Beyond species: why ecological interactions interaction networks vary through space and time

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- **Abstract**: Community ecology is tasked with the considerable challenge of predicting the 20 structure, and properties, of emerging ecosystems. It requires the ability to understand how 21 and why species interact, as this will allow the development of mechanism-based predictive 22 models, and as such to better characterize how ecological mechanisms act locally on the 23 existence of inter-specific interactions. Here we argue that the current conceptualization of 24 species interaction networks is ill-suited for this task. Instead, we propose that future research must start to account for the intrinsic variability of interaction species interactions, then scale 26 up from here onto complex networks. This can be accompslihed accomplished simply by 27 recognizing that there exists intra-specific variability, in traits or properties related to the establishment of species interactions. By shifting the scale towards population-based 29 processes, we show that this new approach will improve our predictive ability and mechanistic understanding of how species interact over large spatial or temporal scales.

32 Introduction

Interactions between species are the driving force behind ecological dynamics within communities (Berlow et al. 2009). Likely for this reason more than any other, the structure of communities have been described by species interaction networks for over a century (Dunne 2006). Formally an ecological network is a mathematical and conceptual representation of 36 both species, and the interactions they establish. Behind this conceptual framework is a rich 37 and expanding literature whose primary focus has been to quantify how numerical and 38 statistical properties of networks relate to their robustness (Dunne et al. 2002), productivity 39 (Duffy et al. 2007), or tolerance to extinction (Memmott et al. 2004). Although this approach 40 classically focused on food webs (Ings et al. 2009), it has proved particularly successful 41 because it can be applied equally to all types of ecological interactions (Kéfi et al. 2012). This body of literature generally assumes that, short of changes in local densities due to ecological dynamics, networks are inherently static objects. This assumption calls into 44 question the relevance of network studies at biogeographic scales. More explicitly, if two 45 species are known to interact at one location, it is often assumed that they will interact 46 whenever and wherever they co-occur (see e.g. Havens 1992); this neglects the fact that local environmental conditions, species states, and community composition can intervene in the 48 realization of interactions. More recently, however, it has been established that networks are 49 dynamic objects that have structured variation in α , β , and γ diversity, not only with regard to the change of species composition at different locations but also to the fact that the same species will interact in different ways over time or across their area of co-occurrence (Poisot et al. 2012). Of these sources of variation in networks, the change of species composition has been addressed explicitly in the context of networks (Gravel et al. 2011, Dáttilo et al. 2013) and within classical meta-community theory. However, because this literature still tends to assume that interactions happen consistently between species wherever they co-occur, it is ill-suited to address network variation as a whole and needs be supplemented with new concepts and mechanisms. 58

Within the current paradigm, interactions are established between species and are an immutable "property" of a species pair. Starting from empirical observations, expert knowledge, or literature surveys, one could collect a list of interactions for any given species pool. Several studies used this approach to extrapolate the structure of networks over time and space (Havens 1992, Piechnik et al. 2008, Baiser et al. 2012) by considering that the network at *any* location is composed of *all* of the potential interactions known for this species pool. This stands in stark contrast with recent results showing that (i) the identities of interacting species vary over space and (ii) the dissimilarity of interactions is not related to the dissimilarity in species composition (Poisot et al. 2012). The current conceptual and operational tools to study networks therefore leaves leave us poorly equipped to understand the causes of this variation. In this paper, we propose a general research agenda to understand to shift the research agenda towards understanding the mechanisms involved in the variability of co-occurring species interactions.

In contrast to the current paradigm, we propose that future research on interaction networks
should be guided by the following principleprinciples: the existence of an interaction between
two species is the result of a stochastic process involving (i) local traits distributions, (ii) local
abundances, and (iii) higher-order effects by the local environment or species acting "at a
distance" on the interaction; regionally, the observation of interactions results is the result of
the accumulation of local observations. This approach is outlined in Box 1. Although this
proposal is a radical yet intuitive change in the way we think about ecological network
structure, we demonstrate in this paper that it is well supported by empirical and theoretical
results alike. Furthermore, our new perspective is well placed to open the door to novel
predictive approaches integrating a range of key ecological mechanisms. Notably, we propose
in Box 2 that this approach facilitates the study of indirect interactions, for which predictive
approaches have long proved elusive (Tack et al. 2011).

Since the next generation of predictive biogeographic models will need to account for species interactions (Thuiller et al. 2013), it is crucial not to underestimate the fact that these

interactions they are intrinsically variable and exhibit a geographic variability of their own. Indeed, investigating the impact of species interactions on species distributions only makes 87 sense under the implicit assumption that species interactions themselves vary over 88 biogeographical scales. Models of species distributions will therefore increase their predictive 89 potential ability if they account for the variability of ecological interactions. In turn, tighter 90 coupling between species-distribution and interaction-distribution models will provide mode 91 accurate predictions of the properties of emerging ecosystems (Gilman et al. 2010, Estes et al. 2011) and the spatial variability of properties between existing ecosystems. By paying more 93 attention to the variability of species interactions, the field of biogeography will be able to re-visit classical observations typically explained by species-level mechanisms; for example, how does community complexity and function vary along latitudinal gradients, is there information hidden in the co-occurrence or avoidance of species interactions, etc. This predictive effort is made all the more important as both the phenology (Parmesan 2007) and ranges (Devictor et al. 2012) of species occupying different positions in their interactions networks are affected differently by climate change. Predicting that species will move and 100 change while interactions remain the same is probably a very conservative approach to 101 estimating the changes to come, and building explicitly on biological mechanisms is one 102 possible way to overcome this limitation. 103

In this paper, we outline the mechanisms that are involved in the variability of species
interactions over time, space, and environmental gradients. We discuss how they will affect
the structure of ecological networks, and how these mechanisms can be integrated into new
predictive and statistical models (**Box 1**). Most importantly, we show that this approach
integrates classical community ecology thinking and biogeographic questions (**Box 2**) and will
ultimately result in a better understanding of the structure of ecological communities.

