

1 **Do closely related species interact with similar partners?**

2 **Testing for phylogenetic signal in bipartite interaction networks**

3
4 Running title: Measuring phylogenetic signal in interactions

5
6 Benoît Perez-Lamarque ^{1,2}, Odile Maliet ¹, Benoît Pichon ^{3,1}, Marc-André Selosse ^{2,4,5},
7 Florent Martos ², and Hélène Morlon ¹

8
9 ¹ *Institut de biologie de l'École normale supérieure (IBENS), École normale supérieure, CNRS,*
10 *INSERM, Université PSL, 46 rue d'Ulm, 75 005 Paris, France*

11 ² *Institut de Systématique, Évolution, Biodiversité (ISYEB), Muséum national d'histoire*
12 *naturelle, CNRS, Sorbonne Université, EPHE, UA, CP39, 57 rue Cuvier 75 005 Paris, France*

13 ³ *Institut d'écologie et des sciences de l'environnement (iEES), Sorbonne Université, CNRS,*
14 *UPEC, CNRS, IRD, INRA, 75 005 Paris, France*

15 ⁴ *Department of Plant Taxonomy and Nature Conservation, University of Gdansk, Wita*
16 *Stwosza 59, 80-308 Gdansk, Poland*

17 ⁵ *Institut universitaire de France (IUF), Paris, France*

18
19 Corresponding author: Benoît Perez-Lamarque (benoit.perez@ens.psl.eu)

20
21 Supplementary data: [https://github.com/BPerezLamarque/Phylosignal_network/](https://github.com/BPerezLamarque/Phylosignal_network/blob/master/Supplementary_figures_phylo_signal_network.pdf)
22 [blob/master/Supplementary_figures_phylo_signal_network.pdf](https://github.com/BPerezLamarque/Phylosignal_network/blob/master/Supplementary_figures_phylo_signal_network.pdf)

24 **Abstract**

25 Whether interactions between species are conserved on evolutionary time-scales has
26 spurred the development of both correlative and **process**-based approaches for testing
27 phylogenetic signal in interspecific interactions: do closely related species interact with
28 similar partners? Here we use simulations to test the statistical performances of the
29 two approaches that are the most widely used in the field: Mantel tests and the
30 Phylogenetic Bipartite Linear Model (PBLM). Mantel tests investigate the correlation
31 between phylogenetic distances and dissimilarities in sets of interacting partners,
32 while PBLM is a **process**-based approach that relies on strong assumptions on how
33 interactions evolve. We find that PBLM often detects phylogenetic signal when it
34 should not. Simple Mantel tests instead have **infrequent false positives** and moderate
35 statistical power; however, they often artifactually detect that closely related species
36 interact with dissimilar partners. Partial Mantel tests, which are used to partial out the
37 phylogenetic signal in the number of partners, actually fail at correcting for this
38 confounding effect, and we instead **recommend to evaluate the significance of Mantel**
39 **tests with network permutations constraining the number of partners**. We also explore
40 the ability of simple Mantel tests to analyze clade-specific phylogenetic signal. We
41 provide general guidelines and an application on an interaction network between
42 orchids and mycorrhizal fungi.

43

44 **Keywords:** ecological network, phylogenetic signal, Mantel tests, clade-specific signal,
45 species interactions, mycorrhizal symbiosis.

46 Introduction

47

48 Species in ecological communities engage in numerous types of interspecific
49 interactions, such as pollination, mycorrhizal symbioses, herbivory, and parasitism
50 (Bascompte et al. 2003; Fontaine et al. 2011; Martos et al. 2012; Bascompte and Jordano
51 2013), which are often summarized using bipartite interaction networks (Bascompte &
52 Jordano 2013; Fig. 1). Understanding the processes that shape these interaction
53 networks, including the role of evolutionary history, is a major focus of ecology and
54 evolution (Rezende et al. 2007; Futuyma and Agrawal 2009; Vázquez et al. 2009;
55 Gómez et al. 2010; Krasnov et al. 2012; Elias et al. 2013; Rohr and Bascompte 2014;
56 Braga et al. 2021). One way to assess the role of evolutionary history in shaping
57 contemporary interactions is to test for phylogenetic signal in species interactions, *i.e.*
58 whether closely related species interact with similar sets of partners (Peralta 2016).

59

60 Testing for phylogenetic signal in a trait **for a given clade**, *i.e.* whether a trait is
61 phylogenetically conserved, is mainstream (Felsenstein 1985; Blomberg et al. 2003;
62 Münkemüller et al. 2012). One approach (the ‘correlative’ approach) is to perform a
63 Mantel test between phylogenetic and trait distances (Mantel 1967); another approach
64 (the ‘**process**-based’ approach) relies on trait evolution models such as Pagel’s λ (Pagel
65 1999) or Blomberg’s K (Blomberg et al. 2003). The **process**-based approach has a higher
66 ability to detect an existing phylogenetic signal (power) and a lower propensity to infer
67 a phylogenetic signal when it should not (**false positive**; Harmon & Glor 2010): The
68 correlative approach should therefore only be used when the **process**-based approach
69 is not applicable, *e.g.* if the ‘trait’ data is expressed in terms of pairwise distances.

70

71 Testing for phylogenetic signal in species interactions falls in the category of
72 cases where the ‘trait’ data are pairwise distances, here the between-species
73 dissimilarity in sets of interacting species. Simple Mantel tests have therefore been
74 widely used in this context (*e.g.* Cattin *et al.* 2004; Rezende *et al.* 2007; Elias *et al.* 2013;

75 Fontaine & Thébault 2015). Partial Mantel tests have also been used to test whether the
76 phylogenetic signal reflects more the identity of the interacting partners than their
77 number, as similarity in the number of partners can increase the value of similarity
78 metrics (“phylogenetic signal in the number of partners”; Rezende *et al.* 2007;
79 Jacquemyn *et al.* 2011; Aizen *et al.* 2016). Mantel tests, that are easy and fast to run and
80 that do not rely on strong hypotheses, have therefore been vastly used to test for
81 phylogenetic signal in species interactions in empirical networks (Cattin *et al.* 2004;
82 Rezende *et al.* 2007; Jacquemyn *et al.* 2011; Elias *et al.* 2013; Fontaine and Thébault
83 2015). Besides these correlative approaches, several **process**-based approaches have
84 been developed (Ives and Godfray 2006; Rafferty and Ives 2013; Hadfield *et al.* 2014;
85 Li *et al.* 2020). The first of **these approaches**, the Phylogenetic Bipartite Linear Model
86 (PBLM, Ives & Godfray 2006) has been widely used to test for phylogenetic signal in
87 species interactions in a variety of networks, *e.g.* in host-parasite, plant-fungus, and
88 pollination networks (Ives and Godfray 2006; Martos *et al.* 2012; Martín González *et al.*
89 *et al.* 2015; Xing *et al.* 2020). In short, PBLM assumes that interaction strengths between
90 species from the two guilds are determined by (unobserved) traits that evolve on the
91 two phylogenies each following a simplified Ornstein-Uhlenbeck process (Blomberg
92 *et al.* 2003). PBLM performs a phylogenetic regression to infer the Ornstein-Uhlenbeck
93 parameters, which are then interpreted in terms of phylogenetic signal (Ives &
94 Godfray 2006). Other models have been developed more recently (Rafferty and Ives
95 2013; Hadfield *et al.* 2014; Li *et al.* 2020), including the phylogenetic generalized linear
96 mixed model (PGLMM; Rafferty and Ives 2013) that uses linear mixed models to infer
97 phylogenetic signals in both the number of partners and species interactions. Yet, the
98 higher computational requirements of these methods have prevented their
99 widespread use on empirical networks. PBLM thus remains the method frequently
100 used in empirical studies (*e.g.* Xing *et al.* 2020; Corro *et al.* 2021).

101

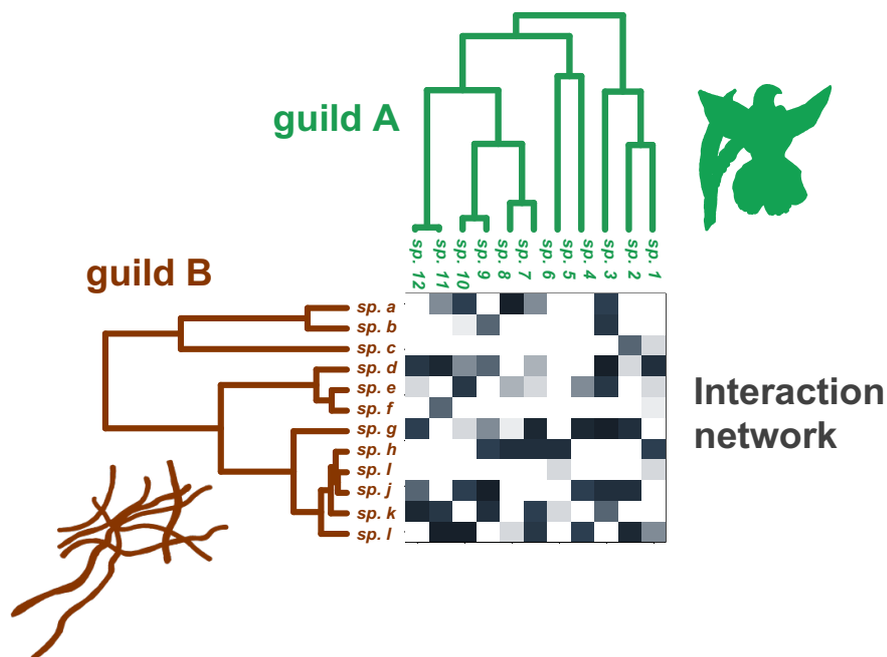
102 Mantel tests and PBLM sometimes provide contradictory conclusions on
103 empirical data and this is difficult to interpret because the statistical performances of

104 the two approaches have never been compared (Peralta 2016). Importantly, the
105 statistical performances of PBLM have not been tested. Here, we use simulations to
106 perform a comparative analysis of the statistical performances of these approaches.
107 We consider both weighted and unweighted bipartite interaction networks between
108 species from two guilds A and B (Fig. 1). Our results lead us to propose **alternative**
109 **approaches** for measuring phylogenetic signal in interaction networks. We also
110 investigate the ability of Mantel tests to detect the presence of phylogenetic signal in
111 the different clades of a phylogenetic tree, as phylogenetic signal may be localized.
112 Finally, we provide general guidelines and illustrate them on an orchid-fungus
113 mycorrhizal network identified across the oceanic island of Réunion (Martos et al.
114 2012).

115 **Figure 1: Illustration of the data used to test for phylogenetic signal in species**
116 **interactions**

117 Toy example of an interaction network between orchids (in green) and
118 mycorrhizal fungi (in brown) with associated phylogenetic trees. The bipartite
119 interaction network between two guilds A (here the orchids) and B (the fungi) is
120 represented by a matrix which elements indicate either whether or not species interact
121 (*i.e.* 1 if they do and 0 otherwise, 'unweighted' or 'binary' network) or the frequency
122 of the interaction ('weighted' network; for example here we indicated the number of
123 times a given pairwise interaction has been observed using shades of gray from white
124 (no interaction) to dark gray (many interactions)). Each guild is also characterized by
125 a rooted phylogenetic tree, used to compute phylogenetic distances between pairs of
126 species.

