

# Inferring biotic interactions from proxies

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**Inferring biotic interactions from functional, phylogenetic and geographical proxies remains one great challenge in ecology. We propose a conceptual framework to infer the backbone of biotic interaction networks within regional species pools. First, interacting groups are identified to order links and remove forbidden interactions between species. Second, additional links are removed by examination of the geographical context in which species co-occur. Third, hypotheses are proposed to establish interaction probabilities between species. We illustrate the framework using published food-webs in terrestrial and marine systems. We conclude that preliminary descriptions of the web of life can be made by careful integration of data with theory.**

## Why infer interactions?

Even if serious gaps in knowledge of biodiversity remain, much progress has been made in determining how many different types of organisms exist (the Linnaean shortfall [1]), what evolutionary relationships connect different lineages to a common ancestor (the Darwinian shortfall [2]), and where different species are distributed (the Wallacean shortfall [3]). Much less is known about the types of interactions that exist among species (the Eltonian shortfall [4]) and the importance of such interactions for the maintenance and evolution of life on earth. One reason why the Eltonian shortfall is so prevalent is that detection of direct and indirect interactions requires significant empirical efforts, even in simple systems. For example, identifying direct species interactions within an ecosystem with only seven species would require accounting for 42 potential links and up to 13 650 links if indirect interactions are considered [5]. Given that most systems have more than seven species, documenting all biotic interactions at any site (let alone across the world) would be a considerable undertaking.

Furthermore, the widespread idea that biotic interactions affect ecological processes mainly at local scales of resolution and extent [6–8] has discouraged researchers working at biogeographical scales from addressing them: why deal with biotic interactions if they do not contribute to understanding the bigger ecological picture? Recent evidence, however, has shown that biotic interactions can affect species ranges [9–11] and co-distribution of species [12] at regional to continental scales, thus potentially mediating biological responses to environmental changes [13]. There is a demand for predictions about the dynamics and functioning of novel ecosystems emerging from differential responses of species to global changes [14,15]. Because determining the nature of all interactions among species is currently beyond reach, the development of a methodological framework for inferring interactions from proxies is timely.

Can biotic interactions be inferred? It has been argued that the study of pairwise interactions between species has failed to provide general principles about the dynamics and organization of communities [16], but the recent upsurge of interest in ecological networks [17] is gradually changing this perspective. We propose that, if appropriately analyzed, existing data on functional traits, geographical distributions, and phylogenies provide a starting point for making predictions about the likelihood of pairwise biotic interactions among species and the general properties of the interaction networks. The proposed framework is complementary to empirical approaches for network construction (wherein links are established only after observations are made [18]), and can be useful when empirical data on interactions are not accessible. Our approach generates the backbone of an interactions network that can be used to remove forbidden interactions and to identify plausible links among species in a given regional species pool.