110 The dynamic nature of ecological interaction networks

Recent studies on the sensitivity of network structure to environmental change provide some context for the study of dynamic networks. Menke et al. (2012) showed that the structure of a 112 plant-frugivore network changed along a forest-farmland gradient. At the edges between two 113 habitats, species were on average less specialized and interacted more evenly with a larger 114 number of partners than they did in habitat cores. Differences in network structure have also 115 been observed within forest strata that differ in their proximity to the canopy and visitation by 116 birds (Schleuning et al. 2011). Tylianakis et al. (2007) reports a stronger signal of spatial 117 interaction turnover when working with quantitative rather than binary interactions, 118 highlighting the importance of *measuring* rather than assuming (or simply reporting) the 119 existence of interactions. Eveleigh et al. (2007) demonstrated that outbreaks of the spruce 120 budworm were associated with changes in the structure of its trophic network, both in terms 121 of species observed and their interactions. Poisot et al. (2011) used a microbial system of hosts 122 and pathogens to study the impact of productivity gradients on realized infection; when the 123 species were moved from high to medium to low productivity, some interactions were lost and 124 others were gained. As a whole, these results suggest that the existence, and properties, of an 125 interaction are not only contingent on the presence of the two species involved but may also 126 require particular environmental conditions, including the presence or absence of species not directly involved in the interaction.

We argue here that there are three broadly-defined classes of mechanisms that ultimately
determine the realization of species interactions. First, individuals must be in high enough
local relative abundances species must be locally abundant enough for their individuals to
meet; this is the so-called "neutral" perspective of interactions. Second, there must be
phenological or trait matching between individuals, such that an interaction will actually
occur given that the encounter takes place. Finally, the realization of an interaction is
regulated by the interacting organisms' surroundings and should be studied in the context of
indirect interactions.

Population dynamics and neutral processes

Over the recent years, the concept of neutral dynamics has left a clear imprint on the analysis 138 of ecological network structure, most notably in bipartite networks (Blüthgen et al. 2006). 139 Re-analysis of several host-parasite datasets, for example, showed that changes in local species abundances triggers variation in parasite specificity (Vazquez et al. 2005). More generally, it is possible to predict the structure of trophic interactions (Canard et al. 2012) and host-parasite 142 communities (Canard et al. 2014) given only minimal assumptions about the distribution of 143 species abundance. In this section, we review recent studies investigating the consequences of 144 neutral dynamics on the structure of interaction networks and show how variations in 145 population size can lead directly to interaction turnover. 146

147 The basic processes

As noted previously, for an interaction to occur between individuals from two populations, these individuals must first meet, then interact. Assuming that two populations occupy the same location and are active at the same time of the day/year, then the likelihood of an 150 interaction is roughly proportional to the product of their relative abundance (Vázquez et al. 151 2007). This means that individuals from two large populations are more likely to interact than 152 individuals from two small populations, simply because they tend to meet more often. This 153 approach can also be extended to the prediction of interaction strength (Blüthgen et al. 2006, 154 Vázquez et al. 2007), i.e. how strong the consequences of the interaction will be. The neutral 155 perspective predicts that locally-abundant species should have more partners and that 156 locally-rare species should appear more specialized. In a purely neutral model (i.e. 157 interactions happen entirely by chance, although the determinants of abundance can still be 158 non-neutral), the identities of species do not matter, and it becomes easy to understand how 159 the structure of local networks can vary since species vary regionally in abundance. Canard et 160 al. (2012) proposed the term of "neutrally forbidden links" to refer to interactions that are

phenologically feasible but not realized because of the underlying population size
distribution. The identity of these neutrally forbidden links will vary over time and space,
either due to stochastic changes in population sizes or because population size responds
deterministically (*i.e.* non-neutrally) to extrinsic drivers.

166 Benefits for network analysis

It is important to understand how local variations in abundance, whether neutral or not,
cascade up to affect the structure of interaction networks. One approach is to use simple
statistical models to quantify the effect of population sizes on local interaction occurrence or
strength (see *e.g.* Krishna et al. 2008). These models can be extended to remove the
contribution of neutrality to link strength, allowing us to work directly on the interactions as
they are determined by traits (**Box 1**). Doing so allows us to compare the variation of neutral
and non-neutral components of network structure over space and time. To achieve this goal,
however, it is essential that empirical interaction networks (i) are replicated and (ii) include
independent measurements of population sizes.

An additional benefit of such sampling is that these data will also help refine neutral theory. 176 Wootton (2005) made the point that deviations of empirical communities from neutral 177 predictions were most often explained by species trophic interactions which are notoriously, 178 albeit intentionally, absent from the original formulation of the theory (Hubbell 2001). 179 Merging the two views will increase our explanatory power, and provide new ways to test 180 neutral theory in interactive communities; it will also offer a new opportunity, namely to 181 complete the integration of network structure with population dynamics. To date, most 182 studies have focused on the effects of a species' position within a food web on the dynamics of 183 its biomass or abundance (Brose et al. 2006, Berlow et al. 2009, Stouffer et al. 2011, Saavedra 184 et al. 2011). Adopting this neutral perspective brings things full circle since the abundance of 185 a species will also dictate its position in the network: changes in abundance can lead to 186 interactions being gained or lost, and these changes in abundance are in part caused by 187

existing interactions (**Box** 2). For this reason, there is a potential to link species and
interaction dynamics and, more importantly, to do so in a way which accounts for the
interplay between the two. From a practical point of view, this requires repeated sampling of a
system through time, so that changes in relative abundances can be related to changes in
interaction strength (Yeakel et al. 2012). Importantly, embracing the neutral view will force us
to reconsider the causal relationship between resource dynamics and interaction strength
since, in a neutral context, both are necessarily interdependent.