127



128

129 **Methods**

130

131 **Simulating bipartite interaction networks with or without phylogenetic signal in** 132 **species interactions**

133

134 We used *BipartiteEvol*, an individual-based eco-evolutionary model (Figure 2;
135 see Maliet *et al.* 2020 for a complete description of the model), to generate interaction
136 networks with or without phylogenetic signal between two guilds interacting in a
137 mutualistic, antagonistic, or neutral way. In short, each individual from guild A (resp.
138 B) is characterized by a multidimensional continuous trait and interacts with one
139 individual from guild B (resp. A). The effect of this interaction on the fitness of each
140 individual from guilds A or B is determined by the distance in trait space of the two
141 interacting individuals, according to a classical trait matching expression
142 parametrized by two parameters α_A and α_B (Supplementary Methods 1, Maliet *et al.*
143 2020). These parameters determine the nature and specificity of the interaction:
144 positive α_A and α_B correspond to mutualistic interactions, negative α_A and positive α_B
145 to antagonistic interactions (with guild A representing hosts/preys and guild B
146 parasites/predators), high $|\alpha|$ values to scenarios with strong fitness effects (*i.e.* highly
147 specialized interactions), and $|\alpha|$ values close to 0 to more neutral scenarios (Figure 2).
148 *BipartiteEvol* simulates the random death of individuals that are replaced by new ones
149 proportionally to their fitness. At birth, new individuals have a probability μ to mutate,
150 leading to new trait values slightly different from the parental ones (Figure 2). Such
151 mutations can lead to the formation of new species. Networks simulated using
152 *BipartiteEvol* show typical structural properties observed in empirical networks,
153 including significant nestedness and/or modularity according to the sets of simulated
154 parameters (Maliet *et al.* 2020). For instance, networks simulated with antagonistic
155 interactions ($\alpha_A < 0$) tend to be significantly modular, while networks simulated with
156 neutral or mutualistic interactions ($\alpha_A = 0$ or $\alpha_A > 0$) tend to be nested. Here, instead of

157 using the species delineation of the original *BipartiteEvol* model (Maliet et al. 2020), we
158 considered that each combination of traits corresponds to a distinct species. This
159 increased our ability to generate phylogenetic signal in the simulated networks, and
160 we show that it does not affect their overall structure (see below).

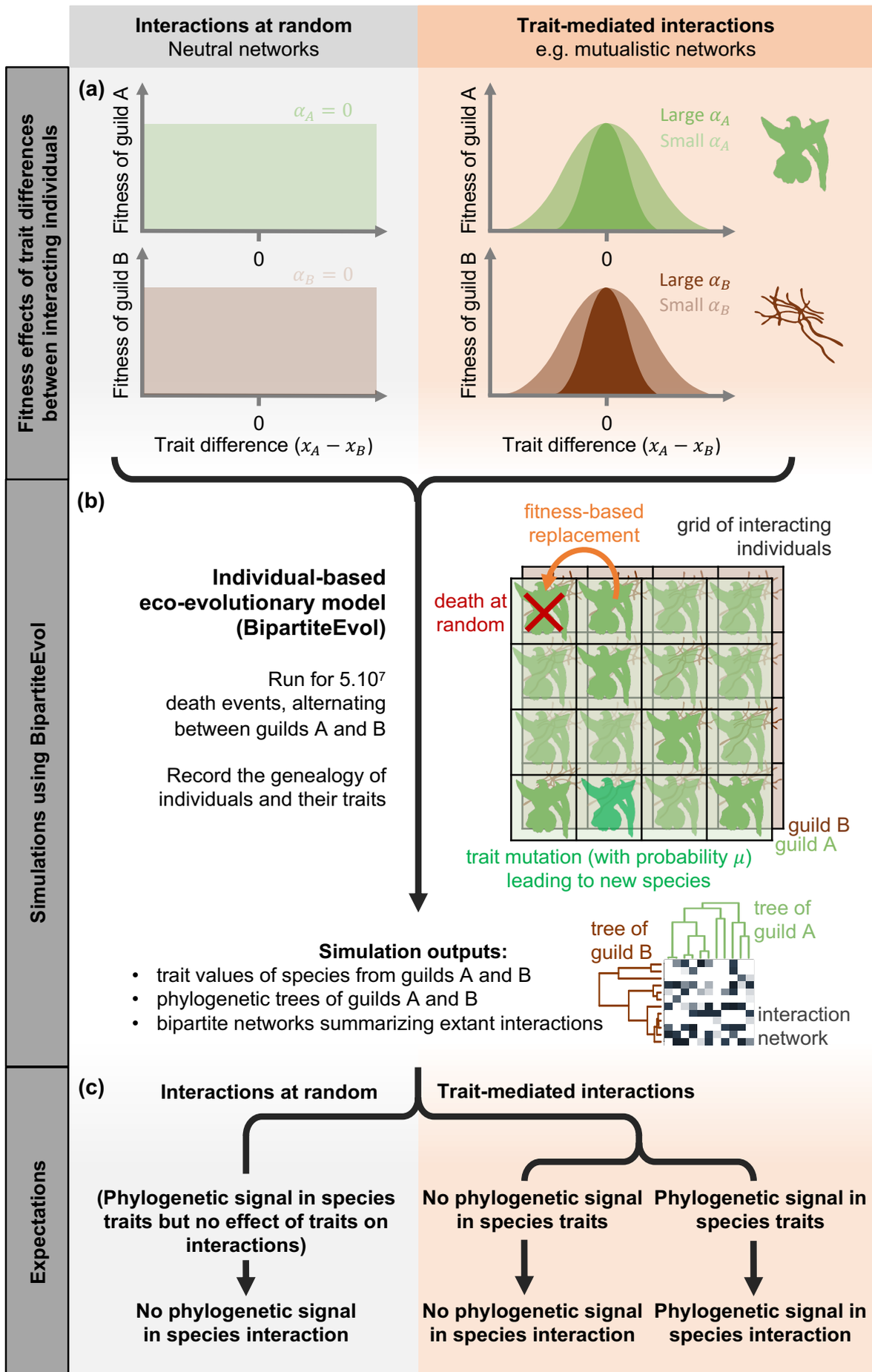
161

162 **Figure 2: Simulation scheme used to generate interaction networks with or without**
163 **phylogenetic signal in species interactions:**

164 (a) The fitness of a given individual is either affected by its trait value and that of the
165 individual it interacts with (right; “mutualistic or antagonistic interactions”) or not
166 (left; “neutral interactions”). In the first case, fitness depends on the trait matching
167 between the pair of interacting individuals ($x_A - x_B$), where x_A (resp. x_B) are the trait
168 values of the individual from guild A (resp. B). The strength of the effect of the traits
169 on fitness is modulated by the parameters α_A and α_B ($|\alpha|$ values close to 0 tends toward
170 neutrality, where interactions happen at random).

171 (b) *BipartiteEvol* assumes that pairs of individuals from guilds A and B interact on a
172 grid of a given size. Each cell of the grid contains one pair of individuals. The process
173 starts with one monomorphic species in each guild. At each time step, one individual
174 of guild A is killed at random and replaced by another individual proportionally to its
175 fitness in the cell. The new individual can mutate with probability μ , in which case the
176 new trait value is drawn from a normal distribution centered on the parental trait. A
177 mutation generates a new species. The same events are applied to an individual from
178 guild B, and the process is repeated a large number of time steps. This simulation
179 model outputs a list of species of guilds A and B with associated trait values,
180 phylogenetic trees, and the bipartite interaction network of extant species.

181 (c) A phylogenetic signal in species interactions can occur only when trait values
182 modulate interactions (for large α) and there is a phylogenetic signal in trait values.
183 When interactions are completely or quasi random ($\alpha = 0$ or low α), there cannot be a
184 phylogenetic signal in species interactions.



186 Under the *BipartiteEvol* model, closely related species tend to interact with
187 similar sets of partners, *i.e.* there is a phylogenetic signal in species interactions, if and
188 only if: (1) closely related species have similar traits, *i.e.* there is a phylogenetic signal
189 in species traits, and (2) these traits determine who interacts with whom, *i.e.* $\alpha \neq 0$
190 (Figure 2). Similarly, a **negative** phylogenetic signal in species interactions, *i.e.* the
191 tendency for closely related species to associate with dissimilar partners, is expected if
192 there is a **negative** phylogenetic signal in species traits, *i.e.* closely related species have
193 dissimilar traits, and $\alpha \neq 0$.

194 We used the *sim.BipartiteEvol* function from the R-package RPANDA (Morlon
195 et al. 2016; R Core Team 2022) to simulate a total of 2,400 interaction networks. To
196 obtain a wide range of network sizes, we considered a total number of 500, 1,000, 2,000,
197 3,000, 4,000, or 5,000 pairs of interacting individuals per simulation. For each size, we
198 simulated the evolution of 100 neutral networks ($\alpha_A=0$; $\alpha_B=0$), 120 mutualistic
199 networks (**i**: $\alpha_A=1$; $\alpha_B=1$; **ii**: $\alpha_A=0.1$; $\alpha_B=0.1$; **iii**: $\alpha_A=0.01$; $\alpha_B=0.01$; **iv**: $\alpha_A=1$; $\alpha_B=0.1$; **v**: $\alpha_A=1$;
200 $\alpha_B=0.01$; and **vi**: $\alpha_A=0.1$; $\alpha_B=0.01$) and 180 antagonistic networks (**i**: $\alpha_A=-1$; $\alpha_B=1$; **ii**: $\alpha_A=-$
201 0.1 ; $\alpha_B=0.1$; **iii**: $\alpha_A=-0.01$; $\alpha_B=0.01$; **iv**: $\alpha_A=-1$; $\alpha_B=0.1$; **v**: $\alpha_A=-1$; $\alpha_B=0.01$; **vi**: $\alpha_A=-0.1$; $\alpha_B=1$;
202 **vii**: $\alpha_A=-0.1$; $\alpha_B=0.01$; **viii**: $\alpha_A=-0.01$; $\alpha_B=1$; **ix**: $\alpha_A=-0.01$; $\alpha_B=0.1$). Each individual was
203 characterized by a six-dimensional trait, and trait mutation occurred at birth with
204 probability $\mu=0.01$. Upon mutation, the new trait values were drawn independently in
205 a normal distribution centered on the parental traits and with a variance of 1. We
206 followed the interacting individuals during $5 \cdot 10^7$ death events. At the end, we
207 extracted for each guild a species tree from its genealogy by randomly selecting one
208 individual per species (Fig. S1), we also recorded the number of individuals belonging
209 to each species, and counted the number of occurrences of each interspecific
210 interaction; we then reconstructed the corresponding weighted interaction network.

211 First, we evaluated whether these simulations generated realistic networks by
212 comparing their structure with that of empirical networks. Empirical networks were
213 gathered from the Web of Life database (web-of-life.es (Fortuna et al. 2014)) and the
214 database of Michalska-Smith & Allesina (2019). We compared the structures of

215 simulated *versus* empirical networks in terms of connectance, nestedness, and
216 modularity (Supplementary Methods 2).

