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The ecological impact of pest-induced tree dieback on insect biodiversity in Yunnan pine plantations, China

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ABSTRACT

China has recently announced a reform of forestry policy, with a major goal being to transform from plantation to heterogeneous forests, which have higher resistance to pests and disease and house more biodiversity. One driver of reform is increased intensity and frequency of pest-induced tree-dieback events. To inform management, we ask what effects these events have on insect biodiversity in Pinus yunnanensis monocultures in Yunnan province, the province with one of the highest proportions of forest cover in China. We sampled aerial arthropods (mostly insect) biodiversity along gradients of Pinus yunnanensis dieback severity using Malaise traps and used metabarcoding to characterise the insect community. We used MS-GDM ('multi-site generalized dissimilarity modelling of zeta diversity'), zeta-decline analysis, and iNEXT ('Interpolation and extrapolation for species diversity') to assess community change as functions of forest-structure covariates. Metabarcoding of Malaisetrapped insects reveals that bark-beetle induced forest dieback does not result in detectable differences in species diversity but does result in compositional change, with the biggest turnover occurring between 0% and infested-0%-open-canopy forests and 20%-infested-20%-open-canopy forests. Zeta-decline analysis found that the insect community in low-infestation forests is characterized by a stochastic assembly, while in highinfestation forests, the community structure is consistent with niche assembly. Our results thus suggest that bark-beetle dieback mimics natural forest-gap dynamics, consistent with the interpretation of bark beetles as a keystone species in European conifer forests, where it has been proposed that forest heterogeneity can be created efficiently by allowing natural disturbances, including bark-beetle outbreaks, to proceed naturally, without being mitigated by deadwood removal and dense replanting. In Yunnan's situation, and given predicted increases in bark-beetle dieback severity and frequency, this strategy should probably be supplemented with anthropogenic treatments, such as deadwood enhancement and planting of multiple tree species, to accelerate the succession of plantations into heterogeneous forests.

1. Introduction

The largest reforestation programmes in the world are China's Natural Forest Protection Program (NFPP) and Grain for Green Program (GFGP), which were implemented after widespread flooding in 1998

(Liu et al., 2008; Vina et al., 2016; Xu et al., 2006; Yin et al., 2009). The NFPP protects native forests in the upstream watersheds of the Yangtze and Yellow rivers (Liu et al., 2008; Ren et al., 2015), and the GFGP controls soil erosion by paying farmers to plant trees on sloping land that had been used for food production (Delang & Yuan, 2015; Liu et al.,

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2008; Ma et al., 2017; Xu et al., 2006; Zhai et al., 2014). The GFGP reforested 9.06 million ha of cropland between 1999 and 2014, and not surprisingly, the GFGP has primarily established low-diversity tree plantations ('plantations' hereafter), rather than restoring native forest (Hua et al., 2018, 2016; Zhai et al., 2014).

Studies have previously shown that these plantations support lower levels of bird, bee, and general insect diversity than do native forests in the same locations (Hua et al., 2016; Wang et al., 2019). These findings complement those of Cao et al. (2019), who recently calculated that plantations in China return a lower net value of ecosystem services relative to native forests, even after counting income from timber sales. Plantations require a high initial outlay for tree planting, some nonnative tree species like *Eucalyptus* require more water input than do native tree species, and more management effort is required to protect plantations from pest attack (Brockerhoff et al., 2013). In contrast, income from timber sales is low. Thus, to better protect and restore terrestrial biodiversity, studies have recommended that reforestation policy in China should prioritize the conservation and restoration of native forest over plantations (Hua et al., 2016; Wang et al., 2019).

These initial results in China are consistent with those from a larger body of research on forest biodiversity and ecosystem functioning in Central Europe, where professional silviculture has long promoted plantations, which have now grown to be dominated by dense tree stands with few canopy gaps and low volumes of deadwood (Doerfler et al., 2018; Thorn et al., 2018, 2019). Such forests support a low diversity of plants and animals, especially of saproxylic species (Thorn et al., 2018, 2019) and are more vulnerable to large-scale bark-beetle (Curculionidae, Scolytinae) outbreaks because the forests are even-aged and thus grow to provide an extensive and continuous cover of the large trees that are ideal hosts for bark beetles (Seidl et al., 2016).

Bark-beetle outbreaks are now a primary killer of coniferous forest in central Europe (Thorn et al., 2019), as well as North America (Robertson et al., 2009) and China (Gan, 2015; He & Zhang, 2004). Moreover, climate change is increasing the frequency of bark-beetle outbreaks (Carroll et al., 2004; Esper et al., 2007; Sambaraju et al., 2012; Seidl et al., 2017). For instance, more frequent and severe droughts and high temperatures impede pines from producing enough toxic resin to disable attacking beetles (Erbilgin et al., 2017; Kichas et al., 2020; Raffa & Berryman, 1983), and consequently, bark-beetle populations can more easily grow to outbreak levels (Cullingham et al., 2011).

Bark-beetle outbreaks leave many standing dead trees, leading to an overall increase in deadwood amount and stand structural heterogenity (Swanson et al., 2011). Forest managers often carry out salvage logging by removing infected trees in order to stop the expansion of the beetles (Stadelmann et al., 2013) and to recover the economic value of wood (Lindenmayer et al., 2008). The removal of infected trees has a negative impact not only on bark beetles but also on other species associated with dead wood (Thorn et al., 2018) but can have positive effects by on species that are normally associated with open areas (Rost & Clavero, 2012).