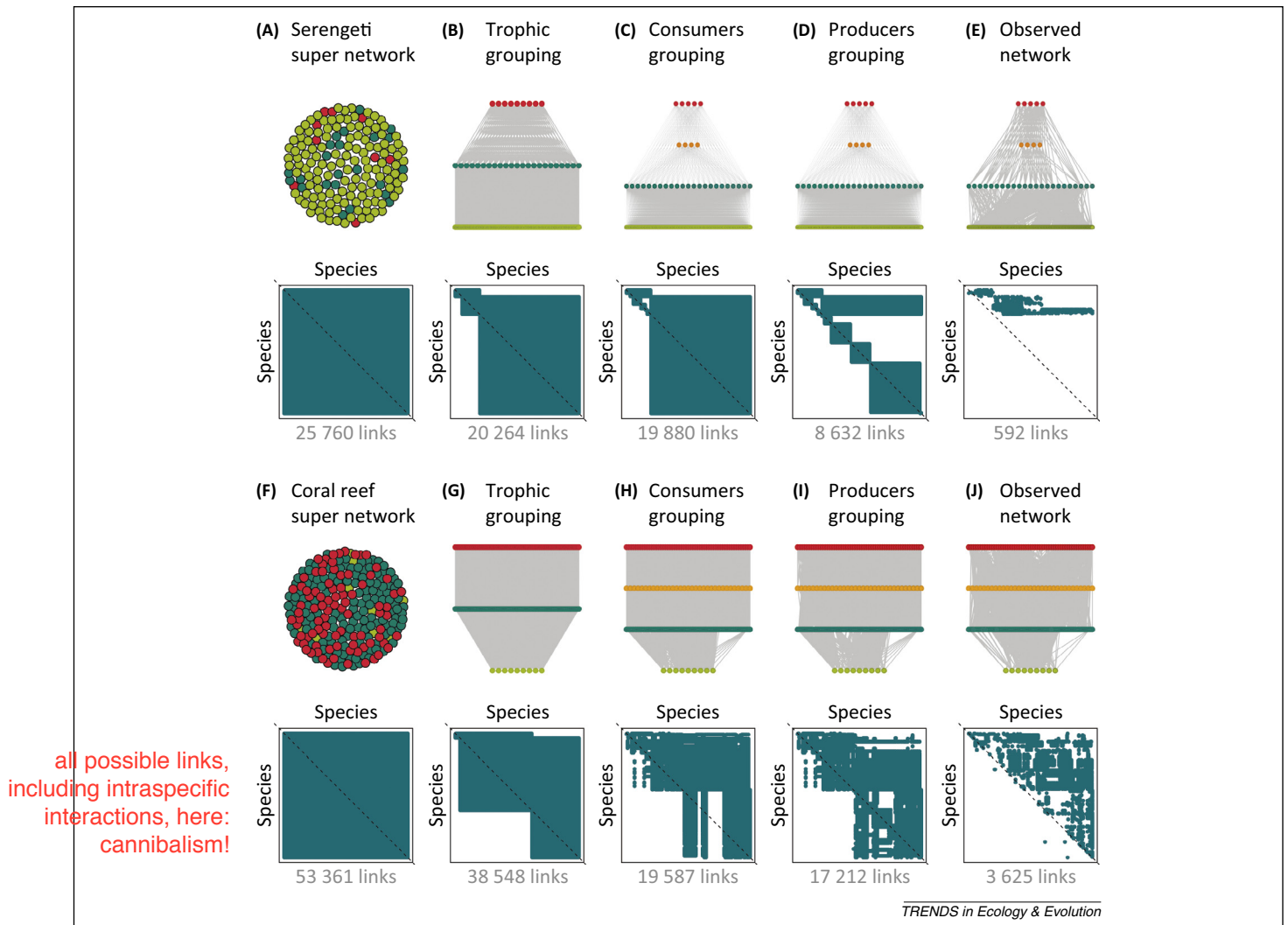
## Which interactions to infer?

There are many different ways to describe a biotic interaction. Interactions may vary in their type (e.g., antagonistic or facilitative), their strength (e.g., weak or strong

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**Figure 1.** Inferring biotic interactions from proxies and testing them with two examples of food-webs from the Serengeti grassland ecosystem (A–E) and from Cuban coral reefs (F–J). The examples differ in the numbers of species and geographic realm but are comparable in scales of extent because data were pooled across large regions (>25 000 km<sup>2</sup>). Whereas the first example comprises 161 species inhabiting the Serengeti National Park, (north of Tanzania), the second comprises 265 species found in all Cuban coral reefs within the 100 m depth platform surrounding the island. By applying three nested successive constraints the number of potential links was reduced by more than two-thirds. The first step involved assigning species into three major trophic groups (B,G): carnivores (red circles), herbivores (dark-green circles) and primary producers (light-green circles). The second step involved classification of consumer species based on their diets (C,H). The third step involved refinement of groups by accounting for the geographic location of species or by establishing where producer species feed in environmental space (D,I). The trophic structure of the two examples is reversed, with more diversity of primary producers in the Serengeti. Even so, the ability to remove forbidden links and match empirical food-webs is similar (i.e., 66.49% for the Serengeti and 67.74% for the Cuban coral reef).

