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Forest decline differentially affects trophic guilds of canopy-dwelling beetles

Aurélien Sallé¹ · Guilhem Parmain² · Benoît Nusillard² · Xavier Pineau¹ · Ravène Brousse¹ ·
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Abstract

• **Key message** Decline can affect the structure, resources, and microclimates of the forest canopy, and potentially have cascading effects on canopy-dwelling species. Our survey shows that an oak decline can promote saproxylic beetles, especially xylophagous ones, and generalist phyllophagous weevils. However, it negatively affects specialist leaf-eating species and has no effect on seed-eating weevils.

• **Context** Decline in a context of climate change is expected to induce considerable changes in forest structure, potentially affecting habitat opportunities and trophic resources for numerous species. Nonetheless, the consequences of decline in forest biodiversity have rarely been studied.

• **Aims** We aimed to characterize the impact of oak decline on different guilds of canopy-dwelling beetles.

• **Methods** Beetles were sampled for three consecutive years in oak stands exhibiting different levels of decline. Several guilds were considered: (i) Buprestidae, (ii) other saproxylic beetles split into wood-boring species and non-wood-boring species, (iii) seed-eating weevils, and (iv) specialist and generalist leaf-eating weevils.

• **Results** Overall, decline had positive effects on the abundance and biomass of beetles, though contrasting variations were observed at the species or guild levels. Wood-boring species, especially the main oak-associated buprestids, and other saproxylic species benefitted from decline conditions. However, at odds with the insect performance hypothesis, decline had a positive effect on generalist leaf-eating species, a negative effect on specialist leaf-eating species, and a null effect on seed-eating species.

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This article is part of the topical collection on *Entomological issues during forest diebacks*

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• **Conclusion** The increase in species richness for saproxylic and leaf-eating beetle communities suggests that decline might promote forest biodiversity. Our results call for further studies to thoroughly assess the functional outcomes of forest decline and to suggest management strategies for conservation biologists.

Keywords *Agrilus* · *Quercus* · Saproxylic beetles · Phyllophagous beetles · Seminiphagous beetles

1 Introduction

Global change can dramatically affect the organization and functioning of forest ecosystems by promoting the introduction and establishment of invasive species (Liebhold et al. 2017), by intensifying land use at the landscape level (Seibold et al. 2019), and through the direct and indirect effects of climate change on forest health (Seidl et al. 2017). Climate change already challenges the ability of European forests to adapt (Allen et al. 2010; Carnicer et al. 2011), and unprecedented forest declines are expected in response to the predicted increase in frequency and severity of droughts and heat waves (Allen et al. 2010; IPCC 2013).

Forest decline generally consists in a progressive loss of vigor of the trees, over several years, in response to multiple, successive, or concomitant driving factors (Manion 1981). These factors include (i) predisposing factors such as site conditions that constantly affect the stands, (ii) inciting factors such as defoliation or droughts that trigger declines, and (iii) contributing factors such as secondary pests and pathogens, which aggravate the deleterious effects of inciting factors, ultimately killing trees (Sinclair 1967; Manion 1981; Thomas et al. 2002; Sallé et al. 2014). The gradual loss of tree vigor progressively affects all forest compartments but the canopy is certainly the first to exhibit conspicuous modifications as decline progresses. The crown of a declining tree is characterized by an accumulation of dead branches, cavities, and fruiting bodies of saprotrophic or pathogenic fungi (Houston 1981; Ishii et al. 2004). Therefore, a forest decline generates novel structures and favors the accumulation of uncommon ones for healthy trees, and consequently tends to increase the structural complexity of the canopy at stand, tree, and branch scales (Ishii et al. 2004). Crowns of declining trees also exhibit reduced foliage density, which in turn can considerably alter microclimates within and beneath the canopy (Houston 1981; Ishii et al. 2004). Such profound structural modifications affect habitat opportunities and trophic resources, with likely marked cascading effects on canopy-dwelling communities.

The tree canopy and the soil are the two key compartments supporting forest biodiversity and their contribution is tremendous (Stork and Grimbacher 2006). Compared to tropical forests, temperate forests have less vertical stratification and a more marked seasonality with leaf fall, so temperate forest canopies probably shelter a lower proportion of specific taxa

(Ulyshen 2011). Canopy functional biodiversity in temperate forests has therefore received relatively little attention to date (Ulyshen 2011). However, the studies conducted in temperate forests (e.g., Bouget et al. 2011; Vodka and Cizek 2013; Plewa et al. 2017) have shown a clear vertical stratification of insect assemblages, just as in tropical forests, with 20–40% of all forest insect species strictly associated with canopies (Bouget et al. 2011). In addition to these specialist species, many Arthropods also rely on the canopy for a part of their life cycle, for maturation feeding and mating on foliage, such as *Agrilus* spp. for instance (Ulyshen 2011; Sallé et al. 2014). However, canopies are still relatively unknown biotic frontiers. These crown ecosystems harbor poorly understood, rarely described (both in terms of composition and abundance) insect communities (Bouget et al. 2011). They potentially shelter an underestimated pool of not only rare or patrimonial species (Plewa et al. 2017) but also native and invasive pests.

Canopy modifications in response to decline may change resource availability and microclimates and may create novel colonization opportunities, thus modulating in different ways the community dynamics of canopy-dwelling insects, depending on their functional guilds. Changes in foliage quality during plant stress may influence the performance of leaf feeders, but the magnitude and orientation of the herbivore response likely depend on both stress intensity and the feeding strategy of the herbivore (Larsson 1989; Herms and Mattson 1992). In addition, the decrease in the number of living branches in the canopy of declining trees may also negatively affect leaf-, seed-, and flower-feeding species (Martel and Mauffette 1997). A survey of Lepidopteran communities in declining maple stands indicated that exposed caterpillars became more abundant while the density of semi-concealed or endophagous species decreased (Martel and Mauffette 1997). This suggests that phyllophagous or seminiphagous insects with an intimate relationship with their host-tree, like specialist species with an endophytic larval development, may be negatively affected by the decrease in foliage quantity or quality and/or the change in microclimate, while these modifications might promote generalist species (Martel and Mauffette, 1997). Conversely, saproxylic beetles are likely to show a marked positive response to forest decline, both in terms of abundance and species richness. Saproxylic beetles form a functional guild associated with dead and decaying wood, related microhabitats, and other saproxylic taxa (Stokland et al. 2012). This guild also includes xylophagous species developing on weakened

trees and acting as secondary pests, like the buprestids (Coleoptera: Buprestidae), which are contributing agents during declines (Sallé et al., 2014; Tiberi et al. 2016). The weakened trees and the accumulation of dead wood and related microhabitats typical of declining stands should promote the abundance and diversity of this functional guild.

Our investigation focused on oak forests, which have at least two relevant characteristics for our study purpose. First, oak forests have regularly undergone periods of decline throughout Europe during the last centuries (e.g., Delatour, 1983; Oszako, 2000; Thomas et al., 2002; Sonesson and Drobyshev, 2010; Denman et al., 2014). Moreover, the frequency and intensity of declines have recently increased, and extended canopy modifications have already been documented in Mediterranean oak forests (Allen et al. 2010; Carnicer et al. 2011; Millar and Stephenson 2015). Second, oak forests host a species-rich insect fauna (Southwood 1961). We sampled the communities of leaf-dwelling weevils (Coleoptera: Curculionidae) and saproxylic beetles for three consecutive years in oak stands exhibiting different levels of tree decline.

Firstly, we hypothesized that saproxylic beetles, especially xylophagous species, would be favored by decline. Secondly, we expected to find contrasted responses to decline intensity for leaf-feeding weevils, dependent on their relationship with the host plant. More specifically, we hypothesized that weevils with endophytic larvae would be negatively affected by decline while species feeding on foliage only during the adult stage would be favored. Finally, we hypothesized that seminiphagous weevils would be negatively affected by the reduced amount of acorns in declining stands. Consequently, our objectives were (i) to describe the canopy-dwelling communities of buprestid beetles, other saproxylic species and weevils, and (ii) to evaluate how the local intensity of forest decline was modifying the diversity of these communities.

2 Material and methods

2.1 Study area

The study was conducted in the two adjacent state forests of Vierzon and Vouzeron, with a surface area of 5300 ha and 2200 ha, respectively, located in the center of France 200 km south of Paris (47° 26' 89" N, 02° 10' 74" E). The Vierzon forest is dominated by oaks (mostly *Quercus petraea* and *Quercus robur*) at 70% and conifers (mostly *Pinus sylvestris* and *Pinus nigra*) at 30% in both pure and mixed stands. The Vouzeron forest is dominated by conifers (mostly *Pinus sylvestris* and *Pinus nigra*) at 65% in pure stands or mixed with *Q. robur* and *Q. petraea*. The oaks, especially *Q. robur* in the Vierzon forest, have suffered from regular declines (documented in 1920, 1940, and 1982 (Douzon 2006)). The last severe oak decline occurred between 2000 and 2010, which

was followed by a sanitation cutting of 100,000 m³ of oak over 1000 ha (Douzon, 2006). Several factors were implicated in these successive periods of decline. The prominent predisposing factor was edaphic. In most areas, the water table is shallow and variable, and therefore inadequate for the development of *Q. robur*, which was, however, extensively planted in this forest (Douzon 2006; Marçais and Desprez-Loustau 2014). The prominent inciting factors were severe droughts and defoliation caused by powdery mildew (Douzon, 2006; Marçais and Desprez-Loustau 2014). Finally, the most frequently observed contributing biotic agents were *Agrilus biguttatus* Fabricius and *Armillaria mellea* (Vahl ex Fr.) P. Kumm. (Douzon 2006).

2.2 Site selection and characterization

Overall, 14 stands dominated by mature oaks were monitored during our study in the two forests (Table 1). In 2016, 13 plots were set up in 11 stands (Table 1). Plots were homogeneous areas (approx. 2000 m²) within stands in terms of tree composition and dendrometric parameters. Most plots were located in different stands (Table 1). They were located at a minimum of 100 m from the others, but in most cases, several kilometers from each other. In 2017 and 2018, 12 plots located in 11 stands were selected in the two forests (Table 1). Some of the original plots were changed in 2018 because the stands had been either cleared or harvested selectively (Table 1). We selected one tree at the center of each plot, on which we suspended one trap for beetle collection (see below “Beetle sampling”).