Our study region of Yunnan province, southwestern China, has one of the highest proportions of forest cover in China (Ren et al., 2015; SFA, 2016), and Pinus yunnanensis plantations account for 28.2% of this forest cover (YNFA, 2018), 80% of which is monoculture (Cai et al., 2006). Most of the Pinus yunnanensis forest has grown up on land where primary evergreen broadleaved forests had been destroyed (Deng et al., 2014). The provincial forestry bureau carries out salvage logging by cutting and removing 'snags' (upright dead trees) to control local outbreaks of three species of pine shoot beetles of genus Tomicus (Coleoptera: Curculionidae: Scolytinae) (Gan, 2015; Kirkendall et al., 2008; Lu & Zhang, 2000; Lu et al., 2014; Wang et al., 2015). In addition, locals remove logs for firewood, leading to very low volumes of dead wood in Pinus yunnanensis plantations, despite massive shoot beetle outbreaks affecting over 200,000 ha of pine plantations in Yunnan (Ji et al., 2007; Lieutier et al., 2003). This combination in Yunnan of GFGP-financed plantation dominance, Tomicus outbreaks, and salvage logging results in conifer

forests similar to those in Central Europe: structurally simple, even-aged tree cover that, despite cut-and-removal of infested trees, remains vulnerable to bark-beetle outbreaks (Cai et al., 2006) and supports low levels of native biodiversity compared to native forest.

However, China announced its intention to implement a new forest restoration plan in 2019 (Xinhua News Agency, 2019), with a major policy goal being to transform plantations into heterogeneous forests that have higher resistance to pests and disease.

In this study, we used Malaise traps to sample aerial insect biodiversity (dominated by Diptera and Hymenoptera) along gradients of Pinus yunnanensis dieback severity. The initial goal of our study was to study the ecological impact of bark-beetle-induced dieback on flying insect diversity. In particular, we were interested in whether patterns of forest insect diversity in Yunnan plantations are similar to those in Central Europe, where impacts of tree-dieback on habitat structure and salvage logging have been extensively studied (Doerfler et al., 2018; Hilmers et al., 2018; Müller et al., 2010; Seibold et al., 2016a, 2016b, 2018; Thorn et al., 2018). If similar, then this increases our confidence in applying lessons learned there to Yunnan and neighboring provinces (e. g. the efficacy of deadwood enrichment as a means of promoting saproxylic taxa; Doerfler et al., 2018; Seibold et al., 2016a, 2016b). Secondly, given China's recent forest-policy reform announcement, our results serve as a baseline survey of aerial insect biodiversity in Yunnan's Pinus yunnanensis plantations, to allow comparison with future forests in which, we presume, China will promote, or at least allow, the accumulation of greater structural and age heterogeneity and more deadwood.

We characterized the Malaise-trap samples using DNA metabarcoding, which combines DNA barcoding with high-throughput DNA sequencing to generate large sample X species tables that can be used to test the effects of candidate environmental variables on biodiversity. We did not specifically collect saproxylic taxa because the current cut-andremove policy means that there are no *Pinus yunnanensis* forests in Yunnan with high volumes of deadwood to act as a contrast. Metabarcoding has been tested against morphologically identified samples and been shown to be a reliable and efficient method of characterizing the species compositions of bulk samples of insects and invertebrates generally (Aylagas et al., 2018; Cordier et al., 2017; Edwards et al., 2014; Ji et al., 2013; Lejzerowicz et al., 2015; Pawlowski et al., 2016; Wang et al., 2019; Yu et al., 2012). Accessible explanations of metabarcoding are available in Bush et al. (2017), Ji et al. (2013), Piper et al. (2019), Yang et al. (2020), and Zinger et al. (2019).

2. Methods

2.1. Field sampling and environmental variables

Following the distribution of *Pinus yunnanensis* in Yunnan province, southwest China, we sampled in five counties across the elevational range of optimal growth (1800–3000 m, Table 1) (Deng et al., 2013). In each county, we sampled in six *P. yunnanensis*-dominated forest stands of at least 1 Ha extent along a gradient of bark-beetle-induced dieback severity: two sites each in low, medium, and high severity (Fig. 1) (sampling locations and elevations in Table S1). Severity was judged by local forestry officials, who are charged with responding to bark-beetle outbreaks, using a method defined in 2006 by the then-State Administration of Forestry (now National Forestry and Grassland Administration) (SFA, 2006). All sampling plots are reported to have been attacked the first time in the 1980s (Zhao & Långström, 2012). Our goal with this initial blocking was only to maximize coverage of the local gradient of dieback severity.

We set out and retrieved Malaise traps during 4–14 July 2016, placing two traps 10 m apart in each forest site (pairs were pooled at DNA extraction before downstream processing) for a total of 60 samples (5 counties X 6 sites/county X 2 traps/site). We used absolute ethanol as the killing and preserving agent, and the traps were left out for seven

Table 1
Environmental covariates and definitions.