217 Second, we separated the 2,400 simulated networks between those for which
218 we should expect a phylogenetic signal in species interactions and those for which we
219 should not (Figure 2). We did not expect phylogenetic signal in species interactions in
220 neutral networks and in non-neutral networks with no phylogenetic signal in species
221 traits. Conversely, we expected phylogenetic signal in non-neutral networks with
222 phylogenetic signal in species traits. We evaluated phylogenetic signal in species traits
223 using two approaches. First, for simplicity and consistency with the rest of the paper,
224 we used Mantel tests (Pearson correlation) between phylogenetic distances and trait
225 distances computed as the Euclidian distances between trait values for each species
226 pair. Second, given that process-based approaches usually perform better (Harmon
227 and Glor 2010), we used a multivariate extension of Pagel's λ (Pagel 1999)
228 implemented in R (Goolsby 2015); we assessed the significance of the phylogenetic
229 signal in species traits with likelihood ratio tests comparing the inferred Pagel's λ
230 model to a null model where $\lambda=0$ (*i.e.* no phylogenetic signal).

231

232 **Computing phylogenetic signal in species interactions**

233

234 We computed phylogenetic signal in species interactions in the simulated
235 networks using Mantel tests and PBLM, as well as the computationally-intensive
236 PGLMM for the smallest networks. Complete descriptions of these methods are
237 available in Supplementary Methods 3. Mantel tests, PBLM, and PGLMM rely on
238 different strategies to evaluate the significance of the phylogenetic signal, and it could
239 be argued that results of these tests are not directly comparable. Our approach is to
240 follow the methodologies traditionally used in empirical studies and compare their
241 conclusions (detection or not of a phylogenetic signal).

242

243 Mantel tests: We evaluated the phylogenetic signal in species interactions in guilds A
244 and B separately using simple Mantel tests between phylogenetic and ecological (set
245 of interacting partners) distances. Ecological distances were measured both without
246 accounting for evolutionary relatedness of the interacting partners, using (weighted or
247 unweighted) Jaccard, and accounting for relatedness using (weighted or unweighted)
248 UniFrac distances (Supplementary Methods 3 (Lozupone et al. 2011)). Accounting for
249 evolutionary relatedness of the interacting partners can be particularly relevant for
250 organisms with uncertain species delineations (e.g. microorganisms delineated using
251 only molecular data (Martos et al. 2012; Sanders et al. 2014)). We used Pearson,
252 Spearman, and Kendall correlations (R) by extending the *mantel* function in the R-
253 package *ecodist* (Goslee and Urban 2007); the significance of each correlation was
254 evaluated using 10,000 permutations, except for the computationally intensive Kendall
255 correlation (100 permutations only). For each network, we considered that there was
256 a significant **positive** phylogenetic signal (resp. **negative** phylogenetic signal) if the
257 correlation coefficient (R) was higher (resp. lower) than >95% of the randomized
258 correlations; we computed the p-value of each one-tailed Mantel test as the fraction of
259 the randomized correlations above (resp. below) the original value.

260

261 PBLM: To estimate phylogenetic signal based on PBLM, we modified the function *pblm*
262 from the R-package *picante* (Kembel et al. 2010) to more efficiently perform matrix
263 inversions and handle large interaction networks. In short, the parameters d_A and d_B
264 of the Ornstein-Uhlenbeck processes of PBLM were estimated using generalized least
265 squares (Ives & Godfray 2006). d_A and d_B are interpreted as a measure of phylogenetic
266 signal in species interactions: if $d=0$, there is no effect of the phylogeny (similar as
267 evolution on a star phylogeny, *i.e.* no phylogenetic signal); $0 < d < 1$ generates stabilizing
268 selection (*i.e.* phylogenetic signal) and $d > 1$ disruptive selection (*i.e.* **negative**
269 phylogenetic signal). We followed Ives & Godfray (2006; Supplementary Methods 3)
270 by considering that the phylogenetic signal is significant when the mean square error
271 (MSE) of the model is smaller than that obtained using star phylogenies (MSE_{star}); we

272 also used a more stringent criterion by considering that the signal is significant when
273 the MSE is at least 5% lower than MSE_{star} . Finally, we applied the bootstrapping
274 method of Ives & Godfray (2006; Supplementary Methods 3) to the smallest networks.
275 A single PBLM inference can take several days to run (time measured on an Intel 2.8
276 GHz MacOSX laptop) on networks of intermediate sizes (*e.g.* between 50 and 100
277 species per guild), which prevented us from applying the bootstrap approach to large
278 networks; we therefore **only** tested this approach on networks simulated with 500
279 individuals (*i.e.* a total of 400 networks).

280

281 PGLMM: We performed analyses of the statistical performances of PGLMM (Rafferty
282 and Ives 2013) using the function *pglmm* in the R-package *phyr* (Li et al. 2020).
283 Following the procedure used in Lajoie and Kembel (2021), we fitted for each network
284 different models accounting or not for phylogenetic signals in both the number of
285 partners and in the species interactions in both clades, using restricted maximum
286 likelihood and evaluating significance with likelihood ratio tests. Because fitting these
287 models can require large amount of memory (*e.g.* >80 Gb for some networks with >50
288 species per guild) **and long computation time (Fig. S2)**, we **only** tested this approach
289 on networks simulated with 500 individuals. We fitted the PGLMM using either a
290 Gaussian or a Poisson distribution of abundances for weighted networks, and a
291 binomial distribution (presence/absence data) for unweighted networks (Li et al. 2020).

292

293 **Confounding effect of the phylogenetic signal in the number of partners**

294

295 To test the performances of the partial Mantel test at measuring phylogenetic
296 signal in species interactions while controlling for the number of partners
297 (Supplementary Methods 3), we first performed partial Mantel tests between
298 phylogenetic and ecological distances, while controlling for **pairwise** differences in **the**
299 **number of partners**, on the networks simulated with *BipartiteEvol*. There is no reason
300 that *BipartiteEvol* simulations generate a phylogenetic signal in the number of partners,

301 and we verified this by performing Mantel tests between phylogenetic distances and
302 pairwise differences in the number of partners. These analyses thus assess whether
303 partial Mantel tests lose power compared to simple Mantel tests in the absence of
304 phylogenetic signal in the number of partners. If they do not suffer power loss, partial
305 Mantel tests applied to *BipartiteEvol* simulations should be significant when simple
306 Mantel tests are significant.

307

308 Second, we assessed whether partial Mantel tests successfully correct for
309 phylogenetic signal in the number of partners using networks simulated under a
310 process that generate phylogenetic conservatism in the number, but not the identity,
311 of interacting partners (*i.e.* partial Mantel tests should not be significant when applied
312 to such networks). To simulate network with only phylogenetic conservatism in the
313 number of partners in guild A, we first simulated phylogenetic trees for guilds A and
314 B using *pbtree* (R-package *phytools*; Revell 2012) with a number of species uniformly
315 sampled between 40 and 150 by guild. Next, we simulated the number of partners of
316 the species from guild A using an Ornstein-Uhlenbeck process with an attraction
317 toward 0, a variance of 0.1 (noise of the Brownian motion), and a selection strength (a_A)
318 ranging from 5 (strong stabilizing effect, weak phylogenetic signal) to 0 (Brownian
319 motion, strong phylogenetic signal). We computed the number of partners per species
320 by calibrating the simulated values between 1 and the number of species in guild B
321 and taking the integer part. For each a_A value (5, 1, 0.5, 0.05, or 0), we performed 100
322 simulations using *mvSIM* (R-package *mvMORPH*; Clavel *et al.* 2015). Finally, for each
323 species in A, we attributed the corresponding number of partners in B at random to
324 obtain binary networks. We checked that our simulations indeed generated a signal in
325 the number of partners by performing simple Mantel tests between phylogenetic
326 distances and pairwise differences in the number of partners. Finally, we performed
327 on each simulated network a partial Mantel test between phylogenetic and ecological
328 distances, while controlling for pairwise differences in number of partners.

329

330 Given the poor performances of partial Mantel tests (see Results), we tested
331 three alternative approaches to partial out the confounding effect of the number of
332 partners in measures of phylogenetic signals. First, we tested whether using sequential
333 Mantel tests would provide a good alternative: based on simple Mantel tests, we
334 consider that there is a phylogenetic signal in the identity of the partners if there is a
335 phylogenetic signal in species interactions and no phylogenetic signal in the number
336 of partners. Second, we tested the use of methods that directly partition ecological
337 distance metrics into a part due to the dissimilarity in the number of partners and a
338 part due to the dissimilarity in the identity of the partners, *i.e.* “species turnover”
339 (Baselga 2010). We used the betapart R-package (Baselga and Orme 2012) to extract the
340 part of the unweighted Jaccard distances due to species turnover and tested its
341 correlation with phylogenetic distances using a simple Mantel test. Third, we designed
342 specific network permutations to test for the significance of the Mantel correlation
343 between phylogenetic distances and ecological distances while accounting for the
344 number of partners. To measure whether the phylogenetic signal observed in guild A
345 is not due to a phylogenetic signal in the number of partners, instead of shuffling the
346 distance matrix as in a regular Mantel test (Supplementary Methods 3), we
347 randomized the interaction network by keeping constant the number of partners per
348 species from guild A while permuting the partner identities. Because this third
349 approach requires recomputing the ecological distances for each permutation, it is
350 much slower than regular Mantel tests (Fig. S2) and we thus used only 1,000
351 permutations. We applied these three methods to all our simulated networks.

352

353 **Effect of phylogenetic uncertainty, sampling asymmetry, and network** 354 **heterogeneity on measures of phylogenetic signal in species interactions**

355

356 Unlike simulations, such as those provided by *BipartiteEvol*, empirical bipartite
357 networks suffer from (i) uncertainty in the phylogenetic reconstructions, *e.g.* in the
358 microbial partners’ tree when studying host-associated microbiota, which often

359 prevents accounting for evolutionary relatedness (*i.e.* using UniFrac distances), (ii)
360 sampling asymmetry, *i.e.* one side of the network is more thoroughly sampled than
361 the other, and (iii) network heterogeneity, *i.e.* different sub-clades in the network have
362 different levels of phylogenetic signal. We performed additional analyses to
363 investigate the effect of these aspects on phylogenetic signals in species interactions
364 measured using simple Mantel tests.

