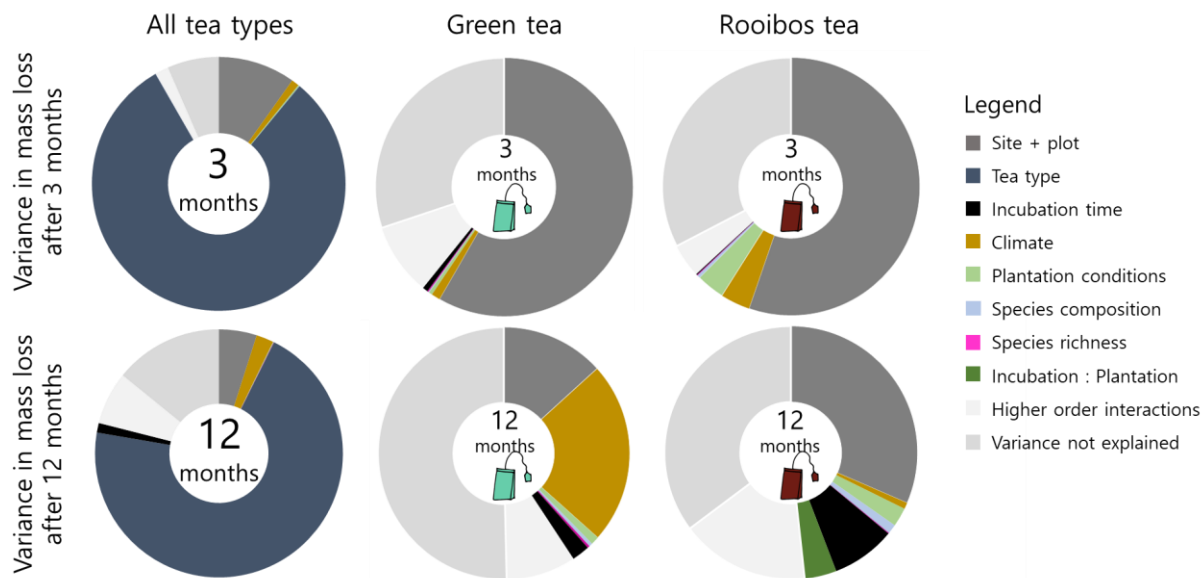


Figure 5. Percentage of variation explained by site, litter type, climate (MAT+MAP), tree composition (Focal tree identity + mixture composition), species richness (SR), plantation conditions (age + density), and the shared effects

339 between incubation period, climate, and plantation conditions for the total mass loss (left), green tea mass loss
340 (middle) and rooibos tea mass loss (right) for incubation periods (top: 3 months, bottom: 12 months).

4 Discussion

The impact of forest silvicultural management practices, such as the selection of a certain species composition and planting density, on biogeochemical processes and their potential co-benefits for carbon sequestration is still not well understood. Our study of the mass loss of two standardized litter types in planted juvenile forests across biomes aimed to examine the effects of overstory composition (tree identity, mixture composition and diversity), plantation conditions (density and age) relative to climate (temperature and precipitation) on the litter decomposition process over time.

4.1 Litter quality and climate effects on mass loss over time

As generally acknowledged (Coûteaux et al., 1995; Aerts, 1997), litter decomposition is largely influenced by litter quality and climate. We observed two times faster decomposition of high-quality green tea compared to low-quality rooibos tea along the investigated gradients of tree diversity located in three continents (Figure S3). The higher water-soluble fraction and nutrient content in green tea compared to rooibos result in faster mass loss due to leaching and decomposer activity during early stages of decomposition (Berg and McClaugherty, 2008; Ristok et al., 2017; Fanin et al., 2020). Hence, our results (80% of variance explained by tea type in our study; Figure 5) further illustrates the importance of litter quality for decomposition at a multi-continental scale (Djukic et al., 2018; Kwon et al., 2021). After litter quality, climate was the most important driver of mass loss in our study, explaining 0.9 % of variation after 3-months and 2 % after 12-months (Figure 5). Accordingly, biome significantly affected decomposition dynamics with mass loss increasing from boreal < Mediterranean < temperate < warm, temperature humid biomes (Figure S5, Table S3), as already illustrated by Djukic et al. (2018), corresponding to the

anticipated increase in mass loss with temperature (Bradford et al., 2016), as long as moisture does not become limiting (Petraglia et al., 2019). It is important to note that the impact of temperature on mass loss depends greatly on soil moisture conditions (Petraglia et al., 2019), but we did not consider this factor specifically in our study. Moreover, differences in microclimate (e.g., through shading and interception) and local differences in the environment (e.g., micro-relief, litter traits on the forest floor) all play an important role in decomposition (Joly et al., 2017), yet they are often neglected when evaluating broader climatic gradients.

4.2 Tree identity, composition, and diversity effects

In our study, we found significant overstory tree identity effects on mass loss of low-quality litter during early stages of decomposition (after 3 months of incubation - conducted in the growing season): rooibos tea decomposed faster under gymnosperm compared to angiosperm overstories (Table 2, Figure 2). Most studies report opposite trends: for instance, Chomel et al. (2015) observed slower decomposition of cellulose in a spruce plantation than in a poplar plantation; Joly et al. (2017) reported a negative correlation between litter decomposition and the relative basal area of evergreen trees; and recent findings of Fanin et al. (2020) showed that rooibos tea decomposed slower under coniferous species than under broadleaved species. These studies were executed in mature forest stands, whereas our study was carried out in young plantations (maximum 12 years old) where microbial and invertebrate communities had less time to adapt to the prevailing tree species (and its litter input) and also microclimatic conditions still change considerably in the first years after planting (Zhang et al., 2022). The higher mass loss of low-quality litter after three months of incubation, when placed under young gymnosperm canopies in our study could be related to higher shading provided by gymnosperm species due

to their dense canopies. This could have improved moisture retention (Petraglia et al., 2019) or provided a more buffered temperature regime (Zhang et al., 2022), both of which promote mass loss. This mechanistic explanation is thus in contrast to our expectation that denser canopies may reduce soil moisture due to higher rainfall interception than in the more open canopies of the angiosperm species (cf Hypothesis 1). These identity effects became subtler with progressing decomposition. We assume that this reflects the decreasing impact of climate on decomposition processes as the proportion of lignin increases with time (Berg, 2000), explaining the lack of effects after 12 months of incubation. Additionally, we cannot exclude the effect of differences in the decomposer communities between angiosperm and gymnosperm overstories, which could affect decomposition differentially through decomposition stages.

We did not find a significant effect of tree species richness on the decomposition of standardized substrates (Table 2, Figure 3). This is in line with other studies that identified a predominant role of identity over diversity in belowground ecosystem functioning (Schwarz et al., 2015; Dawud et al., 2016, 2017; Joly et al., 2017; Zhou et al., 2020). We did, however, find a significant effect of tree composition on mass loss of low-quality rooibos tea after 12 months: more diverse neighborhoods (e.g., mixtures of angiosperm and gymnosperm species) promote the decomposition of low-quality litter after accounting for the positive impact of the focal species being a gymnosperm, which is in line with our hypothesis 1. Taken together, these results suggest that tree species diversity and specifically mixtures of angiosperms and gymnosperms can have positive effects on the decomposition of low-quality litter, as mentioned in previous research (Gartner and Cardon, 2004; Handa et al., 2014; Joly et al., 2017): (1) the presence of tree species with high-quality litter (which are more likely to be present in mixtures) promotes the breakdown

of low quality litter via nutrient transfer, improved water retention or other positive interactions with recalcitrant litter (Porre et al., 2020), and (2) more diverse substrates support a larger decomposer community, which is more likely to process low-quality litter (Vogel et al., 2013). A probable mechanism is that specific decomposers break down specific litter components, such as lignin or phenolics (Ristok et al., 2017), and the diversity of these different decomposer groups increases the overall decomposition process through niche partitioning. This implies that, beyond the overarching importance of focal tree identity, the surrounding environment and its diversity and composition could also affect decomposition (Hättenschwiler, 2005). Such diversity effects are often context dependent and cannot be generalized across continents and over different species compositions (Scherer-Lorenzen et al., 2007); this may explain the subtle differences found in our study, which could become more pronounced with time (Li et al., 2019; Xu et al., 2020).

4.3 Plantation conditions effects on mass loss

We did not find any effect of planting density on mass loss, irrespective of tea type, during early stages of forest development, leading to the rejection of hypothesis 3. This suggests that the density at which young trees were planted did not affect nutrient availability or the forest microclimate to the extent that it affected decomposition. In other studies of mature forests, variables such as canopy density and packing had some explanatory power for decomposition (Jucker et al., 2015; Trogisch et al., 2016), similarly we expect density effects in our experiments may strengthen as the experiments age. Furthermore, we found that mass loss of low-quality litter after 12 months marginally decreases with stand age ($P=0.09$), whereas we found no effect of stand age on green tea decomposition. We did, however, find a significant interaction effect

between age and tree identity during decomposition after 3 months for both rooibos and green tea: the impact of stand age on litter decomposition was more negative under gymnosperm overstories. Hence, we reject hypothesis 2, that tree species identity effects become more pronounced with the age of the stand, as we observed a reversal of the impact tree identity. The possible beneficial microclimate effect of young gymnosperm stands promoting decomposition (by providing a more buffered temperature regime (Zhang et al., 2022) or higher soil moisture due to shading (Petraglia et al., 2019)) compared to angiosperm stands is likely to disappear with stand age. In addition, the soil biochemical composition is likely to change over time under gymnosperm species, negatively affecting acidity status, nutrient availability, and microbial communities (Coûteaux et al., 1995). This illustrates that plantation age can alter identity effects and that caution is needed when generalizing our findings to mature stands where belowground functioning, canopy closure and microclimate may be very different (Trogisch et al., 2016; Joly et al., 2017; Zhang et al., 2022). In addition, the interaction effect between tree identity and stand age was not observed for the 12-month incubation period (Table 2). This suggests that earlier stages of litter decomposition may be better suited to understanding the relative importance of overstory-related decomposition drivers compared to mid-stages of decomposition; as decomposition progresses (and thus relative higher lignin content and less mass remains) overstory effects become subtler. This corresponds with García-Palacios et al. (2016) who also illustrated a shift in control from biotic (overstory) to abiotic (edaphic and climatic) factors with ongoing litter decay.

4.4 Limitations and scope for further research

This study was carried out in young plantations (maximum 12 years old) across boreal, temperate, Mediterranean, and subtropical biomes. As a consequence, some tree community effects on decomposition processes may emerge in the future as ecosystem functioning may become more strongly controlled by biotic factors during later stand development (Jucker et al., 2020; Xu et al., 2020). For example, decomposer community (Eisenhauer et al., 2012) and soil chemistry (Dhiedt et al., 2022), microclimatic conditions (Mayer et al., 2017), and litter production (in absolute quantities) change over time, and such differences can vary among biomes. Moreover, no information on edaphic properties could be included in our research, despite its explanatory importance for context-dependent effects on litter decomposition (Fanin et al., 2020; Desie et al., 2021). Previous research has indicated that the local environment (including edaphic properties, microrelief and microclimate (Seidelmann et al., 2016; Joly et al., 2017)), the quantity and quality of the litter (Briones, 2014), the composition of the local soil community (Hättenschwiler, 2005; Briones, 2014), as well as the interactions between litter quality and decomposing environment (Veen et al. 2018) all affect decomposition. Such interactions are very complex, and future large-scale studies should take into account the ambient soil conditions, litter quality, and microclimate (Makkonen et al., 2012). Nevertheless, our study using standardized litters across a multi-continent set of tree diversity experiments provides a first indication of the relative importance of the surrounding tree community and its diversity, relative to macroclimate.