Environmental covariates	Definitions	Range _{(of}		
Elevation	Recorded by GPS at the plot center.	1757–3052		
		m		
Height	Mean height of 40 trees in the quadrat,	4.61-13.00		
	where the trees are the first 10 trees north, south, east, and west of the quadrat centre.	m		
DBH	Mean diameter at breast height of 40 trees	7.9-23.6 cm		
	in the quadrat, using the same trees used			
	for the height measurement.			
Canopy openness	Mean proportion of sky visible in the	0.01 - 0.68		
	quadrat, measured by spherical			
	densiometer (Paletto & Tosi, 2009).			
	Measurements were taken at quadrat			
	centre and each corner, and averaged.			
Infestation rate	The percentage of trees that are infested.	0–0.79		
	Trees with one or more bark beetle			
	emergence holes on the north, south, east,			
	and west sides of their trunks were scored			
	as infected, using the same trees used for			
	the height measurement.			
Stumps	The total number of tree stumps in the	0–22		
	quadrat.			

days each. For efficiency, we set out all traps in one county and then moved the next day to another county. Retrieval followed the same schedule, and we replaced the ethanol in each trap with fresh ethanol for transport at ambient temperature to our laboratory, where samples were stored at $-80\,^{\circ}\text{C}$ until DNA extraction. In addition, in each of the 30 sampling sites, we centered a 30 X 30 m quadrat over the Malaise traps

and measured six environmental covariates: elevation, mean tree height, mean diameter at breast height (DBH), percentage canopy openness, mean infestation rate (the percentage of trees with one or more bark beetle emergence holes on all four cardinal sides), and stump number (details in Table 1).

2.2. DNA extraction and PCRs

Before DNA extraction, the storage ethanol was decanted, and the sample was air-dried on single-use filter papers. To reduce PCR dominance by large-biomass individuals (Elbrecht et al., 2017), we used two legs from all individuals larger than a housefly and whole bodies of everything smaller. Tissue was digested using a modified non-destructive protocol from Gilbert et al. (2007) and Nielsen et al. (2019) in one 50-ml falcon tube per sample, followed by DNA extraction with the DNEasy Blood & Tissue Kit (Qiagen GmbH, Germany). After extraction, we pooled the DNA from the paired Malaise traps, leaving us with 30 samples, one per site.

We used *mlCOlintF–Fol-degen-rev* primers (Leray et al., 2013; Yu et al., 2012), which amplify a 313-bp fragment of the COI barcode, and we followed the DAMe metabarcoding protocol (Alberdi et al., 2018; Bohmann et al., 2018; Zepeda-Mendoza et al., 2016), which is a codesigned wet-lab and bioinformatic pipeline that combines qPCR-optimized PCR conditions, multiple, independent PCR replicates per sample, twin-tagging, and negative and positive controls to (i) remove sequence-to-sample misassignment due to tag-jumping (Schnell et al., 2015), (ii) reduce sequence dropout and taxonomic bias in amplification, and (iii) reduce erroneous sequences. Twin-tagging means that the same tag is used on both the forward and reverse primers in a reaction

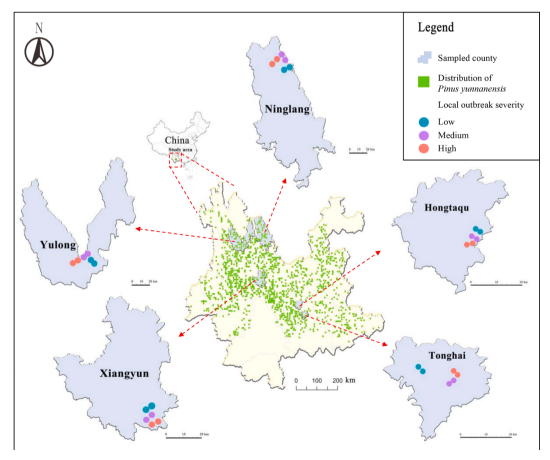


Fig. 1. Map of study area, Yunnan province, southwest China. Blue insets are the five sampled counties. Green squares indicate *Pinus yunnanensis* forest distribution. Coloured circles indicate sampling sites, stratified by severity of bark-beetle outbreak, as judged subjectively by local forestry officials. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

(F1-R1, F2-R2,...), and multiple, independent PCR replicates per sample means that a different twin tag is used for each of the six PCRs per sample, which lets them be distinguished in bioinformatic processing. The DAMe logic is that tag-jump events can be filtered out by removing reads carrying non-twinned tags (e.g. F1-F2, F3-F5) and that nearly all erroneous sequences (indels, substitutions, chimeras) can be filtered out by removing sequences that appear in only one (or a low number of) PCR replicate(s) at a low copy number, while true sequences are more likely to appear in multiple PCRs at higher copy numbers. Extensive testing with a recently updated version of DAMe (now called Begum) using mock samples finds that erroneous sequences can be nearly eliminated at the cost of only a small rise in drop-outs, and a detailed explanation of the protocol can be found there (Yang et al., 2020).