Traits matching in space and time

Once individuals meet, whether they will interact is widely thought to be the product of an array of behavioral, phenotypic, and cultural aspects that can conveniently be referred to as a 197 "trait-based process". Two populations can interact when their traits values allow it, e.g. viruses are able to overcome host resistance, predators can capture the their preys, trees provide enough shading for shorter grasses to grow. Non-matching traits will effectively 200 prevent the existence of an interaction, as demonstrated by Olesen et al. (2011). Under this 201 perspective, the existence of interactions can be mapped onto trait values, and interaction 202 networks will consequently vary along with variation in local trait distribution. In this 203 section, we review how trait-based processes impact network structure, how they can create 204 variation, and the perspective they open for an evolutionary approach. 205

206 The basic processes

There is considerable evidence that, at the species level, interaction partners are selected on
the grounds of matching trait values. Random networks built on these rules exhibit realistic
structural properties (Williams and Martinez 2000, Stouffer et al. 2005). Trait values,
however, vary from population to population within species; it is therefore expected that the

local interactions will be contingent upon traits spatial distribution the spatial distribution of 211 the traits (??). The fact that a species' niche can appear large if it is the aggregation of narrow 212 but differentiated individual or population niches is now well established (Bolnick et al. 2003, 213 Devictor et al. 2010a) and has also reinforced the need to understand intra-specific trait 214 variation to describe the structure and dynamics of communities (Woodward et al. 2010, 215 Bolnick et al. 2011). Nevertheless, this notion has yet to percolate into the literature on 216 network structure despite its most profound consequence: a species appearing generalist at 217 the regional scale can easily be specialized in *each* of the patches it occupies. This reality has 218 long been recognized by functional ecologists, which are now increasingly predicting the 219 variance in traits of different populations within a species (Violle et al. 2012). 220

Empirically, there are several examples of intraspecific trait variation resulting in extreme 221 interaction turnover. A particularly spectacular example was identified by Ohba (2011) who 222 describes how a giant waterbug is able to get hold of, and eventually consume, juveniles from 223 a turtle species. This interaction can only happen when the turtle is small enough for the 224 morphotraits of the bug to allow it to consume the turtle, and as such will vary throughout the 225 developmental cycle of both species. Choh et al. (2012) demonstrated through behavioral assays that prey which evaded predation when young were more likely to consume juvenile predators than the "naive" individuals; their past interactions shaped behavioral traits that 228 alter the network structure over time. These examples show that trait-based effects on 229 networks can be observed even in the absence of genotypic variation (although we discuss this 230 in the next section). 231

From a trait-based perspective, the existence of an interaction is an emergent property of the trait distribution of local populations: variations in one or both of these distributions, regardless of the mechanism involved (development, selection, plasticity, environment), are likely to alter the interaction. Importantly, when interaction-driving traits are subject to environmental forcing (for example, body size is expected to be lower in warm environments, Angilletta et al. (2004)), there can be covariation between environmental conditions and the

occurrence of interactions. Woodward et al. (2012) used macrocosms to experimentally
demonstrate that changes in food-web structure happen at the same time as changes in species
body mass distribution. Integrating trait variation over gradients will provide more predictive
power to models of community response to environmental change.

Benefits for network analysis

Linking spatial and temporal trait variation with network variation will help identify the mechanistic basis of network dissimilarity. From a sampling point of view, having enough data requires that, when interactions are recorded, they are coupled with trait measurements. Importantly, these measurements cannot merely be extracted from a reference database because interactions are driven by local trait values and their matching across populations from different species. Within our overarching statistical framework (**Box 1**), we expect that (i) network variability at the regional scale will be dependent on the variation of populations' 249 traits, and (ii) variation between any series of networks will depend on the covariance between 250 species traits. Although it requires considerably larger quantities of data to test, this approach 251 should allow us to infer a priori network variation. This next generation of data will also help 252 link variation of network structure to variation of environmental conditions. Price (2003) 253 shows how specific biomechanical responses to water input in shrubs can have pleiotropic 254 effects on traits involved in the interaction with insects. In this system, the difference in 255 network structure can be explained because (i) trait values determine the existence of an 256 interaction, and (ii) environmental features determine trait values. We have little doubt that 257 future empirical studies will provide similar mechanistic narratives. 258

At larger temporal scales, the current distribution of traits also reflects past evolutionary
history (Diniz-Filho and Bini 2008). Recognizing this important fact offers an opportunity to
approach the evolutionary dynamics and variation of networks. Correlations between
different species' traits, and between traits and fitness, drive coevolutionary dynamics
(Gomulkiewicz et al. 2000, Nuismer et al. 2003). Both of these correlations vary over space

and time (Thompson 2005), creating patchiness in the processes and outcomes of coevolution.

Trait structure and trait correlations are also disrupted by migration (Gandon et al. 2008,

Burdon and Thrall 2009). Ultimately, understanding of how ecological and evolutionary trait

dynamics affect network structure will provide a mechanistic basis for the historical signal

found in contemporary network structures (Rezende et al. 2007, Eklof et al. 2011, Baskerville

et al. 2011, Stouffer et al. 2012).

Beyond direct interactions

In this section, we argue that, although networks are built around observations of direct interactions like predation or pollination, they also offer a compelling tool with which to address indirect effects on the existence and strength of interactions. Any direct interaction 273 arises from the "physical" interaction of only two species, and, as we have already detailed, these can be modified by local relative abundances and/or species traits. Indirect interactions, on the other hand, are established through the involvement of another party than the two focal species, either through cascading effects (herbivorous species compete with insect laying eggs on plants) or through physical mediation of the environment (bacterial exudates increase 278 the bio-availability of iron for all bacterial species; plants with large foliage provide shade for 279 smaller species). As we discuss in this section, the fact that many (if not all) interactions are 280 indirectly affected by the presence of other species (i) has relevance for understanding the 281 variation of interaction network structure and (ii) can be studied within the classical 282 network-theory formalism. 283

284 The basic processes

Biotic interactions themselves interact (Golubski and Abrams 2011); in other words,
interactions are contingent on the occurrence of species other than those interacting. Because

the outcome of an interaction ultimately affects local abundances (over ecological time scales)
and population trait structure (over evolutionary time scales), all interactions happening
within a community will impact one another. This does not actually mean pairwise
approaches are bound to fail, but it does clamor for a larger scale approach that accounts for indirect effects.

