# The role of evolutionary modes for trait-based cascades in mutualistic networks

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## Abstract

- 2 The erosion of functional diversity may foster the collapse of ecological systems. Functional diversity is ultimately
- 3 determined by the distribution of species traits. As species traits are a legacy of species evolutionary history, one might
- 4 expect that the mode of trait evolution influences community resistance under the loss of functional diversity. In this
- 5 paper, we investigate the role of trait evolutionary dynamics on the robustness of mutualistic networks undergoing
- 6 the following scenarios of species loss: i) random extinctions, ii) loss of functional distinctiveness and iii) extinctions
- <sup>7</sup> biased towards larger sizes. We simulated networks defined by models of single trait complementary and evolutionary
- 8 modes where traits can arise in recent diversification events with weak phylogenetic signal, in early diversification
- 9 events with strong phylogenetic signal, or as a random walk through evolutionary time. Our simulations show that
- mutualistic networks are especially vulnerable to extinctions based on trait distinctiveness and more robust to random
- extinction dynamics. The networks show an intermediate level of robustness against size-based extinctions. Despite

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the small range of variation in network robustness, our results show that the mode of trait evolution matters for network robustness in all three scenarios. Networks with low phylogenetic signal are more robust than networks with high phylogenetic signal across all scenarios. As a consequence, our results predict that mutualistic networks based upon current adaptations are more likely to cope with extinction dynamics than those networks that are based upon conserved traits.

# Introduction

Understanding how ecological systems respond to disturbances is a central and long-standing issue in theoretical and applied ecology (May 1972, 2001; Pimm 1984; Neutel et al. 2002; Allesina & Tang 2012; Myers et al. 2015; Pires et al. 2015). Given the pace of anthropogenic-induced mass extinction, with species dying out three orders of magnitude faster than the extinction background 21 rate inferred from fossil record (Pimm et al. 2014; Ceballos et al. 2015), the need to understand and 22 predict species extinctions has become a fundamental task to mitigate human impact on ecosystems (Vieira & Almeida-Neto 2014; Ceballos et al. 2015). Ecologists have long acknowledged that the species loss may trigger cascading effects in ecological communities, which might bring other species to extinction and even entire ecosystems to collapse (Estes et al. 1998; Jackson et al. 2001; 26 Colwell et al. 2012; Säterberg et al. 2013; Vieira et al. 2013; Brodie et al. 2014; Vieira & Almeida-Neto 2014). However, to date, most studies that have examined the magnitude of biodiversity loss usually ignore co-extinction processes (Dunn et al. 2009; Vieira & Almeida-Neto 2014; but see Silva et al. 2007 and Strona & Bradshaw 2018).

Mutualistic networks are formed by sets of interacting species, generating mutual benefits for par-31 ticipant species (Bronstein 2001; Bascompte & Jordano 2007, 2014). Mutualistic networks include a wide range of taxonomic groups and interaction types, such as interactions between flowering plants and their animal pollinators and seed dispersers (e.g., Bascompte & Jordano 2007; Muller-Landau & Hardesty 2005; Vizentin-Bugoni et al. 2014), animal cleaning associations (e.g., Wicksten 1998; Guimarães et al. 2007; Sazima et al. 2010) and many forms of human-microbe interactions (Dethlefsen et al. 2007). Mutualistic interactions provide an important model system for 37 understanding properties of ecological communities given their paramount role in shaping ecoevolutionary dynamics, biodiversity patterns, ecosystem functioning (Ferriere & Legendre 2013; 39 Bascompte & Jordano 2014; Scheuning et al. 2015, Guimarães Jr 2020) and, consequently, for their 40 importance to the development of conservation strategies (Kiers et al. 2010; Brodie et al. 2014). Although mutualists can be flexible with regards to their partners (Bascompte and Jordano 2014), extinctions in mutualistic systems have the potential to accelerate biodiversity loss and ecosystem disruption (Kiers et al. 2010).

Among the factors that are recognized as important drivers of mutualistic network organization, species functional traits play a crucial role. Functional traits are behavioral, morphological or ecological characteristics associated with organismal fitness, biotic interactions and/or an ecosystem function of interest (Schmitz et al. 2015; Lefcheck et al. 2015). Functional traits are critical to network organization because they can directly constrain or enable the likelihood of an interaction among two or more individuals, imposing thresholds on trait values for feasible interactions (Santamaría & Rodríguez-Gironés 2007; Vizentin-Bugoni et al. 2014; Minoarivelo & Hui 2016; Bastazini et al. 2017, Guimarães Jr 2020).

