1 Multi-dimensionality of tree communities structure host-parasitoid networks and their

2 phylogenetic composition

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38 Data availability

- 39 Data is available on Science Data Bank at https://www.scidb.cn/en/s/Ujq22u, and will be available
- 40 on the BEF-China project database at https://data.botanik.uni-halle.de/bef-china/datasets.

41 Conflict of interest statement

42 The authors have no conflict of interest.

43 Author contributions

- 44 C.D.Z., A.L. and M.Q.W. conceived the idea for the manuscript; C.D.Z., A.M.K., S.K.G., P.F.G.
- 45 and M.Q.W. designed the study. S.K.G., P.F.G., J.J.Y., G.A.C., Z.Q.N., M.S., J.T.C., Y.L., Q.S.Z.,
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- 50

51 Abstract

52 Environmental factors can influence ecological networks, but these effects are poorly understood 53 in the realm of the phylogeny of host-parasitoid interactions. Especially, we lack a comprehensive 54 understanding of the ways that biotic factors, including plant diversity, tree identity, genetic 55 diversity, overall community composition of higher trophic levels, and abiotic factors such as 56 microclimate, interact to determine host-parasitoid network structure and host-parasitoid 57 community dynamics. To address this, we leveraged a five-year dataset of trap-nesting bees and 58 wasps and their parasitoids collected in a highly-controlled, large-scale subtropical tree 59 biodiversity experiment. We tested for effects of tree species richness, tree phylogenetic and 60 functional diversity, and taxonomic and phylogenetic composition on taxonomic, phylogenetic, 61 and network composition of both host and parasitoid communities. We show that multiple 62 components of tree diversity, tree composition, and canopy cover impacted both, taxonomic and 63 phylogenetic composition of hosts and parasitoids. Generally, top-down control was stronger than 64 bottom-up control via phylogenetic association between hosts and parasitoids, reflecting 65 non-randomly structured interactions between phylogenetic trees of hosts and parasitoids. Further, 66 host-parasitoid network structure was influenced by tree species richness, tree phylogenetic 67 diversity, and canopy cover. Our study indicates that the composition of higher trophic levels and 68 corresponding interaction networks are determined by habitat structure and heterogeneity, which 69 is maintained by trees and especially via phylogenetic links in species-rich ecosystems.

70 Introduction

71 Understanding the ecological consequences of biodiversity loss is an increasingly important task

72 in ecology, given the ongoing biodiversity crisis (Isbell et al. 2022). Representing the

73 interdependencies among organisms, ecological networks reflect whether and how species interact

74 with each other across trophic levels, playing an indispensable role in assessing ecosystem

stability and integrity (De Ruiter et al. 1995, Harvey et al. 2017). Changes in network structure

- visually coincide with variations in diversity and community of trophic levels, which could be in
- turn affected by the changes in producers via trophic cascades (Barnes et al. 2018, Gonzalez et al.

78 2020). However, we still lack a generalizable framework for how these networks and especially 79 their phylogenetic interdependencies respond to basal changes of biodiversity loss in ecosystems, 80 such as changes in the tree diversity of forests (Tylianakis et al. 2008, Grossman et al. 2018). To 81 better understand species codependence and its role for biodiversity conservation, we must further 82 study the complex dynamics of networks from multiple angles (Tittensor et al. 2014, Brondizio et 83 al. 2019). Interactions can be viewed from a top-down or bottom-up perspective. Higher trophic 84 levels can impact lower trophic levels through antagonistic species interactions, while lower 85 trophic levels can provide resources to support higher trophic levels, Previous studies have shown 86 there might be asymmetric effects between top-down and bottom-up control by different 87 mediators (Vidal and Murphy 2018), with both factors shaping multitrophic communities 88 together(Hunter et al. 1992). Moreover, the diversity and community of higher trophic levels 89 could be driven by environmental microclimate, which could be determined by plant structuring, 90 such as canopy cover (Fornoff et al. 2021, Perlík et al. 2023). Understanding how these forces 91 respond to changing biodiversity is therefore imperative to predict how ongoing environmental 92 changes will impact the functioning and stability of ecosystems (Hines et al. 2019).

93 A crucial network that unites bottom-up and top-down processes in many ecosystems and is 94 prone to strong alterations due to environmental change is the interaction network between 95 parasitoids and their hosts (Tylianakis et al. 2006, Jeffs and Lewis 2013). Insect parasitoids attack 96 and feed on and eventually kill their insect hosts (Godfray and Godfray 1994). Parasitoids are 97 thought to be particularly sensitive to environmental changes, because species in higher trophic 98 levels usually have smaller population sizes and the potential specialization of their hosts may 99 cascade up to impact the parasitoids' presence or absence. Therefore, insect host-parasitoid 100 systems are ideal for studying the relationships between habitat-level changes and species 101 interactions (Jeffs and Lewis 2013). Previous studies on host-parasitoid interactions mainly 102 focused on the influence of abiotic factors, such as elevation and habitat structure (e.g. Valladares 103 et al. 2012, Maunsell et al. 2015, Grass et al. 2018), or sometimes interaction structure (e.g. 104 Cagnolo et al. 2011). However, the role of multiple components of plant diversity in modifying 105 host-parasitoid interaction networks, as key biotic determinants of overall ecosystem structure,