We used qPCR on a subset of samples to optimize PCR annealing temperature, cycle number, and initial DNA template concentration, as recommended by Murray et al. (2015) and Bohmann et al. (2018). Afterwards, for each sample, we ran 6 independent PCRs with 6 different twin-tags, under the following qPCR-optimised conditions: initial denaturation 95 °C for 5 min, followed by 27 cycles of 95 °C for 10 s, 45.5 °C for 45 s, 72 °C for 1 min, and finishing at 72 °C for 10 mins. All PCRs were performed in 20 µl reactions containing 0.6 U Ex Taq HS DNA polymerase, 1 X Ex Taq Buffer (Mg²⁺ plus), 0.2 mM dNTP Mixture (TaKaRa, Biotechnology Co. Ltd, Dalian, China), 0.4 µM of each primer, 1 μl DMSO, 0.1 μg/μl BSA (Bovine Serum Albumin Solution, TaKaRa Biotechnology Co. Ltd, Dalian, China), and 2 µl genomic DNA. We visualized the PCR products on 2% agarose gels. The PCR plate also included three extraction blanks and a row of PCR blanks. Finally, we included a positive control containing seven insect species from France. The 30 samples were combined into six, approximately equimolar pools for bead purification (Agencourt AMPure XP kit, Beckman Coulter, Inc., USA) and subsequent library preparation using the NEXTflex Rapid DNA-Seq Kit for Illumina (Bioo Scientific Corp., Austin, USA). The six libraries were sequenced on the Illumina MiSeq platform (300PE) at the Southwest Biodiversity Institute Regional Instrument Center in Kunming.

2.3. Bioinformatic processing

Raw MiSeq data were first trimmed for remnant Illumina adapters with *AdapterRemoval* 2.2.0 (Schubert et al., 2016), followed by Schirmer et al.'s (2015) recommended pipeline: we trimmed low-quality ends using *sickle* 1.33 (Joshi & Fass, 2011), denoised reads using the *Bayes-Hammer* module in *SPAdes* 3.10.1 (Nikolenko et al., 2013), and merged read pairs using *PandaSeq* 2.11 (Masella et al., 2012). In all cases, we used default parameters.

Sequence were demultiplexed to sample and filtered for tag-jumps using a modified version of DAMe that ignores heterogeneity spacers in the primers (github.com/shyamsg/DAMe, accessed 10 October 2020). We then filtered out putatively erroneous sequences by keeping only those that appeared in > 2 of the 6 PCRs per sample, at a minimum copy number of 30 per PCR, which is the stringency level that minimized false negatives and maximized true positives in the positive control. We further filtered by removing sequences ≤ 300 bp length and using the *de* novo chimera search function in vsearch 2.4.3 (Rognes et al., 2016). After filtering, sequences were clustered into 97% similarity Operational Taxonomic Units (OTUs) using SUMACLUST 1.0.20 (Mercier et al., 2013), from which we created a Sample X OTU table, and we used the R package 'lulu' 0.1.0 (Frøslev et al., 2017) with default parameters to combine likely 'parent' and 'child' OTUs that had failed to cluster. Finally, we assigned taxonomies to the remaining OTUs with the RDP Classifier function (Wang et al., 2007) on the Midori metazoan mitochondrial gene website (Leray et al., 2018). OTUs assigned to Arthropoda with < 80% probability were removed. No OTUs remained in the extraction-blank and PCR negative controls, and the positive control and samples shared no OTUs. We also tried assigning taxonomies on BOLD (Ratnasingham & Hebert, 2007), but only a few OTUs received hits, due

to a lack of samples from this region.

2.4. Statistical analyses

All statistical analyses were carried out in R 3.6.3. Read numbers per OTU per sample were transformed to presence/absence (1/0). We first used the 'boral' 1.6.1 R package (Hui, 2016) to cluster sites by community composition. Boral is a Bayesian, model-based ordination method that allows the selection of an appropriate error distribution. We used a binomial error distribution and no row effect to fit the model since we were using presence/absence data. For the same reason, we used 'mvabund' 3.12.3 (Wang et al., 2012) to test for the effects of environmental covariates on community composition.

Because the boral ordination showed that the dominant driver of change in community composition is geographic distance, which is not surprising given the large spatial extent of our sampling (Fig. 1), we followed up with Multi-Site Generalized Dissimilarity Modelling (MS-GDM), using the 'zetadiv' 1.2.0 package (Latombe et al., 2017). Classical GDMs try to identify the dominant drivers of change in community composition by using a combination of pairwise (i.e. between-two-sites) differences in geographic distance and in environmental-covariate values to explain pairwise differences in community composition (Ferrier et al., 2007). However, pairwise differences in composition (e.g. 1-Jaccard) are dominated by the contributions of the many species that are present in just two samples (i.e. rare species), resulting in GDMs that more heavily weight the variables that explain turnover in rare species, such as geographic distance.

To identify the environmental variables that are more important for explaining the distributions of widespread species (i.e. those present in multiple samples), Latombe et al. (2017) combined GDMs with the concept of zeta diversity (Hui & McGeoch, 2014) to create MS-GDMs. Zeta diversity is a generalization of pairwise beta diversity and is the mean number of species shared by *i* number of sites, where *i* is known as the zeta order. Zeta diversity order 4, for instance, is the mean number of species shared by 4 sites (in a dataset of 100 sites, there are \sim 3.9 million combinations of 4 sites). Zeta diversities can be converted to multi-site equivalents of the pairwise Jaccard dissimilarity and used as response variables in an MS-GDM (Latombe et al., 2017, 2019), with the six environmental covariates as candidate predictors (Table 1), rescaled between 0 and 1. We also used zeta diversity to ask if the insect communities in low- and high-infestation forests show evidence for different assembly mechanisms, by using the 'zetadiv' package to calculate zeta diversity decline and species retention rates for low- and for highinfestation forests. Finally, we partitioned variation in zeta diversity into environmental, distance, indistinguishable, and unexplained components.