Species traits may also affect extinction probability, as taxa with some specific traits, such as large body size, and narrow niche breadth are especially more prone to extinction (Purvis et al. 54 2000; Cardillo et al. 2005; Reynolds et al. 2005; but see Chichorro et al. 2019). The robustness of ecological networks, i.e., the system's tolerance to species loss, has been traditionally evaluated based on scenarios where secondary extinctions are driven by species specialization (i.e., number 57 of interacting partners) and/or on stochastic processes (Solé & Montoya 2001; Dunne et al. 2002; 58 Memmot et al. 2004; Burgos et al. 2007; Rezende et al. 2007a; Pocock et al 2012 but see Vidal et al. 2013). These studies help us to broaden our understanding of ecological resistance. A next step in the analysis of network vulnerability is to explore the role of how ecological and evolutionary factors affect the likelihood of species becoming extinct (Bastazini et al. 2019; but see Vieira et al. 2013; Astegiano et al. 2015). For example, as trait redundancy may play an important role in 63 network robustness, extinctions are expected to have a small effect on robustness if all species 64 are functionally similar (higher trait resemblance), but a large effect if species have different trait values (Fonseca & Ganade, 2001).

As species traits are largely a legacy of their evolutionary history (Grafen 1989; Diniz-Filho et al. 2012; Mouquet et al. 2012), it is expected that the mode of evolution, i.e., how traits arise along the phylogenetic history of a clade (Burin et al. 2021), may play a pivotal role in ecological dynamics, and consequently, the robustness of networks that are losing functional diversity. Furthermore, recent evidence suggests that the loss of functional trait diversity takes a larger toll in ecological communities than taxonomic loss alone, making them more likely to collapse (Galetti et al. 2013; Brodie et al. 2014; Valiente-Banuet et al. 2015; Bastazini et al. 2019; Crooke et al. 2020).

Here, we theoretically explore how different modes of trait evolution may affect the robustness of mutualistic networks undergoing three distinct extinction scenarios of species loss: i) random extinctions, which serves as a baseline scenario; ii) loss of functional distinctiveness (i.e., species disappearing sequentially as a function of their functional distinctiveness); and iii) size trait (i.e. species with larger size-related traits disappearing first). As phylogenetically related species tend to interact with a similar set of species (Rezende et al. 2007a), we predicted that networks formed by species with higher levels of phylogenetic signal in traits would be more robust to secondary extinction, as a result of a more cohesive and redundant structure within the network.

# 82 Modeling Approach and Statistical Analysis

83 Eco-evolutionary Dynamics

We modeled the evolutionary dynamics of bipartite mutualistic networks, formed by two sets of interacting species (Fig. 1). We first produced simulated ultrametric phylogenetic trees of different sizes for each set of species, resulting from a uniform birth-death process (Nee et al. 1994). The size of simulated phylogenetic trees ranged from 10 to 20 species, which generated networks that varied in size, ranging from 20 to 40 species.

Secondly, we simulated the evolution of a single trait using a family of power transformations to the branch lengths of simulated phylogenetic trees (Grafen 1989). These transformations were achieved by raising the height of each phylogenetic tree to a different power, denoted by  $\rho$  (Grafen 1989). The range of powers used in these transformations simulates different evolutionary models (Fig. 1). When the height of a phylogenetic tree is raised to the power of 1, it simulates trait evolution under Brownian motion, as if evolution of traits followed a random walk through evolutionary time (Diniz-Filho et al. 2012), which implies that traits divergence increases linearly with time. Power values smaller than 1 compress deeper branch lengths, and expand them near the tips of the tree, simulating a recent diversification of traits with low phylogenetic signal (Diniz-Filho et al. 2012), while  $\rho$  values larger than 1 increase branch lengths near the root of the tree and simulate early diversification of traits with high phylogenetic signal (Diniz-Filho et al. 2012).

Networks were then generated using the single-trait complementarity model, given by equations 1 and 2, proposed by Santamaría & Rodríguez-Gironés (2007) which assumes that interactions between species can be described by a single trait. This approach emulates ecological systems such as pollination networks formed by flowering plants and birds, in which species interactions can be predicted by flowers' corola length and hummingbird's bill length (Vizentin-Bugoni et al. 2014, 2020; see also: Garibaldi et al. 2015, Stang et al. 2009, Donoso et al. 2017). Following Santamaría & Rodríguez-Gironés (2007) approach, a mean trait value and its variability characterize each species in the network and a pair of species is more likely to interact if their trait values overlap.