106 remains relatively little explored. Recent studies mainly focused on basic diversity associations 107 (Ebeling et al. 2012, Schuldt et al. 2019, Guo et al. 2021). These studies have demonstrated both 108 direct and indirect effects of plant diversity on both hosts and parasitoids, possibly via increased 109 niche space and resource availability (Guo et al. 2021). Nevertheless, how these patterns 110 propagate to their interaction networks is still unclear. Moreover, the effects of changing plant 111 diversity are not always obvious when looking at plant species richness, and other diversity 112 components such as plant phylogenetic diversity have been shown to better predict 113 diversity-dependent bottom-up effects on host-parasitoid networks (e.g. Staab et al. 2016). It is 114 especially important to take phylogenetic dependences into account when it comes to the 115 phylogenetic structure within and across trophic levels (Webb et al. 2002, Emerson and Gillespie 116 2008). This makes it vital to account for multiple dimensions of biodiversity and relevant trophic 117 interactions (e.g. taxonomic, phylogenetic, functional; Peralta et al. 2015, Volf et al. 2017, Wang 118 et al. 2020). These components might jointly affect host-parasitoid networks in a system with high 119 species diversity. Forests have garnered special attention lately, because they represent complex 120 and large ecosystems susceptible to global change (add citation here). Understanding how such 121 relationships modulate the effects of tree diversity loss on the structure and interaction strength of 122 host-parasitoid networks clearly requires further study (Staab et al. 2016, Fornoff et al. 2019). 123 Here, we leverage standardized trap nests for solitary cavity-nesting bees, wasps, and their 124 parasitoids in a large-scale subtropical forest biodiversity experiment to test how multiple 125 dimensions of tree diversity and community composition influence host-parasitoid network 126 structure. A multi-faceted approach is particularly important when considering that associations 127 between trophic levels might be non-random and phylogenetically structured (Volf et al. 2018, 128 Wang et al. 2020). We aimed to quantify the strength and stability of associations between hosts 129 and parasitoids to both discern the primary components of the diversity and composition of tree 130 communities that affect higher trophic level interactions We expected that (a) multiple tree community metrics, such as taxonomic, phylogenetic and functional diversity and phylogenetic 131 132 and species community composition can structure host and parasitoid community compositions, 133 especially via phylogenetic processes, as species interactions often show phylogenetic

134 conservatism (e.g. Pellissier et al. 2013, Peralta et al. 2015). Further, we hypothesized that (b)

135 host-parasitoid networks will be more complex and stable with increasing tree species richness

due to potential links from higher richness of hosts and parasitoids promoted by more tree species,

137 and (c) both, community and interaction network changes can also be related to abiotic factors,

138 such as canopy cover, which might play a role in structuring hymenopteran communities (Haddad

- tal. 2011, Fornoff et al. 2021). By better understanding these dynamics, we can begin building a
- 140 generalized framework for understanding host-parasitoid interactions in forest ecosystems.

141 **Results**

148

142 Overall, 34,398 brood cells were collected from 13,267 tubes across five years of sampling

143 (2015, 2016, 2018, 2019, and 2020). Six families of hosts and seventeen families of

parasitoids were identified. Among them, we found 56 host species (12 bees and 44 wasps)

and 50 parasitoid species (38 Hymenoptera and 12 Diptera). The full species list and their

Host species composition was significantly related to the species and phylogenetic

abundances are given in Table S1.

147 Community composition of hosts and parasitoids

149 composition of the tree and parasitoid communities (NMDS scores), as well as to canopy 150 cover, tree MPD, elevation, and eastness (Fig. 1a, Table 1, Table S2). Parasitoid species 151 composition was significantly associated with host phylogenetic diversity, tree functional 152 diversity, tree MPD, eastness, and elevation, and was significantly related to tree species 153 composition, host species composition, and canopy cover (Fig. 1b, Table 1, Table S3). Host 154 phylogenetic composition was affected by tree species composition, tree MPD, tree functional 155 diversity, canopy cover, eastness, elevation and was especially obviously affected by 156 parasitoid species and phylogenetic composition (Fig. 1c, Table 1, Table S4). For parasitoid 157 phylogenetic composition, significant relationships were found with tree species and 158 phylogenetic composition, host species composition, tree functional diversity, canopy cover, 159 elevation, and eastness (Fig. 1d, Table 1, Table S5).