5. Conclusions

Our findings highlight the important role of local conditions shaped by the forest overstory composition and stand structure as determinants of litter decomposition dynamics in young

472 plantations. Overstory diversity had no clear effect on mass loss, irrespective of tree type,
473 whereas tree species identity (and composition) affected decomposition of low-quality litter
474 directly via overstory litter quality and quantity and indirectly via changes in the micro-
475 environment. Moreover, these direct and indirect effects of tree species composition on litter
476 loss were dependent on the age of the stand. In these young stands, both planting density and
477 overstory composition had little effect on litter decomposition compared to litter quality and
478 macroclimate, suggesting limited consequences of management decisions related to planting
479 density or species choice at early stages of stand development. To further disentangle these
480 effects, it will be important for future studies to include a direct quantification of the micro-
481 environmental conditions at the site of decomposition. Further, we suggest not only relying on
482 standard substrates for such studies, which can only test the role of the environmental and
483 edaphic controls of decomposition, but combining them with the decomposition of plot-specific
484 litters that would capture the influence of tree diversity and species composition through their
485 effects on litter quality and quantity.

Acknowledgements

We are grateful to UNILEVER for sponsoring the Lipton tea bags and to ILTER Initiative Grant for supporting the work within the TeaComposition Initiative. We thank Stef Haesen for his assistance with the climate data and Ilié Storms for his feedback on manuscript. JZ worked on this paper with a postdoc grant from the Federal Belgian Science Office (Belspo) in the framework of the Forbio Climate project. NE and OF acknowledge support from the German Centre for Integrative Biodiversity Research Halle-Jena-Leipzig, funded by the German Research Foundation (FZT 118, 202548816). HB, ST, BY, StS and HF acknowledge support from the German Research Foundation (DFG FOR 891/3). We acknowledge the support of Agence nationale de la recherche (ANR), in the University framework of the Chinese Academy of Sciences (UCAS). MW acknowledges funding by the Swedish Energy Agency (36654-1, 36654-2, 36654-3project DiPTiCC (16-CE32-0003). MSL acknowledges support and site maintenance of the BIOTREE experiment by the Federal Forestry Office Thüringer Wald — Bundesforstamt Thüringer Wald, Bad Salzungen. We also thank Martin Mörsdorf (Freiburg) for assistance in field work.

Data accessibility

Data for this paper will be accessible at (doi: ...).

6. References

- Abatzoglou, J.T., Dobrowski, S.Z., Parks, S.A., Hegewisch, K.C., 2018. TerraClimate, a high-resolution global dataset of monthly climate and climatic water balance from 1958-2015. *Scientific Data* 5, 1–12. doi:10.1038/sdata.2017.191
- Aerts, R., 1997. Climate, Leaf Litter Chemistry and Leaf Litter Decomposition in Terrestrial Ecosystems: A Triangular Relationship. *Oikos* 79, 439. doi:10.2307/3546886
- Aranda, I., Forner, A., Cuesta, B., Valladares, F., 2012. Species-specific water use by forest tree species: From the tree to the stand. *Agricultural Water Management* 114, 67–77. doi:10.1016/j.agwat.2012.06.024
- Augusto, L., De Schrijver, A., Vesterdal, L., Smolander, A., Prescott, C., Ranger, J., 2015. Influences of evergreen gymnosperm and deciduous angiosperm tree species on the functioning of temperate and boreal forests. *Biological Reviews* 90, 444–466. doi:10.1111/brv.12119
- Barton, K., 2022. MuMIn: Multi-Model Inference. R package version 1.46.0.
- Bates, D., Maechler, M., Bolker, B., Walker, S., 2015. Fitting linear mixed-effects models using lme4. *J Stat Softw* 67, 1–48.
- Berg, B., McClaugherty, C., 2008. Plant litter: decomposition, humus formation, carbon sequestration. Springer-Verlag, Berlin, Germany. 1–10.
- Binkley, D., Giardina, C., 1998. Why do tree species affect soils? The Warp and Woof of tree-soil interactions. *Biogeochemistry* 42, 89–106.
- Blair, J.M., Parmelee, R.W., Beare, M.H., 1990. Decay Rates , Nitrogen Fluxes , and Decomposer Communities of Single- and Mixed-Species Foliar Litter. *JSTOR* 71, 1976–1985.

524 Bradford, M.A., Berg, B., Maynard, D.S., Wieder, W.R., Wood, S.A., 2016. Understanding the
 525 dominant controls on litter decomposition. *Journal of Ecology* 104, 229–238.
 526 doi:10.1111/1365-2745.12507

527 Briones, M.J.I., 2014. Soil fauna and soil functions: a jigsaw puzzle. *Frontiers in Environmental*
 528 *Science* 2, 7. doi:10.3389/fenvs.2014.00007

529 Bueis, T., Bravo, F., Pando, V., Turrión, M.B., 2018. Local basal area affects needle litterfall ,
 530 nutrient concentration , and nutrient release during decomposition in *Pinus halepensis*
 531 Mill . plantations in Spain. *Annals of Forest Science* 75.

532 Chomel, M., Guittonny-Larchevêque, M., DesRochers, A., Baldy, V., 2015. Home Field
 533 Advantage of Litter Decomposition in Pure and Mixed Plantations Under Boreal Climate.
 534 *Ecosystems* 18, 1014–1028. doi:10.1007/s10021-015-9880-y

535 Cornwell, W.K., Cornelissen, J.H.C., Amatangelo, K., Dorrepaal, E., Eviner, V.T., Godoy, O.,
 536 Hobbie, S.E., Hoorens, B., Kurokawa, H., Pérez-Harguindeguy, N., Quested, H.M., Santiago,
 537 L.S., Wardle, D.A., Wright, I.J., Aerts, R., Allison, S.D., van Bodegom, P., Brovkin, V., Chatain,
 538 A., Callaghan, T. V., Díaz, S., Garnier, E., Gurvich, D.E., Kazakou, E., Klein, J.A., Read, J.,
 539 Reich, P.B., Soudzilovskaia, N.A., Vaieretti, M.V., Westoby, M., 2008. Plant species traits
 540 are the predominant control on litter decomposition rates within biomes worldwide.
 541 *Ecology Letters* 11, 1065–1071. doi:10.1111/j.1461-0248.2008.01219.x

542 Coûteaux, M.-M., Bottner, P., Berg, B., 1995. Litter decomposition, climate and litter quality.
 543 *Trends in Ecology & Evolution* 10, 63–66. doi:10.1016/S0169-5347(00)88978-8

544 Davidson, E.A., Janssens, I.A., 2006. Temperature sensitivity of soil carbon decomposition and
 545 feedbacks to climate change. *Nature* 440, 165–173. doi:10.1038/nature04514

546 Dawud, S.M., Raulund-Rasmussen, K., Domisch, T., Finér, L., Jaroszewicz, B., Vesterdal, L., 2016.
 547 Is Tree Species Diversity or Species Identity the More Important Driver of Soil Carbon
 548 Stocks, C/N Ratio, and pH? *Ecosystems* 19, 645–660. doi:10.1007/s10021-016-9958-1
 549 Dawud, S.M., Raulund-Rasmussen, K., Ratcliffe, S., Domisch, T., Finér, L., Joly, F.-X.,
 550 Hättenschwiler, S., Vesterdal, L., 2017. Tree species functional group is a more important
 551 driver of soil properties than tree species diversity across major European forest types.
 552 *Functional Ecology* 31, 1153–1162. doi:10.1111/1365-2435.12821
 553 De Schrijver, A., Frenne, P., Staelens, J., Verstraeten, G., Muys, B., Vesterdal, L., Wuyts, K.,
 554 Nevel, L., Schelfhout, S., Neve, S., 2012. Tree species traits cause divergence in soil
 555 acidification during four decades of postagricultural forest development. *Global Change*
 556 *Biology* 18, 1127–1140.
 557 Desie, E., Muys, B., Jansen, B., Vesterdal, L., Vancampenhout, K., 2021. Pedogenic threshold in
 558 acidity explains context-dependent tree species effects on soil carbon. *Frontiers in Forests*
 559 *and Global Change*. doi:10.3389/ffgc.2021.679813
 560 Desie, E., Vancampenhout, K., Heyens, K., Hlava, J., Verheyen, K., Muys, B., 2019. Forest
 561 conversion to conifers induces a regime shift in soil process domain affecting carbon
 562 stability. *Soil Biology and Biochemistry* 136, 107540. doi:10.1016/J.SOILBIO.2019.107540
 563 Dhiedt, E., Baeten, L., De Smedt, P., Verheyen, K., 2022. Little effect of tree species richness on
 564 within- and between-plot variability in soil chemical properties in a young plantation
 565 forest. *European Journal of Soil Science* 73, 1–8. doi:10.1111/ejss.13164
 566 Djukic, I., Kepfer-Rojas, S., Schmidt, I.K., Larsen, K.S., Beier, C., Berg, B., Verheyen, K., Caliman,
 567 A., Paquette, A., Gutiérrez-Girón, A., Humber, A., Valdecantos, A., Petraglia, A., Alexander,

568 H., Augustaitis, A., Saillard, A., Fernández, A.C.R., Sousa, A.I., Lillebø, A.I., da Rocha Gripp,
 569 A., Francez, A.J., Fischer, A., Bohner, A., Malyshev, A., Andrić, A., Smith, A., Stanisci, A.,
 570 Seres, A., Schmidt, A., Avila, A., Probst, A., Ouin, A., Khuroo, A.A., Verstraeten, A., Palabral-
 571 Aguilera, A.N., Stefanski, A., Gaxiola, A., Muys, B., Bosman, B., Ahrends, B., Parker, B.,
 572 Sattler, B., Yang, B., Juráni, B., Erschbamer, B., Ortiz, C.E.R., Christiansen, C.T., Carol Adair,
 573 E., Meredieu, C., Mony, C., Nock, C.A., Chen, C.L., Wang, C.P., Baum, C., Rixen, C., Delire, C.,
 574 Piscart, C., Andrews, C., Rebmann, C., Branquinho, C., Polyanskaya, D., Delgado, D.F.,
 575 Wundram, D., Radeideh, D., Ordóñez-Regil, E., Crawford, E., Preda, E., Tropina, E., Groner,
 576 E., Lucot, E., Hornung, E., Gacia, E., Lévesque, E., Benedito, E., Davydov, E.A., Ampoorter,
 577 E., Bolzan, F.P., Varela, F., Kristöfel, F., Maestre, F.T., Maunoury-Danger, F., Hofhansl, F.,
 578 Kitz, F., Sutter, F., Cuesta, F., de Almeida Lobo, F., de Souza, F.L., Berninger, F., Zehetner, F.,
 579 Wohlfahrt, G., Vourlitis, G., Carreño-Rocabado, G., Arena, G., Pinha, G.D., González, G.,
 580 Canut, G., Lee, H., Verbeeck, H., Auge, H., Pauli, H., Nacro, H.B., Bahamonde, H.A.,
 581 Feldhaar, H., Jäger, H., Serrano, H.C., Verheyden, H., Bruelheide, H., Meesenburg, H.,
 582 Jungkunst, H., Jactel, H., Shibata, H., Kurokawa, H., Rosas, H.L., Rojas Villalobos, H.L.,
 583 Yesilonis, I., Melece, I., Van Halder, I., Quirós, I.G., Makelele, I., Senou, I., Fekete, I., Mihal,
 584 I., Ostonen, I., Borovská, J., Roales, J., Shoqeir, J., Lata, J.C., Theurillat, J.P., Probst, J.L.,
 585 Zimmerman, J., Vijayanathan, J., Tang, J., Thompson, J., Doležal, J., Sanchez-Cabeza, J.A.,
 586 Merlet, J., Henschel, J., Neiryneck, J., Knops, J., Loehr, J., von Oppen, J., Þorlákssdóttir, J.S.,
 587 Löffler, J., Cardoso-Mohedano, J.G., Benito-Alonso, J.L., Torezan, J.M., Morina, J.C.,
 588 Jiménez, J.J., Quinde, J.D., Alatalo, J., Seeber, J., Stadler, J., Kriiska, K., Coulibaly, K.,
 589 Fukuzawa, K., Szlavec, K., Gerháto, K., Lajtha, K., Käppeler, K., Jennings, K.A., Tielbörger,