In their definition,  $V_i$  and  $W_j$  is the central trait value for species i in one set (e.g., flowering plants) and species j in the other set (e.g., pollinators), respectively, and  $\delta V_i$  and  $\delta W_j$  are the range of variability of each trait for species i and j. Then, the value of each cell in the bi-adjacency matrix, corresponding to this pair of species  $I_{ij}$  will be

$$I_{ij} = 1if|V_i - W_j| < 0.5x(\delta V_i + \delta W_j)$$

$$I_{ij} = 0$$
 otherwise

which means that a pair of species interact if the cell value is equal to one and they do not interact if it is equal to zero. The variability represented by  $\delta V_i$  and  $\delta W_j$  were defined as random variables with uniform distributions in the intervals 0-0.25 (Santamaría & Rodríguez-Gironés 2007). To ensure all species within the network interacted at least with one species from the other set of species, we assigned a random interaction to species that did not have any overlapping trait (Fig. 1).

## 118 Co-extinction analyses

We estimated network robustness (R) based on the area below the Attack Tolerance Curve (ATC; 119 Albert & Barabási 2002; Memmott et al. 2004; Burgos et al. 2007). The ATC is a quantitative 120 description of the network robustness measuring its ability to maintain its structural connectivity 121 as species go extinct. The ATC is contained in the unit square and starts at a value 1 in the y-axis, 122 when no species in one set of species are eliminated and all the species in the other set survive. As 123 species are eliminated, the curve decreases monotonically to 1 in the x -axis as no species in one 124 set survives because all the species in the other set went extinct (for further details see Burgos et al. 2007). R values closer to 1 indicate higher network robustness, i.e., the system is more tolerant to 126 species extinctions. We used three distinct species elimination scenarios. First, we removed species 127 based on their trait distinctiveness, which means that at each time step, the species with the most 128 distinct trait value is eliminated (Bastazini et al. 2019). We estimated the functional distinctiveness of each species, following the approach proposed in Bastazini et al. (2019), using an analogous metric used in phylogenetic studies (Redding et al. 2008). Therefore, we built a functional den-131 drogram based on species trait resemblance and then calculated the functional distinctiveness of each species, defined as the sum of all edge lengths between the species and the root of the dendrogram, with each edge length divided by the number of species in the cluster it subtends (Bastazini et al. 2019). In the second elimination scenario, we simulated species extinctions based on size trait values, eliminating species with larger sizes, as empirical evidence suggest that larger species have a higher chance of dying out (Cardillo et al. 2005, Reynolds et al. 2005; Donoso et al. 2017; but see Chichorro et al. 2019). At last, we eliminated species at random from the higher trophic level, which serves as a baseline scenario to compare the effects of the two functional extinction scenarios.

We compared four evolutionary modes under four distinct scenarios. The four models compared are related to the evolutionary mode generating traits in the species belonging to the higher trophic level (e.g., pollinators, seed dispersers): i) Traits of species diversified recently, presenting low 143 phylogenetic signal ( $\rho = 1e-04$ ); ii) Trait evolution follows a Brownian process, ( $\rho = 1.0$ ); and two 144 models, where traits diversified in the beginning of the evolutionary process, with strong phyloge-145 netic signal ( $\rho = 2.0$  and 5.0, modes iii and iv, respectively). These models were compared in four distinct scenarios according with the evolution of traits in species in the set of species belonging to 147 the lower trophic level (e.g., flowering plants): A) a random combination of evolutionary modes, 148 where phylogenetic signal varies from low ( $\rho = 1e-04$ ) to high phylogenetic signal ( $\rho = 5.0$ ); B) 149 late diversification of traits ( $\rho = 1e-04$ ); C) trait evolution following a Brownian process, ( $\rho = 1.0$ ); 150 and D) early diversification of traits ( $\rho = 5.0$ ). 151