- 160 An effect of host composition on the composition of the parasitoid communities was
- 161 further indicated by a significant parafit test (p = 0.032), suggesting nonrandom associations
- 162 in the phylogenetic structure of parasitoid and host communities (Fig. 2, Fig. S3).
- 163 Host-parasitoid network associations

164 The linear regression model results showed that host vulnerability, linkage density, and

- 165 robustness of parasitoids were significantly negatively related to tree species richness, while
- 166 remaining unaffected by other environmental covariables (Table 2, Fig. 3; except for
- 167 elevation, which was marginally significantly related to robustness). Interaction evenness was
- 168 significantly negatively associated with canopy cover, and interaction evenness was also
- negatively related to eastness (Fig. 4c; Table 2).. Generality was only marginally associated
- 170 with canopy cover, and was not related to tree species richness or the other environmental
- 171 factors. In the alternative models (tree species richness replaced by tree MPD), vulnerability
- and linkage density were significantly positively related to tree MPD (Fig. 4a, 4b; Table S6),
- 173 while parasitoid robustness was negatively related to tree MPD (Fig. S4, Table S6). The
- 174 results of other network metrics (generality and interaction evenness) were consistent with
- those of the primary models
- 176 The results of the null model analysis suggested that our metrics calculated by the
- 177 observed network were significantly different from a random distribution (72, 71, and 77 out
- 178 of 85 values for generality, vulnerability, and linkage density, respectively; all values for
- 179 robustness, and interaction evenness), strongly demonstrating that interactions between
- 180 species were not driven by random processes.
- 181

182 Discussion

Our study demonstrates that tree species richness and phylogenetic diversity play key roles in
modulating communities of hosts and parasitoids and their interactions with each other. These
interactions are further structured by the phylogenetic associations between hosts and

parasitoids. Moreover, canopy cover partly determined host-parasitoid networks, supporting a recent finding that the structure of host-parasitoid networks is also mediated by changes in microclimate to some extent (Fornoff et al. 2021). These patterns were highly associated with multiple tree diversity metrics (tree taxonomic, phylogenetic and functional diversity), and compositional changes which are key to understanding how host-parasitoid interactions may be impacted by biodiversity loss through trait- and phylogeny-based processes.

192 Community composition associations

193 Host species composition was influenced by several factors, including the taxonomic and

194 phylogenetic composition of trees and parasitoids, tree diversity (species richness and MPD),

and other biotic factors. The effects of tree diversity and composition on host species

196 composition agree with previous studies where solitary bee and wasp species composition

197 were related to plant community structure (e.g. Loyola and Martins 2008). It seems likely that

198 these results are based on bee linkages to pollen resources and predatory wasp linkages to the

diverse of food sources, which may themselves be closely linked to tree species richness

200 (Reitalu et al. 2019, Staab and Schuldt 2020).

201 We also found that tree MPD, FD, and species composition affect parasitoid species 202 composition, as many studies have found significant relationships between plant and 203 parasitoid diversity in forests, including tree phylogenetic diversity (Staab et al. 2016), 204 functional diversity (Rodriguez et al. 2019), and structural diversity (Schuldt et al. 2019). 205 Similar to predators, parasitoids might also be more active or efficient with increasing tree 206 community dissimilarity due to higher prey resources or lower intraguild parasitism caused by more diverse habitats (Finke and Denno 2002). On the other hand, our results also show that 207 208 host species composition and parasitoid species composition relate to each other and their 209 phylogenetic compositions, which are structured by tree communities to some extent. This 210 pattern could propagate to the adjacent two higher trophic interactions through both top-down 211 and bottom-up control.

212 Both host and parasitoid phylogenetic composition were related to tree species 213 composition. This pattern has important implications for cascading effects among trophic 214 levels, in that producer communities could structure a higher trophic level community via an 215 intermediate trophic level. As previous studies usually found weak effects of plants on higher 216 trophic levels (e.g. Cappelli et al. 2022). However, while both host and parasitoid 217 phylogenetic composition was related to tree MPD, only parasitoids responded to tree 218 phylogenetic composition. This may be because there are many caterpillar-hunting wasps in 219 our host communities, and the community composition of caterpillars were usually correlated 220 with tree phylogenetic communities(Wang et al. 2019). Therefore, the prev highly associated 221 with tree phylogenetic composition (e.g. caterpillars) might indirectly determine predatory 222 wasp (host) phylogenetic composition, as recently found for plants-caterpillars-spiders (Chen 223 et al. 2023). This could be further tested by collecting the food directly used by the wasps 224 (caterpillars). For parasitoids, tree phylogenetic composition might drive the process of 225 community assembly through trophic cascades (e.g. from plants to parasitoids via herbivores 226 and host wasps) (Webb et al. 2002, Cavender-Bares et al. 2009). Additionally, parasitoid 227 phylogenetic composition can be influenced by tree structural diversity (e.g. host availability 228 in plots with higher heterogeneity; Schuldt et al. 2019), which can be determined by 229 conserved traits across phylogenies (Webb et al. 2002). The phylogenetic associations 230 between hosts and parasitoids exhibited a nonrandom structure (significant *parafit* correlation) 231 between the phylogenetic trees of the host and their parasitoids (see also Peralta et al. 2015). 232 Interestingly, we found that only host phylogenetic composition was affected by parasitoid 233 phylogenetic composition (and not vice-versa). This asymmetry suggests that top-down 234 control (parasitoids to hosts) was stronger than bottom-up control (hosts to parasitoids), 235 supporting prior hypotheses, demonstrating the strong control of parasitoids (Vidal and 236 Murphy 2018). This pattern might reflect disproportionate interactions between hosts and 237 parasitoids, with one trophic level more strongly influencing the other in some instances, but 238 further study is needed.