interactions), or their symmetry (e.g., symmetric or asymmetric). An important step for inferring biotic interactions is to determine what information is to be inferred. We propose building interaction networks bottom-up, in other words predicting the links among species and then exploring the collective properties of the networks (e.g., connectance, degree distribution, or distribution of motifs) that emerge from these links (see also [13]). In the proposed framework, four key metrics are the focus of inference:

(i) *The probability of occurrence of a link*

The first step is to infer whether an interaction between any given pair of species is possible. Many species co-occur without interacting with each other [4] and, although detection of links is the focus of much research in community ecology, predicting if a link is absent is of equal importance. Forbidden interactions, such as small fish eating big fish or grassland herbivores eating leaves in trees, can be

easily identified – thus helping to remove links from a full set of candidate links. Such a procedure of pruning forbidden links from potential networks of interactions helps to limit the scope of inference to possible interactions alone (Figure 1). After removing the forbidden links from the tree, the next step is to calculate the probability with which possible links occur using a combination of data that can include traits, phylogenies, and geographical/environmental features [19].

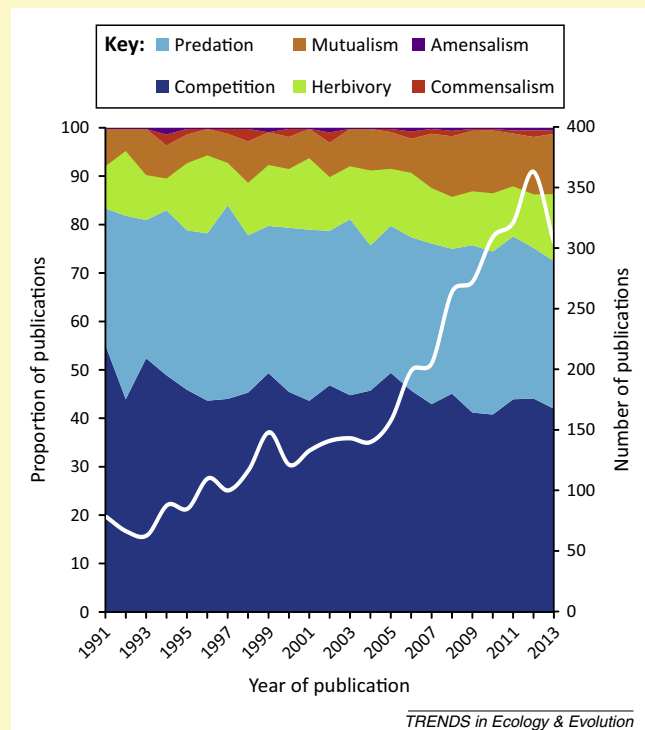
(ii) *The type of interaction*

Once a link between two species is deemed probable, the next step is to determine the expected type of interaction involved (e.g., antagonistic, facilitative, direct, or indirect). Such inference is not always straightforward. Of the myriad of interactions in nature, not all are equally prevalent and not all are equally detectable. Variation in the prevalence and