We compared the effects of the evolutionary modes, within each scenario of species extinction, using a Bayesian analysis of variance, based on Jeffreys non-informative priors (Kinas & Andrade 2010). Based on the posterior distribution we calculated 95% Bayesian Credible intervals for each scenario. The posterior distributions of parameters are defined as:

$$p(\mu_g | Data) = St\left(n - G, \overline{y_g}, \frac{S_e}{\sqrt{n_g}}\right)$$

$$p\left(\sigma_{g}^{2} \mid Data\right) = GInv\left(\frac{n-G}{2}, \frac{n-G}{2}S_{e}^{2}\right)$$

where G is the fixed factor representing the evolutionary modes,  $\mu_g$  is the mean robustness response for each scenario, and  $\sigma^2$  is the variance (for further details see Kinas & Andrade 2010). The posterior distribution is simulated first sampling  $\sigma_g^2$ . Then, sampling values are taken from

a multivariate normal distribution with  $\mu_g$  equal to the mean robustness, with covariance matrix giving by

$$\sigma = \sigma_g^2 D_{v_g}$$

$$V_g = \frac{1}{n_g}$$

Due to the theoretical predictions of the association between network structure and robustness

(Bascompte 2009, Bascompte & Jordano 2007, 2014 and references therein), we also evaluated the correlation between network robustness and nestedness and modularity. To do so, we ran another 163 set of simulations (with 1,000 iterations), across four scenarios, depending on the strength of phy-164 logenetic signal of the species in the higher trophic level ( $\rho$  = 1e-04, 1, 2 and 5). Grafen's  $\rho$  varied 165 randomly across the lower trophic level, in each scenario (from 1e-04 to 5). We estimated nested-166 ness using the nested overlap and decreasing fill (NODF) index proposed by Almeida-Neto et al. (2008), and modularity using QuaBiMo algorithm that computes modules, based on a hierarchical 168 representation of species link weights (as we are simulating qualitative networks, all interactions 169 have the same weight) and optimal allocation to modules (Dormann and Strauss 2014). 170 All numerical simulations and statistical analyses were performed in the R environ-171 ment (R Core Team 2012) and the simulation code is available at github (https: 172 //github.com/bastazini/The-role-of-evolutionary-modes-for-trait-173

driven-coextinctions-in-mutualistic-networks-network).

## Results

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The three scenarios of species extinction were stochastically different, leading to different dynamics of co-extinctions. Networks under a process of random extinctions were more robust than networks experiencing trait-driven cascades (Fig. 2). Despite the smaller differences, networks losing species based on trait distinctiveness were less robust than networks losing species based on size (Fig.2;  $p(\mu_{Size\ trait} > \mu_{Trait\ distinctiveness}) = 0.99$ ).

Our simulations show that the mode of trait evolution matters for network robustness losing functional diversity (Fig. 3). Networks with strong phylogenetic signals were less robust to species

extinctions in all three scenarios (Fig. 3). Species traits evolving under Brownian motion led to

intermediate levels of robustness in networks undergoing functional attacks (Fig. 3). Our simulations also showed that the phylogenetic signals between interacting species influenced network robustness under functional attacks (Fig. 4). Furthermore, networks losing species with strong phylogenetic signals were less robust to primary species loss in most of the combinations of phylogenetic signals (Fig. 4). The only exception is when the phylogenetic signal in species traits in the lower trophic level is also strong. In this case, there is a large superposition of the posterior distribution in both scenarios of functional attack (Fig. 4).

The association between network robustness and structure behaved similarly across all scenarios, independently of the strength of phylogenetic signal (Fig. 5). Robustness was positively correlated with nestedness (mean correlation  $\pm$  SD = 0.62  $\pm$  0.10; Fig. 5), and negatively correlated with modularity (-0.63  $\pm$  0.11; Fig 5).

## Discussion

The recent merging of functional and phylogenetic ecology has been contributing to our under-196 standing of the mechanisms underlying species interaction networks (Rezende et al. 2007a,b; 197 Peralta 2016; Bastazini et al. 2017) and the impact of environmental changes on natural commu-198 nities (e.g., Rezende et al. 2007a; Díaz et al. 2013; Astegiano et al. 2015; Bastazini et al. 2019). Here we evaluated trait-based cascades using minimal model systems of phylogenetically structu-200 red mutualistic networks. Although species' traits can drive organismal and organism-environment 201 interactions in a myriad of complex manners, we explored two specific dimensions of functional 202 diversity expected to have strong consequences to ecological dynamics: body size, a dimension of functional diversity with ubiquitous effect on ecological systems (Purvis et al. 2000; Cardillo et al. 204 2005; Reynolds et al. 2005; Seguin et al. 2014; Terzopoulou et al., 2015; Verde Arregoitia, 2016; 205 Chichorro et al. 2019) and trait distinctiveness, a facet of functional diversity that warrants unique 206 and/or rare biological interactions and ecosystem functions (Violle et al. 2017). Our results show 207 that extinction cascades based on trait distinctiveness have a more detrimental effect on network robustness, especially when traits evolve under strong phylogenetic signal. 209