The level of decline was evaluated yearly, during winter, at two embedded spatial scales. (i) At the tree scale: individual decline status was assessed for all bearing-trap trees. (ii) At the plot scale: decline status was assessed for trees surrounding the bearing-trap tree, which encompassed only the five closest oak trees in 2016 and 2017, and all the oaks located within a radius of 20 m around the bearing-trap tree in 2018. Decline level was evaluated following the protocol designed by the French Forest Health Service (Département de la Santé des Forêts, DSF) (Nageleisen 2005). In brief, crown transparency, the proportion of branches without leaves, the proportion of dead branches and leaf distribution in the canopy were evaluated. Based on these criteria, each tree was given a decline index ranging from 0 (no decline) to 5 (dead tree) (Table 2). Trees with an index value equal to or below two were considered healthy. Trees with an index value equal to or above three were considered in decline. For the plot scale, the proportion of declining trees (with a decline index equal to or above three) was calculated, following a routine DSF procedure (Table 2).

Table 1 characteristics of the monitored oak stands

Stand ID	Latitude	Longitude	Mean \pm SE stand height ¹ (m)	Monitoring year	Composition	Mean \pm SE oak DBH ² (cm)	Oak density ³ (n/ha)	Tree density ³ (n/ha)	Total basal area ³ (m ² /ha)
19	47.2579	2.1812	15 \pm 1	2018	Oak with a beech understorey	51.8 \pm 2.4	151	151	33
24	47.2681	2.1733	20 \pm 2	2016–2017	Pure oak	65.0 \pm 3.5	88	88	30
35–1	47.2980	2.1752	17 \pm 1	2016–2017–2018	Oak with scattered beech and Scots pine	54.2 \pm 3.4	111	207	47
35–2	47.2974	2.1743	17 \pm 1	2016–2017–2018	Oak with scattered beech and Scots pine	51.7 \pm 3.5	119	167	30
55	47.2567	2.1616	21 \pm 2	2016–2017–2018	Mixed stand of oak and beech	69.0 \pm 8.3	16	111	33
70	47.2886	2.1551	17 \pm 2	2016–2017–2018	Oak with scattered beech and Scots pine	47.1 \pm 5.0	64	119	21
71	47.2877	2.1536	15 \pm 1	2018	Oaks with scattered beech and Scots pine	39.7 \pm 3.7	119	223	26
81–1	47.2666	2.1967	23 \pm 2	2016	Oak with a hornbeam understorey	37.4 \pm 1.3	175	175	20
81–2	47.2672	2.1980	23 \pm 2	2016–2017–2018	Oak with a hornbeam understorey	37.4 \pm 1.3	175	175	20
133	47.2918	2.1434	18 \pm 2	2016–2017	Oak with scattered beech and Scots pine	75.0 \pm 2.6	56	80	28
161	47.2626	2.1243	22 \pm 1	2016–2017	Pure oak	64.6 \pm 5.6	80	88	31
179	47.2654	2.1189	19 \pm 2	2016–2017–2018	Pure oak	59.3 \pm 3.6	151	151	45
236	47.2675	2.0807	15 \pm 3	2016–2017–2018	Mixed stand of oak and beech	40.5 \pm 4.8	167	422	51
249	47.2656	2.0674	16 \pm 1	2018	Oak with scattered beech	48.2 \pm 1.3	199	223	38
290	47.26119	2.0328	16 \pm 1	2016–2017–2018	Oak with scattered wild service trees	38.6 \pm 1.7	183	207	24
299	47.2569	2.0257	21 \pm 3	2016–2017–2018	Pure oak	47.8 \pm 8.1	64	64	14

¹ Mean stand height was measured on three trees.

² DBH was measured on all trees within a radius of 20 m around the tree carrying traps.

³ Densities and total basal area were assessed through the inventory and measurement of all trees within a radius of 20 m around the tree carrying traps.

Table 2 Decline index of trees carrying traps and percentage of surrounding declining trees in the monitored oak plots for the three survey years

Stand ID	Decline index of trees with traps			Percentage of surrounding declining trees		
	2016	2017	2018	2016	2017	2018
19	NA	NA	3	NA	NA	89
24	4	3	NA	17	33	NA
35-1	2	1	1	17	0	19
35-2	0	1	1	17	33	14
55	3	3	3	83	67	100
70	3	3	3	83	71	100
71	NA	NA	3	NA	NA	87
81-1	1	1	1	17	17	14
81-2	2	NA	NA	0	NA	NA
133	3.5	2	NA	33	57	NA
161	2	2	NA	33	29	NA
179	2	4	2	17	57	61
236	3	3	3	33	33	45
249	NA	NA	3.5	NA	NA	60
290	2	2	2	33	14	43
299	3.5	2	2	50	57	50

2.3 Beetle sampling

2.3.1 Optimization of the sampling protocol

We used multi-funnel traps (Lindgren traps, Chemtica Internacional, San Jose, Costa Rica), each with 12 fluon-coated funnels, to collect the insects. To optimize our protocol for sampling canopy-dwelling beetles, we tested two trap heights and two trap colors. In 2016, we assessed whether trap height significantly affected the composition or relative abundance of the captured species. To do so, we suspended two green traps at different heights in the same tree, one approximately 15 m from the ground and the other 10 m from the ground. Thirteen pairs of traps were considered.

In 2017 and 2018, we also compared the trapping efficiency of green and purple multi-funnel traps. Green multi-funnel traps have successfully captured a wide array of buprestid species in North America and Europe (Petrice and Haack 2015; Rassati et al. 2019), but Brown et al. (2017) showed that some European *Agriilus* species might be more attracted to purple than to green. In ten trees, one green and one purple trap were both suspended roughly 15 m from the ground.

2.3.2 Routine sampling protocol

For the sake of consistency, and because it was the best sampling design (see “Results”), only green multi-funnel traps, suspended at 15 m from the ground (i.e., among the lower branches of the canopy) were considered to assess the effect of decline on canopy-dwelling beetles. One trap was set up

within each monitored plot. The collectors were filled with a 50% (v/v) monopropylene glycol solution diluted with water and a drop of detergent. No lure was added to the traps. The traps were emptied every month and the captured species were recorded. In 2016, the traps were installed in June and collection continued until September. In 2017, the traps were installed in April and collection continued until October. In 2018, the traps were installed in April and collection continued until September.

2.4 Beetle identification and ecological trait assignment

Three beetle groups were considered for analysis: (i) oak-associated buprestid beetles, i.e., xylophagous species specifically attracted by green Lindgren traps; (ii) other saproxylic beetles, excluding “tourist” species associated to conifer tree species, split into two feeding guilds, i.e., the xylophagous species guild (including wood-boring sensu stricto and saproxylophagous species) and the non-xylophagous species guild (including saprophagous, zoophagous, and mycetophagous species); and (iii) oak-associated phytophagous weevils, which were split into two feeding guilds, i.e., the leaf-eating (phyllophagous, i.e., folivore) species guild and the seed- and fruit-eating (seminiphagous, i.e., acorn borers) species guild (Sallé et al. 2020). In the phyllophagous guild, we considered species whose adults feed on oak leaves (mainly leaf chewers) as generalist species, and species whose both larvae and adults feed on oak leaves (larvae dwelling in foliar tissues) as specialist species.

Certain saproxylic families were removed from the data set (Latridiidae, Leioididae, Malachiidae, Cantharidae, Corylophidae, Cryptophagidae, Ptiliidae, Staphylinidae) because they are often difficult to identify at the species level and due to the lack of specialists able to check species identifications. The French Frisbee database was used as the reference list of feeding guilds for saproxylic beetle species from the 39 recorded beetle families (Bouget et al. 2019). Most of the beetle specimens were identified by several of the authors (GP, BN, XP, RB, TFG, and RL). The remaining families were identified by specialists, as mentioned in the Acknowledgments. For each guild, we computed the number of species (richness), number of individuals (abundance), and beetle biomass caught at the trap level and cumulated over all the sampling season per year. Biomass, actually dry weight (in mg), was assessed through the following formula: $Biomass = 3.269 + L^{2.463}$, where “L” is the body length in millimeters (Ganihar 1997 in Seibold et al. 2019). The abundance of several dominant species of oak-associated buprestids and phytophagous weevils was also analyzed.

2.5 Data analysis

All analyses were performed in R, version 3.5.1 (R Core Team 2018). Trap color and trap height effects on catches of buprestids, other saproxylic beetles, and phytophagous weevils were assessed with Wilcoxon signed-ranked tests. To analyze the effect of the decline level at tree scale on cumulative number of detected species, we rarefied species richness to the same sample size (interpolated rarefaction sampling without replacement; `specaccum` function, `vegan` R-package). To rank the effect of decline level at tree or plot scale on variations in average univariate metrics (mean values per trap of guild richness, species abundance, guild abundance, guild biomass), we used the differences in the Akaike information criterion (AICc) scores to compare the fit between the generalized linear mixed models including separately each of the two explanatory variables and their fit with the null model. To assess the significance of the estimates of the best decline features for each response variable, the error structure of the generalized linear mixed-effects models was adjusted to better fit the data. To do so, `glmm` were fitted for the negative binomial family, the Gaussian family, the log-normal family (i.e., log-transformed response), and the Poisson family (functions `glmer.nb`, `glmer`, `lmer`, `lme4` R-package). To account for repeated measures and configuration of sampling design, plot and year were added as nested random effects on the intercept in mixed models. To rank the effect of decline level at tree or tree-group scale on variations in species composition (including singletons), we performed a Canonical Analysis of Principal coordinates (function `capscale`, `vegan` R-package, CAP, Anderson and Willis 2003). Based on Bray-Curtis distance matrices, we carried out inertia partitioning on all the

explanatory environmental variables, as collinearity among predictor variables is not a problem in CAP. We calculated total constrained inertia, the total inertia explained by each variable, the latter’s statistical significance (permutation tests—100 runs), and the relative individual contribution of each variable to constrained inertia.

We used the `IndVal` method to identify beetle species indicating tree decline level (healthy vs. declining) (Dufrêne and Legendre 1997, `indicspecies` R-package). This method calculates the association value (`IndVal` index) between the species and a group of sites, based on between-group variations in occurrence (fidelity) and abundance (specificity), and then tests the statistical significance of this relationship with a permutation test. *P*-values were corrected for multiple testing. Only species shown to be significant in the permutation test with an indicator value above 25%, occurring in more than 10% of the samples and with more than 10 individuals sampled were considered here.

3 Results

For all guilds, the number of individuals captured was higher in the upper traps (Fig. 1). For both guilds of leaf-dwelling species (i.e., Agrilinae and phytophagous weevils), green traps were markedly more attractive than purple traps, while no difference between traps was detected for the other saproxylic species (Fig. 2).