Moreover, the species and phylogenetic composition of hosts and parasitoids was also related to abiotic factors, especially to canopy cover, which has been considered especially important (Sobek et al. 2009, Fornoff et al. 2021). In future studies, it will be useful to incorporate other, more direct metrics of microclimate, such as local temperature and humidity, to determine the proximal drivers of these microclimatic effects (Ma et al. 2010, Fornoff et al. 2021).

245

246 Community-level host-parasitoid networks

247 Tree community species richness did not significantly influence the diversity of hosts targeted 248 by parasitoids (generality), but caused a significant increase in the diversity of parasitoids 249 per host species (vulnerability). This is likely because niche differentiation often influences 250 network specialization via potential higher resource diversity in plots with higher tree 251 diversity (Lopez-Carretero et al. 2014). For the significant relationship between vulnerability 252 and tree species richness, a potential explanation is that host-parasitoid interactions could be 253 driven through bottom-up effects. Moreover, higher trophic levels will particularly benefit 254 from tree diversity, inducing parasitoid species increases more than host diversity with 255 increasing tree species richness (Guo et al. 2021). These will further increase vulnerability at 256 community level. According to the enemies hypothesis (Root 1973), which posits a positive 257 effects of plant richness on natural enemies, the higher trophic levels in our study (e.g. 258 predators and parasitoids) would benefit from tree diversity and regulate herbivores thereby 259 (Staab and Schuldt 2020). Indeed, previous studies at the same site found that bee parasitoid 260 richness and abundance were positively related to tree species richness, but not their bee hosts 261 (Fornoff et al. 2021, Guo et al. 2021). Because our dataset considered all hosts and reflects an 262 overall pattern of host-parasitoid interactions, the effects of tree species richness on generality 263 might be more complex and difficult to predict, as we found that neither tree species richness 264 nor tree MPD were related to generality. Thus, our results again indicate that top-down

265 control imposed by higher trophic levels is more significant than bottom-up control with an266 increasing number of plant species.

267 Linkage density was positively related to tree species richness, supporting the food web 268 theory, which predicts in our case that network complexity (linkage density) depends on the 269 number of plants (Melián and Bascompte 2002). Although trees were not directly included as 270 a trophic level in our networks, potential network complexity increased with tree species 271 richness, likely enabling higher network stability/resistance (Ebeling et al. 2011). For 272 example, a network might be more sensitive to extinctions because of key species loss due to 273 lower linkage density and lower redundancy (e.g. Naeem and Li 1997). However, parasitoid 274 robustness was unexpectedly negatively related to tree species richness. It was expected that 275 higher trophic levels would be more robust, less influenced by perturbations from lower 276 trophic levels, when plant diversity is higher, as more potential interactions at lower trophic 277 levels should theoretically increase redundancy and resilience of connected higher levels 278 (Blüthgen and Klein 2011, Fornoff et al. 2019). Dilution effects may explain this, as plots 279 with higher richness held fewer individuals of a given tree species. If there are strong prey 280 item (caterpillars, grasshoppers, etc.) preferences for one species, there may be fewer or they 281 may be more densely aggregated and less likely to be encountered by parasitoids. This 282 increased stochasticity in parasitoid wasps could benefit hosts by reducing parasitism pressure 283 overall, weakening top-down controls.

284 Similar to tree species richness, tree MPD was also positively correlated with 285 vulnerability and linkage density, meaning that the mean number of parasitoids per host 286 species and number of links within the host-parasitoid system can also be promoted by tree 287 MPD, in agreement with several recent studies (Pellissier et al. 2013, Staab et al. 2020, Wang 288 et al. 2020). Our results suggest that the specialization and complexity of higher trophic levels 289 can also be affected by plant phylogenetic diversity. This pattern can be traced to the effects 290 of habitat heterogeneity caused by tree species richness and MPD on higher trophic levels via 291 bottom-up control. The effects of tree MPD were consistent with effects of tree species 292 richness on robustness of parasitoids to host loss. This result suggests that higher trophic

levels are sensitive to changes in both plant phylogenetic relatedness and general species
dissimilarity via trophic interactions, even the hosts are not all directly interacting with plants,
bees excluded. Therefore, it may be that stronger linkages would be found when exclusively
exploring such plant-herbivore-parasitoid systems.