Our results suggest the loss of functional distinctiveness is more detrimental to mutualistic networks than the loss of species with larger size trait values, corroborating empirical results that
demonstrate that the loss of more functionally distinct species have a large effect on network robustness (Bastazini et al. 2019). This effect is due to the fact that the role of "functionally unique"
species cannot be compensated for by the remaining species in the network (Bastazini et al. 2019;
Crooke et al. 2020). Distinct species are irreplaceable components of ecological networks, and yet,

still largely ignored in current conservation frameworks (Crooke et al. 2020). Our results support the importance of targeted conservation efforts on species that have unique roles in ecological systems (Crooke et al. 2020).

As phylogenetically related species tend to interact with a similar set of species (Rezende et al. 219 2007a), we expected that networks exhibiting strong phylogenetic signal would be more robust, as 220 a result of higher trait similarity among species. Contrary to our expectations, this was not the case, 221 and in some situations, strong phylogenetic signal was even associated with reduced robustness. 222 Robustness may be especially reduced when phylogenetic signal in the other set of species is low 223 or when the evolution of traits follows a random walk through evolutionary time in both scenarios of trait based cascades. Scenarios where traits evolve under Brownian motion or traits with weak 225 phylogenetic signal in the lower trophic level suggest that there is a strong coupled phylogenetic 226 response in the set of interacting species, as both scenarios show a proportional response of net-227 work robustness with increasing phylogenetic signal. However, when species in the lower trophic 228 level present low phylogenetic signal, network robustness decreases, whereas in the scenario where species in the lower trophic level have strong phylogenetic signal there is not such a clear trend. 230 Rezende et al. (2007a) suggest that ecological communities in which species interactions present a 231 strong phylogenetic component are more likely to suffer co-extinctions following an initial extinc-232 tion event. Our results corroborate this notion showing that strong phylogenetic signal amplifies 233 the cascading effects of co-extinctions in mutualistic systems.

Although body size has been found to be a fundamental trait capable of predicting species respon-235 se to environmental gradients (Seguin et al. 2014; Fritschie & Olden 2016) and their extinction 236 risk (Purvis et al. 2000; Cardillo et al. 2005; Reynolds et al. 2005; Terzopoulou et al., 2015; Verde Arregoitia, 2016; Chichorro et al. 2019), its effects depend on the threat and responses and can be 238 fairly inconsistent, as size is represented by different aspects among taxonomic groups (Chichorro 239 et al. 2019). As our simulations are independent of taxonomic identity (and therefore generalist), 240 our finding that trait distinctiveness was more important for robustness further supports the in-241 clusion of other traits, or other facets of functional diversity, rather than size-related traits alone. Therefore the use of size traits as an "all-encompassing trait", or "key trait" might be misleading 243 in interaction networks or extinction risk studies. Furthermore, empirical evidence from threate-244 ned birds and mammals (Crooke et al. 2020) show that species are more ecologically distinct on 245 average which, together with our simulation results, reinforces the need for targeted conservation 246 efforts on species based on their functional distinctiveness (Crooke et al. 2020). 247

our simulations support previous findings that robustness should increase with nestedness and

decrease with modularity, that network structure can affect its dynamics (Bascompte 2009, Bascompte & Jordano 2007, 2014), and that species phylogenetic relationship affects the degree of ecological network nestedness (Rezende et al. 2007b). Indeed, we found that nestedness had a positive association with the robustness of the system to loss of species or connections. That is likely because a more cohesive structure of nested networks is more redundant, has more alternative states and provides pathways for the persistence of rare species compared to modular ones, and it will not collapse as easily (Bascompte 2009, Bascompte & Jordano 2007, 2014).