To compare alpha diversity across infestation levels (Species richness, Shannon and Simpson diversities), we used the sample-based rarefaction-extrapolation approach in the 'iNEXT' 2.0.12 package (Hsieh et al., 2016). Significant differences in estimated alpha diversity were judged by non-overlapping confidence intervals, which is considered slightly conservative (MacGregor-Fors & Payton, 2013). In case we had oversplit some biological species into multiple OTUs, leading to artefactual differences in species richness, we also carried out a phylogenetic-diversity (PD) analysis because a single species split into multiple OTUs should cluster on a phylogenetic tree and thus contribute less to PD than two OTUs from two different biological species. Our protocol followed that of Wang et al. (2019), in which we aligned the OTU sequences, built a maximum-likelihood (ML) phylogenetic tree (details in S2), and estimated PD with the 'iNextPD' 0.3.2 package (Hsieh & Chao, 2017). We omitted two OTUs because they produced long branches.

3. Results

3.1. Bioinformatic processing and taxonomic composition

The six libraries yielded 11,128,217 paired-end reads. After removing a very large number of tag-jumped, paired-end reads (7,526,449), followed by DAMe filtering (retaining 1,217,449 sequences in ≥ 2 of the 6 PCRs per sample at ≥ 30 copies per PCR), and removal of chimeras and OTUs not assigned to Arthropoda, we ended up with 1,107,100 reads, clustered into 880 97% OTUs, for downstream analysis. Mean reads per OTU was 1,258 (Range =66–54,775; SD =2930), and mean reads per sample was 36,903 (n =30; range 3,881–72,575; SD =17,396). These 880 OTUs were assigned to 35.8% Diptera, 21.7% Lepidoptera, 19.1% Hymenoptera, 9.7% Coleoptera, 7% Hemiptera, and 6.7% other orders.

Read depth varied across samples (Fig. S3A), and we found a positive correlation between read depth and species richness (Pearson, p < 0.001). Thus, to test the robustness of our results, we removed eight samples that had < 25,000 reads, which removed the positive correlation (Pearson, p = 0.68, Fig. S3B), reran the analyses below (3.2–3.5), and as we report below and in Supplementary Information (S4, S6, S9, S10), found essentially the same results. We report the full-dataset results in the Main Text.

3.2. Boral ordination

Boral ordination (Fig. 2) clustered the 30 sites by the five counties in

3

Ľo M which we sampled (Fig. 1) and arranged the clusters by elevation (latent variable 1) and tree-infestation rate (latent variable 2). Mvabund analysis confirmed the same effects (Table 2) and found no evidence for an interaction effect. Boral results without low-read-depth samples in Fig. S4.

3.3. Multi-site generalized dissimilarity modelling

We carried out MS-GDM to identify the main drivers of change in community composition after controlling for geographical distance. Initially, we ran the model with five environmental covariates from Table 1 (omitting elevation), plus geographic distances between sites, because geographic distance and elevation are correlated. At zeta order 2 (equivalent to the Jaccard index, which is pairwise and thus dominated by rare species), distance is the dominant driver of compositional turnover, followed by the local environmental variables canopy openness and DBH (Fig. 3A, Order2). Distance is largely linear in its effects,

Table 2Myabund analysis. Testing for the effects of Elevation, Infestation rate, and their interaction on community composition.

	Res.Df	Df.diff	Score	Pr(>score)
Intercept	29			
Elevation	28	1	100.1	0.001
Infestation rate	27	1	136.8	0.044
Elevation:Infestation rate	26	1	131.1	0.935

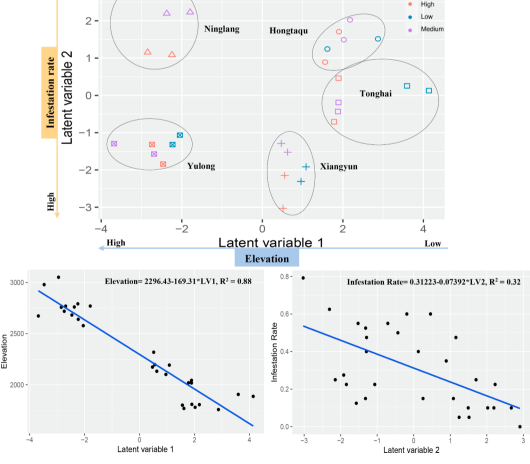


Fig. 2. 'Boral' ordination of beta diversity by disturbance type. Color codes for outbreak severity as in Fig. 1. Symbols (and surrounding ovals) indicate the five counties, and points represent samples. Latent variable 1 predicts elevation (2296.43–169.31*LV1, $R^2 = 88.0\%$, df = 28, p = 4.42e-15), and latent variable 2 predicts tree-infestation rate (0.31–0.07*LV2, $R^2 = 31.5\%$, df = 28, p = 0.0007). Boral residuals in Fig. S5.