The occurrence or absence of a biotic interaction can either affect either the realization of 292 other interactions (thus affecting the "interaction" component of network β -diversity) or the 293 presence of other species. There are several well-documented examples of one interaction 294 allowing new interactions to happen f, e.g. opportunistic pathogens have a greater success of 295 infection in hosts which are already immunocompromised by previous infections, Olivier 296 2012), or conversely preventing them (, e, g, a) a resident symbiont decreases the infection 297 probability of a new pathogen (Heil and McKey 2003, Koch and Schmid-Hempel 2011 op. 298 @hei03). In both cases, the driver of interaction turnover is the patchiness of species 299 distribution; the species acting as a "modifier" of the probability of interaction is only 300 partially present throughout the range of the other two species, thus creating a mosaic of 301 different interaction configurations. Variation in interaction structure can happen through both cascading and environmental effects: Singer et al. (2004) show that caterpillars change 303 the proportion of different plant species in their diet when parasitized in order to favor low 304 quality items and load themselves with chemical compounds which are toxic for their 305 parasitoids. However, low quality food results in birds having a greater impact on caterpillar 306 populations (Singer et al. 2012). It is noteworthy that in this example, the existence of an a 307 parasitic interaction will affect both the strength, and impact, of other interactions. In terms 308 of their effects on network β -diversity, indirect effects are thus likely to act on components of 309 dissimilarity. A common feature of the examples mentioned here is that pinpointing the exact 310 mechanism through which interactions affect each other often requires a good working 311 knowledge of the system's natural history.

Benefits for network analysis

As discussed in previous sections, improved understanding of why and where species interact should also provide a mechanistic understanding of observed species co-occurrences. 315 However, the presence of species is also regulated by indirect interactions. Recent 316 experimental results showed that some predator species can only be maintained if another 317 predator species is present, since the latter regulates a competitively superior prey and allows 318 for prey coexistence (Sanders and Veen 2012). These effects involving several species and 319 several types of interactions across trophic levels are complex (and for this reason, have been 320 deemed unpredictable in the past, Tack et al. (2011)), and can only be understood by 321 comparing communities in which different species are present/absent. Looking at figure ??, it 322 is also clear that the probability of having an interaction between species i and j ($P(L_{ij})$) is 323 ultimately constrained by the probability of simultaneously observing that individuals of 324 species i and j together will meet assuming random movement, i.e. $P(i \cap i)$. Thus, the existence 325 of any ecological interaction will be contingent upon other ecological interactions driving local 326 co-occurrence (Araújo et al. 2011). Based on this argument, ecological networks cannot be limited to a collection of pairwise interactions. Our view of them needs be updated to account for the importance of the context surrounding these interactions (**Box** 2). From a biogeographic standpoint, it requires us to develop a theory based on interaction 330 co-occurrence in addition to the current knowledge encompassing only species co-occurrence. 331 Araújo et al. (2011) and Allesina and Levine (2011) introduced the idea that competitive 332 interactions can leave a signal in species co-occurrence network. A direct consequence of this 333 result is that, for example, trophic interactions are constrained by species' competitive 334 outcomes before they are ever constrained by e.g. predation-related traits. In order to fully 335 understand interactions and their indirect effects, however, there is a need to develop new 336 conceptual tools to represent effects that interactions have on one another. In a graph 337 theoretical perspective, this would amount to establishing edges between pairs of edges, a task 338 for which there is limited conceptual or methodological background. 339

Conclusions Conclusions

Overall, we argue here that the notion of "species interaction networks" shifts our focus away from the level of organization at which most of the relevant biogeographic processes happen — populations. In order to make reliable predictions about the structure of networks, we need to understand what triggers variability of ecological interactions. In this contribution, we have outlined that there are several direct (abundance-based and trait-based) and indirect (biotic 345 modifiers, indirect effects of co-occurrence) effects to account for. We expect that the relative 346 importance of each of these factors and how precisely they affect the probability of 347 establishing an interaction are likely system-specific; nonetheless, we have proposed a unified 348 conceptual approach to understand them better. 349 At the moment, the field of community ecology is severely data-limited to tackle this 350 perspective. Despite the existence of several spatially- or temporally-replicated datasets (e.g. 351 Schleuning et al. 2011 2012 Menke et al. 2012), it is rare that all relevant information has 352 been measured independently. It was recently concluded, however, that even a reasonably 353 small subset of data can be enough to draw inferences at larger scales (Gravel et al. 2013). 354 Paradoxically, as tempting as it may be to sample a network in its entirety, the goal of 355 establishing global predictions might be better furthered by extremely-detailed 356 characterization of a more modest number of interactions (Rodriguez-Cabal et al. 2013). 357 Assuming that there are indeed statistical invariants in the rules governing interactions, this 358 information will allow us to make verifiable predictions on the structure of the networks. 359 Better still, this approach has the potential to substantially strengthen our understanding of 360 the interplay between traits and neutral effects. Blüthgen et al. (2008) claim that the impact of 361 traits distribution on network structure can be inferred simply by removing the impact of neutrality (population densities), based on the idea that many rare links were instances of sampling artifacts. As illustrated here (e.g, **Box** 2), their approach is of limited generality, as the abundance of a species itself can be directly driven by factors such as trait-environment 365 matching. In addition, there are virtually no datasets that follow a collection of interacting 366