365

366 First, we tested the effect of phylogenetic uncertainty in the partners' tree on the
367 measure of phylogenetic signal when evolutionary relatedness is accounted for (*i.e.*
368 using UniFrac distances). We performed these analyses to assess whether accounting
369 for the partners' evolutionary relatedness remains advantageous (see Results) when
370 phylogenetic uncertainty is high. To add some variability in the phylogenetic tree of
371 guild B (resp. A) used to compute the UniFrac distances between species pairs from
372 guild A (resp. B), we first simulated, on the original partners tree, the evolution of a
373 short DNA sequence and then reconstructed the tree from the simulated DNA
374 alignment using neighbor-joining (*nj* function, R-package APE (Paradis et al. 2004)).
375 We used *simulate_alignment* (R-package HOME; Perez-Lamarque & Morlon 2019) to
376 simulate sequences of length 75, 150, 300, 600, or 1,200 base-pairs, with 30% of variable
377 sites, and a substitution rate of 1.5 (shorter fragments should result in noisier
378 phylogenies). *To assess the uncertainty of these reconstructed phylogenetic trees
379 compared with the original trees, we computed the correlations between the pairwise
380 phylogenetic distances in both trees.*

381

382 Second, we tested the influence of sampling asymmetry on measures of
383 phylogenetic signal. Empirical networks are often an incomplete representation of the
384 actual interactions between two guilds because they are under-sampled, and
385 frequently, in an asymmetrical way. For instance, by sampling targeted species from
386 guild A, observed networks are constituted by few species from guild A which have
387 the complete set of their partners and by often more species from guild B which have

388 an incomplete set of their partners (as they likely interact with unsampled species from
389 guild A). We tested the influence of such sampling asymmetry by selecting only 10%
390 of the most abundant species from guild A in each simulated network (while retaining
391 at least 10 species) and computed phylogenetic signal in these asymmetrically-
392 subsampled networks.

393

394 Third, both Mantel tests and PBLM neglect the heterogeneity within networks.
395 Indeed, a non-significant phylogenetic signal at the level of the entire network can
396 potentially hide a sub-clade of species presenting significant phylogenetic signal.
397 Alternatively, a phylogenetic signal in the entire network may be driven by only two
398 sub-clades of guilds A and B, while the other sub-clades present no significant
399 phylogenetic signal. To explore the potential heterogeneity of the phylogenetic signal
400 within one guild, one possibility is to apply Mantel tests to the sub-networks formed
401 by a given sub-clade (*e.g.* Song *et al.* 2020). For each node of the tree of guild A having
402 at least 10 descendants, we estimated the clade-specific phylogenetic signal using a
403 Mantel test investigating whether closely related species from this sub-clade of A tend
404 to interact with similar partners (and *vice-versa* for guild B). Using UniFrac distances,
405 we performed the Mantel tests with 100,000 permutations, and introduced a
406 Bonferroni correction for multiple testing to keep a global risk of false positives of 5%.
407 To test this approach, we simulated networks with known sub-clade signal by
408 artificially combining networks simulated under neutrality with networks simulated
409 with the mutualistic parameters \mathbf{v} (see Results). We grafted each “mutualistic”
410 phylogenetic tree from guilds A and B within a “neutral” phylogenetic tree by
411 randomly selecting a branch, such that it creates a separate module with strong
412 phylogenetic signal. Such simulations could correspond to the evolution of a different
413 niche, *e.g.* terrestrial *versus* epiphytic plants associating with different mycorrhizal
414 fungi (Martos *et al.* 2012). We then performed our clade-specific analysis of
415 phylogenetic signal and investigated in which nodes we recovered a significant
416 phylogenetic signal.

417

418 **General guidelines and illustration with application on the orchid-fungus**
419 **mycorrhizal network from La Réunion**

420

421 We used our results and other empirical considerations to provide general
422 guidelines for testing for phylogenetic signal in interaction networks. We illustrated
423 these guidelines by applying them in a network between orchids and mycorrhizal
424 fungi from La Réunion island (Martos et al. 2012). This network encompasses 70 orchid
425 species (either terrestrial or epiphytic species) and 93 molecularly-identified fungal
426 partners (defined according to 97% sequence similarity; Martos *et al.* 2012). We
427 gathered the maximum-likelihood plant and fungal phylogenies on TreeBASE (Study
428 Accession S12721), calibrated the orchid phylogeny using a relaxed clock with *chronos*
429 (Paradis 2013), and set the divergence between Orchidoideae and Epidendroideae at
430 65 million years (Givnish et al. 2015). To obtain a species-level orchid phylogeny,
431 missing species were grafted in the phylogeny by arbitrarily adding 10 million-year-
432 old polytomies in the corresponding unresolved genera, namely *Habenaria*, *Benthamia*,
433 *Cynorkis*, *Phaius*, *Liparis*, *Bulbophyllum*, and *Polystachya*.

434 Results

435

436 Expected phylogenetic signal in species interactions in *BipartiteEvol* networks

437

438 *BipartiteEvol* simulations resulted in interaction networks with a large range of
439 sizes for guilds A and B, from less than 50 to more than 250 species (Fig. S3). These
440 simulated networks had similar structural properties as empirical networks, including
441 in terms of connectance, nestedness, and modularity (Fig. S4). This means that
442 networks simulated using *BipartiteEvol* are realistic and cover a large range of
443 structures encountered in natural interaction networks.

444 Using Mantel tests, we found a significant phylogenetic signal in species traits
445 for most antagonistic and neutral simulations (Fig. S5A). In contrast, for many
446 mutualistic simulations, closely related species often did not tend to have similar traits,
447 except when $\alpha_B=0.01$ (*i.e.* mutualistic parameters **iii**, **v**, and **vi**; Fig. S5A). When α_B were
448 higher (*i.e.* mutualistic parameters **i**, **ii**, and **iv**), we suspect stabilizing selection to
449 occur and erase the phylogenetic signal in the traits (Maliot et al. 2020): we therefore
450 do not expect phylogenetic signal in species interactions for these simulations, which
451 represent ~40% of the mutualistic simulations. In addition, we found a **negative**
452 phylogenetic signal in species traits (**suggesting that closely related species have**
453 **dissimilar traits**) in less than 1% of the simulations (Fig. S5A). Given that we do not
454 expect *BipartiteEvol* to generate a **negative** phylogenetic signal in species traits and
455 given that the **risk of false positives of a Mantel test** is 5%, these 1% of networks with
456 a **negative** phylogenetic signal in species traits are likely false-positives. We removed
457 them when evaluating the performance of the different approaches and we therefore
458 do not expect **negative** phylogenetic signal in species interactions for the networks we
459 tested, *i.e.* **closely related species should not tend to associate with dissimilar partners**.
460 Results were similar with Pagel's λ , with a significant phylogenetic signal in species
461 traits for almost all antagonistic and neutral simulations, and in ~65% of the

462 mutualistic simulations (Fig. S5B). Mantel tests and Pagel's λ lead to identical
463 conclusions for >95% of the simulated networks.

464

465 **Computing phylogenetic signal in species interactions in *BipartiteEvol* networks**

466

467 Using Mantel tests, as expected, we did not find a significant phylogenetic
468 signal in species interactions for most neutral networks or for networks with no signal
469 in species traits (Figs. 3 & S6): the **false positive** rate was below 5%, corresponding to
470 the **risk of false positives** of the test (Table S1), with one notable exception for small
471 networks when using weighted Jaccard distances and Pearson correlations (~8% of
472 **false positives**). Conversely, we detected a significant unexpected **negative**
473 phylogenetic signal in more than 10% of the simulated networks, in particular in the
474 small ones (Figs. 3 & S6).

475

476 Many mutualistic or antagonistic networks where we expected a phylogenetic
477 signal in species interactions (*i.e.* non-neutral networks with signal in species traits)
478 presented no significant signal with Mantel tests (Figs. 3 & S6), in particular those
479 simulated with low α_A and α_B values (*e.g.* antagonism **vii**), where non-neutral effects
480 were weak. Mantel tests measuring phylogenetic signal in species interactions were
481 most often not significant unless the phylogenetic signal in species traits was strong
482 ($R > 0.6$; Fig. S7). Even when the phylogenetic signal in species traits was very strong
483 ($R > 0.9$), the phylogenetic signal in species interactions was not significant in many
484 networks. In mutualistic networks, phylogenetic signals in species interactions were
485 present only when there was a large asymmetry in the effects of trait matching on the
486 fitnesses of the species from guilds A or B (case **v**: $\alpha_A = 1$; $\alpha_B = 0.01$), *i.e.* when only one
487 guild was specialized. Conversely, in antagonistic networks, phylogenetic signals
488 were found mainly when trait matching had a strong impact on the fitness of guild B
489 (the obligate parasites/predators; $\alpha_B \geq 0.1$). Additionally, when phylogenetic signal was

490 significant in one guild, it was generally also significant in the other; in antagonistic
491 networks, the signal was usually higher in guild A compared to guild B (Fig. S6).

492

493

494 **Figure 3: Statistical performances of the simple Mantel tests and the Phylogenetic**
495 **bipartite linear model (PBLM; Ives & Godfray, 2006)**

496 For each panel, the simulations are divided between networks where phylogenetic
497 signal in species interactions is expected (*i.e.* networks (i) simulated with an effect of
498 the traits on individual fitness - antagonistic and mutualistic simulations - and (ii)
499 presenting traits that are phylogenetically conserved according to a Mantel test – see
500 Fig. S5A) and networks where phylogenetic signal in species interactions is not
501 expected (*i.e.* neutral simulations ($\alpha = 0$) or simulated networks where we observed
502 no phylogenetic signal in the traits). Results are similar when the expectations are
503 based on Pagel’s λ to measure the phylogenetic signals in species traits (Fig. S11).

504

505 **a-d:** Phylogenetic signals in species interactions estimated using simple Mantel tests
506 with Pearson correlation (R) in the guilds A (a, c) and B (b, d). The different panels in
507 rows correspond to the 2 tested ecological distances: weighted Jaccard (a, b) or
508 weighted UniFrac (c, d) distances. One-tailed Mantel tests between phylogenetic
509 distances and ecological distances were performed using 10,000 permutations. In each
510 panel, the bars indicate the percentage of simulated networks that present a significant
511 positive correlation (in green; $p\text{-value} > 0.05$ for the test of phylogenetic signal), a
512 significant negative correlation (in red; $p\text{-value} > 0.05$ for the test of **negative**
513 phylogenetic signal), or no significant correlation (in yellow; both $p\text{-values} > 0.05$).
514 Significant phylogenetic signals (resp. **significant negative** phylogenetic signals) are
515 shaded from light green (resp. red) to dark green (resp. red) according to the strength
516 of the signal: we arbitrarily considered a “low signal” when $R < 0.05$ (resp. $R > -0.05$), an
517 “intermediate signal” when $0.05 < R < 0.15$ (resp. $-0.05 > R > -0.15$), and a “strong signal”
518 when $R > 0.15$ (resp. $R < -0.15$).

519

520 **e:** Phylogenetic signals estimated using PBLM. For a given combination of parameters,
521 the bar indicates the percentage of simulated networks that present no significant (in
522 yellow; $MSE \geq MSE_{star}$) or a significant (green; $MSE < MSE_{star}$) phylogenetic signal.
523 Phylogenetic signals are shaded from light green to dark green according to the
524 strength of the signal: we arbitrarily considered a “low signal” when $d_A < 0.05$ and
525 $d_B < 0.05$, an “intermediate signal” when $d_A > 0.05$ or $d_B > 0.05$, and a “strong signal” when
526 $d_A > 0.15$ or $d_B > 0.15$. PBLM were run on the weighted networks.