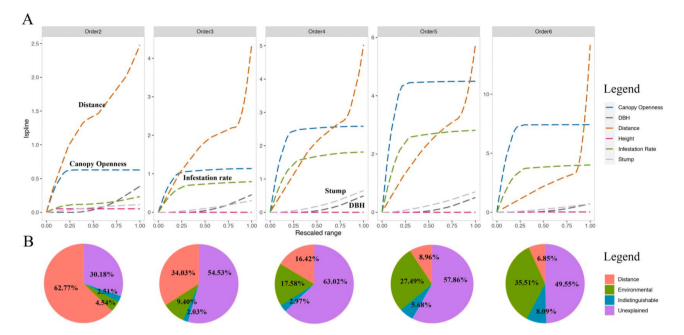


Fig. 3. Multi Site Generalised Dissimilarity Modelling (MS-GDM) analysis. A. Contributions of five environmental covariates and distance to explaining zeta diversity and B. and variation partitioning. Environmental covariates were rescaled between 0 and 1. The vertical axes indicate the relative contributions of each environmental variable, at each zeta order. Geographic distance is most important at low zeta orders, which are dominated by rare species, and as zeta order increases (increasing the importance of common species), canopy openness and then infestation rate become increasingly more important, with most of the compositional change occurring in the first 20% of change in those two covariates. Overall when zeta order > 4, environmental covariates explain more compositional change than distance.

meaning that changes in community composition occur along the full range of distance, as rare species turnover from site to site and county to county. In contrast, for canopy openness, most compositional turnover occurs in the first 20% of its range, in the transition from closed to partially open-canopy forest, and for DBH, most change occurs in the last 20% of its range, in the transition to sites with the largest trees. MS-GDM with low-read-depth samples removed in Fig. S6.

By definition, as zeta order rises, common species increasingly dominate the analysis, and starting at zeta order 4 (Fig. 3A, Order4), the distance variable starts to be less important than the local variables of infestation rate and canopy openness, which both exert their effects primarily in the first $\sim 20\%$ of their ranges. That is, most of the compositional change occurs in the transition from 0% to infested-0%-open-canopy forests to 20%-infested-20%-open-canopy forests. At higher zeta orders, distance explains even less of the change in composition, except at very large distances, since common species are by definition more widespread.

We then re-ran the MS-GDM with elevation included, which returned similar results: as zeta order increases, the five environmental covariates other than elevation (Table 1) explain an increasingly larger proportion of total variation, while distance and elevation become less important (Fig. S7).

3.4. Zeta diversity decline and retention ratios

Another application of zeta diversity is to infer the relative roles of niche partitioning and stochastic assembly in community assembly (McGeoch et al., 2019). Zeta diversity declines as zeta order increases, since fewer and fewer species are shared amongst more and more sites. Steeper rates of decline indicate greater numbers of rarer species over more common species. Here, we asked how infestation affects community assembly over the infestation-rate gradient. To simplify the comparison, we divided the sites roughly evenly into 'low' (\leq 0.25, n = 14) and 'high' (>0.25, n = 16) infestation-rate categories (Figs. 4 and S8), and tested the goodness of zeta-diversity decline functional forms using

the Akaike information criterion (AIC).

In the low-infestation forests, zeta-diversity decline is both steeper and better fit by an exponential function than by a power-law function ($-7.54_{AIC_exp} < 10.67_{AIC_pl}$) (Fig. 4A). This is consistent with low-infestation forests being characterized by a stochastic assembly process. In the extreme form, there is no niche partitioning; all the species have equal probability of occurring at any given site despite environmental variation across sites, and across-species variation in occupancy and turnover arises only stochastically, due to, for instance, random dispersal governing establishment (Hui & McGeoch, 2014; McGeoch et al., 2019). Consistent with this, the zeta-ratio analysis shows fewer common species and generally low retention of species when a new site is sampled (McGeoch et al., 2019, Fig. 4B).

In contrast, in the high-infestation forests, zeta-diversity decline is relatively shallower and better fit by power-law function ($8.52_{AIC_exp} > 0.80_{AIC_pl}$) (Fig. 4A), which obtains when the probability that a species occurs in a newly sampled site increases with that species' overall occupancy, which in turn is consistent with community assembly being driven by niche differentiation. Each OTU has a species-specific probability of occurring at a site due to environmental conditions at that site (Hui & McGeoch, 2014; McGeoch et al., 2019). The zeta-ratio analysis shows that species in the high-infestation forests generally have higher occupancy, even beyond order 6 (Fig. 4B), which is the number of sites per county (Fig. 2) and which thus shows that high-infestation forests share species across large geographic distances, apparently because of shared environmental conditions (Fig. 3A). (See Fig. S9 for the same analysis with low-read-depth samples removed).

3.5. Alpha diversity

The iNEXT and iNextPD analyses found no evidence for a difference in species or phylogenetic diversity between low- and high-infestation forests (Fig. 5, iNEXT analyses with low-read-depth samples removed in Fig. S10).