590 K., Hoshizaki, K., Green, K., Yé, L., Pazianoto, L.H.R., Dienstbach, L., Williams, L., Yahdjian,
 591 L., Bringham, L.M., van den Brink, L., Rustad, L., Zhang, L., Morillas, L., Xiankai, L., Carneiro,
 592 L.S., Di Martino, L., Villar, L., Bader, M.Y., Morley, M., Lebouvier, M., Tomaselli, M.,
 593 Sternberg, M., Schaub, M., Santos-Reis, M., Glushkova, M., Torres, M.G.A., Giroux, M.A.,
 594 de Graaff, M.A., Pons, M.N., Bauters, M., Mazón, M., Frenzel, M., Didion, M., Wagner, M.,
 595 Hamid, M., Lopes, M.L., Apple, M., Schädler, M., Weih, M., Gualmini, M., Vadeboncoeur,
 596 M.A., Bierbaumer, M., Danger, M., Liddell, M., Mirtl, M., Scherer-Lorenzen, M., Růžek, M.,
 597 Carbognani, M., Di Musciano, M., Matsushita, M., Zhiyanski, M., Puşcaş, M., Barna, M.,
 598 Ataka, M., Jiangming, M., Alsafran, M., Carnol, M., Barsoum, N., Tokuchi, N., Eisenhauer,
 599 N., Lecomte, N., Filippova, N., Hölzel, N., Ferlian, O., Romero, O., Pinto, O.B., Peri, P.,
 600 Weber, P., Vittoz, P., Turtureanu, P.D., Fleischer, P., Macreadie, P., Haase, P., Reich, P.,
 601 Petřík, P., Choler, P., Marmonier, P., Muriel, P., Ponette, Q., Guariento, R.D., Canessa, R.,
 602 Kiese, R., Hewitt, R., Rønn, R., Adrian, R., Kanka, R., Weigel, R., Gatti, R.C., Martins, R.L.,
 603 Georges, R., Meneses, R.I., Gavilán, R.G., Dasgupta, S., Wittlinger, S., Puijalon, S., Freda, S.,
 604 Suzuki, S., Charles, S., Gogo, S., Drollinger, S., Mereu, S., Wipf, S., Trevathan-Tackett, S.,
 605 Löfgren, S., Stoll, S., Trogisch, S., Hoeber, S., Seitz, S., Glatzel, S., Milton, S.J., Dousset, S.,
 606 Mori, T., Sato, T., Ise, T., Hishi, T., Kenta, T., Nakaji, T., Michelin, T.S., Camboulive, T.,
 607 Mozdzer, T.J., Scholten, T., Spiegelberger, T., Zechmeister, T., Kleinebecker, T., Hiura, T.,
 608 Enoki, T., Ursu, T.M., di Cella, U.M., Hamer, U., Klaus, V.H., Rêgo, V.M., Di Cecco, V., Busch,
 609 V., Fontana, V., Piscová, V., Carbonell, V., Ochoa, V., Bretagnolle, V., Maire, V., Farjalla, V.,
 610 Zhou, W., Luo, W., McDowell, W.H., Hu, Y., Utsumi, Y., Kominami, Y., Zaika, Y., Rozhkov, Y.,
 611 Kotroczó, Z., Tóth, Z., 2018. Early stage litter decomposition across biomes. *Science of the*

612 Total Environment 628–629, 1369–1394. doi:10.1016/j.scitotenv.2018.01.012
 613 Eisenhauer, N., Reich, P.B., Scheu, S., 2012. Increasing plant diversity effects on productivity
 614 with time due to delayed soil biota effects on plants. Basic and Applied Ecology 13, 571–
 615 578. doi:10.1016/j.baae.2012.09.002
 616 Fanin, N., Bezaud, S., Sarneel, J.M., Cecchini, S., Nicolas, M., Augusto, L., 2020. Relative
 617 Importance of Climate, Soil and Plant Functional Traits During the Early Decomposition
 618 Stage of Standardized Litter. Ecosystems 23, 1004–1018. doi:10.1007/s10021-019-00452-z
 619 Finzi, A.C., Canham, C.D., Breemen, N. Van, 1998. CANOPY TREE–SOIL INTERACTIONS WITHIN
 620 TEMPERATE FORESTS: SPECIES EFFECTS ON pH AND CATIONS. Ecological Applications 8,
 621 447–454.
 622 Fox, J., Weisberg, S., Price, B., Friendly, M., Firth, D., Taylor, S., 2022. Package ‘ effects .’
 623 Fraser, L.H., Henry, H. Al, Carlyle, C.N., White, S.R., Beierkuhnlein, C., Cahill, J.F., Casper, B.B.,
 624 Cleland, E., Collins, S.L., Dukes, J.S., Knapp, A.K., Lind, E., Long, R., Luo, Y., Reich, P.B.,
 625 Smith, M.D., Sternberg, M., Turkington, R., 2013. Coordinated distributed experiments: An
 626 emerging tool for testing global hypotheses in ecology and environmental science.
 627 Frontiers in Ecology and the Environment 11, 147–155. doi:10.1890/110279
 628 García-Palacios, P., Shaw, E.A., Wall, D.H., Hättenschwiler, S., 2016. Temporal dynamics of biotic
 629 and abiotic drivers of litter decomposition. Ecology Letters 19, 554–563.
 630 doi:10.1111/ele.12590
 631 Gartner, T.B., Cardon, Z.G., 2004. Decomposition dynamics in mixed-species leaf litter. Oikos
 632 104, 230–246. doi:10.1111/j.0030-1299.2004.12738.x
 633 Gessner, M.O., Swan, C.M., Dang, C.K., McKie, B.G., Bardgett, R.D., Wall, D.H., Hättenschwiler,

634 S., 2010. Diversity meets decomposition. *Trends in Ecology & Evolution* 25, 372–380.
 635 doi:10.1016/J.TREE.2010.01.010

636 Gottschall, F., Davids, S., Newiger-Dous, T.E., Auge, H., Cesarz, S., Eisenhauer, N., 2019. Tree
 637 species identity determines wood decomposition via microclimatic effects. *Ecology and*
 638 *Evolution* 9, 12113–12127. doi:10.1002/ece3.5665

639 Handa, I.T., Aerts, R., Berendse, F., Berg, M.P., Bruder, A., Butenschoen, O., Chauvet, E.,
 640 Gessner, M.O., Jabiol, J., Makkonen, M., McKie, B.G., Malmqvist, B., Peeters, E.T.H.M.,
 641 Scheu, S., Schmid, B., Van Ruijven, J., Vos, V.C.A., Hättenschwiler, S., 2014. Consequences
 642 of biodiversity loss for litter decomposition across biomes. *Nature* 509, 218–221.
 643 doi:10.1038/nature13247

644 Hättenschwiler, S., 2005. Effects of Tree Species Diversity on Litter Quality and Decomposition,
 645 in: *Forest Diversity and Function*. Springer-Verlag, Berlin/Heidelberg, pp. 149–164.
 646 doi:10.1007/3-540-26599-6_8

647 Hobbie, S.E., Ogdahl, M., Reich, P.B., Oleksyn, J., Hale, C., Zytowski, R., Karolewski, P., 2006.
 648 Tree species effects on decomposition and forest floor dynamics in a common garden.
 649 *Ecology* 87, 2288–2297. doi:10.1890/0012-9658(2006)87[2288:TSEODA]2.0.CO;2

650 Hoeber, S., Fransson, P., Weih, M., Manzoni, S., 2020. Leaf litter quality coupled to *Salix* variety
 651 drives litter decomposition more than stand diversity or climate. *Plant and Soil* 453, 313–
 652 328. doi:10.1007/s11104-020-04606-0

653 Jewell, M.D., Shipley, B., Low-Décarie, E., Tobner, C.M., Paquette, A., Messier, C., Reich, P.B.,
 654 2017. Partitioning the effect of composition and diversity of tree communities on leaf litter
 655 decomposition and soil respiration. *Oikos* 126, 959–971. doi:10.1111/oik.03868

656 Joly, F.-X., Milcu, A., Scherer-Lorenzen, M., Jean, L.-K., Bussotti, F., Dawud, S.M., Müller, S.,
 657 Pollastrini, M., Raulund-Rasmussen, K., Vesterdal, L., Hättenschwiler, S., 2017. Tree species
 658 diversity affects decomposition through modified micro-environmental conditions across
 659 European forests. *New Phytologist* 214, 1281–1293. doi:10.1111/nph.14452
 660 Jucker, T., Bouriaud, O., Coomes, D.A., 2015. Crown plasticity enables trees to optimize canopy
 661 packing in mixed-species forests. *Functional Ecology* 29, 1078–1086. doi:10.1111/1365-
 662 2435.12428
 663 Jucker, T., Koricheva, J., Finér, L., Bouriaud, O., Iacopetti, G., Coomes, D.A., 2020. Good things
 664 take time—Diversity effects on tree growth shift from negative to positive during stand
 665 development in boreal forests. *Journal of Ecology* 108, 2198–2211. doi:10.1111/1365-
 666 2745.13464
 667 Keuskamp, J.A., Dingemans, B.J.J., Lehtinen, T., Sarneel, J.M., Hefting, M.M., 2013. Tea Bag
 668 Index: A novel approach to collect uniform decomposition data across ecosystems.
 669 *Methods in Ecology and Evolution* 4, 1070–1075. doi:10.1111/2041-210X.12097
 670 Kwon, T.O., Shibata, H., Kepfer-Rojas, S., Schmidt, I.K., Larsen, K.S., Beier, C., Berg, B., Verheyen,
 671 K., Lamarque, J.F., Hagedorn, F., Eisenhauer, N., Djukic, I., 2021. Effects of Climate and
 672 Atmospheric Nitrogen Deposition on Early to Mid-Term Stage Litter Decomposition Across
 673 Biomes. *Frontiers in Forests and Global Change* 4, 1–18. doi:10.3389/ffgc.2021.678480
 674 Lado-monserrat, L., Lidon, A., Bautista, I., 2015. Litterfall , litter decomposition and associated
 675 nutrient fluxes in *Pinus halepensis* : influence of tree removal intensity in a Mediterranean
 676 forest. *European Journal of Forest Research* 134, 833–844. doi:10.1007/s10342-015-0893-
 677 z

678 Li, Y., Bruelheide, H., Scholten, T., Schmid, B., Sun, Z., Zhang, N., Bu, W., Liu, X., Ma, K., 2019.
 679 Early positive effects of tree species richness on soil organic carbon accumulation in a
 680 large-scale forest biodiversity experiment. *Journal of Plant Ecology* 12, 882–893.
 681 doi:10.1093/jpe/rtz026

682 Lin, H., Li, Y., Bruelheide, H., Zhang, S., Ren, H., Zhang, N., Ma, K., 2021. What drives leaf litter
 683 decomposition and the decomposer community in subtropical forests – The richness of the
 684 above-ground tree community or that of the leaf litter? *Soil Biology and Biochemistry* 160,
 685 108314. doi:10.1016/j.soilbio.2021.108314

686 Makkonen, M., Berg, M.P., Handa, I.T., Hättenschwiler, S., van Ruijven, J., van Bodegom, P.M.,
 687 Aerts, R., 2012. Highly consistent effects of plant litter identity and functional traits on
 688 decomposition across a latitudinal gradient. *Ecology Letters* 15, 1033–1041.
 689 doi:10.1111/j.1461-0248.2012.01826.x

690 Maxwell, T.L., Augusto, L., Bon, L., Courbineau, A., Altinalmazis-Kondylis, A., Milin, S., Bakker,
 691 M.R., Jactel, H., Fanin, N., 2020. Effect of a tree mixture and water availability on soil
 692 nutrients and extracellular enzyme activities along the soil profile in an experimental
 693 forest. *Soil Biology and Biochemistry* 148. doi:10.1016/j.soilbio.2020.107864

694 Mayer, M., Matthews, B., Rosinger, C., Sandén, H., Godbold, D.L., Katzensteiner, K., 2017. Tree
 695 regeneration retards decomposition in a temperate mountain soil after forest gap
 696 disturbance. *Soil Biology and Biochemistry* 115, 490–498.
 697 doi:10.1016/j.soilbio.2017.09.010

698 Messier, C., Bauhus, J., Sousa-Silva, R., Auge, H., Baeten, L., Barsoum, N., Bruelheide, H.,
 699 Caldwell, B., Cavender-Bares, J., Dhiedt, E., Eisenhauer, N., Ganade, G., Gravel, D.,

700 Guillemot, J., Hall, J.S., Hector, A., Hérault, B., Jactel, H., Koricheva, J., Kreft, H., Mereu, S.,
 701 Muys, B., Nock, C.A., Paquette, A., Parker, J.D., Perring, M.P., Ponette, Q., Potvin, C., Reich,
 702 P.B., Scherer-Lorezen, M., Schnabel, F., Verheyen, K., Weih, M., Wollni, M., Zemp, D.C.,
 703 2022. For the sake of resilience and multifunctionality, let's diversify planted forests!
 704 Conservation Letters 15, 1–8. doi:10.1111/conl.12829

705 Naeem, S., Thompson, L.J., Lawler, S.P., Lawton, J.H., Woodfin, B.R.M., 1994. of Ecosystems.
 706 Nature 368, 19–22.

707 Nakagawa, S., Schielzeth, H., 2013. A general and simple method for obtaining R² from
 708 generalized linear mixed-effects models. Methods in Ecology and Evolution 4, 133–142.