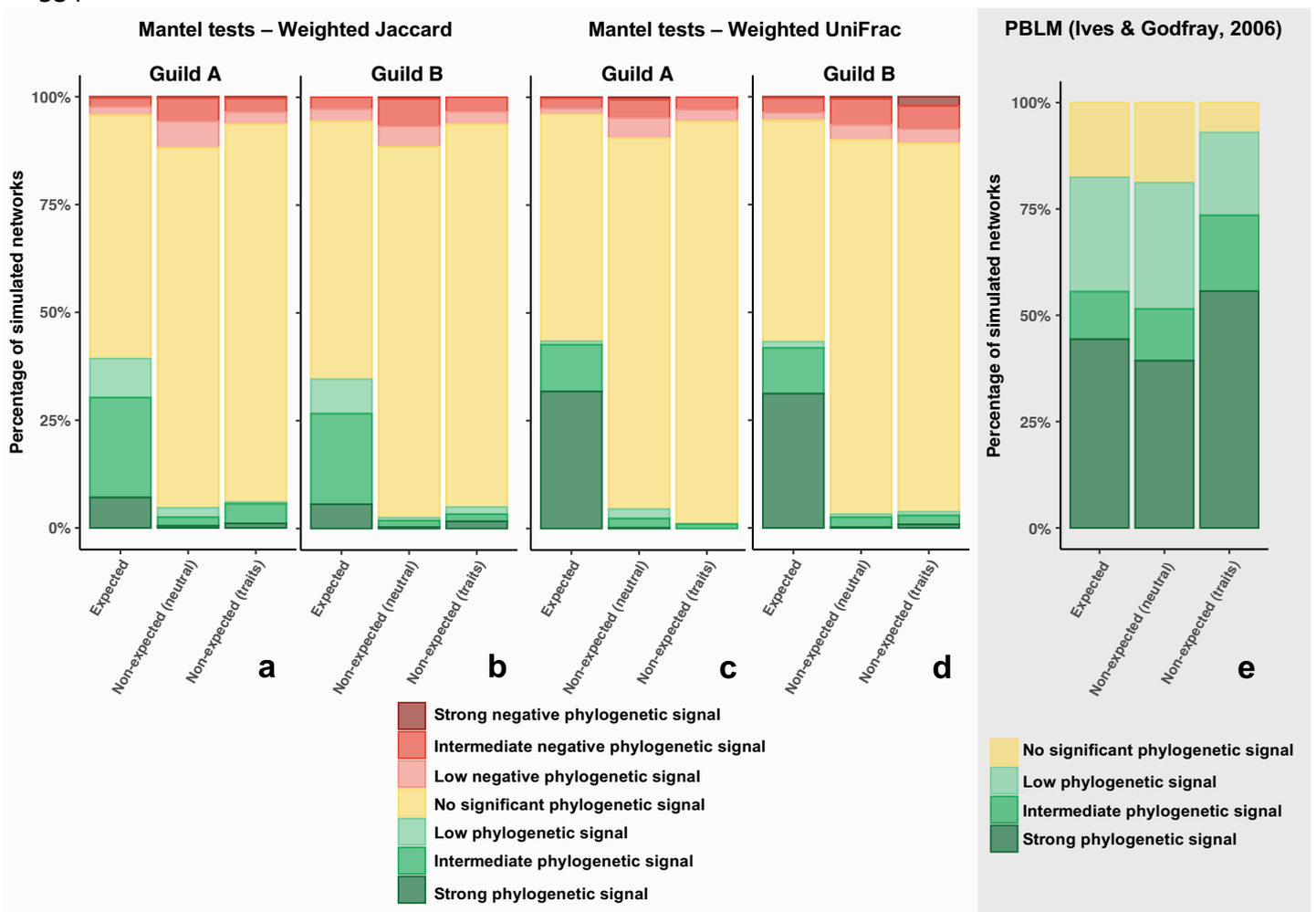
527

528 In each panel, the first bar indicates the statistical power of the test, whereas the second
529 and third bars indicate the **false positive** rate of the test. Note that the strength the
530 phylogenetic signals (based on the R and d values) are not directly comparable.

531

532 Results discriminating the simulated networks of different sizes and with different sets
533 of parameters are available in Figures S6 & S8.

534



535 The statistical power of Mantel tests measuring phylogenetic signal in species
536 interactions seems to be modulated by network size, as phylogenetic signals were less
537 often significant but generally stronger in smaller networks (Fig. S6). Moreover,
538 Mantel tests based on Pearson correlations had higher power than Spearman and
539 Kendall correlations (Fig. S6) and weighted UniFrac distances **performed better than**
540 other ecological distances in terms of power **in the context of these simulations** (Fig.
541 S6; Table S2).

542
543 When using mean square errors to evaluate the significance of PBLM, we found
544 a significant phylogenetic signal in species interactions in most of the simulated
545 networks including when we did not expect any (Fig. 3e). The strength and the
546 significance of the inferred phylogenetic signals were independent of the strength of
547 the phylogenetic signal in species traits (Fig. S7). The propensity of PBLM to detect
548 phylogenetic signal decreased in large unweighted networks, but the **false positives**
549 remained >30%, including when using a more stringent significance cutoff (Figs. S8).
550 Similar results were obtained when bootstrapping to evaluate the significance (Fig.
551 S9). PGLMM on weighted networks with a Gaussian or Poisson distribution had
552 slightly lower but still **frequent false positives** (>25% or 20%, respectively) and
553 intermediate statistical power (<50%) when measuring phylogenetic signals in species
554 interactions (Fig. S10). PGLMM also often artifactually detected phylogenetic signals
555 in the number of partners (Fig. S10). Conversely, PGLMM on unweighted networks
556 never detected any significant signal (Fig. S10).

557
558 We inferred similar statistical performances of both Mantel tests and PBLM
559 when we used Pagel's λ to evaluate phylogenetic signal in species traits (Figs. S7 and
560 S11).

561

562 **Confounding effect of the phylogenetic signal in the number of partners**

563

564 As expected, tests of phylogenetic signal in the number of partners were non-
565 significant in the large majority of the *BipartiteEvol* networks, especially the larger ones
566 (Fig. S12). We did however observe significant correlations between ecological
567 distances and differences **in the number of partners** (Fig. S13). Partial Mantel tests
568 testing for phylogenetic signal in species interactions while accounting for
569 phylogenetic signal in the number of partners had similar **false positive rates** and
570 power as simple Mantel tests (Figs. S6 & S14; Table S2). Sequential Mantel tests
571 decreased the statistical power by less than 2% (Table S2). **Partitioning the ecological**
572 **distances before running the Mantel test reduced the power by only 5% and resulted**
573 **in less than 1.5% of false positives, but also resulted in an artefactual detection of**
574 **negative phylogenetic signal in 9% of the simulations (Table S2; Fig. S15). Finally,**
575 **Mantel tests with network permutations keeping the number of partners constant**
576 **increased the statistical power by ~5% (Table S2), but resulted in an artefactual**
577 **detection of (positive or negative) phylogenetic signal in ~10% of the simulations when**
578 **using Jaccard distances (Fig. S16).**

579

580 Networks simulated with phylogenetic conservatism in the number, but not the
581 identity of partners covered a realistic range of sizes (Fig. S17). As expected, Mantel
582 tests revealed significant phylogenetic signals in the number of partners in >60% of
583 these networks, with an increasing percentage of significant tests with decreasing α_A
584 (*i.e.* increasing conservatism in the number of partners; Fig. S18). We found significant
585 correlations between differences in the number of partners and ecological distances in
586 most of these simulated networks (Fig. S19), **generating a confounding effect**. As a
587 result, simple Mantel tests testing for phylogenetic signal in species interactions
588 without accounting for phylogenetic signal in the number of partners were frequently
589 significant (>30%; Fig. S20; Table S3). Partial Mantel tests controlling for differences **in**
590 **the number of partners** slightly decreased the proportion of false-positives, but it

591 remained **very** high (>25% of **false positives**; Fig. S21). In addition, partial Mantel tests
592 detected a spurious significant **negative** phylogenetic signal in species interactions in
593 >15% of the networks (Fig. S21). Conversely, only few networks with a significant
594 simple Mantel test in species interactions did not produce a significant simple Mantel
595 test in the number of partners, such that sequential Mantel tests had only ~7% of **false**
596 **positives** (Table S3). **Partitioning the ecological distances before running the Mantel**
597 **test (Fig. S22) or using Mantel tests with network permutations keeping the number of**
598 **partners constant (Fig. S23) had even lower false positive rates (~4% and ~5%**
599 **respectively; Table S3). However, partitioning the ecological distances led to an**
600 **artefactual detection of negative phylogenetic signals in more than 30% of the**
601 **simulated networks (Fig. S23).**

602

603 **Effect of phylogenetic uncertainty, sampling asymmetry, and network** 604 **heterogeneity on measures of phylogenetic signal in species interactions**

605

606 **As expected, phylogenetic uncertainty in the partner's tree increased when the**
607 **length of the simulated DNA sequence used for phylogenetic reconstruction decreased**
608 **(Fig. S24), resulting in a decreased** statistical power of Mantel tests using UniFrac
609 distances (Fig. S25). However, even when the simulated DNA sequences were the
610 shortest (75 base pairs), resulting in very noisy reconstructed partners' trees (Fig. S26),
611 the statistical power of the Mantel tests using UniFrac distances **decreased by less than**
612 **5% (Fig. S25).**

613

614 Our results on the statistical performance of tests of phylogenetic signal were
615 similar when considering sampling asymmetry (Figs. S27-30): PBLM spuriously
616 detected phylogenetic signal when it should not, and Mantel tests had decent
617 statistical performances, especially when using weighted UniFrac distances. In
618 addition, the correlations of the Mantel tests in guild A were generally higher when
619 significant (Fig. S29).

620

621 Our clade-specific tests of phylogenetic signal using Mantel tests while
622 correcting for multiple testing recovered a significant phylogenetic signal in 82% of the
623 nodes where mutualism originated (Fig. S31), as well as in most of the ascending
624 nodes. Conversely, we did not find spurious phylogenetic signals in nodes with only
625 neutrally-evolving lineages (Fig. S31).

626

627 **General guidelines and illustration with application on the orchid-fungus** 628 **mycorrhizal network from La Réunion**

629

630 Figure 4 provides general guidelines based on our results and empirical
631 considerations for accurate tests of phylogenetic signal in interaction networks. We
632 applied these guidelines on the orchid-fungus mycorrhizal network from La Réunion
633 (available in Martos et al. (2012)). First (step 1), simple Mantel tests of phylogenetic
634 signal in species interactions for fungi and orchids revealed a significant but low
635 phylogenetic signal ($R < 0.10$) on the orchid side using Jaccard distances; however, the
636 significance disappeared with UniFrac distances (Table S4). Similarly, marginally not-
637 significant and low phylogenetic signals were detected in the mycorrhizal fungi side
638 ($R < 0.04$; Table S4). Next (step 2), **results were qualitatively similar when using Mantel**
639 **tests with permutations keeping the number of partners constant, suggesting that the**
640 **phylogenetic signal in species interaction did not result from a phylogenetic signal in**
641 **the number of partners.** Our investigation of clade-specific phylogenetic signals in
642 species interactions in orchids (option 1) revealed a significant phylogenetic signal in
643 Angraecinae, a sub-tribe composed of 34 epiphytic species (Mantel test: $R = 0.37$;
644 Bonferroni-corrected p -value = 0.016; Fig. 5) interacting with 53 fungi, suggesting that
645 closely related Angraecinae tend to interact with more similar mycorrhizal fungi.
646 When we checked the robustness of the significant phylogenetic signal detected in
647 Angraecinae (option 2) by subsampling the Angraecinae clade down to 10 species, we
648 still recovered a significant signal in species interactions in both cases (Fig. S32).