species through both space and time in a replicated fashion. This type of data, although 367 exceedingly tedious to collect, would provide important indications of which mechanisms 368 should be explored to improve our understanding the variability of species interactions. 369 With the accumulation of data Assuming that suitable and accessible empirical data will 370 inevitably accumulate in the coming years, these approaches will rapidly expand our ability to 371 predict the re-wiring of networks under environmental change. The effect of environmental 372 changeis expected to occur because (i) population sizes will be affected by the change and (ii) 373 either. There are two broad mechanisms linking network structure to environmental change: 374 changes in population sizes due to modification of demographic parameters, and plastic or 375 adaptive responses will shift or disrupt the resulting in shifted or disrupted trait distributions. 376 The framework proposed in **Box 1** predicts interaction probabilities under different scenarios. 377 Ultimately, being explicit about the trait-abundance-interaction feedback will provide a better 378 understanding of short-term and long-term dynamics of interaction networks. We illustrate 379 this in Fig. ??. The notion that population sizes have direct effects on the existence of an 380 interaction stands opposed to classical consumer-resource theory, which is one of the bases of 381 network analysis. Considering this an opposition, however, is erroneous. Consumer-resource 382 theory considers a strong effect of abundance on the intensity of interactions (**Box 2**), and itself is a source of (quantitative) variation. Furthermore, these models are entirely determined by variations in population sizes in the limiting case where the coefficient of interactions are 385 similar. As such, any approach seeking to understand the variation of interactions over space 386 ought to consider that local densities are not only a consequence, but also a predictor, of the 387 probability of observing an interaction. The same reasoning can be held for local trait 388 distributions, although over micro-evolutionary time-scales. While trait values determine 389 whether two species are able to interact, they will be modified by the selective effect of species 390 interacting. Therefore, conceptualizing interactions as the outcome of a probabilistic process 391 process regulated by local factors, as opposed to a constant, offers the unprecedented 392 opportunity to investigate feedbacks between different time scales. This is especially 393 important since all of the mechanisms mentioned above are also likely to change rapidly over 394

- spatial scales. The situation in which the phenologies of populations are synchronized locally 395 but not regionally (as shown by Singer and McBride 2012) is an excellent example of when we 396 must integrate these mechanisms into our interpretation of spatial and temporal dynamics. 397
- Over the past decade, many insights have been gained by looking at the turnover of different 398 facets of biodiversity (taxonomic, functional, and phylogenetic) through space (Devictor et al. 399 2010b, Meynard et al. 2011). Here, we propose that there is another oft-neglected side of 400
- biodiversity: species interactions. The perspective we bring forth allows us to unify these 401 dimensions and offers us the opportunity to describe the biogeographic structure of all 402 components of community and ecosystem structure simultaneously.
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Boxes

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Box 1: A mathematical framework for population-level interactions

We propose that the occurrence (and intensity) of ecological interactions at the population level relies on several factors, including relative local abundances and local trait distributions. It is important to tease apart these different factors so as to better disentangle neutral and 414 niche processes. We propose that these different effects can adequately be partitioned using 415 the model a model for the adjacency matrix A, where the probability of an interaction between 416 species i and j is

$$\mathbf{A}_{ij} \propto [\mathcal{N}(i,j) \times \mathcal{T}(i,j)] + \epsilon$$
,

where \mathcal{N} is a function giving the probability that species i and j interact based only on their 418 relative local abundances, (that is, the probability of encounter), and T is a function giving the 419 per encounter probability that species i and j interact based on their trait values. The term ϵ 420 accounts for all higher-order effects, such as indirect interactions, local impact of 421 environmental conditions on the interaction, and impact of co-occurring species. Both of these 422 functions can take any form needed. In several papers, $\mathcal{N}(i,j)$ was expressed as $\mathbf{n}_i \times \mathbf{n}_j$, where 423 **n** is a vector of relative abundances (Canard et al. 2014). The expression of \mathcal{T} can in most 424 cases be derived from mechanistic hypotheses about the observation. For example, Gravel et 425 al. (2013) used the niche model of Williams and Martinez (2000) to predict interactions with 426 the simple rule that T(i, j) = 1 if i can consume j based on allometric rules, and 0 otherwise. 427 Following Rohr et al. (2010), the expression of T can be based on latent variables rather than 428 actual trait values. This simple formulation could be used to partition, at the level of individual interactions, the relative importance of density-dependent and trait-based processes using variance decomposition. Most importantly, it predicts (i) how each of these 431 components will vary over space and (ii) how the structure of the network will be affected by, 432 for example, changes in local abundances or trait distributions. The results provided by this 433 framework will only be as good as the empirical data used, and there is a dire need for a 434 methodological discussion about how "predictor" variables (traits, population sizes, etc.) 435 should be measured in the field, in a way that is not biased by the observation of the 436 interactions. This will prove challenging for some types of interactions; e.g. estimating the 437 population size of parasites is often contingent upon catching and examining hosts. 438 Understanding non-independence between these variables in a system-specific way is a 439 crucial point. 440

This model can further be extended in a spatial context, as

$$\mathbf{A}_{ijx} \propto [\mathcal{N}_x(i_x, j_x) \times \mathcal{T}_x(i_x, j_x)] + \epsilon_{ijx}$$

in which i_x is the population of species i at site x. In this formulation, the ϵ term could include 442 the spatial variation of interaction between i and j over sites, and the covariance between the 443 observed presence of this interaction and the occurrence of species i and j. This can, for 444 example, help address situations in which the selection of prey items is determined by traits, 445 but also by behavioral choices. Most importantly, this model differs from the previous one in 446 that each site x is characterized by a set of functions \mathcal{N}_x , \mathcal{T}_x that may not be identical for all 447 sites considered. For example, the same predator may prefer different prey items in different 448 locations, which will require the use of a different form for \mathcal{T} across the range of locations. 449 (???Gravel et al. (2013) show that it is possible to derive robust approximation for the T450 function even with an incomplete set of data, which gives hope that this framework can be 451 applied even when all species information is not known at all sites (which would be an 452 unrealistic requirement for most realistic systems). Both of these models can be used to partition the variance from existing data or to test which trait-matching function best 454 describes the observed interactions. They also provide a solid platform for dynamical simulations in that they will allow re-wiring the interaction network as a function of trait 456 change and to generate simulations that are explicit about the variability of interactions.