Overall, for the assessment of decline effects on canopy-dwelling beetles, the compiled data set of 27,627 individual specimens included 266 beetle species, 10,440 individuals and ten species of oak-associated buprestid beetles; 8280 individuals and 21 species of oak-associated phytophagous weevils (4 seminiphagous, 10 specialist phyllophagous, 5 generalist phyllophagous, and 2 flower-eating (anthophagous) species); 3008 individuals and 102 species of non-xylophagous saproxylic beetles; and 5899 individuals and 133 species of xylophagous saproxylic beetles (Table 6 in annex). This corresponds to 14,490 individuals and 223 beetle species found in the 23 traps hanging from non- or slightly-declining trees, and 13,137 individuals and 194 species found in the 14 traps hanging from declining trees. On the whole, cumulative species-richness estimates at a standardized sample size did not display any significant contrast between decline levels at the tree scale, either for the whole beetle community or for individual guilds (Fig. 7 in annex).

We detected many significant effects of decline level on guild metrics (mean abundance, biomass, and richness per trap) and on species mean abundances (Table 3). Most of these effects were positive except at the plot scale for (i) a negative effect of decline level on the mean abundance of xylophagous beetles (Table 3 and Fig. 3), and (ii) a negative effect of decline level on the mean abundance of two specialist

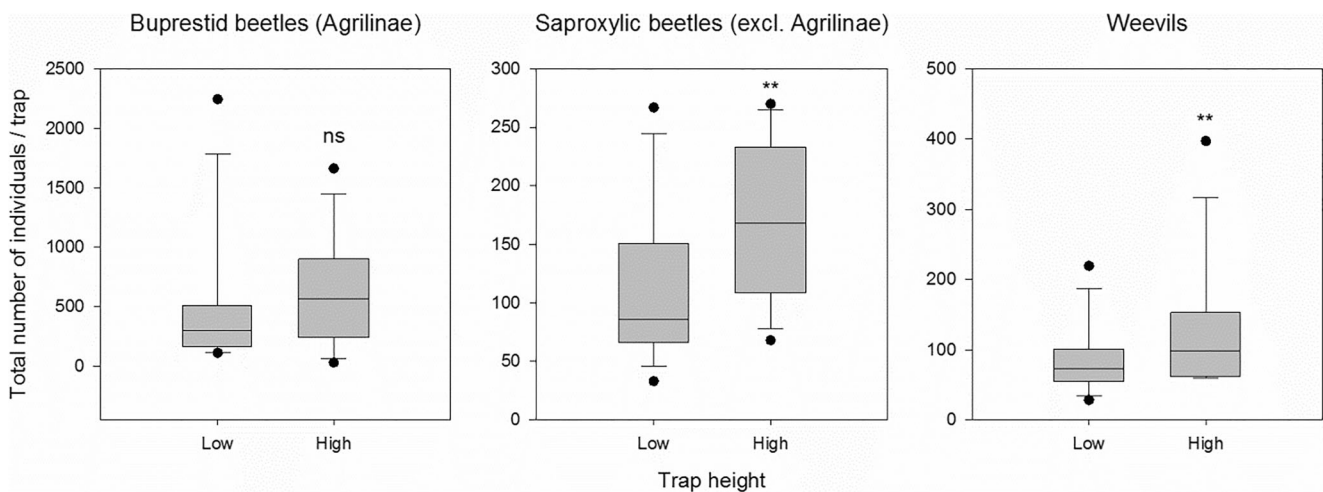


Fig. 1 Effect of trap height (10 m vs. 15 m above the ground) on the number of oak-associated Agrilinae (i.e., *Agrilus* sp., *Coraeus* sp., and *Meliboeus* sp.), other saproxylic beetles, and phytophagous weevils (i.e., phyllophagous and seminiphagous species) captured per trap. $P < 0.01$: **

phyllophagous species, i.e., *Archarius pyrrhoceras* Marsham and *Orchestes quercus* L. (Table 3). We measured significant positive effects of decline level at the tree scale on the species richness of xylophagous beetles (Fig. 3) and on the biomass and abundance of buprestids (Fig. 4); and at plot scale, on the biomass of non-xylophagous saproxylic beetles (Fig. 3), on species richness of buprestids (Fig. 4) and phyllophagous weevils (Fig. 5), and on the abundance of generalist phyllophagous weevils (Fig. 5). Five of the six buprestid species tested responded positively in abundance to decline intensity (at tree scale: *Agrilus angustulus* Illiger, *A. biguttatus*, *Agrilus laticornis* Illiger, *Agrilus obscuricollis* Kiesenwetter, *Agrilus sulcicollis* Lacordaire; and at the plot scale: *Coraeus undatus* Fabricius), as well as one of the two tested generalist phyllophagous weevil (*Phyllobius pyri* L.) (Table 3). Seminiphagous species were not significantly affected by

decline level at any scale, either at species or guild level (Table 3, Fig. 5). Specialist phyllophagous weevils responded to decline intensity at the species level but not at the guild level (Table 3, Fig. 5). When all the sampled species were pooled, we also observed significant positive effects of decline at the tree scale on the biomass and abundance of all beetles (Table 3, Fig. 6).

From CAP analyses, we estimated low but significant contributions of decline level to variations in the community composition of most of the beetle groups, i.e., phytophagous weevils, buprestids, and xylophagous saproxylic, though not for the non-xylophagous saproxylic guild (Table 4). Significant effects on community composition were mainly related to decline level at the plot scale, except for xylophagous saproxylic beetles, which were affected at the tree scale. A larger portion of inertia was explained by sampling year.

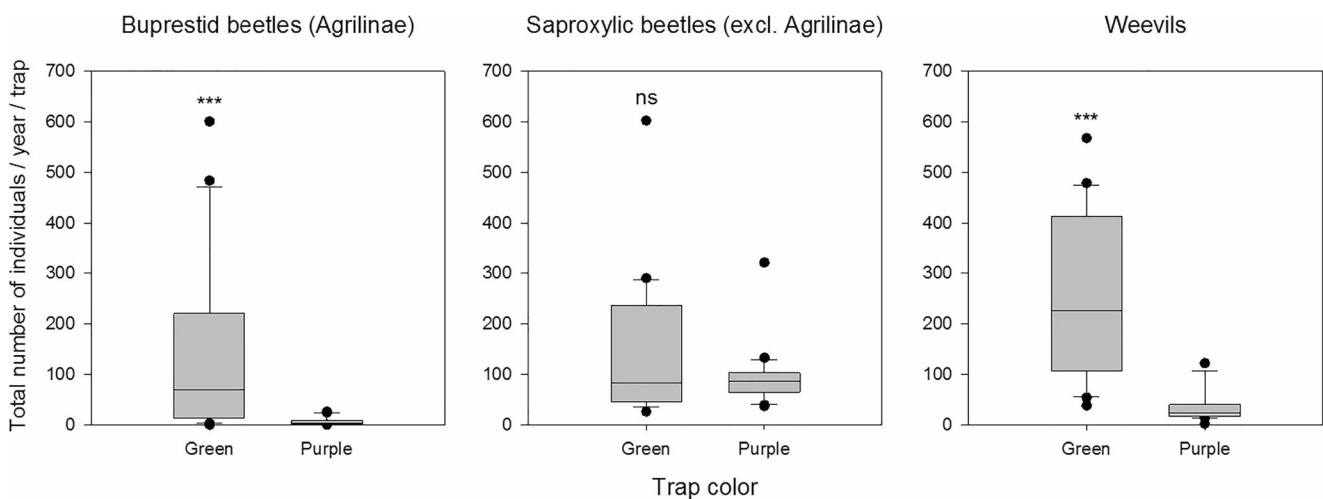


Fig. 2 Effect of trap color (green vs. purple) on the number of oak-associated Agrilinae (i.e., *Agrilus* sp., *Coraeus* sp., and *Meliboeus* sp.), other saproxylic beetles, and phytophagous weevils (i.e., phyllophagous and seminiphagous species) captured per trap. $P < 0.01$: **, $P < 0.001$: ***

Table 3 Effect of decline on biomass, abundance, and species richness for the different groups and guilds of beetles, and on abundance for the main species collected in the canopy

	Variable (mean value per trap)	Best ecological model	Delta[AICc] ^e	Effect of decline		
				Estimate	SE	
Oak-associated buprestid beetles	Biomass ^c	Tree	-6	1.26**	0.44	
	Abundance ^a	Tree	-5	1.38**	0.48	
	Species richness ^d	Plot	-2	0.14*	0.06	
	<i>Agrilus angustulus</i> ^{c,f}	Tree	-4	1.49*	0.59	
	<i>Agrilus biguttatus</i> ^a	Tree	-2	2.03*	0.93	
	<i>Agrilus hastulifer</i> ^a	Tree	0	1.84	1.05	
	<i>Agrilus laticornis</i> ^a	Tree	-2	0.98*	0.46	
	<i>Agrilus obscuricollis</i> ^a	Tree	-2	1.87*	0.93	
	<i>Agrilus sulcicollis</i> ^a	Tree	-9	1.88***	0.51	
Saproxylic beetles (excl. buprestids)	<i>Coraeus undatus</i> ^a	Plot	-7	0.94***	0.28	
	Biomass ^c	Tree	0	0.71	0.46	
	Xylophagous	Abundance ^c	Plot	-2	-0.39*	0.16
		Species richness ^a	Tree	-1	0.29*	0.14
	Non-xylophagous	Biomass ^c	Plot	-3	0.31**	0.10
		Abundance ^a	Plot	+1	0.12	0.10
Oak-associated weevils	Species richness ^b	Plot	0	3.41	1.84	
	All phytophagous	Biomass ^c	Plot	0	0.19	0.12
		Abundance ^c	Tree	+2	0.10	0.10
		Species richness ^b	Plot	0	0.64	0.37
	All phyllophagous	Biomass ^c	Tree	-1	0.23	0.13
		Abundance ^c	Tree	+2	-0.10	0.26
		Species richness ^c	Plot	-3	0.10*	0.04
	Generalist phyllophagous	Biomass ^c	Plot	-1	0.59	0.32
		Abundance ^c	Plot	-5	0.77**	0.23
		Species richness ^c	Tree	+5	0.04	0.12
	Specialist phyllophagous	<i>Phyllobius pyri</i> ^a	Plot	-6	1.69**	0.56
		<i>Polydrusus cervinus</i> ^a	Plot	+2	0.27	0.28
		Biomass ^c	Tree	+2	-0.10	0.04
		Abundance ^c	Tree	+1	-0.12	0.10
		Species richness ^d	Plot	0	0.14	0.09
		<i>Archarius pyrrhoceras</i> ^a	Plot	-3	-0.79*	0.37
	Seminiphagous	<i>Orchestes quercus</i> ^a	Plot	-1	-0.27*	0.13
		Biomass ^c	Tree	+1	-0.38	0.57
Abundance ^a		Plot	+2	-0.14	0.21	
Species richness ^d		Plot	+2	0.06	0.10	
<i>Curculio glandium</i> ^a		Plot	+2	-0.15	0.22	
All beetles	Biomass ^c	Tree	-10	0.80***	0.22	
	Abundance ^b	Tree	-3	298.2*	121.10	
	Species richness ^b	Tree	+1	3.70	2.33	

biomass = dry weight (in mg)

^a Generalized linear mixed-effects models fitted for the negative binomial family.