We are aware that there are shortcomings to our simulations. First, single trait models may show 256 a poor fit to empirical data (Santamaría & Rodríguez-Gironés 2007; but see Pires et al. 2011). 257 However, trait matching seems to be common in many mutualistic interactions (Garibaldi et al. 258 2015, Stang et al. 2009, Vizentin-Bugoni et al. 2014, Donoso et al. 2017). Additionally, in our 259 simulations phylogenetic signal is associated with evolutionary process and rate. However, it is 260 important to note that in some situations, this may not be the case, or that this association may be 261 complex (Revell et al. 2008). Other scenarios involving more complex relationships between phy-262 logenetic signal and evolutionary process and rate could bring further insights. Finally, we stress 263 that our framework shares a common shortcoming with similar studies, which assume that a spe-264 cies cannot establish new interactions ("rewire") in the absence of original mutualistic partners 265 (Vizentin-Bugoni et al. 2020), when secondary extinctions can take place every time a species has 266 no surviving partner (Dunne et al. 2002, Memmot et al. 2004, Burgos et al. 2007, Vieira et al. 2013, Astegiano et al. 2015, Bastazini et al. 2019). Although experimental studies have suggested that 268 rewiring may promote higher resistance in seed dispersal networks (Timoteo et al. 2016, Costa 269 et al. 2018), it should not be common in mutualistic networks with strong trait coupling such as 270 the ones simulated here. That is because trait mismatch prevents new interactions (Santamaría & 271 Rodríguez-Gironés 2007, Bascompte 2009, Vizentin-Bugoni et al. 2014). We still lack a deep understanding of the underlying mechanisms driving rewiring in mutualistic networks. For example, 273 different factors such as spatiotemporal co-occurrence, environmental gradients, and species traits 274 and abundances may determine the probability of species to rewire (Vizentin-Bugoni et al. 2020). 275 The inability to correctly account for the factors determining network rewiring or simulations based on an unconstrained rewiring process could lead to an overestimation of network robustness (Costa et al. 2018), which is undesirable from a conservation point of view. 278

## 279 Conclusions

Over the past years, ecologists have greatly advanced our understanding of how mutualistic net-

work robustness is associated with phylogenetic patterns (Rezende et al. 2007a; Vieria & Almeida-Neto 2013; Emer et al. 2019; Bastazini et al. 2019). However, these studies are usually concerned 282 with primary extinctions and the phylogenetic information of just one trophic level or set of species 283 (Vieria & Almeida-Neto 2013; Bastazini et al. 2019). Our simulations demonstrate that cascading 284 effects of co-extinction may spread across taxonomically related species, increasing the erosion 285 of species diversity (Rezende et al. 2007a). Moreover, we show that the interaction between phylogenies of each partite of interacting species may influence network robustness and thus should 287 be considered in studies investigating the association between phylogeny and network robustness. 288 Our models integrating phylogenies of each set of mutualistic species suggest that networks are 289 the most susceptible to collapse when 1) they suffer a targeted attack on more functionally distinct, 290 rather than just larger species, and when 2) species have a strong phylogenetic signal in traits. As a 291 consequence, our results predict that mutualistic networks molded by more recent adaptations are 292 more likely to cope with extinction dynamics than those networks that are based upon conserved 293 traits under trait-based cascades. Despite its simplicity, our in silico approach reveals the import-294 ance of considering phylogenetic patterns to predict co-extinctions, providing a step forward in 295 understanding cascading effects in natural communities, and developing better conservation stra-296 tegies. 297

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- Fig. 1. Examples of some of the possible eco-evolutionary dynamics of bipartite mutualistic net-
- works we adopted in our simulations (six out of the 16 possible combinations). Species interactions
- are denoted in the bi-adjacency matrices along with their evolutionary trees and traits (black cir-
- cles). The size of each circle corresponds to trait values. Graphen's pdefine the tempo and mode

of trait evolution. A single-trait complementarity model defines the probability of interaction of

two species. We assigned a random interaction to species that did not have any overlapping trait,

to ensure all species interacted with at least one from the other set of species. More details are in

the main text.

Fig. 2. Robustness (95% Credible Interval) for different species elimination schemes, based on

trait distinctiveness, trait size and random extinctions.

Fig. 3. Robustness (95% Credible Interval) for each species elimination schemes, under different

<sub>89</sub> phylogentic signal in traits.

Fig. 4. Robustness (95% Credible Interval) for the different evolutionary modes under distinct

extinction scenarios in response to each level of phylogenetic signal ( $\rho$ ) in the other partite (in this

case a theoretical lower trophic level).

Fig. 5. Correlation network, based on Pearson correlation, depicting the association between met-

rics of network structure (modularity and nestedness) and network robustness across the three

scenarios of species loss (Trait size, trait distinctiveness and Random extinctions).

# **Higher trophic level**