709 Paquette, A., Hector, A., Castagneyrol, B., Vanhellemont, M., Koricheva, J., Scherer-Lorezen,
 710 M., Verheyen, K., 2018. A million and more trees for science. Nature Ecology & Evolution 2,
 711 763–766. doi:10.1038/s41559-018-0544-0

712 Peng, Y., Holmstrup, M., Kappel, I., Ruggiero, L., Schelfhout, S., Zheng, H., Hed, P., Yue, K.,
 713 Vesterdal, L., 2022. Tree species identity is the predominant modulator of the effects of
 714 soil fauna on leaf litter decomposition. Forest Ecology and Management 520.
 715 doi:10.1016/j.foreco.2022.120396

716 Petraglia, A., Cacciatori, C., Chelli, S., Fenu, G., Calderisi, G., Gargano, D., Abeli, T., Orsenigo, S.,
 717 Carbognani, M., 2019. Litter decomposition: effects of temperature driven by soil moisture
 718 and vegetation type. Plant and Soil 435, 187–200. doi:10.1007/s11104-018-3889-x

719 Porre, R.J., van der Werf, W., De Deyn, G.B., Stomph, T.J., Hoffland, E., 2020. Is litter
 720 decomposition enhanced in species mixtures? A meta-analysis. Soil Biology and
 721 Biochemistry 145, 107791. doi:10.1016/j.soilbio.2020.107791

722 Prescott, C.E., 2002. The influence of the forest canopy on nutrient cycling. *Tree Physiology* 22,
 723 1193–1200. doi:10.1093/treephys/22.15-16.1193

724 Prescott, C.E., Vesterdal, L., 2013. Tree species effects on soils in temperate and boreal forests:
 725 Emerging themes and research needs. *Forest Ecology and Management* 309, 1–3.
 726 doi:10.1016/j.foreco.2013.06.042

727 Reich, P.B., Oleksyn, J., Modrzynski, J., Mrozinski, P., Hobbie, S.E., Eissenstat, D.M., Chorover, J.,
 728 Chadwick, O.A., Hale, C.M., Tjoelker, M.G., 2005. Linking litter calcium, earthworms and
 729 soil properties: a common garden test with 14 tree species. *Ecology Letters* 8, 811–818.

730 Ristok, C., Leppert, K.N., Franke, K., Scherer-Lorenzen, M., Niklaus, P.A., Wessjohann, L.A.,
 731 Bruelheide, H., 2017. Leaf litter diversity positively affects the decomposition of plant
 732 polyphenols. *Plant and Soil* 419, 305–317. doi:10.1007/s11104-017-3340-8

733 Scherer-Lorenzen, M., Bonilla, J.L., Potvin, C., 2007. Tree species richness affects litter
 734 production and decomposition rates in a tropical biodiversity experiment. *Oikos* 116,
 735 2108–2124. doi:10.1111/j.2007.0030-1299.16065.x

736 Schwarz, B., Dietrich, C., Cesarz, S., Scherer-Lorenzen, M., Auge, H., Schulz, E., Eisenhauer, N.,
 737 2015. Non-significant tree diversity but significant identity effects on earthworm
 738 communities in three tree diversity experiments. *European Journal of Soil Biology* 67, 17–
 739 26. doi:10.1016/J.EJSOBI.2015.01.001

740 Seidelmann, K.N., Scherer-lorenzen, M., Niklaus, P.A., 2016. Direct vs . Microclimate-Driven
 741 Effects of Tree Species Diversity on Litter Decomposition in Young Subtropical Forest
 742 Stands 1–16. doi:10.1371/journal.pone.0160569

743 Silva Pedro, M., Rammer, W., Seidl, R., 2015. Tree species diversity mitigates disturbance

744 impacts on the forest carbon cycle. *Oecologia* 177, 619–630. doi:10.1007/s00442-014-
 745 3150-0
 746 Stoffel, M.A., Nakagawa, S., Schielzeth, H., 2021. partR2 : Partitioning R² in generalized linear
 747 mixed models.
 748 Strukelj, M., Parker, W., Corcket, E., Augusto, L., Khelifa, R., Jactel, H., Munson, A.D., 2021. Tree
 749 species richness and water availability interact to affect soil microbial processes. *Soil*
 750 *Biology and Biochemistry* 155. doi:10.1016/j.soilbio.2021.108180
 751 Tilman, D., Isbell, F., Cowles, J.M., 2014. Biodiversity and ecosystem functioning. *Annual Review*
 752 *of Ecology, Evolution, and Systematics* 45, 471–493. doi:10.1146/annurev-ecolsys-120213-
 753 091917
 754 Trogisch, S., He, J., Hector, A., Scherer-lorenzen, M., 2016. Impact of species diversity , stand
 755 age and environmental factors on leaf litter decomposition in subtropical forests in China
 756 337–350. doi:10.1007/s11104-015-2737-5
 757 Verheyen, K., Vanhellefont, M., Auge, H., Baeten, L., Baraloto, C., Barsoum, N., Bilodeau-
 758 Gauthier, S., Bruelheide, H., Castagneyrol, B., Godbold, D., Haase, J., Hector, A., Jactel, H.,
 759 Koricheva, J., Loreau, M., Mereu, S., Messier, C., Muys, B., Nolet, P., Paquette, A., Parker,
 760 J., Perring, M., Ponette, Q., Potvin, C., Reich, P., Smith, A., Weih, M., Scherer-Lorenzen, M.,
 761 2016. Contributions of a global network of tree diversity experiments to sustainable forest
 762 plantations. *Ambio* 45, 29–41. doi:10.1007/s13280-015-0685-1
 763 Vivanco, L., Austin, A.T., 2008. Tree species identity alters forest litter decomposition through
 764 long-term plant and soil interactions in Patagonia, Argentina. *Journal of Ecology* 96, 727–
 765 736. doi:10.1111/j.1365-2745.2008.01393.x

766 Vogel, A., Eisenhauer, N., Weigelt, A., Scherer-Lorenzen, M., 2013. Plant diversity does not
 767 buffer drought effects on early-stage litter mass loss rates and microbial properties. *Global*
 768 *Change Biology* 19, 2795–2803. doi:10.1111/gcb.12225

769 Wardle, D., Lavelle, P., Cadisch, G., Giller, K., 1997. Linkages between soil biota, plant litter
 770 quality and decomposition, in: *Driven by Nature: Plant Litter Quality and Decomposition*.
 771 CAB International, Wallingford, Oxon, pp. 107–124.

772 Wardle, D.A., Bardgett, R.D., Klironomos, J.N., Setälä, H., van der Putten, W.H., Wall, D.H., 2004.
 773 Ecological linkages between aboveground and belowground biota. *Science (New York,*
 774 *N.Y.)* 304, 1629–33. doi:10.1126/science.1094875

775 Wardle, D.A., Nicholson, K.S., 1996. Synergistic Effects of Grassland Plant Species on Soil
 776 Microbial Biomass and Activity: Implications for Ecosystem-Level Effects of Enriched Plant
 777 Diversity. *Functional Ecology* 10, 410. doi:10.2307/2390291

778 Xu, S., Eisenhauer, N., Ferlian, O., Zhang, J., Zhou, G., Lu, X., Liu, C., Zhang, D., 2020. Species
 779 richness promotes ecosystem carbon storage: Evidence from biodiversity-ecosystem
 780 functioning experiments: Species richness promotes C storage. *Proceedings of the Royal*
 781 *Society B: Biological Sciences* 287. doi:10.1098/rspb.2020.2063rspb20202063

782 Zhang, N., Bruehlheide, H., Li, Y., Liang, Y., Wubet, T., Trogisch, S., Ma, K., 2020. Community and
 783 neighbourhood tree species richness effects on fungal species in leaf litter. *Fungal Ecology*
 784 47, 100961. doi:10.1016/j.funeco.2020.100961

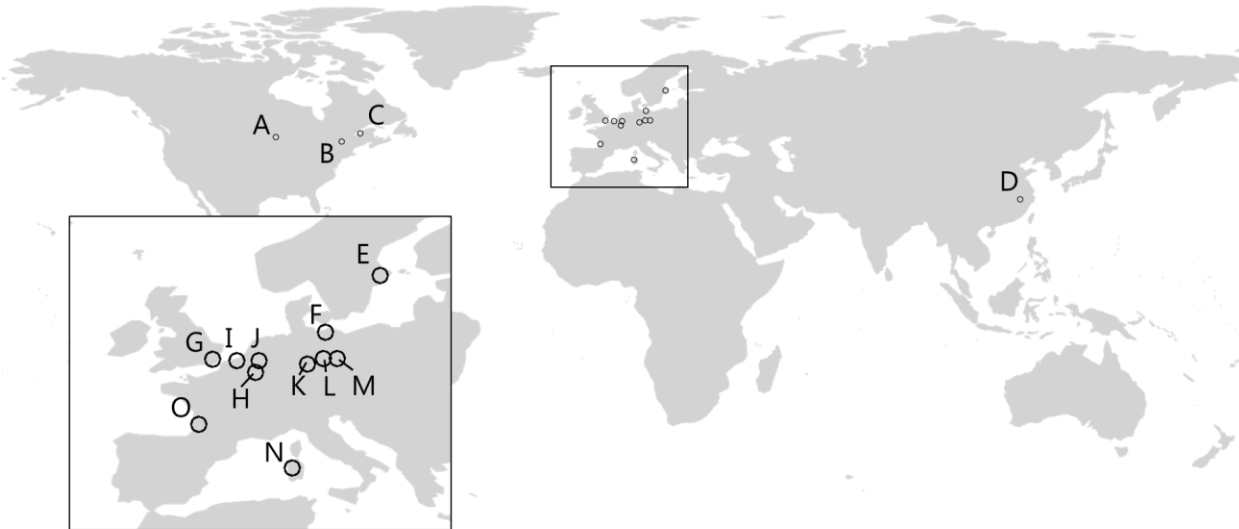
785 Zhang, S., Landuyt, D., Verheyen, K., De Frenne, P., 2022. Tree species mixing can amplify
 786 microclimate offsets in young forest plantations. *Journal of Applied Ecology* 1–12.
 787 doi:10.1111/1365-2664.14158

788 Zhou, S., Butenschoen, O., Barantal, S., Handa, I.T., Makkonen, M., Vos, V., Aerts, R., Berg, M.P.,
789 McKie, B., Van Ruijven, J., Hättenschwiler, S., Scheu, S., 2020. Decomposition of leaf litter
790 mixtures across biomes: The role of litter identity, diversity and soil fauna. *Journal of*
791 *Ecology* 2283–2297. doi:10.1111/1365-2745.13452

792 Zuo, J., Hefting, M.M., Berg, M.P., van Logtestijn, R.S.P., van Hal, J., Goudzwaard, L., Liu, J.-C.,
793 Sass-Klaassen, U., Sterck, F.J., Poorter, L., Cornelissen, J.H.C., 2018. Is there a tree
794 economics spectrum of decomposability? *Soil Biology and Biochemistry* 119, 135–142.
795 doi:10.1016/j.soilbio.2018.01.019

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7. Supporting Information



- | | | |
|--|--|--|
| A. US / IDENT / Cloquet | B. CA / IDENT / Montreal | C. CA / IDENT / Auclair |
| D. CN / BEF-China / Xingangshan | E. SE / ECOLINK-Salix / Uppsala | F. DE / ECOLINK-Salix / Rostock |
| G. UK / Climate Match / Kent | H. BE / FORBIO / Gedinne | I. BE / FORBIO / Zedelgem |
| J. BE / FORBIO / Hechtel-Eksel | K. DE / BIOTREE / Kaltenborn | L. DE / Kreinitz / Zeithain |
| M. DE / MyDiv / Bad Lauchstadt | N. IT / IDENT / Macomer | O. FR / ORPHEE / Pierroton |

Figure S1. Geographical distribution of the site locations. Experimental sites are indicated with a letter code.