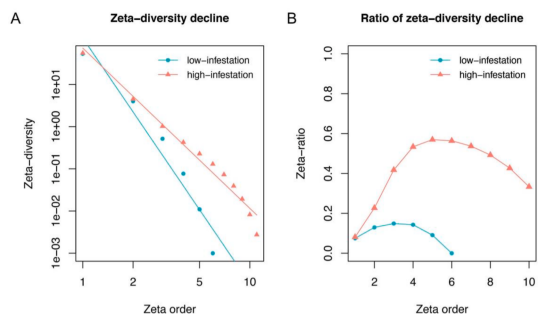


Fig. 4. Community assembly mechanism. A. Comparison of zeta diversity decline and B. retention rate between low- and high-infestation forests. Zeta orders 1 to 11 are shown, as zeta diversity equals zero for orders > 11. High-infestation sites A. are characterized by a power-law decline and B. share more common species, consistent with a niche-differentiated community. Low-infestation sites A. are characterized by an exponential zeta decline and B. share fewer common species, consistent with a stochastic community-assembly process.

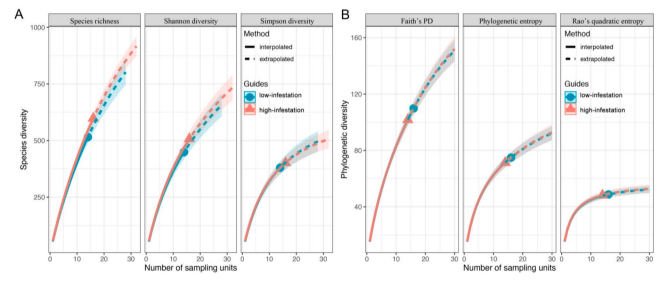


Fig. 5. Alpha diversity analysis by A. iNEXT and B. iNextPD. Sample-size-based rarefaction (solid lines) and extrapolation (dashed lines) sampling curves for three measures of A. species diversity and B. phylogenetic diversity in low-infestation and high-infestation forests. Shaded areas represent 95% confidence intervals. Symbols indicate sample size per forest type. Overlapping confidence intervals indicate no evidence for difference between forest types.

4. Discussion

Metabarcoding of Malaise-trapped insects reveals that bark-beetle induced forest dieback does not result in detectable differences in species richness or phylogenetic diversity (Fig. 5) but does result in compositional change (Figs. 2 and 3). For rarer species, MS-GDM and boral ordination explain this turnover with distance and elevation (Figs. 2 and 3), which are correlated in our sampling design (Fig. 1). For more common species, local-forest environmental variables explain relatively more of the compositional differences, with the biggest compositional change occurring between 0% and infested-0%-opencanopy forests and 20%-infested-20%-open-canopy forests (Figs. 3 and S7). Bark-beetle dieback thus appears to affect the larger insect

community (at least the portion that can be sampled by Malaise traps) by mimicking the transition between closed-canopy forest and structurally heterogeneous forest. That said, at higher zeta orders, just over half of compositional variation across sites remains unexplained (Fig. 3).

The zeta-decline analysis found that low-infestation sites showed evidence of stochastic assembly, while high-infestation forest sites showed evidence of niche partitioning as the dominant community assembly mechanism (Fig. 4). This suggests that the species which colonize the higher-infestation-rate (and higher-energy-availability) sites are adapted to these conditions.

Interestingly, Müller et al. (2010) have also reported that saproxylic beetle composition changes nonlinearly with canopy openness (measured as LiDAR penetration), with rapid compositional change

occurring from closed canopy up to 23% penetration (11–49% 95% CI), after which composition changes slowly. Our results are thus remarkably similar (Figs. 3 and S7), despite differences in geography, dominant tree species, and focal taxa, and we speculate that the driving mechanism is the effect of light availability on understorey vegetation and microclimate. Seibold et al. (2016a) have also reported that canopy openness is a major driver of species assemblage composition of nonsaproxylic epigeal arthropods after intensive logging (see also Bishop et al., 2009; Bouget et al., 2013; Franc et al., 2007). Unfortunately, unlike Müller et al. (2010), we were unable to measure the biodiversity effects of deadwood volume (and thus, the effect of deadwood removal), given that, to our knowledge, there are no Pinus yunnanensis sites in Yunnan with high amounts of deadwood to contrast with lowdeadwood-volume sites. If possible, there would be value in running deadwood enrichment experiments in Yunnan, in order to test the prediction that saproxylic animal and fungal species will benefit (Doerfler et al., 2018; Seibold et al., 2015, 2018). With that important omission, our results in Yunnan seem consistent with Thorn et al.'s (2020) diagnosis of biodiversity decline in European forests, which they attribute to the loss of tree species diversity and the loss of age and structural heterogeneity, which together provide microhabitats for light-demanding plant and insect species. Species, age, and structural heterogeneity also likely contribute to resilience against large-scale bark-beetle outbreaks (Seidl et al., 2016). On the other hand, Trzcinski & Reid (2008) argue that deadwood removal could be effective at preventing the longdistance spread of bark-beetle outbreaks.