Box 2: Population-level interactions in the classical modelling framework

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As noted in the main text, most studies of ecological networks—particularly food 459 webs—regard the adjacency matrix **A** as a fixed entity that specifies observable interactions on 460 the basis of whether two species co-occur or not. Given this assumption, there is a lengthy 461 history of trying to understand how the strength or organization of these interactions 462 influence the dynamic behavior of species abundance (May 1973). Often, such models take the 463 form

$$\frac{dN_i(t)}{dt} = N_i(t) \left(a_i - \sum_{j \neq i} \alpha_{ij} A_{ij} N_j(t) \right),$$

where a_i is the growth rate of species i (and could, in principle, depend on other species' abundances N) and α_{ij} is the strength of the effect of j on i. In this or just about any related 466 model, direct species-species interaction can influence species abundances but their 467 abundances never feedback and influence the per capita interaction coefficients α_{ij} . They do, 468 however, affect the realized interactions, which are defined by $\alpha_{ij}N_i(t)N_j(t)$, something which 469 is also the case when considering more complicated functional responses (Koen-Alonso 2007). 470 More recently, there have been multiple attempts to approach the problem from the other 471 side. Namely, to understand how factors such as species' abundance and/or trait distributions 472 influence the occurrence of the interactions themselves ($\mathbf{Box} 1$). One potential drawback to 473 that approach, however, is that it still adopts the assumption that the observation of any 474 interaction A_{ij} is only an explicit function of the properties of species i and j (traits and 475 co-occurrence).

Since dynamic models demonstrate quite clearly that non-interacting species can alter each 477 others' abundances (e.g. via apparent competition (Holt and Kotler 1987)), this is a deeply-ingrained inconsistency between the two approaches. Such a simplification does 479 increase the analytical tractability of the problem (Allesina and Tang 2012), but there is little, if any, guarantee that it is ecologically accurate. In our opinion, the "higher-effects" term ϵ in the models presented in **Box 1** is the one with the least straightforward expectations, but it may also prove to be the most important if we wish to accurately describe all of these indirect effects.

A similar problem actually arises in the typical statistical framework for predicting 485 interaction occurrence. Often, one attempts to "decompose" interactions into the component 486 that is explained by species' abundances and the component explained by species' traits (e.g., 487 Box 1). Just like how the underlying functions \mathcal{N} and \mathcal{T} could vary across sites, there could 488 also be feedback between species' abundances and traits, in the same way that we have 489 outlined the feedback between interactions and species' abundances. In fact, given the 490 increasing evidence for the evolutionary role of species-species interactions in explaining 491 extant biodiversity and their underlying traits (Janzen and Martin 1982, Herrera et al. 2002), a framework which assumes relative independence of these different phenomenon is likely starting from an overly-simplified perspective. 494

495 Figures

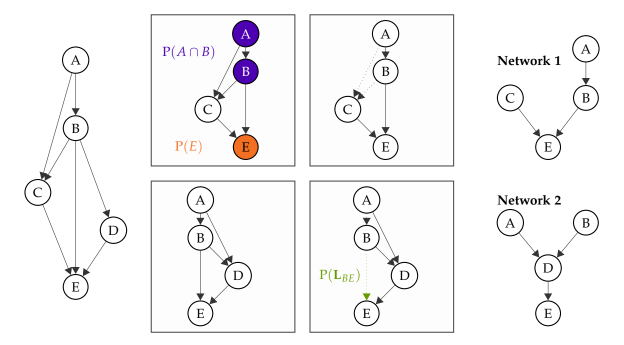


Figure 1: An illustration of the metaweb concept. In its simplest form, a metaweb is the list of all possible species and interactions between them for the system being studied, at the regional level (far left side). Everything that is ultimately observed in nature is a *realisation* of the metaweb (far right side), *i.e.* the resulting network after several sorting processes have occurred (central panel). First, species and species pairs have different probabilities to be observed (top panels). Second, as a consequence of the mechanisms we outline in this paper, not all interactions have the same probability to occur at any given site (bottom panels, see **Box 1**).

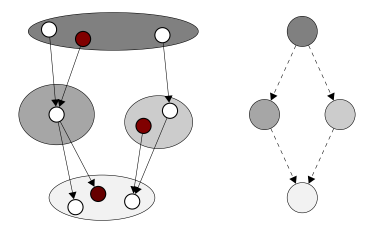


Figure 2: The left-hand side of this figure represents possible interactions between populations (circles) of four species (ellipses), and the aggregated species interaction network on the right. In this example, the populations and species level networks have divergent properties, and the inference on the system dynamics are likely to be different depending on the level of observation. More importantly, if the three populations highlighted in red were to co-occur, there would be no interactions between them, whereas the species-level network would predict a linear chain.

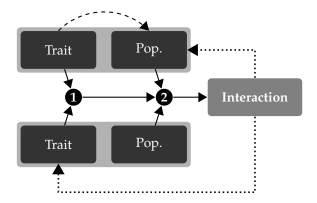


Figure 3: The approach we propose (that populations can interact at the conditions that 1 their trait allow it and 2 they are locally abundant enough for some of their individuals to meet by chance) requires to shift our an increased focus to on population-level processes. A compelling argument to work that supports working at this level of organisation is that eco-evolutionary feedbacks are explicit. All of the components of interaction variability we described are potentially related, either through variations of population sizes due to the interaction itself, or due to selection stemming arising from these variations in population size. In addition, some traits involved in the existence of the interaction may also affect local population abundance.

References

- ⁴⁹⁷ Allesina, S. and Levine, J. 2011. A competitive network theory of species diversity. -
- Proceedings of the National Academy of Sciences of the United States of America 108: 5638.
- Allesina, S. and Tang, S. 2012. Stability criteria for complex ecosystems. Nature 483:
- 500 205-208.
- Angilletta, M. J. et al. 2004. Temperature, Growth Rate, and Body Size in Ectotherms: Fitting
- Pieces of a Life-History Puzzle. Integrative and Comparative Biology 44: 498–509.
- Araújo, M. B. et al. 2011. Using species co-occurrence networks to assess the impacts of
- climate change. Ecography 34: 897–908.
- Baiser, B. et al. 2012. Geographic variation in network structure of a nearctic aquatic food
- web. Global Ecology and Biogeography 21: 579–591.