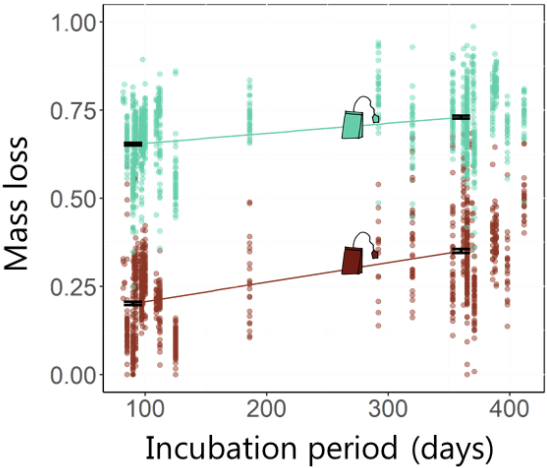


Figure S2. Mass loss (%) as a function of incubation period for the two litter types. Mean and SE are indicated for the 3-month and 12-month periods by the black horizontal errorbar.

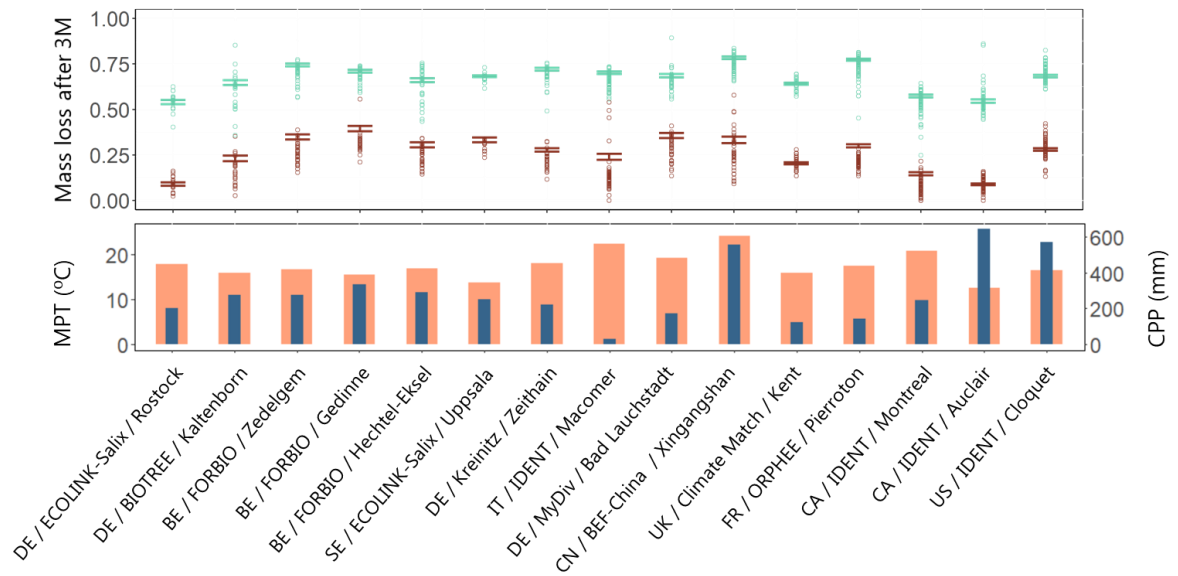


Figure S3. Top: Mass loss (mean and SE) after 3 months of incubation for each site per tea type: green tea is indicated in light green and rooibos tea is indicated in dark red. Bottom: mean period temperature (MPT) (orange wide bars) and cumulative period precipitation (CPP) (blue narrow bars) for the 3 month incubation period per site.

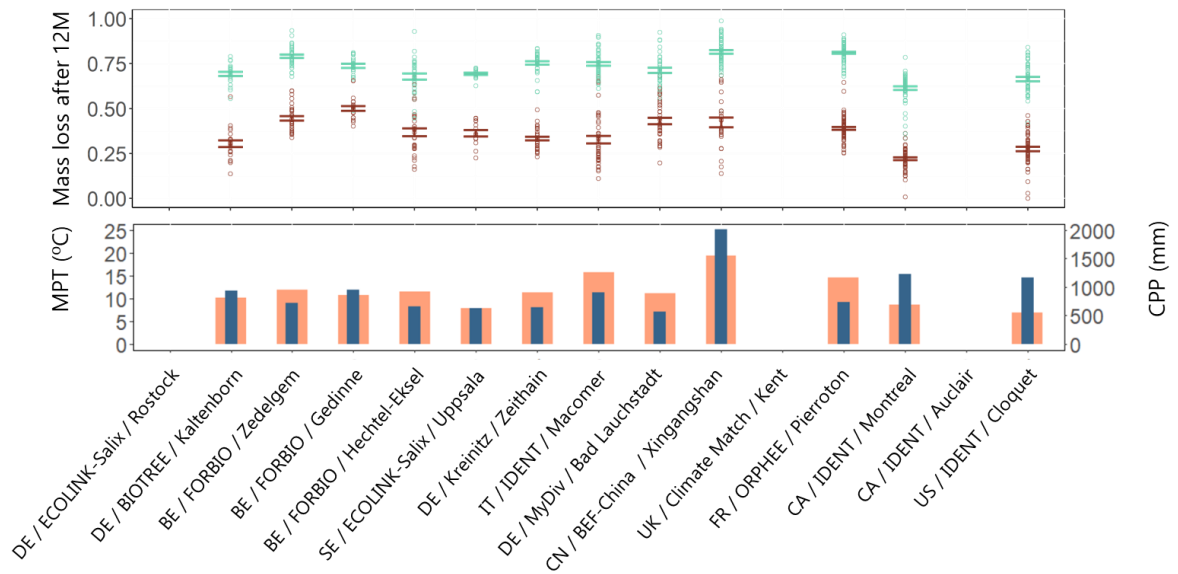


Figure S4. Top: Mass loss (mean and SE) after 12 months of incubation for each site per tea type: green tea is indicated in light green and rooibos tea is indicated in dark red. Bottom: mean period temperature (MPT) (orange wide bars) and cumulative period precipitation (CPP) (blue narrow bars) for the 12 month incubation period per site.

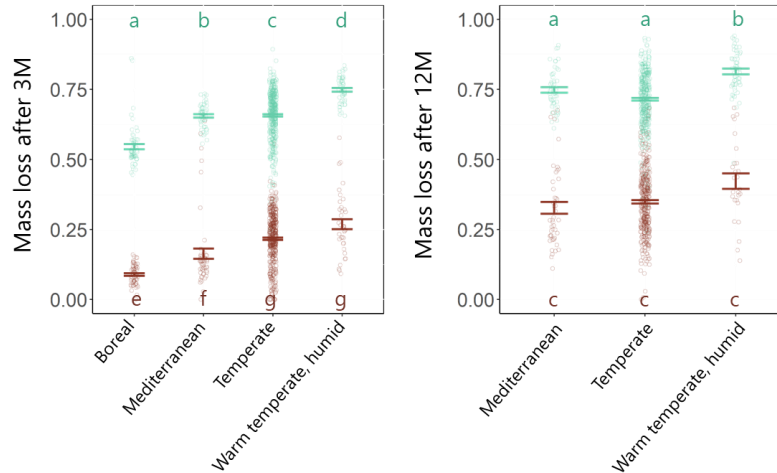


Figure S5. Mass loss (mean and se) after 3 months (left) and 12 months (right) of incubation for each biome and per tea type: green tea is indicated in light green and rooibos tea is indicated in dark red. Results of a multiple comparison test are indicated in the top and bottom of the graph: different letters indicate significant differences ($\alpha<0.05$) between biomes.

Table S1. Tree species of the experiment with their abbreviation and group (Ang = angiosperm; Gym = gymnosperm). Focal tree species are indicated in bold.

Tree species	Abbreviation	Group	Tree species	Abbreviation	Group
<i>Acer davidii</i>	ACDA	Ang	<i>Abies balsamea</i>	ABBA	Gym
<i>Acer monspessulanum</i>	ACMO	Ang	<i>Larix decidua</i>	LADE	Gym
<i>Acer platanoides</i>	ACPL	Ang	<i>Larix x marschlinii</i>	LAEU	Gym
<i>Acer rubrum</i>	ACRU	Ang	<i>Larix kaempferi</i>	LAKA	Gym
<i>Acer saccharum</i>	ACSA	Ang	<i>Larix laricina</i>	LALA	Gym
<i>Acer pseudoplatanus</i>	ACPS	Ang	<i>Picea abies</i>	PIAB	Gym
<i>Aesculus hippocastanum</i>	AEHI	Ang	<i>Picea glauca</i>	PIGL	Gym
<i>Arbutus unedo</i>	ARUN	Ang	<i>Pinus halepensis</i>	PIHA	Gym
<i>Betula alleghaniensis</i>	BEAL	Ang	<i>Picea omorika</i>	PIOM	Gym
<i>Betula papyrifera</i>	BEPA	Ang	<i>Pinus pinaster</i>	PIPI	Gym
<i>Betula pendula</i>	BEPE	Ang	<i>Pinus pinea</i>	PIPIEA	Gym
<i>Carpinus betulus</i>	CABE	Ang	<i>Pinus resinosa</i>	PIRE	Gym
<i>Castanopsis carlesii</i>	CACA	Ang	<i>Picea rubens</i>	PIRU	Gym
<i>Castanopsis eyrei</i>	CAEY	Ang	<i>Pinus strobus</i>	PIST	Gym
<i>Castanea henryi</i>	CAHE	Ang	<i>Pinus sylvestris</i>	PISY	Gym
<i>Castanea sativa</i>	CASA	Ang	<i>Pseudotsuga menziesii</i>	PSME	Gym
<i>Castanopsis sclerophylla</i>	CASC	Ang	<i>Thuja occidentalis</i>	THOC	Gym
<i>Choerospondias axillaris</i>	CHAX	Ang			
<i>Cinnamomum camphora</i>	CICA	Ang			
<i>Cyclobalanopsis glauca</i>	CYGL	Ang			
<i>Cyclobalanopsis myrsinaefolia</i>	CYMY	Ang			
<i>Daphniphyllum oldhamii</i>	DAOI	Ang			
<i>Diospyros glaucifolia</i>	DIGL	Ang			
<i>Fagus sylvatica</i>	FASY	Ang			
<i>Fraxinus excelsior</i>	FREX	Ang			
<i>Fraxinus ornus</i>	FROR	Ang			
<i>Koelreuteria bipinnata</i>	KOBI	Ang			
<i>Liquidambar formosana</i>	LIFO	Ang			
<i>Lithocarpus glaber</i>	LIGL	Ang			
<i>Melia azedarach</i>	MEAZ	Ang			
<i>Nyssa sinensis</i>	NYSI	Ang			
<i>Olea europaea</i>	OLEU	Ang			
<i>Phillyrea angustifolia</i>	PHAN	Ang			
<i>Pistacia lentiscus</i>	PILE	Ang			

<i>Prunus avium</i>	PRAV	Ang				
<i>Quercus acutissima</i>	QUAC	Ang				
<i>Quercus fabri</i>	QUFA	Ang				
<i>Quercus ilex</i>	QUIL	Ang				
<i>Quercus petraea</i>	QUPE	Ang				
<i>Quercus pubescens</i>	QUPU	Ang				
<i>Quercus robur</i>	QURO	Ang				
<i>Quercus rubra</i>	QURU	Ang				
<i>Quercus serrata</i>	QUSE	Ang				
<i>Quercus suber</i>	QUSU	Ang				
<i>Rhus chinensis</i>	RHCH	Ang				
<i>Sapindus mukorossi</i>	SAMU	Ang				
<i>Schima superba</i>	SCSU	Ang				
<i>Sorbus aucuparia</i>	SOAU	Ang				
<i>Quercus rubra</i>	QURU	Ang				
<i>Tilia cordata</i>	TICO	Ang				
<i>Tilia platyphyllos</i>	TIPL	Ang				
<i>Triadica cochinchinensis</i>	TRCO	Ang				
<i>Triadica sebifera</i>	TRSE	Ang				
<i>Salix dasyclados</i> (cv 'Loden')	SADA	Ang				
<i>Salix schwerinii</i> x <i>viminalis</i> (cv 'Tora')	SASV	Ang				
<i>S. schwerinii</i> x <i>viminalis</i> + <i>S. dasyclados</i>	SADASV	Ang				