^b The Gaussian family.

^c The log-normal family (i.e., log-transformed response)

^d The Poisson family with year as a random effect.

^e $\Delta AICc = AICc$ (best ecological model) - $AICc$ (null model).

^f The variable considered for species is abundance.

* $P < 0.05$

** $P < 0.01$

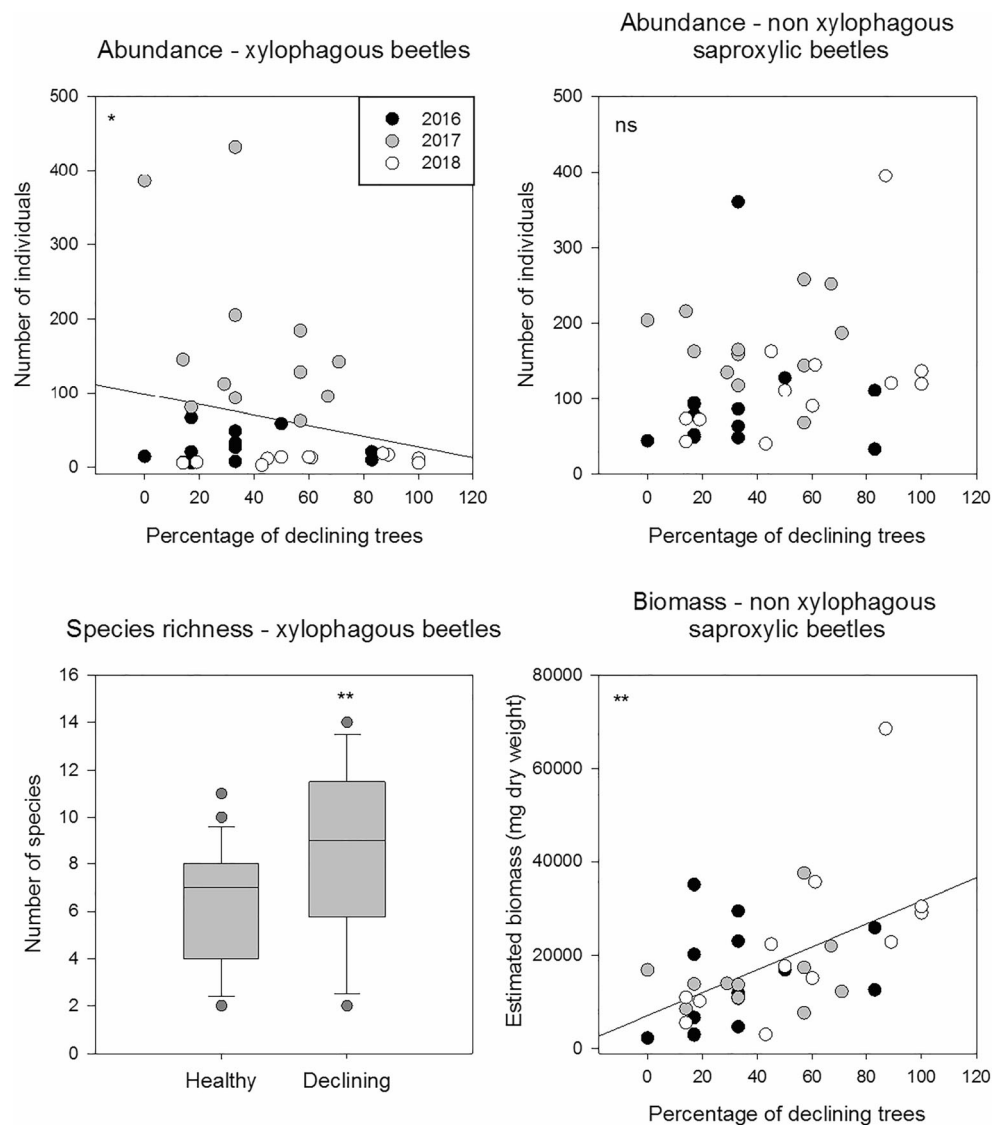
*** $P < 0.001$

The IndVal analysis detected 15 characteristic species in declining trees, and only one in healthy trees (Table 5). The group of species associated with declining trees consisted primarily of xylophagous species, including four species of *Agrilus*, five other xylophagous beetle species, and two saproxylophagous species.

4 Discussion

Oak decline affected the communities of the canopy-dwelling beetles considered in our study differently depending on their feeding guild and/or host specialization. As predicted, the abundance, biomass, and species richness of oak-associated

Fig. 3 Abundance of xylophagous and non-xylophagous saproxylic beetles, species richness of xylophagous beetles, and biomass of non-xylophagous saproxylic beetles depending on decline level at the tree scale (healthy vs. declining) or plot scale (percentage of declining trees). See Table 3 for statistical results; $P < 0.05$: *; $P < 0.01$: **



buprestids increased with the decline severity. The abundance of most major Agrilinae species followed a similar pattern. Consequently, most species contributed to this overall increase. Agrilinae preferentially colonize weakened hosts (e.g., Moraal and Hilszczanski 2000; Jennings et al. 2014; Petrice and Haack 2014; Poole et al. 2019). Their abundance is positively influenced by the availability of fresh snags and coarse woody debris in the environment (Redilla and McCullough 2017) and damaged trees (Brück-Dyckhoff et al. 2019); these features typically occur in declining stands. Several of the species collected, namely *A. biguttatus*, *A. sulcicollis*, *A. angustulus*, and *C. undatus*, can act as major contributing agents during oak declines (Sallé et al. 2014). Consequently, they may also have exerted a positive feedback by further weakening their host trees, thus, contributing to the accumulation of favorable breeding material. Interestingly, three of the four, *A. biguttatus*, *A. sulcicollis*, and *A. angustulus*, were also good indicators of declining trees,

together with *Scolytus intricatus* Ratzeburg and *Gasterocercus depressirostris* Fabricius, other contributing agents of oak decline (Saintonge and Nageleisen 2001; Sallé et al. 2014). However, while the species richness of other xylophagous species also increased in declining stands, their abundance slightly decreased. This might reflect increased competition among xylophagous species in declining stands. However, the variation in abundance of these other xylophagous beetles should be considered with caution since it was mostly driven by variations in the abundance of *Anisandrus dispar* Fabricius. The abundance of this generalist ambrosia beetle might have been loosely connected to oak decline level. The increased availability of resources and habitats in declining stands, especially large woody debris, probably also participated in the observed increase in the biomass of non-xylophagous saproxylic beetles, since the size of saproxylic species tends to increase with the diameter of the available deadwood resources (Brin et al. 2011).

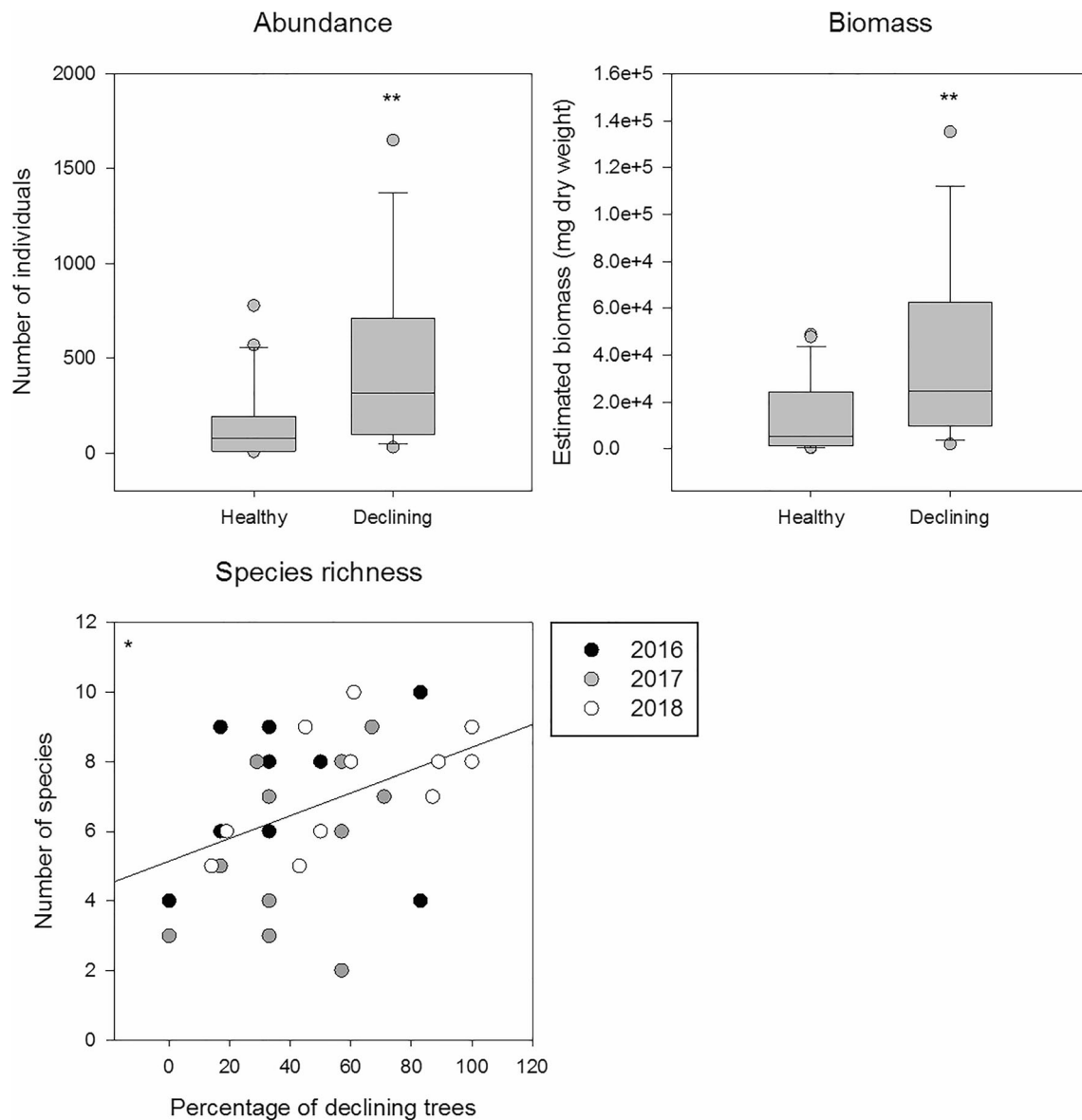
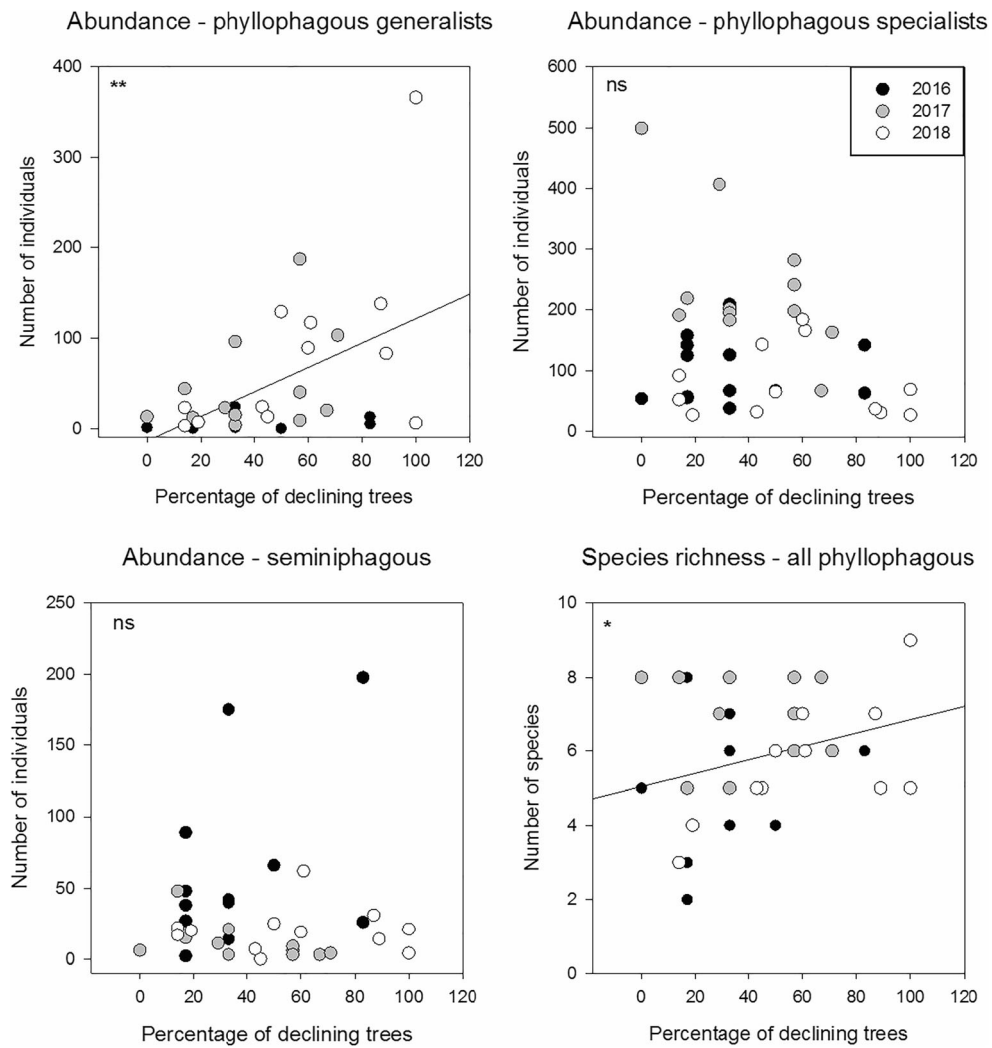