297 Interaction evenness was significantly negatively related to canopy cover, further 298 reinforcing an important role of microclimate (likely temperature and humidity; (Sobek et al. 299 2009, Fornoff et al. 2021). Our results agree with a previous study on ants, where plant-insect 300 interactions were more even with more open canopies (Dáttilo and Dyer 2014). In our case, 301 canopy cover might change hymenopteran species evenness and then further influence 302 interaction evenness, certain host species tended to nest in plots with higher canopy cover, 303 which might decrease the interaction evenness by favoring parasitoids of fewer, more 304 dominant hosts. This pattern would become more significant when more host and parasitoid 305 species are in a plot, given the positive relationship between higher trophic level diversity and 306 canopy cover.

307 Future prospects

308 Overall, our study enables new insights into the dynamics of host-parasitoid interactions 309 under varying environmental conditions, an important step toward building a synthetic model 310 for such biodiversity. A key finding was that although parasitoids and hosts respond to tree 311 species richness, top-down control seems predominant in parasite-host interaction, though 312 whether this holds for others antagonistic interactions requires further investigation. 313 Different trophic levels and functional groups of species responded differently to lower level 314 changes (Fornoff et al. 2021, Guo et al. 2021). This highlights the complexities of building 315 such models and calls for more studies across habitat types and taxa, to test the generality of 316 our findings. Future studies should also consider the role of host/parasitoid functional traits, 317 because they might play a critical role in modifying network structures and ecosystem 318 functioning.

319

320 Materials and methods

321 Study sites design

- 322 This study was conducted in the BEF-China biodiversity experiment, which is the largest tree
- 323 diversity experiment worldwide. The experiment is located in a subtropical forest near
- 324 Xingangshan, Jiangxi province, south-east China (29°08′–29°11′N, 117°90′–117°93′E). The mean
- annual temperature is 16.7°C and mean annual precipitation 1821 mm (Yang et al. 2013). The
- 326 experiment includes two study sites (Site A and Site B), 4 km apart from each other, that were
- 327 established in 2009 (Site A) and 2010 (Site B) respectively. A total of 566 plots (25.8×25.8 m)
- were designed on the two sites, and per plot 400 trees were initially planted in 20 rows and 20
- 329 columns with a planting distance of 1.29 m. A tree species richness gradient (1, 2, 4, 8, 16 and 24
- 330 species) was established at each site, based on a species pool of 40 local, broadleaved tree species
- **331** (Bruelheide et al. 2014).

332 For our study, at both sites (site A and site B) eight plots of each tree species richness
333 level (1, 2, 4, 8) were randomly selected, as well as six and two plots of 16 and 24 mixtures. In

addition, at site B eight additional monocultures were sampled (Fornoff et al. 2021), resulting 48

335 plots in Site B (including 16 monocultures, eight plots for each 2, 4, 8 mixtures and six and two

336plots of 16 and 24 mixtures. In total, 88 study plots were used (40 plots on Site A and 48 plots on

337 Site B, see Fig. S1).

338 Sampling

339 We collected trap nests monthly to sample solitary bees and wasps (Staab et al. 2018) in the 88

340 plots from September to November in 2015 and April to November in 2016, 2018, 2019 and 2020.

- 341 For each plot, we installed two poles with trap nests (11 m apart from each other and 9 m away
- 342 from the nearest adjacent plots) along a SW–NE diagonal centrally per plot (following the design
- of Ebeling et al. 2012). Each pole stood 1.5 m above ground, and each trap nest consisted of two
- 344 PVC tubes (length: 22 cm \times diameter: 12.5 cm) filled with 75 \pm 9 (SD) reed internodes of 20 cm
- length and diameters varying between 0.1 and 2.0 cm (Staab et al. 2014, Fornoff et al. 2021).

346 Every month, we sampled the reeds with nesting hymenopterans and replaced them with

347 internodes of the same diameter. All the samples were reared in glass test tubes under ambient

- 348 conditions until specimens hatched. We identified hatched hosts and parasitoids to species or
- 349 morphospecies (Supplementary Table S1) based on reference specimens (vouchered at the
- 350 Institute of Zoology, CAS, Beijing). We were interested in the general patterns of host-parasitoid
- interactions at the community level, so for the analysis we did not distinguish between the two
- 352 life-history strategies of parasitoids (true parasitoids and kleptoparasitoids, including
- 353 hymenopteran and dipteran parasitoids) because they both have the same ecological result, death
- of host brood cells.

355 DNA extraction and amplification

- 356 All specimens were sequenced for a region of the mitochondrial cytochrome c oxidase subunit I
- 357 (COI) gene (Hebert et al. 2003). We extracted whole-genomic DNA of hosts and parasitoids using
- 358 DNeasy Blood & Tissue Kits (QIAGEN GmbH, Hilden, Germany), following the manufacturer's
- 359 protocols. COI sequences of samples were amplified using universal primer pairs, LCO1490