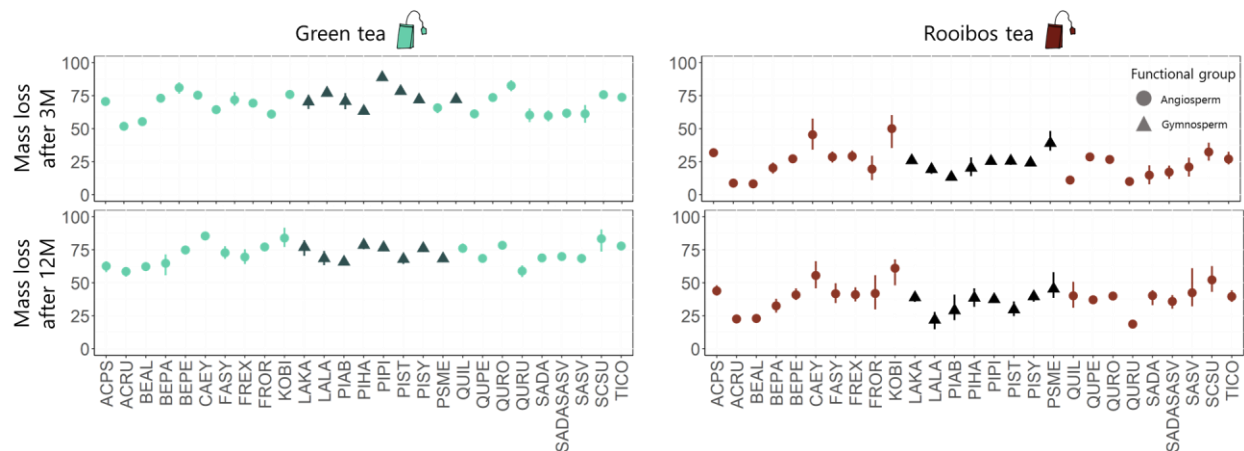


Figure S7: Mass loss (mean and SE) after 3 months (top) and 12 months (bottom) per focal tree species for green tea (left) and rooibos tea (right). Angiosperm species are indicated with light circles and gymnosperm species with dark triangles. Abbreviations per tree species name are found in Table S1.

Table S2: ANOVA output of linear mixed models testing the effect of incubation period, litter type and the interaction. Site and plot were included as random factors.

	Sum Sq	Mean Sq	NumDF	DenDF	F value	p
Incubation period	4.81	4.81	1	2034.5	818.06	< 0.001
Litter type	35.69	35.69	1	1983.6	6070.40	< 0.001
Incubation period : Litter type	0.77	0.77	1	1985.3	131.04	< 0.001

Table S3: ANOVA output of linear mixed models testing the effect of incubation period, litter type, biome and their interactions. Site and plot were included as random factors.

	Sum Sq	Mean Sq	NumDF	DenDF	F value	p
Incubation period	4.63	4.63	1	2020.38	798.27	< 0.001

Litter type	28.14	28.14	1	1983.61	4851.00	< 0.001
Biome	0.33	0.11	3	207.92	19.13	< 0.001
Incubation period: Litter type	0.72	0.72	1	1983.17	125.11	< 0.001
Incubation period: Biome	0.17	0.05	3	1988.65	10.03	< 0.001

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Table S4. P-values for fixed effects in alternative models (including cumulative precipitation and aridity indices $GI = \text{precipitation} / 2 * \text{temperature}$ and $AI = \text{precipitation} / \text{potential evotranspiration}$) for the incubation period and/or excluding BEF-China) accounting for the random effects site and plot (plot nested in site) executed on centered variables. The models included in the main manuscript are highlighted in respectively green and red.

3M	Mass loss ~	Incubation	Identity	Age	Identity*Age	Composition	SR	Density	MPT	CPP	
	GREEN	0.005	0.32	0.86	0.01	0.86	0.50	0.13	0.35	0.29	
	RED	0.30	0.02	0.16	<0.001	0.52	0.59	0.06	0.80	0.20	
	Mass loss ~	Incubation	Identity	Age	Identity*Age	Composition	SR	Density	MPT	CPP	MPT*CPP
	GREEN	0.005	0.33	0.88	0.01	0.89	0.50	0.75	0.71	0.65	0.51
	RED	0.32	0.03	0.18	<0.001	0.54	0.59	0.62	0.51	0.59	0.43
	Mass loss ~	Incubation	Identity	Age	Identity*Age	Composition	SR	Density	MPT	GI = P / 2*T	
	GREEN	0.005	0.32	0.93	0.01	0.89	0.49	0.16	0.28	0.39	
	RED	0.32	0.02	0.21	<0.001	0.51	0.57	0.09	0.90	0.35	
	Mass loss ~	Incubation	Identity	Age	Identity*Age	Composition	SR	Density		GI = P / 2*T	
	GREEN	0.007	0.33	0.98	0.01	0.88	0.42	0.26		0.76	
	RED	0.32	0.02	0.19	<0.001	0.51	0.55	0.07		0.27	
12M	Mass loss ~	Incubation	Identity	Age	Identity*Age	Composition	SR	Density	MPT	AI = P / PET	
	GREEN	0.005	0.33	0.95	0.01	0.90	0.49	0.15	0.30	0.34	
	RED	0.31	0.02	0.18	<0.001	0.52	0.58	0.06	0.98	0.22	
	Mass loss ~	Incubation	Identity	Age	Identity*Age	Composition	SR	Density		AI = P / PET	
	GREEN	0.006	0.33	0.95	0.01	0.89	0.43			0.53	
	RED	0.30	0.02	0.16	<0.001	0.52	0.58	0.05		0.16	
	Mass loss ~	Incubation	Identity	Age	Identity*Age	Composition	SR	Density	MPT	CPP	
	GREEN	0.01	0.79	0.78	0.92	0.36	0.06	0.71	0.01	0.34	
	RED	<0.001	0.22	0.02	0.26	0.02	0.62	0.03	0.49	0.48	
	Mass loss ~	Incubation	Identity	Age	Identity*Age	Composition	SR	Density	MPT	CPP	MPT*CPP
	GREEN	0.008	0.81	0.44	0.26	0.34	0.07	0.60	0.01	0.15	0.26
	GREEN (without BEF-China)	0.09	0.83	0.77	0.81	0.34	0.71	0.37	0.05	0.08	0.49
	RED	<0.001	0.23	0.09	0.24	0.02	0.55	0.26	0.59	0.71	0.35
12M	Mass loss ~	Incubation	Identity	Age		Composition	SR	Density	MPT	GI = P / 2*T	
	GREEN	0.01	0.79	0.76	0.92	0.36	0.06	0.91	0.01	0.39	
	RED	<0.001	0.22	0.03	0.26	0.02	0.63	0.05	0.15	0.52	
	Mass loss ~	Incubation	Identity	Age		Composition	SR	Density		GI = P / 2*T	
	GREEN	0.03	0.84	0.91	0.80	0.33	0.04	0.26		0.65	
	GREEN (without BEF-China)	0.09	0.80	0.57	0.84	0.36	0.74	0.21		0.02	
	RED	<0.001	0.21	0.04	0.22	0.03	0.87	0.05		0.80	
	Mass loss ~	Incubation	Identity	Age		Composition	SR	Density	MPT	AI = P / PET	
	GREEN	0.01	0.79	0.89	0.91	0.36	0.06	0.61	0.01	0.43	
	RED	<0.001	0.22	0.02	0.26	0.02	0.60	0.02	0.17	0.38	
	Mass loss ~	Incubation	Identity	Age		Composition	SR	Density		AI = P / PET	
	GREEN	0.03	0.80	0.91	0.80	0.33	0.04	0.26		0.65	
	GREEN (without BEF-China)	0.18	0.80	0.88	0.79	0.36	0.74	0.57		0.02	
	RED	<0.001	0.21	0.02	0.22	0.03	0.79	0.01		0.42	

Table S6: Results of variance partitioning combining all tea types.

R ² term	estimate for 3 months	estimate for 12 months
Full	0.934461	0.858468
Fixed effects	0.836022	0.80911
Random effects	0.098439	0.049358
Climate (MPT + CPP)	0.009737	0.022611
Plantation conditions (planting density + stand age)	0.001462	0.001005
Species composition (Tree identity + tree composition)	0.00015	0
Species richness	0.000319	0.000528
Tea type	0.806591	0.703487
Incubation period	0.000423	0.012451
Higher order interactions	0.01734	0.069027
Variance not explained	0.065539	0.141532

Table S7: Results of variance partitioning per tea type and per incubation period

R ² term	Green 3 months	Green 12 months	Rooibos 3 months	Rooibos 12 months
Full	0.698917	0.496436	0.673709	0.64771
Fixed effects	0.115483	0.363739	0.120832	0.332876
Random effects	0.583433	0.132697	0.552877	0.314834
Climate	0.010978	0.233291	0.037085	0.008118
Plantation conditions	0.004356	0.010577	0.035503	0.024143
Species composition	0.001116	0.004516	0.003746	0.012015
Species richness	0.001404	0.002817	0.000948	0.000916
Incubation period	0.007021	0.023169	0.001311	0.081743
Higher order interactions	0.090608	0.089369	0.042239	0.205941
Variance not explained	0.301083	0.503564	0.326291	0.35229

Table 1. Basic characteristics of the experimental sites and number of tea bags used in this study.

No.	Country	Name of Experiment	Site	Location		Climate	MAT ^a	MAP ^b	Local weather during experiment				SR gradient ^e	Tree planting year	Tree density (tree m ⁻²)	No. bags 3M	No. bags 12M
				Latitude	Longitude				MPT ^c 3M	MPT 12M	CPP ^d 3M	CPP 12M					
A	Belgium	FORBIO	Gedinne	49.99	4.98	Temperate	10.40	670	15.45	10.81	322.20	897.30	1, 2, 4	2010	0.44	62	52
B	Belgium	FORBIO	Hechtel-Eksel	51.16	5.31	Temperate	8.60	1030	16.99	11.43	278.20	631.80	1, 2, 4	2012	0.44	69	69
C	Belgium	FORBIO	Zedelgem	51.15	3.12	Temperate	10.10	708	16.73	11.90	265.00	688.30	1, 2, 4	2009	0.44	71	71
D	Canada	IDENT	Auclair	48.23	-69.10	Boreal	2.30	1015	12.52	4.88	615.90	1400.90	1, 2, 6	2010	6.25	126	NA ^g
E	Canada	IDENT	Montreal	45.86	-73.93	Temperate	6.20	976	20.78	8.64	236.90	1166.40	1, 2, 4	2009	4.00	120	120
F	China	BEF-China	Xingangshan	29.12	117.91	Warm-temperate, humid	17.10	1777	24.10	19.41	534.60	1903.30	1, 8, 24	2009	0.60	94	101
G	Germany	BIOTREE	Kaltenborn	50.78	10.22	Temperate	7.80	650	15.83	10.22	266.00	885.90	1, 2, 4	2004	0.25/0.44	48	47
H	Germany	ECOLINK-Salix	Rostock	54.06	12.08	Temperate	8.50	590	17.92	10.17	194.20	810.20	1, 2	2014	1.56	36	NA ^g
I	Germany	Kreinitz	Zeithain	51.23	13.15	Temperate	8.40	575	18.04	11.24	212.60	607.30	1, 3, 6	2005	1.25	68	72
J	Germany	MyDiv	Bad Lauchstädt	51.39	11.88	Temperate	9.00	492	19.32	11.17	164.70	538.40	1, 2, 4	2015	1.00	72	71
K	France	ORPHEE	Pierroton	44.74	-0.80	Temperate	12.75	876	17.55	14.63	136.00	705.30	1, 2, 3	2008	0.25	141	141
L	Italy	IDENT	Macomer	40.24	8.70	Mediterranean	13.80	866	22.33	15.74	28.80	867.00	1, 2, 6	2014	4.00	108	108
M	Sweden	ECOLINK-Salix	Uppsala	60.44	18.08	Temperate	5.60	470	13.79	7.93	241.80	591.30	1, 2	2014	1.56	26	31
N	UK	Climate-match	Kent	53.40	-0.30	Temperate	9.30	763	15.88	11.41	116.40	681.2	1, 4	2011	0.25	62	NA ^g
O	USA	IDENT	Cloquet	46.68	-92.52	Temperate	2.60	717	16.48	6.99	545.90	1113.50	1, 2, 6	2010	6.25	123	118

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3 ^a MAT = mean annual temperature in °C from Djukic et al. (2018); ^b MAP = mean annual precipitation in mm from Djukic et al. (2018); ^c MPT = Mean
4 Period Temperature in °C with Period referring to the experimental incubation period; ^d CPP = Cumulative Period Precipitation in mm; 3M = 3
5 months of incubation; 12M = 12 months of incubation; ^e SR gradient = species richness gradient; ^f No. of bags = number of installed tea bags, ^g NA
6 = 12M not included in the experiment.