- Baskerville, E. B. et al. 2011. Spatial Guilds in the Serengeti Food Web Revealed by a Bayesian
- Group Model (LA Meyers, Ed.). PLoS Computational Biology 7: e1002321.
- Berlow, E. L. et al. 2009. Simple prediction of interaction strengths in complex food webs. -
- Proceedings of the National Academy of Sciences of the United States of America 106: 187–91.
- Blüthgen, N. et al. 2006. Measuring specialization in species interaction networks. BMC
- 512 ecology 6: 9.
- Blüthgen, N. et al. 2008. What do interaction network metrics tell us about specialization and
- biological traits? Ecology 89: 3387–99.
- Bolnick, D. I. et al. 2003. The ecology of individuals: incidence and implications of individual
- specialization. The American Naturalist 161: 1–28.
- Bolnick, D. I. et al. 2011. Why intraspecific trait variation matters in community ecology.
- Trends in Ecology and Evolution 26: 183–192.
- Brose, U. et al. 2006. Allometric scaling enhances stability in complex food webs. Ecology
- 520 Letters 9: 1228–1236.
- Burdon, J. J. and Thrall, P. H. 2009. Coevolution of plants and their pathogens in natural
- 522 habitats. Science 324: 755.
- 523 Canard, E. et al. 2012. Emergence of Structural Patterns in Neutral Trophic Networks. PLoS
- 524 One 7: e38295.
- 525 Canard, E. F. et al. 2014. Empirical Evaluation of Neutral Interactions in Host-Parasite
- Networks. The American Naturalist 183: 468–479.
- 527 Choh, Y. et al. 2012. Predator-prey role reversals, juvenile experience and adult antipredator
- behaviour. Scientific Reports in press.
- 529 Dáttilo, W. et al. 2013. Spatial structure of ant–plant mutualistic networks. Oikos: no–no.

- Devictor, V. et al. 2010a. Defining and measuring ecological specialization. Journal of Applied Ecology 47: 15–25.
- Devictor, V. et al. 2010b. Spatial mismatch and congruence between taxonomic, phylogenetic
- and functional diversity: the need for integrative conservation strategies in a changing world.
- Ecology Letters 13: 1030–1040.
- Devictor, V. et al. 2012. Differences in the climatic debts of birds and butterflies at a
- continental scale. Nature Climate Change 2: 121–124.
- Diniz-Filho, J. A. F. and Bini, L. M. 2008. Macroecology, global change and the shadow of
- forgotten ancestors. Global Ecology and Biogeography 17: 11–17.
- Duffy, J. E. et al. 2007. The functional role of biodiversity in ecosystems: incorporating trophic
- complexity. Ecology Letters 10: 522–538.
- Dunne, J. A. 2006. The Network Structure of Food Webs. In: Dunne, J. A. and Pascual, M.
- (eds), Ecological networks: Linking structure and dynamics. Oxford University Press, ppp.
- 543 27-86.
- Dunne, J. A. et al. 2002. Network structure and biodiversity loss in food webs: robustness
- increases with connectance. Ecology Letters 5: 558–567.
- Eklof, A. et al. 2011. Relevance of evolutionary history for food web structure. Proceedings
- of the Royal Society B: Biological Sciences 279: 1588–1596.
- Estes, J. A. et al. 2011. Trophic Downgrading of Planet Earth. Science 333: 301–306.
- Eveleigh, E. S. et al. 2007. Fluctuations in density of an outbreak species drive diversity
- cascades in food webs. Proceedings of the National Academy of Sciences of the United States
- of America 104: 16976–16981.
- 552 Gandon, S. et al. 2008. Host-parasite coevolution and patterns of adaptation across time and

- space. Journal of Evolutionary Biology 21: 1861–1866.
- 6554 Gilman, S. E. et al. 2010. A framework for community interactions under climate change.
- 555 Trends in Ecology and Evolution 25: 325–331.
- Golubski, A. J. and Abrams, P. A. 2011. Modifying modifiers: what happens when
- interspecific interactions interact? Journal of Animal Ecology 80: 1097–1108.
- Gomulkiewicz, R. et al. 2000. Hot spots, cold spots, and the geographic mosaic theory of
- coevolution. The American Naturalist 156: 156–174.
- Gravel, D. et al. 2011. Trophic theory of island biogeography. Ecology Letters 14: 1010–1016.
- Gravel, D. et al. 2013. Inferring food web structure from predator-prey body size
- relationships. Methods in Ecology and Evolution in press.
- Havens, K. 1992. Scale and structure in natural food webs. Science 257: 1107–1109.
- Heil, M. and McKey, D. 2003. Protective ant-plant interactions as model systems in ecological
- and evolutionary research. Annual Review of Ecology, Evolution, and Systematics 34:
- 566 425-553.
- Herrera, C. M. et al. 2002. Interaction of pollinators and herbivores on plant fitness suggests a
- pathway for correlated evolution of mutualism-and antagonism-related traits. Proceedings
- of the National Academy of Sciences 99: 16823–16828.
- Holt, R. D. and Kotler, B. P. 1987. Short-Term Apparent Competition. The American
- 571 Naturalist 130: 412–430.
- Hubbell, S. P. 2001. The Unified Neutral Theory of Biodiversity and Biogeography. Princeton
- 573 University Press.
- Ings, T. C. et al. 2009. Ecological networks–beyond food webs. Journal of Animal Ecology 78:
- 575 253-269.