Fig. 4 Abundance, biomass, and species richness of oak-associated buprestid beetles depending on decline level at the tree scale (healthy vs. declining) or plot scale (percentage of declining trees). See Table 3 for statistical results; $P < 0.05$: *; $P < 0.01$: **

Feeding guilds of phytophagous weevils responded differently to decline severity. The abundance of generalist phyllophagous species, especially *P. pyri*, increased with decline intensity, while the abundance of the two main specialist phyllophagous species, i.e., *O. quercus* and *A. pyrrhoceras*, decreased. These variations support our hypotheses and are congruent with previous observations by Martel and Mauffette (1997) for Lepidopteran communities colonizing maple foliage. They are nonetheless inconsistent with predictions from the insect performance hypothesis by Larsson (1989) concerning the response of folivores with various feeding habits to tree stress. Several factors may have affected the abundance of phyllophagous weevils differently. Environmental constraints can have contrasted effects on both

biochemical and morphological leaf traits (Günthardt-Goerg et al. 2013; Hu et al. 2013). Likewise, the greater exposure of leaves in the opened canopies of declining oaks can alter their phytochemical profile, and may have increased their content in phenolic compounds (Yamasaki and Kikuzawa 2003; Lämke and Unsicker 2018). Such modifications may in turn have different impacts on phytophagous insects depending on their feeding guild and specialization (e.g., Gutbrodt et al. 2011; Forkner et al., 2004). In addition to modifying trophic resources or habitats, crown thinning during a decline can also directly impact larval development by altering the thermal buffering provided by the canopy (Martel and Mauffette 1997; Hardwick et al. 2015; De Frenne et al. 2019). Greater leaf exposure will affect leaf microclimate and may have

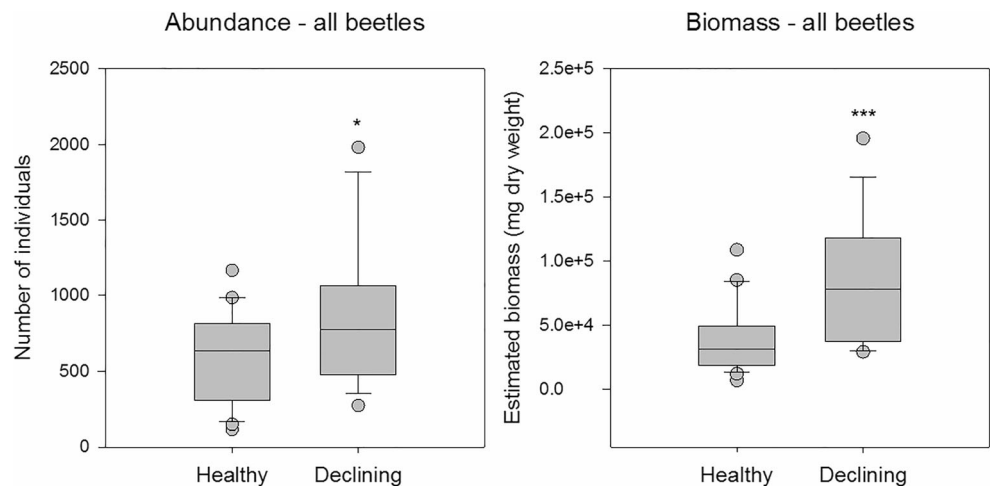
Fig. 5 Abundance of generalist phyllophagous, specialist phyllophagous and seminiphagous weevils, and species richness of all phyllophagous weevils, depending on decline level at the plot scale (percentage of declining trees). See Table 3 for statistical results; $P < 0.05$: *; $P < 0.01$: **



detrimental effects on endophytic larvae (e.g., Pincebourde et al. 2007), like those of *O. quercus* or *A. pyrrhoceras*. Canopy thinning can also affect forest soil microclimates (De Frenne et al. 2013), allowing free-living larvae like those

of *P. pyri* to find optimal microclimatic conditions more easily. Finally, greater leaf exposure may also lead to greater predation pressure on leaf-dwelling endophytic larvae (e.g., Tschanz et al. 2005), which would further explain why

Fig. 6 Abundance and biomass of all the beetle species considered in the analyses, depending on decline at the tree scale (healthy vs. declining). See Table 3 for statistical results; $P < 0.05$: *; $P < 0.01$: **



specialist phyllophagous weevils with endophytic larvae were negatively affected by oak decline. Overall, the negative response of specialist phyllophagous species to decline may relate to a decrease in leaf suitability (phytochemical profile and microclimatic conditions), an increase in predation pressure, or reduced food availability (fewer leaves) (Gely et al. 2019). Conversely, the positive response of generalist phyllophagous species to decline severity could stem from a decrease in interspecific competition with decreasing populations of specialist species (Kaplan and Denno 2007) and from improved conditions for larval development. For seminiphagous weevils, no effect of decline was detected, suggesting either that acorn quantity or quality was not markedly affected by oak decline or that the modifications were not significant enough to impact the species considered.

We considered decline level at two spatial scales: the tree and the plot. Overall, we observed significant responses mainly at the tree scale for xylophagous beetles, including oak-associated buprestids, and mainly at the plot scale for phytophagous and non-xylophagous saproxylic beetles. This might reflect differences among guilds in their dispersal capacity and host-selection behavior. For instance, some xylophagous species might have emerged from the declining trees carrying traps or might have been visually and/or chemically attracted by these declining trees, since weakened hosts often attract secondary pests (e.g., Haack and Benjamin 1982). More specifically, host volatiles such as terpenes or ethanol emitted by weakened trees can be used by these insects to discriminate suitable hosts (e.g., Montgomery and Wargo, 1983; Sánchez-Osorio et al., 2019).

For all the communities we monitored, further experiments would be necessary to identify the main drivers of the variations observed. More specifically, the effect of decline on the abundance of microhabitats and resources such as dead wood, cavities, opportunistic fungi, and acorns should be quantified (Heitzman et al. 2007; Spetich 2007). Likewise, changes in microclimates and predation pressure at the canopy and soil levels during a decline should be characterized. In addition, in our study, we were not able to take into account decline

dynamics, since historical data on decline onset, duration, and intensity at the stand scale was lacking. We considered stands exhibiting different decline levels, which may result from disturbances with different frequency, severity, and/or spatial and temporal extents at the stand scale. Past disturbance regimes can modulate the current taxonomic, functional, and phylogenetic composition forest communities, notably the community of saproxylic beetles (Kozák et al., 2020). Therefore, integrating historical data in future studies would help to disentangle current decline effects from past disturbance legacies.