649 **Figure 4: Recommended guidelines to measure phylogenetic signal in species**
650 **interactions within bipartite ecological networks.**

651 This guideline is composed of two fixed steps followed by two optional ones and can
652 be applied as soon as a bipartite interaction network (with or without abundances)
653 and at least the phylogenetic tree of guild A are available. The phylogenetic tree does
654 not need to be binary, rooted, or ultrametric. For each step, an example of the
655 corresponding function available in the R-package RPANDA is indicated in grey.

656 **Step 1:** The first step consists in testing for phylogenetic signal in species interactions
657 for guild A (*i.e.* whether closely related species from guild A tend to interact with
658 similar partners from guild B) using a one-tailed simple Mantel test. This step requires
659 to pick an ecological distance (*e.g.* UniFrac or Jaccard distances) and a type of
660 correlation (Pearson correlation by default).

661 **Step 2:** Next, to assess whether a phylogenetic signal in species interactions really
662 comes from the identity of species interactions (**and not from a phylogenetic signal in**
663 **the number of partners**), the second step consists in testing whether the phylogenetic
664 signal in guild A **remains significant when the significance of the Mantel correlation is**
665 **evaluated using network permutations keeping the number of partners constant.**

666 **Option 1:** Clade-specific phylogenetic signal in guild A can be tested using simple
667 Mantel tests while correcting for multiple testing (*e.g.* Bonferroni correction). It can be
668 used to test whether some clades present different intensities of phylogenetic signal
669 (*e.g.* because of higher specificity).

670 **Option 2:** The robustness of the findings can be tested by looking at how the
671 conclusions might be affected by phylogenetic uncertainty (*e.g.* using a Bayesian
672 posterior of tree) or sampling bias. The potential effect of sampling bias can be
673 investigated by subsampling all clades to the same number of species.

674 If a phylogenetic tree for guild B is available, all these steps can be replicated to test
675 for phylogenetic signal in species interaction in guild B.

Phylogenetic signal in guild A:

Step 1: Is there a phylogenetic signal in species interactions ?

(simple Mantel test)

(i) choice of ecological distances (Jaccard, UniFrac...)

(ii) with or without interaction abundances

```
RPANDA::phylosignal_network(network, tree_A, tree_B,  
method = "GUniFrac", correlation = "Pearson")
```

↓ Yes

Step 2: Is there still a phylogenetic signal in species interactions when accounting for the signal in the number of partners ?

(simple Mantel test with permutations keeping the number of partners per species constant)

```
RPANDA::phylosignal_network(network,  
tree_A, tree_B, method = "GUniFrac",  
correlation = "Pearson",  
permutations="npartners")
```

↓ No

Option 1: Investigate clade-specific phylogenetic signals (simple Mantel tests with Bonferroni correction)

```
RPANDA::phylosignal_sub_network(network, tree_A, tree_B,  
method = "GUniFrac", correlation = "Pearson")
```

Option 2: Test the robustness of the findings to phylogenetic uncertainty and/or sampling bias

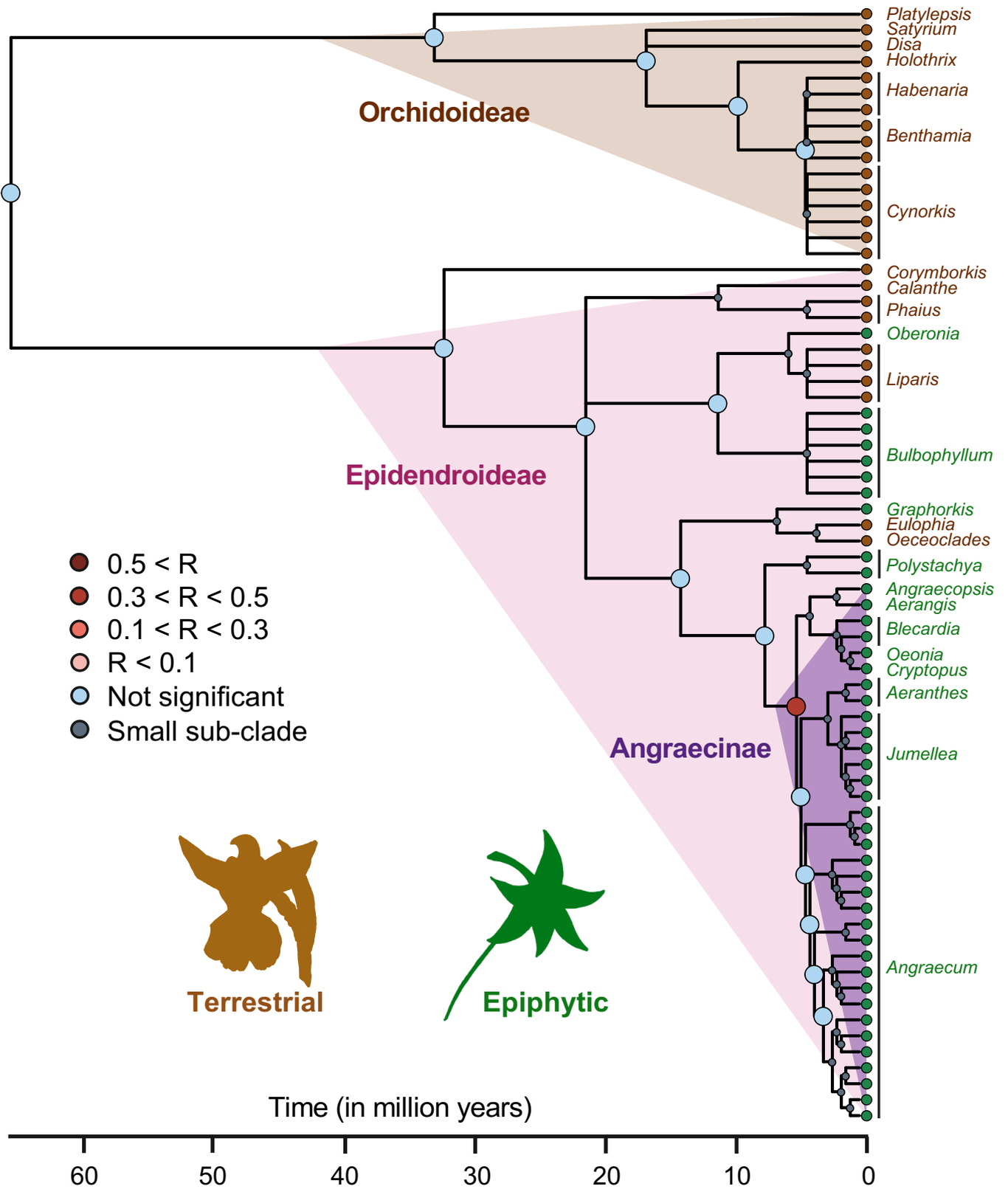
(repeat for guild B)

677 **Figure 5: Empirical application on an orchid-fungus interaction network from La**
678 **Réunion island (Martos *et al.*, 2012): the clade-specific analyses of phylogenetic**
679 **signal in species interactions revealed a significant phylogenetic signal in the**
680 **epiphytic subtribe Angraecinae.**

681 The orchid phylogeny (Martos *et al.*, 2012) is represented with its nodes colored
682 according to the results of the Mantel test performed on the corresponding sub-
683 network: in blue if non-significant, in grey when the node has less than 10 descendent
684 species (the Mantel test was not performed), and in red when the phylogenetic signal
685 is significant. Each one-tailed simple Mantel test was performed using the Pearson
686 correlation and 100,000 permutations and its significance was evaluated while
687 correcting for multiple testing (Bonferroni correction).

688 For each species, its habitat (terrestrial or epiphytic) is indicated at the tips of the tree
689 and the main orchid clades are highlighted in colors. Only the genera are indicated at
690 the tips of the tree (see Supplementary Figure S32 for the species list).

691



693 **Discussion:**

694

695 We used simulations to perform a comparative analysis of the statistical
696 performances of Mantel tests and the Phylogenetic bipartite linear model (PBLM; Ives
697 & Godfray 2006) for testing for phylogenetic signal in species interactions. Our results
698 highlight the weaknesses of PBLM and partial Mantel tests, and advocate for the use
699 of **regular Mantel tests and Mantel tests with network permutations keeping the**
700 **number of partners per species constant.**

701

702 The Phylogenetic bipartite linear model (PBLM) is widely used to test for
703 phylogenetic signal in species interactions, however we found that it has very **frequent**
704 **false positives** (>30%). PBLM assumes that the interaction strength between two
705 species is determined by the product of two unobserved traits evolving on the
706 phylogenies of guilds A and B respectively, according to two independent Ornstein-
707 Uhlenbeck processes with the selection strengths d_A and d_B (Supplementary Methods
708 3). PBLM tests the significance of d_A and d_B , which measure the phylogenetic signal of
709 the unobserved traits. A species with a high trait value will have high interaction
710 strengths with many partner species (*i.e.* it is a generalist species), while a species with
711 a low trait value will have low interaction strengths with most partner species, except
712 with the few species with high trait values (*i.e.* it is a specialist species). Therefore, we
713 suspect d_A and d_B to measure phylogenetic signals in the number of partners rather
714 than in species interactions. However, we also found significant d_A and d_B in the
715 absence of phylogenetic signal in the number of partners, suggesting that PBLM is
716 sensitive to model misspecification (it relies on strong hypotheses on how the number
717 of partners evolves). In any case, our results suggest that PBLM should not be used as
718 a routine for measuring phylogenetic signal in species interactions.

719

720 Other process-based approaches that extend PBLM (Rafferty and Ives 2013;
721 Hadfield et al. 2014; Li et al. 2020) allow to infer parameters thought to reflect the

722 phylogenetic structure of interactions networks, while controlling for phylogenetic
723 signal in the number of partners as well as heterogeneity in sampling effort (Hadfield
724 *et al.*, 2014). Our analyses using the PGLMM approach (Rafferty and Ives 2013) on the
725 smallest simulated networks suggested that it also has frequent false positives and
726 intermediate statistical power when using weighted interactions. It would have been
727 ideal to also test this approach on larger networks, but this was prohibited by their
728 computational cost (Fig. S2). Indeed, fitting PGLMM can require >80 Gb of memory
729 for some networks and our application of the Bayesian approach of Hadfield *et al.*
730 (2014) ran several days (on an Intel 2.8 GHz, MacOSX laptop) without reaching
731 convergence. Because of these high computational demands, these methods are
732 typically not used to measure phylogenetic signal in species interactions in empirical
733 studies, which is either done using Mantel tests or PBLM (see Fontaine and Thébault
734 2015; Xing *et al.* 2020; Corro *et al.* 2021 for recent examples). Future model
735 developments of such approaches would thus benefit from faster inferences and our
736 results highlight the need to thoroughly test these approaches with simulations before
737 they are applied to empirical systems and biological conclusions are drawn.