Table 2. ANOVA output of linear mixed models testing the effect of tree species composition, planting, and climate related variables on mass loss of green and rooibos tea after 3 months and 12 months, respectively. The interaction between age and focal tree identity (angiosperm or gymnosperm) and between mean period temperature (MPT) and cumulative period precipitation (CPP) were included in all models. Site and plot were included as random factors with plot nested in site. Models were executed on centered variables.

	Mass loss of green tea after 3 months							Mass loss of rooibos tea after 3 months					
Fixed effect	Sum Sq	Mean Sq	Num DF	Den DF	F value	P		Sum Sq	Mean Sq	Num DF	Den DF	F value	P
Incubation period length	0.024	0.024	1	503.29	7.83	0.005		0.003	0.003	1	487.30	0.97	0.32
Focal tree identity	0.002	0.002	1	174.13	0.93	0.33		0.019	0.019	1	78.21	4.87	0.03
Age	0.000	0.000	1	9.30	0.02	0.88		0.008	0.008	1	9.30	2.05	0.18
Tree composition	0.000	0.000	2	187.72	0.10	0.89		0.004	0.002	2	94.07	0.60	0.54
Species richness	0.001	0.001	1	84.57	0.43	0.50		0.001	0.001	1	45.32	0.27	0.59
Planting density	0.000	0.000	1	8.93	0.10	0.75		0.001	0.001	1	9.04	0.25	0.62
MPT	0.000	0.000	1	9.12	0.14	0.71		0.001	0.001	1	9.23	0.45	0.51
CPP	0.000	0.000	1	9.14	0.20	0.65		0.001	0.001	1	9.20	0.30	0.59
Focal tree identity: Age	0.020	0.020	1	205.25	6.50	0.01		0.079	0.079	1	126.59	19.63	<0.001
MPT: CPP	0.001	0.001	1	9.01	0.46	0.51		0.002	0.002	1	9.10	0.66	0.43
	Mass loss green tea after 12 months							Mass loss rooibos tea after 12 months					
Fixed effect	Sum Sq	Mean Sq	Num DF	Den DF	F value	P		Sum Sq	Mean Sq	Num DF	Den DF	F value	P
Incubation period	0.037	0.037	1	213.86	7.12	0.008		0.170	0.170	1	139.54	26.85	<0.001
Focal tree identity	0.000	0.000	1	155.04	0.05	0.81		0.008	0.008	1	130.22	1.42	0.23
Age	0.003	0.003	1	6.50	0.66	0.44		0.024	0.024	1	6.05	3.81	0.09
Tree composition	0.011	0.005	2	168.62	1.06	0.34		0.045	0.022	2	146.08	3.61	0.02
Species richness	0.017	0.017	1	40.57	3.27	0.07		0.002	0.002	1	108.45	0.34	0.55
Planting density	0.001	0.001	1	6.26	0.29	0.60		0.009	0.009	1	5.78	1.54	0.26
MPT	0.061	0.061	1	6.08	11.71	0.01		0.002	0.002	1	5.68	0.31	0.59
CPP	0.013	0.013	1	6.31	2.56	0.15		0.000	0.000	1	6.02	0.14	0.71
Focal tree identity: Age	0.000	0.000	1	163.90	0.02	0.87		0.008	0.008	1	134.38	1.38	0.24
MPT: CPP	0.007	0.007	1	6.83	1.48	0.26		0.006	0.006	1	6.76	1.00	0.35