Changes in species richness and abundance led to significant community modifications for both xylophagous beetles and phytophagous weevils, which in turn contributed to a significant modification of the overall beetle community. From a functional standpoint, this type of modification may modulate important processes in forest ecosystems, since saproxylic insects play a significant role in wood decomposition and the nitrogen cycle (Ulyshen 2015). In addition, saproxylic and leaf-dwelling beetles can be important prey for insectivorous vertebrates (e.g., Tillon et al. 2016; Koenig and Liebhold 2017), and changes in beetle community composition may therefore have cascading effects on the food web (e.g., Koenig and Liebhold 2017). From a conservation standpoint, the increase in species richness for the xylophagous and phyllophagous beetle communities suggests that declining stands might enhance forest biodiversity. Decline especially promoted saproxylic species. This community is particularly sensitive to the intensification of management practices involving the extraction of weakened or decaying wood material, and consequently includes several rare and protected species (Grove 2002; Seibold et al. 2015). The accumulation of suitable habitats and resources for this community in declining stands may then counterbalance the adverse effects of intensive management. The increase in abundance and/or biomass of xylophagous and phyllophagous beetles also resulted in an overall increase in beetle abundance and biomass in the declining stands. This also suggests that forest decline may mitigate the reduction in insect biomass recently reported in

Table 4 Canonical analysis of principal coordinates, based on Bray-Curtis distance matrices, ranking the effect of the two spatial levels of decline (plot vs. tree) on variations in species composition

Group	Ecological variable with the best contribution to inertia	Inertia explained by the best ecological variable (and significance)	% inertia explained	Inertia explained by sampling year	% inertia explained by sampling year
Oak-associated buprestid beetles	Plot	0.60*	5.9	2.19***	21.5
Xylophagous beetles (excl. buprestids)	Tree	0.58*	6.7	4.70***	54.5
Non-xylophagous saproxylic beetles	Plot	0.13	1.3	3.14***	30.4
Oak-associated weevils	Plot	0.32*	4.9	1.75***	26.2
All beetles	Plot	0.48**	5.5	2.63***	30.5

* $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$

Table 5 Characteristic species for each tree-decline level, identified using the IndVal approach. We retained only those species significant in the permutation test with an indicator value above 0.25, sampled in more than 10% of traps and with more than ten individuals

Group	Indicator species	Feeding guild	Indicator value	Frequency (%)
Declining trees	<i>Agrilus angustulus</i>	Xylophagous	0.891*	86
	<i>Agrilus sulcicollis</i>	Xylophagous	0.889**	81
	<i>Agrilus hastulifer</i>	Xylophagous	0.798*	65
	<i>Agrilus biguttatus</i>	Xylophagous	0.794**	51
	<i>Trichoferus pallidus</i>	Xylophagous	0.843**	65
	<i>Rhagium sycophanta</i>	Xylophagous	0.573*	16
	<i>Xylotrechus antilope</i>	Xylophagous	0.622*	30
	<i>Scolytus intricatus</i>	Xylophagous	0.683**	30
	<i>Gasterocercus depressirostris</i>	Xylophagous	0.517*	14
	<i>Mordella brachyura</i>	Saproxylophagous	0.804*	57
	<i>Cetonia aurata</i>	Saproxylophagous	0.699**	35
	<i>Opilo mollis</i>	Zoophagous	0.787*	62
	<i>Stenagostus rhombeus</i>	Zoophagous	0.733*	59
	<i>Lygistopterus sanguineus</i>	Zoophagous	0.716**	43
Healthy trees	<i>Calambus bipustulatus</i>	Zoophagous	0.693**	35

* $P < 0.05$ ** $P < 0.01$

European forests, in intensively managed landscapes (Seibold et al. 2019), at least if the increase in resources and structural complexity persists over time (Winter et al. 2015). In this regard, increases in species richness, abundance, and biomass of xylophagous species at the stand scale, as in our study, might prove to be ephemeral (Winter et al. 2015).

In our forests, the CAP sampling year explained a greater percentage of inertia than did decline level for all guilds considered. This strong year effect could not only result from high inter-annual variations in beetle abundance and/or occurrence but may also incorporate multiple methodological factors (i.e., (i) slight variations in sampling periods, (ii) changes in monitored plots, and (iii) modifications in the protocol of decline characterization at the plot level). Marked variations in beetle abundance and biomass occurred on plots and at periods that were consistently monitored throughout the 3 years of survey and between years (i.e., 2016 and 2017) when the protocol of decline characterization was identical (data not shown). This rather supports the hypothesis that the year effect mainly results from marked inter-annual variations in beetle abundance and community composition. Such fluctuations in population and community abundances are commonly observed in temperate forests (e.g., Stange et al. 2011). A longer monitoring period on the same plots would be necessary to identify the factors contributing to the between-year variations we observed.

Green Lindgren traps, placed at the canopy level, have proven to be effective in collecting leaf-dwelling beetles. These traps were specifically designed to collect *Agrilus planipennis* Fairmaire (Francesca et al. 2011) and have also

allowed researchers to collect North American and European Agrilinae species (Petrice and Haack 2015; Rassati et al. 2019). During our survey, all the Agrilinae species associated with oaks in France (i.e., *Agrilus* sp., *Coraebus* sp., and *Meliboeus* sp.) were captured, except for *Agrilus grandiceps hemiphanes* Marseul, a rare Mediterranean species, and *Coraebus florentinus* Herbst. The latter species had previously been collected in the Vierzon forest, and typical shoot browning resulting from its larval activity has already been reported there. The species might have been present but at too low population density for detection, or it might not have been attracted by our traps. We also collected quite diverse communities of phyllophagous and seminiphagous weevil species in our green Lindgren traps, in large amounts for some species. These species were significantly more attracted to green traps than to purple ones, which is congruent with the attraction to green substrates reported for other phytophagous weevils (e.g., Cross et al. 1976; Gadi and Reddy 2014). Overall, this suggests that green Lindgren traps are attractive to phyllobiont species in general, and confirms the tool's utility when investigating canopy-dwelling beetles associated with foliage.

5 Conclusion

Our 3-year survey in a declining forest allowed us to detect significant effects of decline on different canopy-dwelling species and guilds, in spite of strong inter-annual variations and a limited spatial extent, the survey being performed in two

adjacent forests. Overall, decline had a positive effect on the abundance and biomass of beetles, but contrasted variations were observed at the species or guild levels, with positive effects for saproxylic and generalist phylophagous species, null effects for seminiphagous species, and negative effects for specialist phylophagous species. These results call for studies conducted at larger spatial and temporal scales to assess the functional outcomes of the unprecedented level of forest decline expected to affect Europe, and to propose management strategies for conservation biologists.

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Data availability The datasets generated during and/or analyzed during the current study can be accessed at <https://dataverse.harvard.edu/dataverse/BUCHE>

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Reference to pre-print servers (when relevant) The preprint version of this article is available in the BioRxiv server, <https://doi.org/10.1101/2020.02.11.943753>

Annex

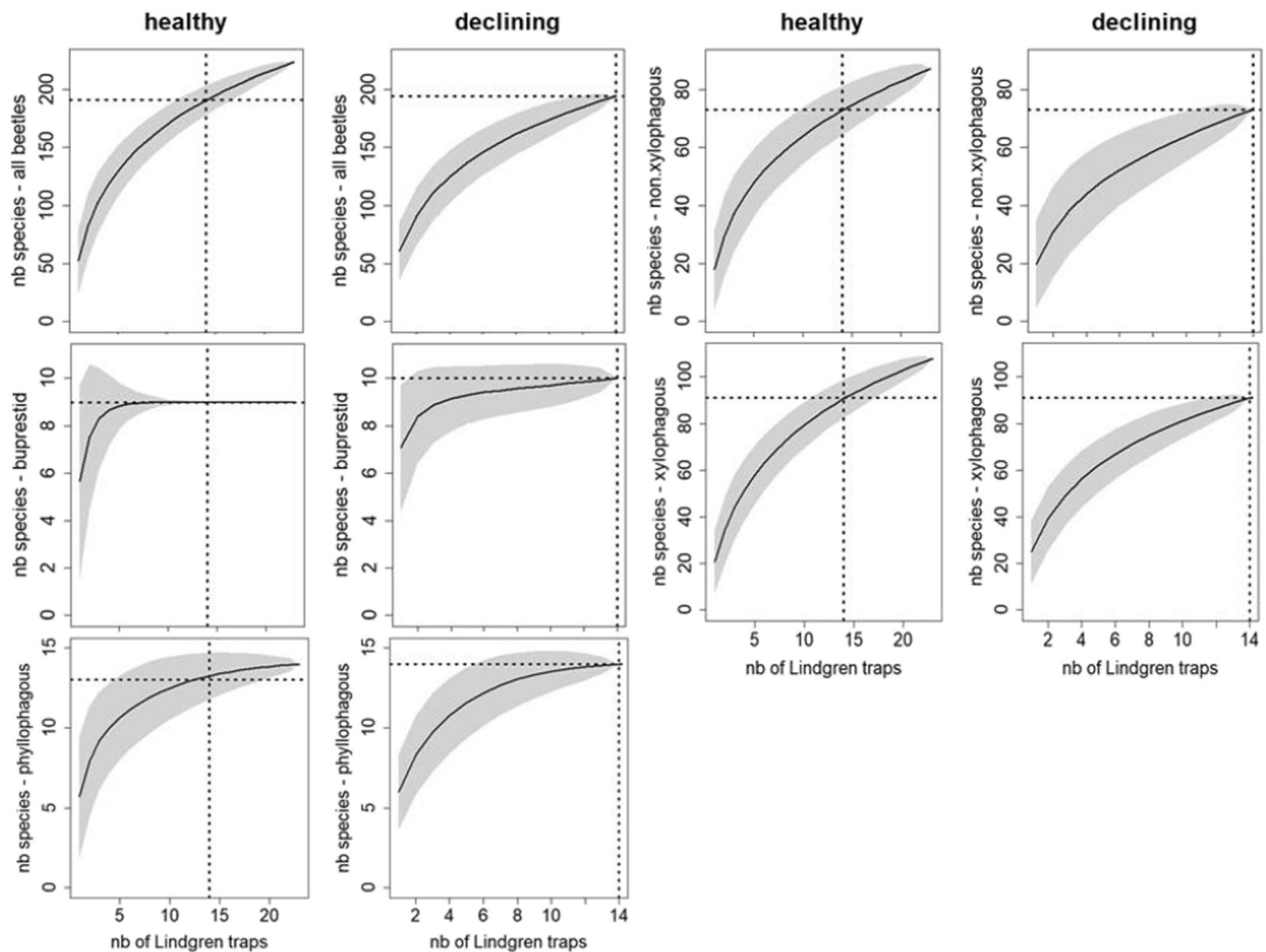


Fig. 7 Cumulative interpolated rarefaction of species richness with sample size (sampling without replacement) by dieback level at tree scale. Vertical dashed line = standard interpolated sample size,

horizontal dashed line = species richness estimate at the standardized sample size, and gray area = standard deviation of species richness estimate

Table 6 list of all the beetle species collected and used for the data analyses with their group, their guild, and their abundance