360 (GGTCAACAAATCATAAAGATATTGG) as the forward primer and HCO2198

- **361** (TAAACTTCAGGGTGACCAAAAAATCA) or HCOout
- **362** (CCAGGTAAAATTAAAATATAAACTTC) as the reverse primer. We carried out polymerase
- 363 chain reactions (PCR) in 96-well plates with 30 μl reactions containing 10 μl ddH2O, 15 μl
- **364** Premix PrimeSTAR HS (TaKaRa), 1 ul of each primer at 10 μM, and 3 μl template genomic DNA
- using a thermo cycling profile. The PCR procedure as follows: 94°C for 1 min; 94°C for 1 min, 45°C
- for 1.5 min and 72°C for 1.5 min, cycle for 5 times; 94°C for 2 min, 58°C for 1.5 min and 72°C for
- 367 1 min, cycle for 36 times; 72°C for 5 min. We performed all PCRs on an Eppendorf Mastercycler
- 368 gradient, which were then visualized on a 1% agarose gel. Samples with clean single bands were
- 369 sequenced after PCR purification using BigDye v3.1 on an ABI 3730xl DNA Analyser (Applied
- Biosystems).

371 Sequence alignment and phylogenetic analysis

372 We applied MAFFT (Misawa, Katoh, Kuma, & Miyata, 2002) to align all sequences, then translated the nucleotides into amino acids via MEGA v7.0 (Kumar, Stecher, & Tamura, 2016) to 373 374 check for the presence of stop codons with manual adjustments. Host and parasitoid sequences 375 were then aligned against the references using a Perl-based DNA barcode aligner (Chesters, 376 2019). 377 We employed two strategies to improve the phylogenetic structure of a DNA barcode 378 phylogeny, which demonstrably improve resulting phylogeny-based diversity indices 379 (Macías-Hernández et al. 2020). These include the integration of 1) molecular sequences of the 380 plot data and 2) phylogenetic relationships from other molecular datasets. Integration was 381 achieved following Wang et al. (2020) and Chesters (2020): reference DNA barcodes of 382 Hymenoptera and Diptera were downloaded from the BOLD API 383 (www.boldsystems.org/index.php/API_Public), which were variously processed (e.g. to retain 384 only fully taxonomically labelled barcodes, to remove low quality or mislabeled entries, and to 385 dereplicate to a single exemplar per species), and then aligned (Chesters 2019). A single outgroup 386 was included for which we selected the most appropriate insect order sister to Diptera and 387 Hymenoptera (Misof et al. 2014), a representative of the order Psocoptera (Psocidae, Psocus 388 *leidvi*). We then constructed a phylogeny of the references and subjects, with references 389 constrained according to the method described earlier (Chesters 2020). A number of backbone 390 topologies were integrated for setting hard and soft constraints, including a 391 transcriptomics-derived topology (Chesters 2020), a mitogenome tree of insects (Chesters 2017), 392 Diptera-specific trees (Wiegmann et al. 2011, Cranston et al. 2012, Ament 2017) and 393 Hymenoptera-specific trees (Peters et al. 2011, Branstetter et al. 2017, Cardinal 2018). The 394 constrained inference was conducted with RaxML version 8 (Stamatakis 2014) under the standard 395 GTRGAMMA DNA model with 24 rate categories. According to the backbone trees used, most 396 taxa present were monophyletic with a notable exception of Crabronidae, for which there is 397 emerging phylogenomic evidence of its polyphyly (Sann et al. 2018).

398 Tree phylogenetic diversity, functional diversity, and environmental covariates

399 The phylogenetic diversity of the tree communities was quantified by calculating wood 400 volume-weighted phylogenetic Mean Pairwise Distance (MPD) (Tucker et al. 2017). Tree 401 wood volume was estimated from data on basal area and tree height (Bongers et al. 2021) 402 measured in the center of each plot. Moreover, to represent variations towards the tips of the 403 phylogeny beyond MPD, we additionally calculated Mean Nearest Taxon Distance (MNTD), 404 which is a measure that quantifies the distance between each species and its nearest neighbor 405 on the phylogenetic tree (Webb 2000). Phylogenetic metrics of trees were calculated based on 406 a maximum likelihood phylogenetic tree available for the tree species in our study area 407 (Purschke et al. 2017). Considering that predatory wasps mainly feed on herbivorous 408 caterpillars, we calculated tree functional diversity to test the indirect effects on 409 hymenopteran communities and relevant network indices. Specifically, seven leaf traits 410 were expressed as Rao's Q (Ricotta and Moretti 2011), including specific leaf area, leaf 411 toughness, leaf dry matter content, leaf carbon content, ratio of leaf carbon to nitrogen, leaf 412 magnesium content, and leaf calcium content. These functional traits were commonly related 413 to higher trophic levels in our study area (Wang et al. 2020, Chen et al. 2023), which are the 414 main food resources of our predatory wasps. All of the traits were measured on pooled 415 samples of sun-exposed leaves of a minimum of five tree individuals per species following 416 standard protocols (Pérez-Harguindeguy et al. 2003). 417 As our analyses mainly compare community patterns among study plots, we additionally 418 considered potential effects of environmental variation by using plot means of slope, 419 elevation, "eastness" (sine-transformed radian values of aspect), and "northness" 420 (cosine-transformed radian values of aspect) as environmental covariates that characterize the 421 heterogeneity of the study plots. We also accounted for the potential effects of canopy cover 422 at plot level for host-parasitoid interactions, as it can structure hymenopteran communities

423 (Perlík et al. 2023). Canopy cover was calculated as in Fornoff et al. (2021) based on

424 above-nest images.