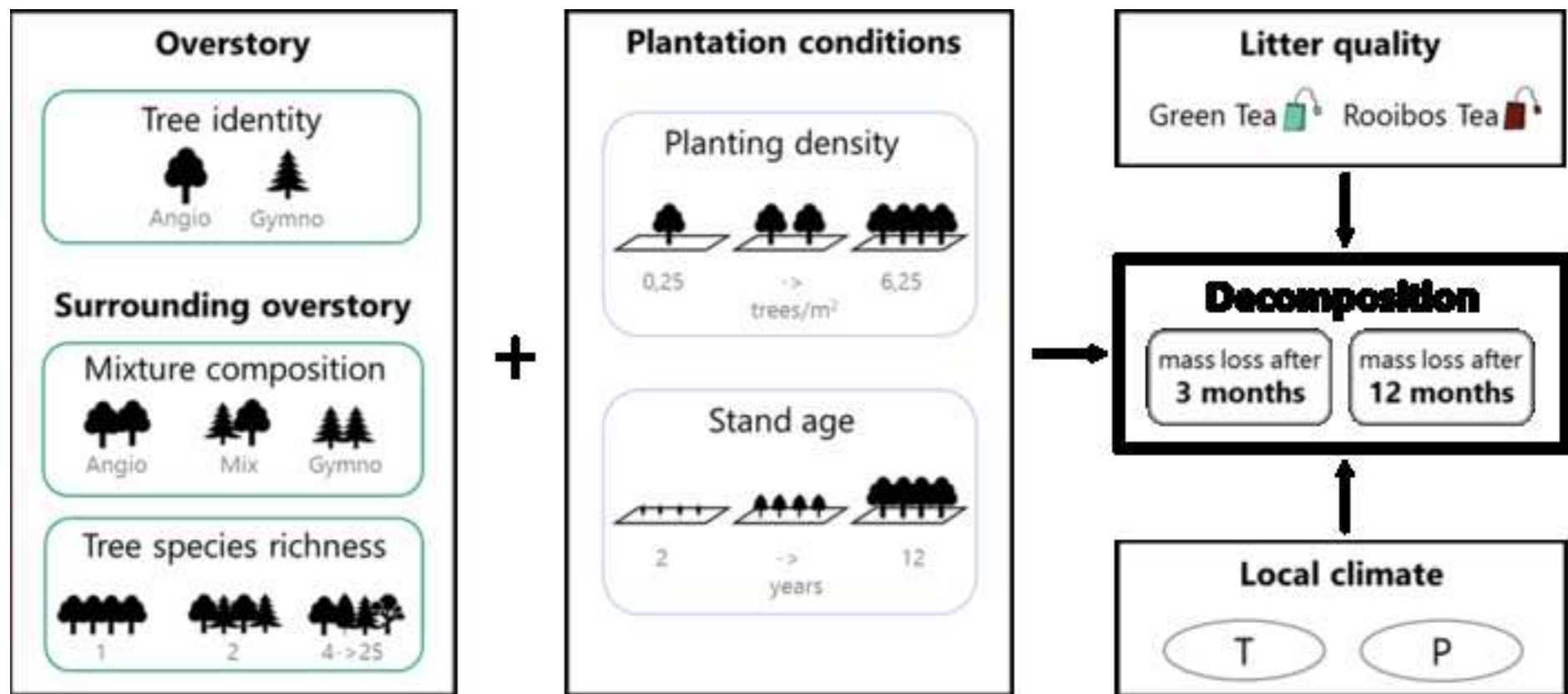


Figure2

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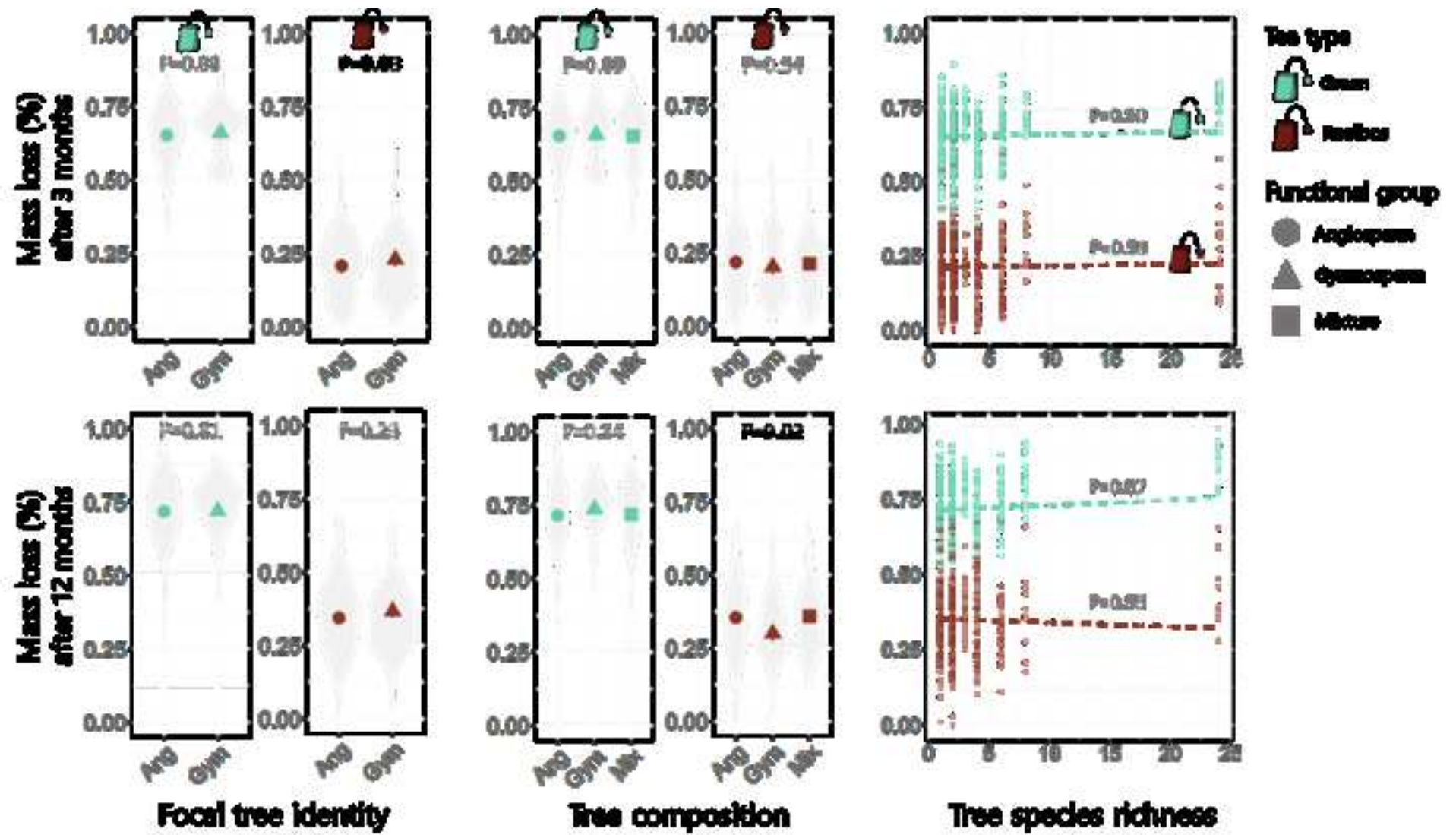
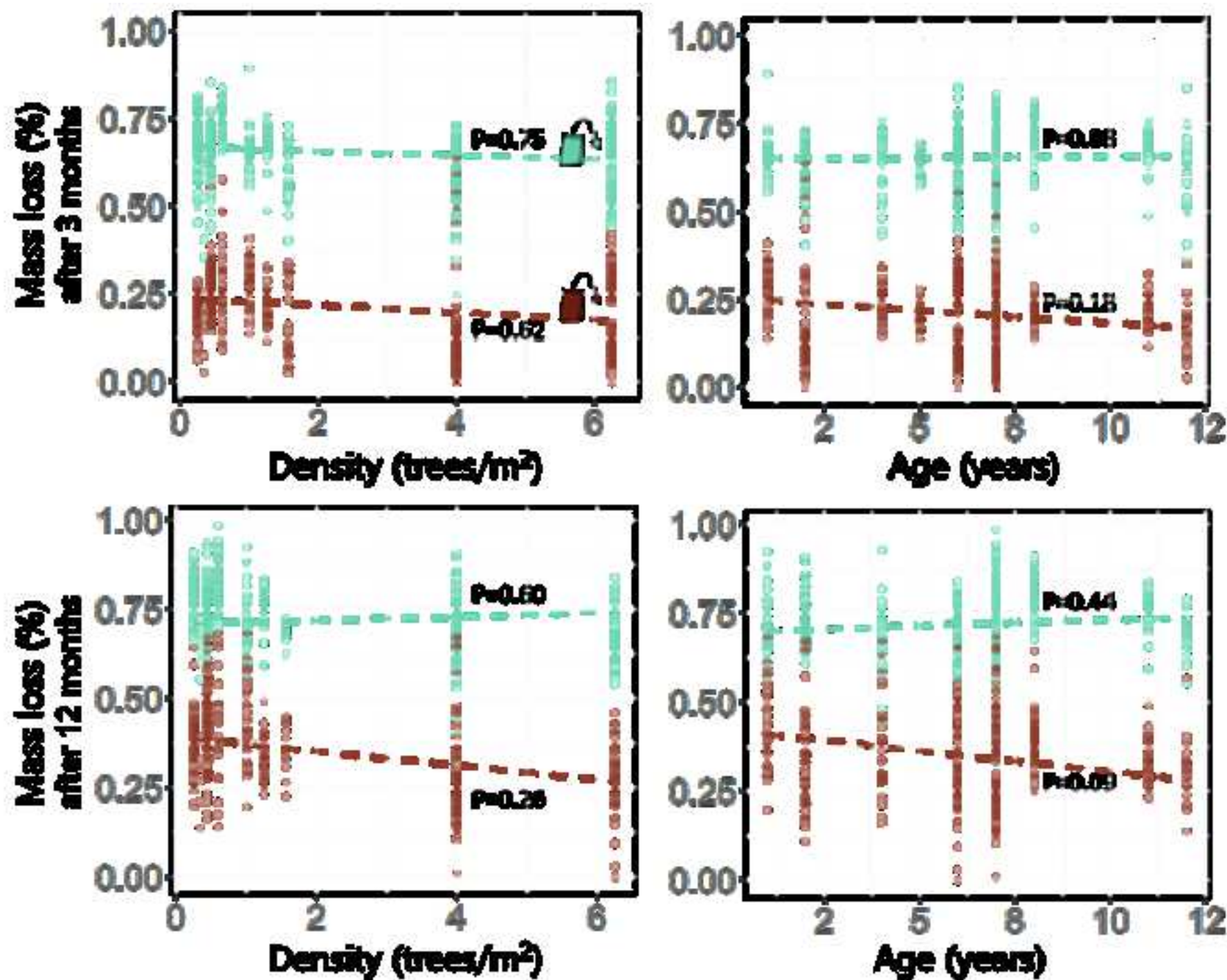
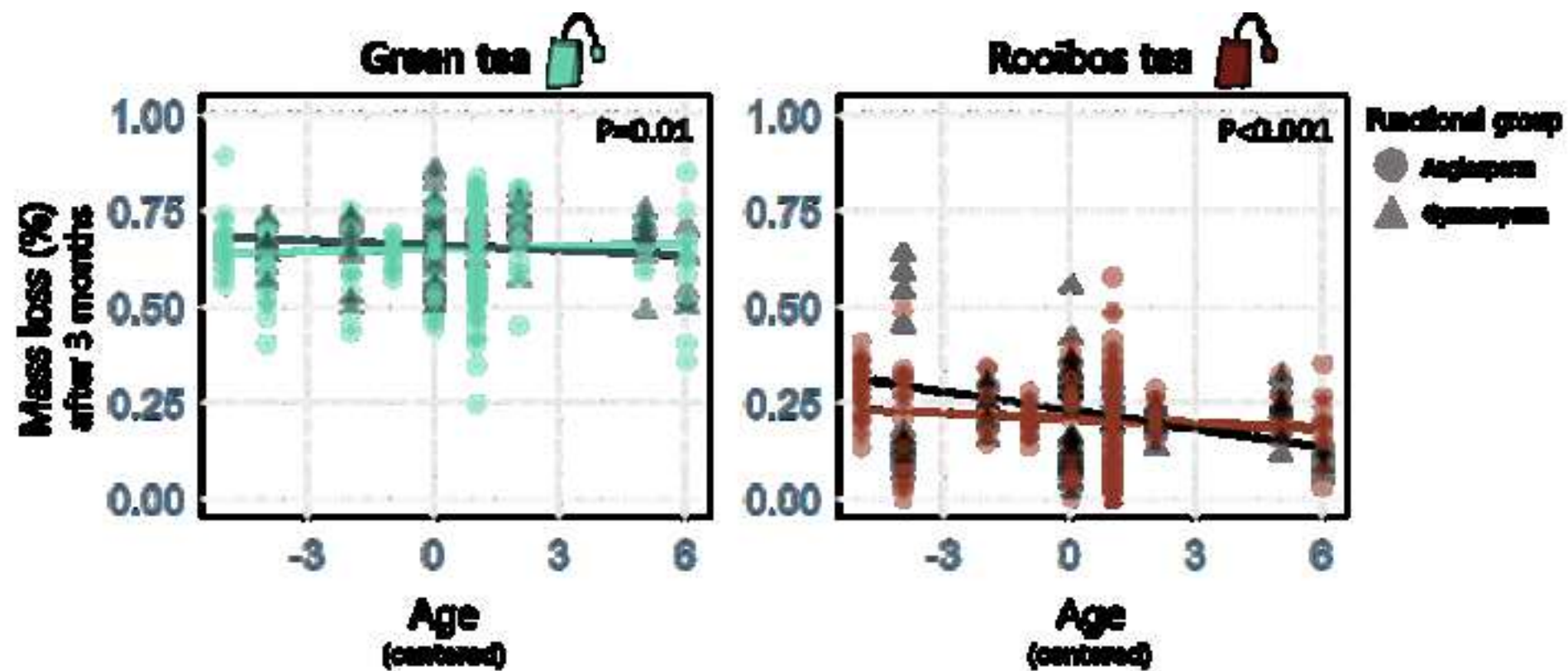
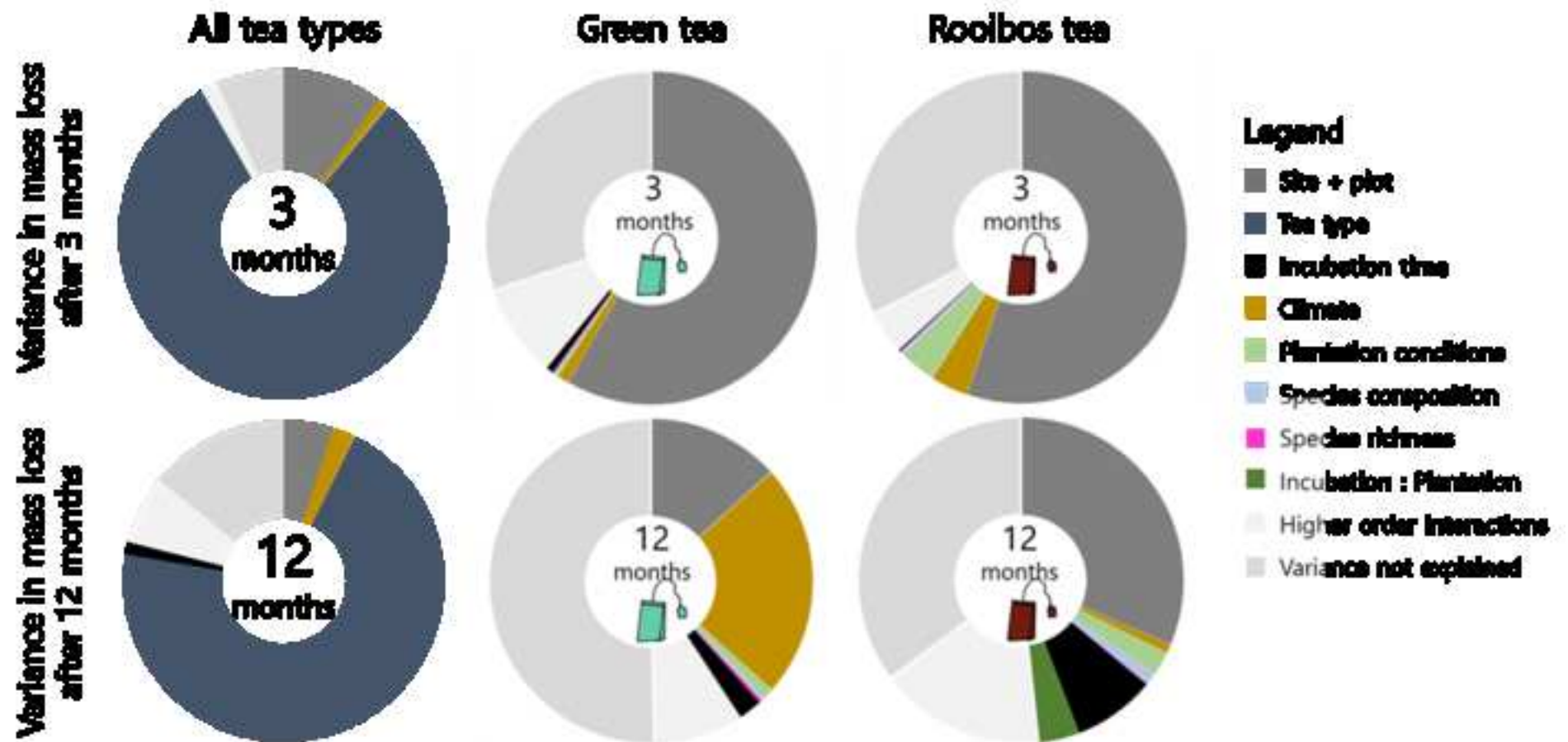


Figure3

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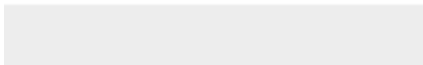
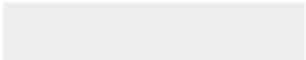








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Declaration of interests

☐The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

☒The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

All authors reports equipment, drugs, or supplies was provided by UNILEVER Lipton Tea Bags. Juan Zuo reports financial support was provided by Belspo. Nico Eisenhauer and Olga Ferlian reports financial support was provided by German Research Foundation FZT 118, 202548816. Helge Bruelheide, Bo Yang, Stefan Trogisch, Heike Feldhaar, Steffen Seitz reports financial support was provided by German Research Foundation FZT 118, 202548816. Martin Weih reports financial support was provided by Swedisch Energy Agency (36654-1, 36654-2, 36654-3 project DiPTiCC 16-CE32-0003. Michael Scherer-Lorenzen reports administrative support was provided by Federal Forestry Office Thüringer Wald. /

Authors' contributions

ID designed the experiment; AP, AS, BY, BM, CB, CM, CN, CAN, HB, HF, HJ, JR, KV, LM, LW, MSL, MW, NB, NE, OF, PBR, QP, StS, ST, SS collected data; ED, ID, JZ assembled the data from the sites; ED, JZ, KVM analyzed the data; ED and JZ wrote the first draft of the manuscript; BM, KV, ID and KVM provided thorough feedback on the manuscript, all coauthors commented on the advanced manuscript.