Group	Guild	Species	Family	Number of individuals	
Oak-associated buprestid beetles (Agrilinae)	NA	<i>Agrilus angustulus</i>	Buprestidae	2089	
	NA	<i>Agrilus biguttatus</i>	Buprestidae	282	
	NA	<i>Agrilus curtulus</i>	Buprestidae	1	
	NA	<i>Agrilus graminis</i>	Buprestidae	65	
	NA	<i>Agrilus hastulifer</i>	Buprestidae	1603	
	NA	<i>Agrilus laticornis</i>	Buprestidae	4029	
	NA	<i>Agrilus obscuricollis</i>	Buprestidae	1433	
	NA	<i>Agrilus sulcicollis</i>	Buprestidae	664	
	NA	<i>Coraeus undatus</i>	Buprestidae	198	
	NA	<i>Meliboeus fulgidicollis</i>	Buprestidae	76	
Oak-associated phytophagous weevils	Anthrophagous	<i>Coeliodes ilicis</i>	Curculionidae	4	
	Anthrophagous	<i>Coeliodes transversealbofasciatus</i>	Curculionidae	2	
	Seminiphagous	<i>Curculio elephas</i>	Curculionidae	67	
	Seminiphagous	<i>Curculio glandium</i>	Curculionidae	926	
	Seminiphagous	<i>Curculio pellitus</i>	Curculionidae	41	
	Seminiphagous	<i>Curculio venosus</i>	Curculionidae	111	
	Generalist	<i>Brachyderes incanus</i>	Curculionidae	74	
	phylophagous				
	Generalist	<i>Phyllobius pyri</i>	Curculionidae	1277	
	phylophagous				
	Generalist	<i>Polydrusus cervinus</i>	Curculionidae	260	
	phylophagous				
	Generalist	<i>Polydrusus marginatus</i>	Curculionidae	4	
	phylophagous				
	Generalist	<i>Strophosoma capitatum</i>	Curculionidae	15	
	phylophagous				
	Specialist	<i>Archarius pyrrhoceras</i>	Curculionidae	425	
	phylophagous				
	Specialist	<i>Attelabus nitens</i>	Curculionidae	7	
	phylophagous				
	Specialist	<i>Curculio villosus</i>	Curculionidae	6	
	phylophagous				
	Specialist	<i>Lasioryhynchites coeruleocephalus</i>	Curculionidae	29	
	phylophagous				
	Specialist	<i>Neocoenorhinidius interpunctatus</i>	Curculionidae	109	
	phylophagous				
	Specialist	<i>Neocoenorhinus minutus</i>	Curculionidae	10	
phylophagous					
Specialist	<i>Orchestes avellanae</i>	Curculionidae	120		
phylophagous					
Specialist	<i>Orchestes irroratus</i>	Curculionidae	10		
phylophagous					
Specialist	<i>Orchestes pilosus</i>	Curculionidae	13		
phylophagous					
Specialist	<i>Orchestes quercus</i>	Curculionidae	4770		
Saproxyllic beetles associated with deciduous trees	Non-xylophagous	<i>Anthribus nebulosus</i>	Anthribidae	16	
	Non-xylophagous	<i>Teredus cylindricus</i>	Bothrididae	1	
	Non-xylophagous	<i>Dromius agilis</i>	Carabidae	7	
	Non-xylophagous	<i>Dromius quadrimaculatus</i>	Carabidae	14	
	Non-xylophagous	<i>Cerylon ferrugineum</i>	Cerylonidae	1	
	Non-xylophagous	<i>Cerylon histeroides</i>	Cerylonidae	1	
	Non-xylophagous	<i>Cis pygmaeus</i>	Ciidae	1	
	Non-xylophagous	<i>Cis villosulus</i>	Ciidae	2	
	Non-xylophagous	<i>Ennearthron cornutum</i>	Ciidae	1	
	Non-xylophagous	<i>Orthocis lucasi</i>	Ciidae	2	
	Non-xylophagous	<i>Clambus armadillo</i>	Clambidae	1	
	Non-xylophagous	<i>Clerus mutillarius</i>	Cleridae	9	
	Non-xylophagous	<i>Opilo mollis</i>	Cleridae	59	
	Non-xylophagous	<i>Tilloidea unifasciata</i>	Cleridae	8	
	Non-xylophagous	<i>Tillus elongatus</i>	Cleridae	3	
	Saproxyllic beetles (excl. Agrilinae) associated with deciduous trees	Non-xylophagous	<i>Aplocnemus impressus</i>	Dasytidae	11
		Non-xylophagous	<i>Aplocnemus nigricornis</i>	Dasytidae	4
Non-xylophagous		<i>Dasytes aeratus</i>	Dasytidae	159	
Non-xylophagous		<i>Dasytes caeruleus</i>	Dasytidae	180	
Non-xylophagous		<i>Dasytes niger</i>	Dasytidae	2	
Non-xylophagous		<i>Dasytes nigrocyaneus</i>	Dasytidae	26	

Table 6 (continued)

Group	Guild	Species	Family	Number of individuals
	Non-xylophagous	<i>Dasytes pauperculus</i>	Dasytidae	116
	Non-xylophagous	<i>Dasytes plumbeus</i>	Dasytidae	8
	Non-xylophagous	<i>Dasytes subaeneus</i>	Dasytidae	1
	Non-xylophagous	<i>Dasytes virens</i>	Dasytidae	1
	Non-xylophagous	<i>Psilothrix viridicoerulea</i>	Dasytidae	1
	Non-xylophagous	<i>Trichoceble floralis</i>	Dasytidae	3
	Non-xylophagous	<i>Attagenus brunneus</i>	Dermestidae	1
	Non-xylophagous	<i>Attagenus pello</i>	Dermestidae	1
	Non-xylophagous	<i>Ctesias serra</i>	Dermestidae	34
	Non-xylophagous	<i>Dermestes lardarius</i>	Dermestidae	4
	Non-xylophagous	<i>Dermestes murinus</i>	Dermestidae	8
	Non-xylophagous	<i>Dermestes undulatus</i>	Dermestidae	69
	Non-xylophagous	<i>Globicornis fasciata</i>	Dermestidae	8
	Non-xylophagous	<i>Globicornis nigripes</i>	Dermestidae	3
	Non-xylophagous	<i>Megatoma undata</i>	Dermestidae	11
	Non-xylophagous	<i>Ampedus balteatus</i>	Elateridae	4
	Non-xylophagous	<i>Ampedus elongatulus</i>	Elateridae	30
	Non-xylophagous	<i>Ampedus nigerrimus</i>	Elateridae	197
	Non-xylophagous	<i>Ampedus pomorum</i>	Elateridae	50
	Non-xylophagous	<i>Ampedus quercicola</i>	Elateridae	13
	Non-xylophagous	<i>Ampedus rufipennis</i>	Elateridae	8
	Non-xylophagous	<i>Ampedus sanguinolentus</i>	Elateridae	59
	Non-xylophagous	<i>Brachygonus megerlei</i>	Elateridae	26
	Non-xylophagous	<i>Calambus bipustulatus</i>	Elateridae	20
	Non-xylophagous	<i>Cardiophorus ruficollis</i>	Elateridae	2
	Non-xylophagous	<i>Denticollis linearis</i>	Elateridae	1
	Non-xylophagous	<i>Elater ferrugineus</i>	Elateridae	1
	Non-xylophagous	<i>Hemicrepidius hirtus</i>	Elateridae	17
	Non-xylophagous	<i>Melanotus villosus</i>	Elateridae	29
	Non-xylophagous	<i>Podeonius acuticornis</i>	Elateridae	2
	Non-xylophagous	<i>Stenagostus rhombeus</i>	Elateridae	37
	Non-xylophagous	<i>Symbiotes gibberosus</i>	Endomychidae	2
	Non-xylophagous	<i>Dacne bipustulata</i>	Erotylidae	43
	Non-xylophagous	<i>Triplax lepida</i>	Erotylidae	65
	Non-xylophagous	<i>Triplax russica</i>	Erotylidae	14
	Non-xylophagous	<i>Tritoma bipustulata</i>	Erotylidae	2
	Non-xylophagous	<i>Gnathoncus nidorum</i>	Histeridae	2
	Non-xylophagous	<i>Laemophloeus monilis</i>	Laemophloeidae	2
	Non-xylophagous	<i>Placonotus testaceus</i>	Laemophloeidae	1
	Non-xylophagous	<i>Lygistopterus sanguineus</i>	Lycidae	28
	Non-xylophagous	<i>Abdera biflexuosa</i>	Melandryidae	10
	Non-xylophagous	<i>Monotoma picipes</i>	Monotomidae	1
Saproxylic beetles (excl. Agrilinae) associated with deciduous trees	Non-xylophagous	<i>Rhizophagus bipustulatus</i>	Monotomidae	3
	Non-xylophagous	<i>Rhizophagus dispar</i>	Monotomidae	1
	Non-xylophagous	<i>Rhizophagus ferrugineus</i>	Monotomidae	2
	Non-xylophagous	<i>Berginus tamarisci</i>	Mycetophagidae	100
	Non-xylophagous	<i>Eulagius filicornis</i>	Mycetophagidae	84
	Non-xylophagous	<i>Litargus balteatus</i>	Mycetophagidae	1
	Non-xylophagous	<i>Litargus connexus</i>	Mycetophagidae	72
	Non-xylophagous	<i>Mycetophagus piceus</i>	Mycetophagidae	2
	Non-xylophagous	<i>Mycetophagus populi</i>	Mycetophagidae	1
	Non-xylophagous	<i>Cryptarcha strigata</i>	Nitidulidae	157
	Non-xylophagous	<i>Cryptarcha undata</i>	Nitidulidae	149
	Non-xylophagous	<i>Cychramus luteus</i>	Nitidulidae	1
	Non-xylophagous	<i>Epuraea sp</i>	Nitidulidae	1
	Non-xylophagous	<i>Soronia grisea</i>	Nitidulidae	295
	Non-xylophagous	<i>Dorcatoma androgyna</i>	Ptinidae	1
	Non-xylophagous	<i>Dorcatoma chrysomelina</i>	Ptinidae	3
	Non-xylophagous	<i>Dorcatoma flavicornis</i>	Ptinidae	1
	Non-xylophagous	<i>Dorcatoma robusta</i>	Ptinidae	1
	Non-xylophagous	<i>Dorcatoma substriata</i>	Ptinidae	2
	Non-xylophagous	<i>Pyrochroa coccinea</i>	Pyrochroidae	58
	Non-xylophagous	<i>Salpingus planirostris</i>	Salpingidae	4
	Non-xylophagous	<i>Salpingus ruficollis</i>	Salpingidae	1
	Non-xylophagous	<i>Uleiota planatus</i>	Silvanidae	1
	Non-xylophagous	<i>Nemozoma elongatum</i>	Trogossitidae	1
	Non-xylophagous	<i>Tenebroides fuscus</i>	Trogossitidae	3
	Non-xylophagous	<i>Colobicus hirtus</i>	Zopheridae	4

Table 6 (continued)