425 Statistical analysis

All analyses were conducted in R 4.1.2 with the packages *ape*, *vegan*, *picante*, *bipartite*, and *caper*(http://www.R-project.org). Prior to analysis, samples from the five years (2015, 2016, 2018, 2019, and 2020) were pooled at the plot level to discern overall and generalizable effects permeating this
system. We excluded three plots with no living trees because of high mortality, resulting in 85
plots in the final analysis.

431 *Composition of trees, hosts and parasitoids*

432 The species and phylogenetic composition of trees, hosts, and parasitoids were quantified at

433 each plot with nonmetric multidimensional scaling (NMDS) analysis based on Morisita-Horn

434 distances. The phylogenetic composition was calculated by mean pairwise distance among the

435 host or parasitoid communities per plot with the R package "*picante*." To test the influence of

436 study plot heterogeneity on these relationships, we fitted their standardized values to the

437 ordination on the basis of a regression with the NMDS axis scores (Quinn and Keough 2002).

438 For the analysis, we considered tree species richness, tree functional and phylogenetic

diversity, canopy cover, and environmental covariates (elevation, eastness, northness, and

slope) as plot characteristics. We assessed the significance of correlations with permutation

441 tests (permutation: n=999).

442 *Phylogenetic match of hosts and parasitoids*

In addition, we used a parafit test (9,999 permutations) with the R package "*ape*" to test whether the associations were non-random between hosts and parasitoids. The species that

445 were not attacked by parasitoids or failed to generate sequences were excluded from the

analyses. For species abundance and composition, see Table S1.

447 *Host-parasitoid interactions*

448 We constructed quantitative host-parasitoid networks with the R package "*bipartite*" for the

449 level of host and parasitoid communities across for each plot of the two sites. All hosts were

- 450 grouped because there were too few abundant bee species to separate them out from other
- 451 hosts. We calculated six indices to quantitatively characterize the structure of the interaction

452 networks: weighted generality (effective number of host species per parasitoid species),

453 weighted vulnerability (Effective number of parasitoid species attacking a host species),

454 robustness (degree of network stability), linkage density (degree of network specialization),

455 and interaction evenness (degree of network evenness). Generality was defined as the

456 weighted mean number of host species per parasitoid species, $G_{qw} = \sum_{j=1}^{J} \frac{A_j}{m} 2^{H_j}$, with A_j

- 457 being the number of interactions of parasitoid species j, m the total number of interactions of
- 458 all species, and H_i the Shannon diversity of interactions of species *j*. Vulnerability was the

459 weighted mean number of parasitoid species per host species, Vulnerability = $\sum_{i=1}^{I} \frac{A_i}{m} 2^{H_i}$

460 (Bersier et al. 2002). Robustness was defined as the area under the extinction curve, reflecting

the degree of decreases of one trophic level with the elimination of the other trophic levels,

462 here using the robustness index for higher trophic levels (i.e. parasitoids). For linkage density,

463 $L_q = 0.5 \left(\sum_{j=1}^{J} \frac{A_j}{m} 2^{H_j} + \sum_{i=1}^{I} \frac{A_i}{m} 2^{H_i} \right)$, we used the realized proportion of possible links

464 between the two trophic levels as the mean number of interactions per species across the 465 entire network (Tylianakis et al. 2007). Interaction evenness was defined as $E_s =$

466 $-\sum_{i}\sum_{j}p_{ij}\ln p_{ij}/\ln IJ$, which is used to describe Shannon's evenness of network interactions 467 (Dormann et al. 2009). To check whether all network indices significantly differ from chance 468 across all study plots, we used Patefield null models (Dormann et al. 2009) to compare

d69 observed indices with simulated values (10,000 times).

470 *Linear mix-effects models* To test the effects of tree species richness, tree phylogenetic, and
471 functional diversity, as well as canopy cover and the other environmental covariates

472 (including slope, elevation, eastness, and northness) on the six network indices, we used

473 linear mixed-effect models. For our analyses, the study sites were considered as random

474 effects, and the others were treated as fixed effects (see above). Given the strong correlation

475 between tree species richness and tree MPD (Pearson's r = 0.74, p<0.01), we excluded tree

476 MPD in the models where tree species richness was a predictor. To evaluate the potential

477 effects caused by tree MPD, we also ran alternative models where tree species richness was

478 replaced with tree MPD. We simplified all models by gradually removing non-significant

479	factors to obtain the most parsimonious model with the lowest AICc. To ensure that the
480	analyses were not strongly affected by multicollinearity, the correlations among all predictors
481	were tested (Fig. S2), and variance inflation factors (VIF) of our statistical models were checked.
482	
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495	

- 496 Table 1 Environmental correlates of dissimilarity matrixes with predictors (NMDS on
- 497 Morisita-Horn dissimilarity) across the study plots. Significant P-values are indicated in bold.
- 498 See Table S2-S5 for the complete information.