- Janzen, D. H. and Martin, P. S. 1982. Neotropical anachronisms: the fruits the gomphotheres ate. Science 215: 19–27.
- Kéfi, S. et al. 2012. More than a meal...integrating non-feeding interactions into food webs. -
- 579 Ecology Letters 15: 291–300.
- Koch, H. and Schmid-Hempel, P. 2011. Socially transmitted gut microbiota protect bumble
- bees against an intestinal parasite. PNAS: 1110474108.
- Koen-Alonso, M. 2007. A process-oriented approach to the multispecies functional response. -
- In: From energetics to ecosystems: the dynamics and structure of ecological systems. Springer,
- 584 ppp. 1–36.
- Krishna, A. et al. 2008. A neutral-niche theory of nestedness in mutualistic networks. Oikos
- 586 117: 1609–1618.
- May, R. M. 1973. Stability in randomly fluctuating versus deterministic environments. -
- American Naturalist: 621–650.
- Memmott, J. et al. 2004. Tolerance of pollination networks to species extinctions. -
- Proceedings of the Royal Society B: Biological Sciences 271: 2605–2611.
- Menke, S. et al. 2012. Plant-frugivore networks are less specialized and more robust at
- forest-farmland edges than in the interior of a tropical forest. Oikos 121: 1553–1566.
- Meynard, C. N. et al. 2011. Beyond taxonomic diversity patterns: how do α , β and γ
- components of bird functional and phylogenetic diversity respond to environmental gradients
- across France? Global Ecology and Biogeography 20: 893–903.
- Nuismer, S. L. et al. 2003. Coevolution between hosts and parasites with partially overlapping
- geographic ranges. Journal of Evolutionary Biology 16: 1337–1345.
- Ohba, S.-y. 2011. Field observation of predation on a turtle by a giant water bug. -

- 599 Entomological Science 14: 364–365.
- Olesen, J. M. et al. 2011. Missing and forbidden links in mutualistic networks. Proceedings.
- ⁶⁰¹ Biological sciences / The Royal Society 278: 725–32.
- Olivier, L. 2012. Are Opportunistic Pathogens Able to Sense the Weakness of Host through
- ⁶⁰³ Specific Detection of Human Hormone? Journal of Bacteriology & Parasitology in press.
- Parmesan, C. 2007. Influences of species, latitudes and methodologies on estimates of
- phenological response to global warming. Global Change Biology 13: 1860–1872.
- Piechnik, D. A. et al. 2008. Food-web assembly during a classic biogeographic study:
- species'"trophic breadth" corresponds to colonization order. Oikos 117: 665–674.
- Poisot, T. et al. 2011. Resource availability affects the structure of a natural
- bacteria-bacteriophage community. Biology Letters 7: 201–204.
- Poisot, T. et al. 2012. The dissimilarity of species interaction networks. Ecology Letters 15:
- 611 1353-1361.
- Price, P. W. 2003. Macroevolutionary Theory on Macroecological Patterns. Cambridge
- 613 University Press.
- Rezende, E. L. et al. 2007. Non-random coextinctions in phylogenetically structured
- mutualistic networks. Nature 448: 925-8.
- Rodriguez-Cabal, M. A. et al. 2013. Node-by-node disassembly of a mutualistic interaction
- web driven by species introductions. Proceedings of the National Academy of Sciences 110:
- 618 16503-16507.
- Rohr, R. P. et al. 2010. Modeling food webs: exploring unexplained structure using latent
- traits. The American naturalist 176: 170–7.
- 621 Saavedra, S. et al. 2011. Strong contributors to network persistence are the most vulnerable to

- extinction. Nature 478: 233–235.
- Sanders, D. and Veen, F. J. F. van 2012. Indirect commensalism promotes persistence of
- secondary consumer species. Biology Letters: 960–963.
- Schleuning, M. et al. 2011. Specialization and interaction strength in a tropical
- plant-frugivore network differ among forest strata. Ecology 92: 26–36.
- Schleuning, M. et al. 2012. Specialization of Mutualistic Interaction Networks Decreases
- toward Tropical Latitudes. Current biology 22: 1925–31.
- Singer, M. C. and McBride, C. S. 2012. Geographic mosaics of species' association: a definition
- and an example driven by plant/insect phenological synchrony. Ecology: 120613103411007.
- 631 Singer, M. S. et al. 2004. Disentangling food quality from resistance against parasitoids: diet
- choice by a generalist caterpillar. The American Naturalist 164: 423–429.
- 633 Singer, M. S. et al. 2012. Tritrophic interactions at a community level: effects of host plant
- species quality on bird predation of caterpillars. The American naturalist 179: 363–74.
- 635 Stouffer, D. B. et al. 2005. Quantitative patterns in the structure of model and empirical food
- 636 webs. Ecology 86: 1301-1311.
- 637 Stouffer, D. B. et al. 2011. The role of body mass in diet contiguity and food-web structure. -
- Journal of Animal Ecology: no-no.
- 639 Stouffer, D. B. et al. 2012. Evolutionary Conservation of Species' Roles in Food Webs. Science
- 640 335: 1489–1492.
- Tack, A. J. M. et al. 2011. Can we predict indirect interactions from quantitative food
- webs?-an experimental approach. The Journal of animal ecology 80: 108-118.
- Thompson, J. N. 2005. The Geographic Mosaic of Coevolution. University Of Chicago Press.

- Thuiller, W. et al. 2013. A road map for integrating eco-evolutionary processes into
- biodiversity models. Ecology Letters 16: 94–105.
- Tylianakis, J. M. et al. 2007. Habitat modification alters the structure of tropical
- 647 host-parasitoid food webs. Nature 445: 202-205.
- Vazquez, D. P. et al. 2005. Species abundance and the distribution of specialization in
- 649 host-parasite interaction networks. Journal of Animal Ecology 74: 946–955.
- Vázquez, D. P. et al. 2007. Species abundance and asymmetric interaction strength in
- ecological networks. Oikos 116: 1120–1127.
- Violle, C. et al. 2012. The return of the variance: intraspecific variability in community
- ecology. Trends in Ecology and Evolution 27: 244–252.
- Williams, R. and Martinez, N. 2000. Simple rules yield complex food webs. Nature 404:
- 655 180–183.
- Woodward, G. et al. 2010. Ecological networks in a changing climate. Advances in Ecological
- 657 Research 42: 71–138.
- 658 Woodward, G. et al. 2012. Climate change impacts in multispecies systems: drought alters
- 659 food web size structure in a field experiment. Philosophical Transactions of the Royal Society
- 660 B: Biological Sciences 367: 2990–2997.
- 661 Wootton, J. T. 2005. Field parameterization and experimental test of the neutral theory of
- biodiversity. Nature 433: 309–12.
- Yeakel, J. D. et al. 2012. Probabilistic patterns of interaction: the effects of link-strength
- variability on food web structure. Journal of The Royal Society Interface: rsif.2012.0481.