Group	Guild	Species	Family	Number of individuals
	Non-xylophagous	<i>Colydium elongatum</i>	Zopheridae	2
	Non-xylophagous	<i>Endophloeus markovichianus</i>	Zopheridae	3
	Xylophagous	<i>Euglenes pygmaeus</i>	Aderidae	1
	Xylophagous	<i>Dissoleucas niveirostris</i>	Anthribidae	1
	Xylophagous	<i>Phaeochrotes pudens</i>	Anthribidae	3
	Xylophagous	<i>Platystomos albinus</i>	Anthribidae	38
	Xylophagous	<i>Pseudeuparius sepicola</i>	Anthribidae	6
	Xylophagous	<i>Rhaphitropis oxyacanthae</i>	Anthribidae	1
	Xylophagous	<i>Tropideres albostris</i>	Anthribidae	41
	Xylophagous	<i>Lyctinae</i>	Bostrichidae	1
	Xylophagous	<i>Xylopertha retusa</i>	Bostrichidae	1
	Xylophagous	<i>Anthaxia salicis</i>	Buprestidae	1
	Xylophagous	<i>Chrysobothris affinis</i>	Buprestidae	1
	Xylophagous	<i>Alosterna tabacicolor</i>	Cerambycidae	1
	Xylophagous	<i>Anaesthetis testacea</i>	Cerambycidae	2
	Xylophagous	<i>Anoplodera sexguttata</i>	Cerambycidae	6
	Xylophagous	<i>Callimus angulatus</i>	Cerambycidae	1
	Xylophagous	<i>Cerambyx cerdo</i>	Cerambycidae	1
	Xylophagous	<i>Cerambyx scopolii</i>	Cerambycidae	8
	Xylophagous	<i>Chlorophorus figuratus</i>	Cerambycidae	3
	Xylophagous	<i>Clytus arietis</i>	Cerambycidae	9
	Xylophagous	<i>Clytus tropicus</i>	Cerambycidae	4
	Xylophagous	<i>Cortodera humeralis</i>	Cerambycidae	8
	Xylophagous	<i>Dinoptera collaris</i>	Cerambycidae	1
	Xylophagous	<i>Exocentrus adpersus</i>	Cerambycidae	3
	Xylophagous	<i>Grammoptera abdominalis</i>	Cerambycidae	5
	Xylophagous	<i>Grammoptera ruficornis</i>	Cerambycidae	9
	Xylophagous	<i>Grammoptera ustulata</i>	Cerambycidae	7
	Xylophagous	<i>Leiopus femoratus</i>	Cerambycidae	1
	Xylophagous	<i>Leiopus linnei</i>	Cerambycidae	5
	Xylophagous	<i>Leiopus nebulosus</i>	Cerambycidae	6
	Xylophagous	<i>Mesosa curculionoides</i>	Cerambycidae	1
	Xylophagous	<i>Mesosa nebulosa</i>	Cerambycidae	27
	Xylophagous	<i>Pedostrangalia revestita</i>	Cerambycidae	1
	Xylophagous	<i>Phymatodes testaceus</i>	Cerambycidae	11
	Xylophagous	<i>Plagionotus detritus</i>	Cerambycidae	8
	Xylophagous	<i>Poecilium alni</i>	Cerambycidae	1
	Xylophagous	<i>Poecilium rufipes</i>	Cerambycidae	22
	Xylophagous	<i>Pogonocherus ovatus</i>	Cerambycidae	2
	Xylophagous	<i>Pseudosphegistes cinerea</i>	Cerambycidae	4
	Xylophagous	<i>Rhagium sycophanta</i>	Cerambycidae	16
	Xylophagous	<i>Rutpela maculata</i>	Cerambycidae	1
	Xylophagous	<i>Tetrops praeustus</i>	Cerambycidae	2
	Xylophagous	<i>Trichoferus pallidus</i>	Cerambycidae	95
	Xylophagous	<i>Xylotrechus antilope</i>	Cerambycidae	66
	Xylophagous	<i>Anisandrus dispar</i>	Curculionidae	1600
	Xylophagous	<i>Cyclorhipidion bodoanum</i>	Curculionidae	24
	Xylophagous	<i>Ermoporicus fagi</i>	Curculionidae	1
	Xylophagous	<i>Gasterocercus depressirostris</i>	Curculionidae	10
	Xylophagous	<i>Hylastinus obscurus</i>	Curculionidae	6
	Xylophagous	<i>Hylesinus varius</i>	Curculionidae	1
	Xylophagous	<i>Magdalis flavicornis</i>	Curculionidae	46
	Xylophagous	<i>Platypus cylindrus</i>	Curculionidae	4
	Xylophagous	<i>Pteleobius kraatzii</i>	Curculionidae	1
	Xylophagous	<i>Scolytus intricatus</i>	Curculionidae	52
	Xylophagous	<i>Scolytus multistriatus</i>	Curculionidae	2
	Xylophagous	<i>Scolytus pygmaeus</i>	Curculionidae	1
	Xylophagous	<i>Scolytus rugulosus</i>	Curculionidae	9
	Xylophagous	<i>Scolytus scolytus</i>	Curculionidae	1
	Xylophagous	<i>Taphrorychus bicolor</i>	Curculionidae	23
	Xylophagous	<i>Taphrorychus villifrons</i>	Curculionidae	2
	Xylophagous	<i>Trypodendron signatum</i>	Curculionidae	2
	Xylophagous	<i>Xyleborinus saxesenii</i>	Curculionidae	291
	Xylophagous	<i>Xyleborus dryographus</i>	Curculionidae	4
	Xylophagous	<i>Xyleborus monographus</i>	Curculionidae	122
	Xylophagous	<i>Xylosandrus germanus</i>	Curculionidae	1
	Xylophagous	<i>Dromaeolus barnabita</i>	Eucnemidae	46
	Xylophagous	<i>Eucnemis capucina</i>	Eucnemidae	4
Saproxylous beetles (excl. Agrilinae) associated with deciduous trees				

Table 6 (continued)

Group	Guild	Species	Family	Number of individuals
Saproxylic beetles (excl. Agrilinae) associated with deciduous trees	Xylophagous	<i>Hylis olexai</i>	Eucnemidae	2
	Xylophagous	<i>Hylis simonae</i>	Eucnemidae	7
	Xylophagous	<i>Isorhipis melasoides</i>	Eucnemidae	1
	Xylophagous	<i>Melasis buprestoides</i>	Eucnemidae	12
	Xylophagous	<i>Microrhagus pygmaeus</i>	Eucnemidae	2
	Xylophagous	<i>Microrhagus pyrenaeus</i>	Eucnemidae	2
	Xylophagous	<i>Dorcus parallelipipedus</i>	Lucanidae	2
	Xylophagous	<i>Platycerus caraboides</i>	Lucanidae	50
	Xylophagous	<i>Lymexylon navale</i>	Lymexylidae	1
	Xylophagous	<i>Anisoxya fuscula</i>	Melandryidae	1
	Xylophagous	<i>Conopalpus brevicollis</i>	Melandryidae	4
	Xylophagous	<i>Conopalpus testaceus</i>	Melandryidae	6
	Xylophagous	<i>Melandrya barbata</i>	Melandryidae	2
	Xylophagous	<i>Phloiotrya tenuis</i>	Melandryidae	7
	Xylophagous	<i>Mordella brachyura</i>	Mordellidae	207
	Xylophagous	<i>Mordella leucaspis</i>	Mordellidae	1
	Xylophagous	<i>Mordellistena humeralis</i>	Mordellidae	4
	Xylophagous	<i>Mordellistena neuwaldeggiana</i>	Mordellidae	25
	Xylophagous	<i>Mordellistena variegata</i>	Mordellidae	11
	Xylophagous	<i>Tolida artemisiae</i>	Mordellidae	2
	Xylophagous	<i>Tomoxia bucephala</i>	Mordellidae	44
	Xylophagous	<i>Ischnomera caerulea</i>	Oedemeridae	223
	Xylophagous	<i>Nacerdes carniolica</i>	Oedemeridae	54
	Xylophagous	<i>Oedemera flavipes</i>	Oedemeridae	1
	Xylophagous	<i>Gastrallus immarginatus</i>	Ptinidae	7
	Xylophagous	<i>Gastrallus laevigatus</i>	Ptinidae	38
	Xylophagous	<i>Gastrallus vavrai</i>	Ptinidae	1
	Xylophagous	<i>Hadrobregmus denticollis</i>	Ptinidae	1
	Xylophagous	<i>Hemicoelus costatus</i>	Ptinidae	4
	Xylophagous	<i>Hemicoelus fulvicornis</i>	Ptinidae	3
	Xylophagous	<i>Hyperisus plumbeum</i>	Ptinidae	6
	Xylophagous	<i>Mesocoelopus niger</i>	Ptinidae	11
	Xylophagous	<i>Oligomerus brunneus</i>	Ptinidae	7
	Xylophagous	<i>Ptilinus pectinicornis</i>	Ptinidae	1
	Xylophagous	<i>Ptinomorphus imperialis</i>	Ptinidae	51
	Xylophagous	<i>Xyletinus pectinatus</i>	Ptinidae	2
	Xylophagous	<i>Cetonia aurata</i>	Scarabaeidae	37
	Xylophagous	<i>Cetonischema speciosissima</i>	Scarabaeidae	22
	Xylophagous	<i>Gnorimus nobilis</i>	Scarabaeidae	1
	Xylophagous	<i>Liocola marmorata</i>	Scarabaeidae	1
	Xylophagous	<i>Potosia fieberi</i>	Scarabaeidae	14
	Xylophagous	<i>Valgus hemipterus</i>	Scarabaeidae	2
	Xylophagous	<i>Anaspis fasciata</i>	Scraptiidae	35
	Xylophagous	<i>Anaspis flava</i>	Scraptiidae	29
	Xylophagous	<i>Anaspis frontalis</i>	Scraptiidae	277
	Xylophagous	<i>Anaspis lurida</i>	Scraptiidae	100
	Xylophagous	<i>Anaspis maculata</i>	Scraptiidae	134
Xylophagous	<i>Anaspis pulicaria</i>	Scraptiidae	11	
Xylophagous	<i>Anaspis regimbarti</i>	Scraptiidae	66	
Xylophagous	<i>Scraptia testacea</i>	Scraptiidae	4	
Xylophagous	<i>Allecula morio</i>	Tenebrionidae	1	
Xylophagous	<i>Gonodera luperus</i>	Tenebrionidae	4	
Xylophagous	<i>Mycetochara maura</i>	Tenebrionidae	10	
Xylophagous	<i>Nalassus laevioctostriatus</i>	Tenebrionidae	2	
Xylophagous	<i>Platydemia violacea</i>	Tenebrionidae	2	
Xylophagous	<i>Prionychus ater</i>	Tenebrionidae	1	
Xylophagous	<i>Prionychus fairmairii</i>	Tenebrionidae	29	

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