738

739 We found that simple Mantel tests have a moderate statistical power and a
740 reasonable false positive rate (<5%) when testing for phylogenetic signal in species
741 interactions. Not surprisingly, these tests have a higher power for larger simulated
742 networks. The fact that Mantel tests have a moderate power for measuring
743 phylogenetic signals in species interactions corroborates the findings about Mantel
744 tests in other contexts (Harmon and Glor 2010; Guillot and Rousset 2013). Hence,
745 although simple Mantel tests might fail at detecting low phylogenetic signal, we can
746 trust their results when they are significant. On the contrary, we found a high
747 proportion of simulated networks (5-10%) presenting a significant negative
748 phylogenetic signal in species interactions, suggesting that closely related species
749 would tend to associate with dissimilar partners. Yet, we did not expect such an
750 outcome in our simulations because we did not observe any negative phylogenetic

751 signal in species traits. False-positives are therefore frequent when testing for **negative**
752 phylogenetic signal using simple Mantel tests and detection of such signal in empirical
753 networks should be interpreted with caution.

754

755 In addition, Pearson correlations performed better than Spearman and Kendall
756 correlations, which is somewhat surprising, as correlations between phylogenetic and
757 ecological distances are not particularly expected to be linear: Spearman and Kendall
758 correlations have less stringent hypotheses, as they only assume monotonicity
759 (Supplementary Methods 3), but they probably lose information. We also reported that
760 using ecological distances that consider interaction abundances, such as weighted
761 **Jaccard or UniFrac** distances, significantly improves the detection of phylogenetic
762 signal. **Using UniFrac distances, which rely on the phylogenetic relatedness of the**
763 **partners, can be particularly relevant when species delineation is somewhat arbitrary,**
764 **e.g. in microbial systems, as it is less sensitive to species delineation than Jaccard**
765 **distances. In addition, results obtained with UniFrac distances were only moderately**
766 **influenced by the phylogenetic uncertainty in the partner's tree, which should thus not**
767 **prevent the use of UniFrac distances. In the context of our *BipartiteEvol* simulations,**
768 **which assume that species interactions are mediated by some phylogenetically-**
769 **conserved traits on both sides of the network, we found that UniFrac distances**
770 **outperform Jaccard distances. We note however that a significant phylogenetic signal**
771 **in UniFrac or Jaccard distances can reflect different evolutionary processes, such as**
772 **one where the traits involved in the interaction are evolutionary conserved on both**
773 **sides of the networks in the case of UniFrac, and on only one side of the network in**
774 **the case of Jaccard (Calatayud et al. 2016). Therefore, choosing between one or the**
775 **other metric (or using both) can also be dictated by the question at stake. Also, if**
776 **communities of interactors differ mainly in terms of recently diverged species, Jaccard**
777 **distances may perform better, as UniFrac distances emphasize differences in long**
778 **branches rather than recent splits (Sanders et al. 2014).**

779

780 We also found that multiple simple Mantel tests combined with a Bonferroni
781 correction perform rather well to investigate clade-specific phylogenetic signals. Such
782 an approach can therefore be valuable for measuring local phylogenetic signal in large
783 “meta-networks”, such as those describing host-microbiota phylosymbiosis (Song et
784 al. 2020), which likely have heterogeneous phylogenetic signals across the network.

785

786 While simple Mantel tests have satisfactory statistical performances, these tests do
787 not control for the potential confounding effect of phylogenetic signal in the number
788 of partners. Partial Mantel tests are frequently used for investigating phylogenetic
789 signal in species interactions while controlling for signal in the number of partners;
790 however, we found that they often detected significant signals in species interactions
791 when we simulated signals in only the number of partners. Thus, partial Mantel tests
792 fail at discerning whether evolutionary relatedness strictly affects the identity of
793 partners, independently of the total number of partners associated with each species
794 (Rezende et al. 2007). This corroborates the poor statistical performances of partial
795 Mantel tests frequently observed in other contexts (Harmon and Glor 2010; Guillot and
796 Rousset 2013). Among the alternative possibilities we tested, using sequential Mantel
797 tests, *i.e.* testing first for phylogenetic signal in species interactions, and if significant
798 testing for phylogenetic signal in the number of partners, has both high statistical
799 power and a low false positive rate. Yet, if both Mantel tests are significant, it does not
800 say whether the signal is entirely due to the signal in the number of partners and
801 therefore, sequential Mantel tests likely have very low power in this case.
802 Alternatively, using methods that can explicitly partition ecological distances into a
803 part due to dissimilarities in the number of partners *versus* the identity of the partners
804 appears promising, although we detected a slight power decrease in our simulations
805 and >30% of artefactual negative phylogenetic signals when partitioning unweighted
806 Jaccard distances. Other partitioning approaches may give better results and should
807 require further attention, as they offer a direct quantification of the contribution of the
808 species identity *versus* the number of partners in the phylogenetic signal (Baselga 2010;

809 Leprieur et al. 2012; Calatayud et al. 2016). Finally, performing a Mantel test with
810 network permutations designed to keep the number of partners associating with each
811 species constant while shuffling their identity has infrequent false positives and does
812 not decrease the statistical power. Therefore, if there is still signal while constraining
813 the number of partners, then we can safely conclude that evolutionary relatedness
814 affects the identity of partners. We thus recommend using such network permutations
815 to correct for the confounding effect of the phylogenetic signal in the number of
816 partners (Figure 4).

817

818 By definition, phylogenetic signals in species interactions measure general patterns
819 that are not informative of the processes at play (Losos 2008). A better understanding
820 of the ecological and evolutionary processes playing a role in the assembly of
821 interaction networks (Harmon et al. 2019) will require developing integrative process-
822 based approaches, for instance inference machineries for eco-evolutionary models
823 such as *BipartiteEvol*. Classical inferences (generalized least-squares or likelihood-
824 based approaches) might be challenging for such complex models (Hadfield et al.
825 2014), but strategies such as machine learning provide promising alternatives.

826

827 In the mycorrhizal network from La Réunion, we found non-significant or weak
828 phylogenetic signals in species interactions at the level of the entire orchid-fungus
829 network, suggesting these interactions are generally poorly conserved over long
830 evolutionary timescales (Jacquemyn et al. 2011; Martos et al. 2012; Perez-Lamarque et
831 al. 2022). Conversely, clade-specific Mantel tests detected a significant phylogenetic
832 signal in the Angraecinae epiphytic clade that is experiencing a radiation in La
833 Réunion island. This signal is likely produced by the different orchids genera in
834 Angraecinae associating with specific fungal clades (Martos et al. 2012). Thus, our
835 results corroborate a trend toward mycorrhizal specialization in epiphytic orchids
836 compared with terrestrial species (Xing et al. 2019), as the epiphytic habitats might

837 require particular adaptations and stronger dependences toward specific mycorrhizal
838 fungi.

839

840 Interaction networks are increasingly being analyzed to unravel the
841 evolutionary processes shaping their structure and to predict their stability. Currently-
842 used tools for measuring phylogenetic signals are clearly misleading. The approach
843 we propose based on Mantel tests **may have a limited statistical power, but it avoids**
844 **false positive, and it is flexible as it allows using different ecological distances and/or**
845 **permutation strategies.** By emphasizing the limits of current tests of phylogenetic
846 signal, we hope to stimulate new developments in the statistical adjustment to
847 empirical data of process-based models for the evolution of interaction networks.

848 **Acknowledgments:**

849 The authors acknowledge M. Elias, E. Thébault, and D. de Vienne for helpful
850 discussions. They also thank I. Overcast, S. Lambert, I. Quintero, C. Fruciano, J. Clavel,
851 and A. Silva for comments on an early version of the manuscript. This work was
852 supported by a doctoral fellowship from the École Normale Supérieure de Paris
853 attributed to BPL and the École Doctorale FIRE – Programme Bettencourt. Funding of
854 the research of FM was from the Agence Nationale de la Recherche (ANR-19-CE02-
855 0002). HM acknowledges support from the European Research Council (grant CoG-
856 PANDA).

857

858 **Author contributions:**

859 All authors designed the study. BPL performed the analyses, BPi performed the
860 analyses on the network structures, and FM gathered the empirical data. BPL and
861 HM wrote the first draft of the manuscript and all authors contributed to revisions.

862

863 **Data accessibility:**

864 The R functions used to measure phylogenetic signals in bipartite interaction
865 networks, including (simple, partial, and clade-specific) Mantel tests and PBLM, are
866 available in the R-package RPANDA (Morlon et al. 2016) (functions
867 *phylosignal_network* and *phylosignal_sub_network*). A tutorial and the simulated
868 networks can be found at https://github.com/BPerezLamarque/Phylosignal_network.
869 Amended functions of *BipartiteEvol* are also included in RPANDA.

870 The scripts for simulating the networks and for measuring the phylogenetic signals in
871 species interactions are available at:

872 https://github.com/BPerezLamarque/Phylosignal_network/tree/master/simulations

873

874

875 Supplementary data (including Supplementary Methods, Tables, and Figures) are
876 available at:

877 https://github.com/BPerezLamarque/Phylosignal_network/blob/master/Supplementa
878 [ry_figures_phylo_signal_network.pdf](https://github.com/BPerezLamarque/Phylosignal_network/blob/master/Supplementary_figures_phylo_signal_network.pdf)

879

880 **Conflict of Interest statement:**

881 The authors declare that there is no conflict of interest with the content of this article.

882 **References:**

883

884 Aizen, M. A., G. Gleiser, M. Sabatino, L. J. Gilarranz, J. Bascompte, and M. Verdú.
885 2016. The phylogenetic structure of plant-pollinator networks increases with habitat
886 size and isolation. *Ecology Letters* 19:29–36.

887 Bascompte, J., and P. Jordano. 2013. Mutualistic networks. *Monographs in*
888 *population biology* (Princeton., Vol. 7). Princeton University Press, Princeton.

889 Bascompte, J., P. Jordano, C. J. Melian, and J. M. Olesen. 2003. The nested assembly of
890 plant-animal mutualistic networks. *Proceedings of the National Academy of Sciences*
891 100:9383–9387.

892 Baselga, A. 2010. Partitioning the turnover and nestedness components of beta
893 diversity. *Global Ecology and Biogeography* 19:134–143.

894 Baselga, A., and C. D. L. Orme. 2012. betapart : an R package for the study of beta
895 diversity. *Methods in Ecology and Evolution* 3:808–812.

896 Blomberg, S. P., T. Garland, and A. R. Ives. 2003. Testing for phylogenetic signal in
897 comparative data: Behavioral traits are more labile. *Evolution* 57:717–745.

898 Braga, M. P., N. Janz, S. Nylin, F. Ronquist, and M. J. Landis. 2021. Phylogenetic
899 reconstruction of ancestral ecological networks through time for pierid butterflies
900 and their host plants. *Ecology Letters* 24:2134–2145.

901 Calatayud, J., J. L. Hórreo, J. Madrigal-González, A. Migeon, M. Á. Rodríguez, S.
902 Magalhães, and J. Hortal. 2016. Geography and major host evolutionary transitions
903 shape the resource use of plant parasites. *Proceedings of the National Academy of*
904 *Sciences* 113:9840–9845.