	Host	Parasitoid	Host	Parasitoid
	species	species	phylogenetic	phylogenetic
	community	community	community	community
Tree phylogeny NMDS1	0.225	0.422	0.386	0.274
Tree phylogeny NMDS2	0.003	0.12	0.128	0.024
Tree composition NMDS1	0.001	0.001	0.001	0.001
Tree composition NMDS2	0.604	0.418	0.433	0.031
Canopy cover	0.001	0.001	0.001	0.004
Tree species richness	0.035	0.122	0.100	0.094
Elevation	0.005	0.007	0.001	0.001
Eastness	0.079	0.045	0.04	0.001
Northness	0.49	0.837	0.821	0.340
Slope	0.031	0.507	0.507	0.959
Tree FD (Rao's Q)	0.094	0.019	0.021	0.031
Tree MPD	0.005	0.021	0.013	0.223
Host phylogeny NMDS1	-	0.016	-	0.584
Host phylogeny NMDS2	-	0.027	-	0.914
Host composition NMDS1	-	0.001	-	0.008
Host composition NMDS2	-	0.169	-	0.138
Parasitoid phylogeny NMDS1	0.001	-	0.001	-
Parasitoid phylogeny NMDS2	0.462	-	0.058	-
Parasitoid composition NMDS1	0.001	-	0.001	-
Parasitoid composition NMDS2	0.014	-	0.001	-

499

- 501 Table 2. Summary results of linear mix-effects models for generality, vulnerability,
- 502 robustness, linkage density, and interaction evenness of host-parasitoid network indices at the
- 503 community level across the tree species richness gradient. Standardized parameter estimates
- 504 (with standard errors, t and P values) are shown for the variables retained in the minimal
- 505 models.

		Est.	SE	t	Р
Generality	Intercept	0.176	0.016	10.96	< 0.001
	Canopy cover	0.033	0.016	2.03	0.046
Vulnerability	Intercept	2.860	0.197	14.49	0.044
	Tree species richness	0.180	0.079	2.29	0.025
Robustness of	Tedamand				
parasitoids	Intercept	0.630	0.007	85.15	< 0.001
	Tree species richness	-0.021	0.007	-2.77	0.007
Linkage density	Intercept	2.033	0.093	21.83	0.029
	Tree species richness	0.106	0.040	2.68	0.009
Interaction	T				
evenness	Intercept	0.511	0.009	59.12	0.025
	Canopy cover	-0.037	0.007	-5.06	< 0.001
	Eastness	-0.018	0.007	-2.50	0.015

506





509 Fig. 1. Ordination plot of the non-metric multidimensional scaling (NMDS) analysis of (a) host taxonomic composition, (b) parasitoid taxonomic composition, (c) host 510 511 phylogenetic composition, and (d) parasitoid phylogenetic composition across the study plots (filled circles) in the BEF-China experiment. Stress = 0.23, 0.23, 0.24 and 512 513 0.20, respectively. Arrows indicate significant (at p < 0.05) correlations of environmental variables with NMDS axis scores. Lengths of arrows are proportional 514 515 to the strength of the correlations. Red crosses refer to the host or parasitoid species in each community. See Table S2-S5 in the Supplementary Material for abbreviations 516 517 and statistical values.

518



- 521 Fig. 2. Dendrogram of phylogenetic congruence for the host species (below) and associated
- 522 parasitoid species (above) recorded in the study. Each rectangle represents a different
- 523 superfamily (for host species) or family (for parasitoid species). H1: Pompilidae H2: Apoidea
- 524 H3: Vespidae; P1: Sarcophagidae P2: Phoridae P3: Bombyliidae P4: Trigonalyidae P5:
- 525 Mutillidae P6: Megachilidae P7: Chrysididae P8: Ichneumonidae P9: Chalcidoidea. The
- 526 trophic network of hosts and parasitoids was non-randomly structured (parafit test: P = 0.032).
- 527 Host and parasitoid species names are given in Fig. S3.







Fig. 3. Community-level relationships of network between tree species richness and (a)
vulnerability, (b) linkage density, and (c) robustness of parasitoids. Values were adjusted for

532 covariates of the final regression model. Regression lines (with 95% confidence bands) show

533 significant (p < 0.05) relationships. Note that axes are on a log scale for tree species richness.

534

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536

537 Fig. 4. Community-level relationships of network between tree mean pairwise phylogenetic538 distance and (a) vulnerability and (b) linkage density and community-level relationships of

ustance and (a) vulnerability and (b) linkage density and community-rever relationships of

network between canopy cover and (c) interaction evenness. Values were adjusted for

540 covariates of the final regression model. Regression lines (with 95% confidence bands) show

541 significant (p < 0.05) relationships.

542

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