Given China's recent announcement that afforestation and reforestation efforts should now aim to create heterogeneous forests that are higher in biodiversity and more resilient to disease and pests, what is the best way to achieve this? Part of the solution is to allow natural disturbances to create forest structural and age heterogeneity, which in turn will benefit light-demanding plants and animals and also provide deadwood volume for saproxylic taxa (Thorn et al., 2020). These natural disturbances include windstorms, bark-beetle outbreaks, and droughtinduced diebacks, as long as dead trees are not subsequently removed and open areas not replanted with plantation trees (Thorn et al., 2020). In particular, bark beetles can be seen as a keystone species (Müller et al., 2008), with their attacks on weak and old trees accelerating the succession of monoculture forests into heterogeneous forests (Cai et al., 2006; Yue et al., 2011). However, extreme climate events are predicted to increase, resulting in a greater rate and severity of natural disturbances (Allen et al., 2010, 2015; Thom et al., 2017; Thom & Seidl, 2016), including an expansion of bark beetles to higher latitudes and elevations (Bentz et al., 2010; Hlásny et al., 2011), which raises the short-term costs of allowing bark beetle outbreaks to proceed unimpeded. Thus, in many areas, anthropogenic treatments could be implemented to accelerate the succession of plantation forests into heterogeneous forests (Baeten et al., 2019; Felipe-Lucia et al., 2018; Schall et al., 2018; Yue et al., 2011), especially given that Pinus yunnanensis covers large areas of poor soil, where tree growth is generally slow and seed sources of other tree species distant.

Methodological considerations. – The combination of metabarcoding and Malaise traps, which preferentially capture species-rich Hymenoptera and Diptera, naturally produces datasets with a large proportion of low-prevalence species. In consequence, we used zeta diversity and MS-GDM to analyse community subsets of increasingly more common species, which showed that local environmental covariates were more important for explaining species distributions at higher orders (Fig. 3). Zeta-decline analysis showed that high-infestation and low-infestation sites differed in their community assembly mechanisms (Fig. 4). In short, removal of the lowest-prevalence species made clearer the effects of forest structure on community composition. In contrast, we failed to find any differences in alpha diversity across low- and high-infestation forests, even for the measures that clearly reached an asymptote: Simpson diversity, Phylogenetic entropy, and Rao's quadratic entropy (Fig. 5). That said, our study is underpowered for comparing alpha

diversities, and we draw only a tentative conclusion on this front. Finally, to test robustness, we reran all analyses after removing the eight lowest-read-depth samples, which removed the correlation between read-depth and species richness, and we recovered the same results (S4, S6, S9, S10).

Another aspect of metabarcoding is that it can be applied to samples from locations where taxonomic coverage is poor, such as arthropods from Southwest China. The resulting OTU dataset, identified only to higher taxonomic ranks, can be used to visualise biodiversity patterns. However, with limited taxonomic information, we are unable to carry out functional (trait-based) analyses to try to explain why particular taxa are favoured or disfavoured under different silvicultural regimes (e.g. Cours et al., 2020; Thorn et al., 2018). We also note that our dataset represents only a single time point, while temporal turnover of forest arthropod communities appears to be high (Barsoum et al., 2019). However, we have shown elsewhere (Zhang et al., 2016) that metabarcoding sample sets taken in rainy season and in dry season are equally able to differentiate forest disturbance gradients, and we have shown in two large studies that Malaise-trapped invertebrates show similar responses to several other methods and taxa in their responses to forest structure and disturbance (Ji et al., 2013; Edwards et al., 2014). Thorn et al. (2018) also found taxonomic congruence in biodiversity response to salvage logging. In the future, one partial way around the lack of taxonomic information for functional inference is to apply joint species distribution modelling to DNA-based time series datasets to infer the relative contributions of environmental covariates and species interactions to changes in species abundances (Abrego et al., 2021).

Conclusion. – Long-term monitoring will be necessary for tracking the biodiversity consequences of conversion from simple to heterogeneous forests and for comparing different anthropogenic treatments. Studies in China, the UK, and Borneo have shown that DNA metabarcoding is an efficient and standardizable tool for measuring how animal biodiversity in forests varies as a function of management and inherent condition (Barsoum et al., 2019; Edwards et al., 2014; Hua et al., 2016; Ji et al., 2013, 2020; Wang et al., 2019; Yang et al., 2014; Yang et al., 2016; Zhang et al., 2016). We also think that there is considerable scope for using remotely sensed measures (multispectral and LiDAR) to efficiently generate environmental covariates for the large-scale mapping and monitoring of pest outbreaks like bark beetles in particular (Ji et al., 2007; Wang et al., 2015) and terrestrial biodiversity in general (Bush et al., 2017).

CRediT authorship contribution statement: D.W.Y., C.LV. designed the study. L.L. and C.LV. designed the environmental covariate measurements. C.Y.Y. and W.C. collected samples and made the environmental covariate measurements. W.C., C.Y.W. and C.Y.Y. performed the molecular experiments. W.C. and X.Y.W. performed the bioinformatic and statistical analyses. D.W.Y., W.C. and Q.Z.W. contributed to the manuscript.

Declaration of Competing Interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: D. W.Y. is a co-founder of NatureMetrics (www.naturemetrics.co.uk), which provides commercial metabarcoding services.

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Data accessibility statement

Sequence data are under GenBank accession number PRJNA668449. Bioinformatic scripts, R scripts are available at https://github.com/CaiWang0503/climtree China.

Appendix A. Supplementary material

Supplementary data to this article can be found online at https://doi.org/10.1016/j.foreco.2021.119173.

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