905 Cattin, M.-F. F., L.-F. Bersier, C. Banašek-Richter, R. Baltensperger, J.-P. P. Gabriel, L.
906 F. Bersler, C. Banašek-Richter, et al. 2004. Phylogenetic constraints and adaptation
907 explain food-web structure. *Nature* 427:835–839.

908 Clavel, J., G. Escarguel, and G. Merceron. 2015. mvMORPH: An R package for fitting
909 multivariate evolutionary models to morphometric data. (T. Poisot, ed.) *Methods in*
910 *Ecology and Evolution* 6:1311–1319.

911 Corro, E. J., F. Villalobos, A. Lira-Noriega, R. Guevara, P. R. Guimarães, and W.
912 Dáttilo. 2021. Annual precipitation predicts the phylogenetic signal in bat–fruit
913 interaction networks across the Neotropics. *Biology Letters* 17.

914 Elias, M., C. Fontaine, and F. J. Frank Van Veen. 2013. Evolutionary history and
915 ecological processes shape a local multilevel antagonistic network. *Current Biology*
916 23:1355–1359.

917 Felsenstein, J. 1985. Phylogenies and the comparative method. *American Naturalist*
918 125:1–15.

919 Fontaine, C., P. R. Guimarães, S. Kéfi, N. Loeuille, J. Memmott, W. H. van der Putten,
920 F. J. F. van Veen, et al. 2011. The ecological and evolutionary implications of
921 merging different types of networks. *Ecology Letters* 14:1170–1181.

922 Fontaine, C., and E. Thébault. 2015. Comparing the conservatism of ecological
923 interactions in plant–pollinator and plant–herbivore networks. *Population Ecology*
924 57:29–36.

925 Fortuna, M. A., R. Ortega, and J. Bascompte. 2014. The Web of Life. *arXiv*
926 Populations and Evolution.

927 Futuyma, D. J., and A. A. Agrawal. 2009. Macroevolution and the biological diversity
928 of plants and herbivores. *Proceedings of the National Academy of Sciences of the*
929 *United States of America*.

930 Givnish, T. J., D. Spalink, M. Ames, S. P. Lyon, S. J. Hunter, A. Zuluaga, W. J. D. D.
931 Iles, et al. 2015. Orchid phylogenomics and multiple drivers of their extraordinary
932 diversification. *Proceedings of the Royal Society B: Biological Sciences* 282:20151553.

933 Gómez, J. M., M. Verdú, and F. Perfectti. 2010. Ecological interactions are
934 evolutionarily conserved across the entire tree of life. *Nature* 465:918–921.

935 Goolsby, E. W. 2015. Phylogenetic comparative methods for evaluating the
936 evolutionary history of function-valued traits. *Systematic Biology* 64:568–578.

937 Goslee, S. C., and D. L. Urban. 2007. The ecodist package for dissimilarity-based
938 analysis of ecological data. *Journal of Statistical Software* 22:1–19.

939 Guillot, G., and F. Rousset. 2013. Dismantling the Mantel tests. *Methods in Ecology*

940 and Evolution 4:336–344.

941 Hadfield, J. D., B. R. Krasnov, R. Poulin, and S. Nakagawa. 2014. A tale of two
942 phylogenies: Comparative analyses of ecological interactions. *The American*
943 *Naturalist* 183:174–187.

944 Harmon, L. J., C. S. Andreazzi, F. Débarre, J. Drury, E. E. Goldberg, A. B. Martins, C.
945 J. Melián, et al. 2019. Detecting the macroevolutionary signal of species interactions.
946 *Journal of Evolutionary Biology* 32:769–782.

947 Harmon, L. J., and R. E. Glor. 2010. Poor statistical performance of the Mantel test in
948 phylogenetic comparative analyses. *Evolution* 64:2173–2178.

949 Ives, A. R., and H. C. J. Godfray. 2006. Phylogenetic analysis of trophic associations.
950 *The American Naturalist* 168:E1–E14.

951 Jacquemyn, H., V. S. F. T. Merckx, R. Brys, D. Tyteca, B. P. A. A. Cammue, O.
952 Honnay, and B. Lievens. 2011. Analysis of network architecture reveals phylogenetic
953 constraints on mycorrhizal specificity in the genus *Orchis* (Orchidaceae). *New*
954 *Phytologist* 192:518–528.

955 Kembel, S. W., P. D. Cowan, M. R. Helmus, W. K. Cornwell, H. Morlon, D. D.
956 Ackerly, S. P. Blomberg, et al. 2010. Picante: R tools for integrating phylogenies and
957 ecology. *Bioinformatics* 26:1463–1464.

958 Krasnov, B. R., M. A. Fortuna, D. Mouillot, I. S. Khokhlova, G. I. Shenbrot, and R.
959 Poulin. 2012. Phylogenetic signal in module composition and species connectivity in
960 compartmentalized host-parasite networks. *American Naturalist* 179:501–511.

961 Lajoie, G., and S. W. Kembel. 2021. Plant-bacteria associations are phylogenetically
962 structured in the phyllosphere. *Molecular Ecology* 30:5572–5587.

963 Leprieur, F., C. Albouy, J. De Bortoli, P. F. Cowman, D. R. Bellwood, and D. Mouillot.
964 2012. Quantifying Phylogenetic Beta Diversity: Distinguishing between ‘True’
965 Turnover of Lineages and Phylogenetic Diversity Gradients. (M. Shawkey, ed.) *PLoS*
966 *ONE* 7:e42760.

967 Li, D., R. Dinnage, L. A. Nell, M. R. Helmus, and A. R. Ives. 2020. phyr: An r package
968 for phylogenetic species-distribution modelling in ecological communities. (S. Price,

969 ed.)Methods in Ecology and Evolution 11:1455–1463.

970 Losos, J. B. 2008. Phylogenetic niche conservatism, phylogenetic signal and the
971 relationship between phylogenetic relatedness and ecological similarity among
972 species. Ecology Letters 11:995–1003.

973 Lozupone, C., M. E. Lladser, D. Knights, J. Stombaugh, and R. Knight. 2011. UniFrac:
974 an effective distance metric for microbial community comparison. The ISME Journal
975 5:169–172.

976 Maliet, O., N. Loeuille, and H. Morlon. 2020. An individual based model for the eco-
977 evolutionary emergence of bipartite interaction networks. (T. Poisot, ed.)Ecology
978 Letters 23:ele.13592.

979 Mantel, N. 1967. The detection of disease clustering and a generalized regression
980 approach. Cancer Research 27:209–220.

981 Martín González, A. M., B. Dalsgaard, D. Nogués-Bravo, C. H. Graham, M.
982 Schleuning, P. K. Maruyama, S. Abrahamczyk, et al. 2015. The macroecology of
983 phylogenetically structured hummingbird-plant networks. Global Ecology and
984 Biogeography 24:1212–1224.

985 Martos, F., F. Munoz, T. Pailler, I. Kottke, C. Gonneau, and M.-A. Selosse. 2012. The
986 role of epiphytism in architecture and evolutionary constraint within mycorrhizal
987 networks of tropical orchids. Molecular Ecology 21:5098–5109.

988 Michalska-Smith, M. J., and S. Allesina. 2019. Telling ecological networks apart by
989 their structure: A computational challenge. (T. Bollenbach, ed.)PLoS Computational
990 Biology 15:1–13.

991 Morlon, H., E. Lewitus, F. L. Condamine, M. Manceau, J. Clavel, and J. Drury. 2016.
992 RPANDA: An R package for macroevolutionary analyses on phylogenetic trees.
993 Methods in Ecology and Evolution 7:589–597.

994 Münkemüller, T., S. Lavergne, B. Bzeznik, S. Dray, T. Jombart, K. Schiffers, and W.
995 Thuiller. 2012. How to measure and test phylogenetic signal. Methods in Ecology
996 and Evolution 3:743–756.

997 Pagel, M. 1999. Inferring the historical patterns of biological evolution. Nature

998 401:877–884.

999 Paradis, E. 2013. Molecular dating of phylogenies by likelihood methods: A
1000 comparison of models and a new information criterion. *Molecular Phylogenetics and*
1001 *Evolution* 67:436–444.

1002 Paradis, E., J. Claude, and K. Strimmer. 2004. APE: Analyses of phylogenetics and
1003 evolution in R language. *Bioinformatics* 20:289–290.

1004 Peralta, G. 2016. Merging evolutionary history into species interaction networks.
1005 *Functional Ecology* 30:1917–1925.

1006 Perez-Lamarque, B., R. Petrolli, C. Strullu-Derrien, D. Strasberg, H. Morlon, M.-A.
1007 Selosse, and F. Martos. 2022. Structure and specialization of mycorrhizal networks in
1008 phylogenetically diverse tropical communities. *bioRxiv* 2022.05.10.491376.

1009 Perez-Lamarque, B., and H. Morlon. 2019. Characterizing symbiont inheritance
1010 during host–microbiota evolution: Application to the great apes gut microbiota.
1011 *Molecular Ecology Resources* 19:1659–1671.

1012 R Core Team. 2022. R: A language and environment for statistical computing. R
1013 Foundation for Statistical Computing, Vienna, Austria.

1014 Rafferty, N. E., and A. R. Ives. 2013. Phylogenetic trait-based analyses of ecological
1015 networks. *Ecology* 94:2321–2333.

1016 Revell, L. J. 2012. phytools: An R package for phylogenetic comparative biology (and
1017 other things). *Methods in Ecology and Evolution* 3:217–223.

1018 Rezende, E. L., J. E. Lavabre, P. R. Guimarães, P. Jordano, and J. Bascompte. 2007.
1019 Non-random coextinctions in phylogenetically structured mutualistic networks.
1020 *Nature* 448:925–928.

1021 Rohr, R. P., and J. Bascompte. 2014. Components of phylogenetic signal in
1022 antagonistic and mutualistic networks. *The American Naturalist* 184:556–564.

1023 Sanders, J. G., S. Powell, D. J. C. Kronauer, H. L. Vasconcelos, M. E. Frederickson,
1024 and N. E. Pierce. 2014. Stability and phylogenetic correlation in gut microbiota:
1025 lessons from ants and apes. *Molecular Ecology* 23:1268–1283.

1026 Song, S. J., J. G. Sanders, F. Delsuc, J. Metcalf, K. Amato, M. W. Taylor, F. Mazel, et al.

1027 2020. Comparative analyses of vertebrate gut microbiomes reveal convergence
1028 between birds and bats. *mBio* 11:1–14.

1029 Vázquez, D. P., N. P. Chacoff, and L. Cagnolo. 2009. Evaluating multiple
1030 determinants of the structure of plant–animal mutualistic networks. *Ecology*
1031 90:2039–2046.

1032 Xing, X., H. Jacquemyn, X. Gai, Y. Gao, Q. Liu, Z. Zhao, and S. Guo. 2019. The impact
1033 of life form on the architecture of orchid mycorrhizal networks in tropical forest.
1034 *Oikos* 128:1254–1264.

1035 Xing, X., Q. Liu, Y. Gao, S. Shao, L. Guo, H. Jacquemyn, Z. Zhao, et al. 2020. The
1036 architecture of the network of orchid–fungus interactions in nine co-occurring
1037 *Dendrobium* species. *Frontiers in Ecology and Evolution* 8:1–10.

1038