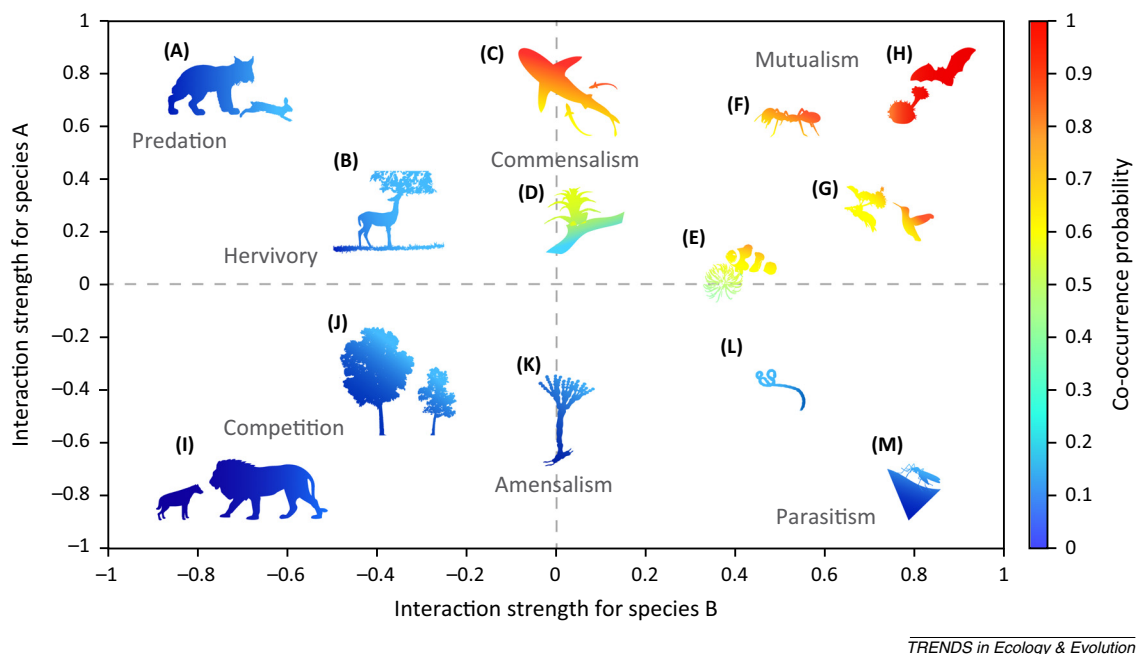
### Box 1. Are all biotic interactions equally prevalent in nature? Can they be predicted from co-occurrence?

Four broad types of biotic interactions can be distinguished based on whether the net effect of the interaction on each interacting species is detrimental for both of them [i.e., competition (–,–)], beneficial for both of them [i.e., mutualism (+,+)], positive for one species and negative for the other [e.g., predation or parasitism (+,–)], or positive for one species and indifferent for the other [i.e., commensalism (+,0)]. Less common is when one interacting species is indifferent while the other is negatively affected [i.e., amensalism (0,–)]. Quantification of the prevalence of the different types of interactions in nature is still lacking, but have they received similar treatment in the literature? A review of the literature of the past two decades reveals that antagonistic interactions (e.g., competition and predation) have been the focus of more than three-quarters of all published studies on biotic interactions (Figure I). These studies often refer to biotic interactions generically, even if only one interaction type – typically competition – is addressed [58].

Why are antagonistic interactions so popular in the ecological literature? (i) Is there evidence that certain types of biotic interactions are more important than others in controlling for the dynamics of species, communities, and ecosystems? (ii) Are antagonistic interactions easier to detect in the field than facilitative interactions? (iii) Are antagonistic interactions more prevalent? Answering these questions requires a comprehensive comparative analysis across all types of biotic interactions. Such analysis is not feasible given the lack of systematic descriptions of biotic interactions across taxa and regions. But simulations may offer an alternative. For example, a recent study provided the first comprehensive simulation of the expected co-occurrence between two species arising from all possible combinations of direct biotic interaction types [20]. The study shows that similar co-occurrences can be achieved by different interactions, leading to the conclusion that co-occurrences alone are not sufficient to provide insight into the biotic interactions generating them (e.g., [20]) (Figure II).



**Figure I.** Published studies on biotic interactions since 1991 categorized by interaction type. Search performed with the Web of Science including the terms 'ecology' AND 'interaction' AND 'interaction type'. Colored sector, proportions by interaction type; white line, total number of publications.



**Figure II.** Interaction strength and probability of co-occurrence. Co-occurrence probability between two different species across biotic interaction space. Biotic interaction space is a representation of all possible types of interactions across two axes, each indicating the direction and the strength of the interaction for each species. Examples are for (A) predation of *Oryctolagus cuniculus* by *Lynx pardinus*, (B) herbivory by *Odocoileus virginianus*, commensalism by (C) *Remora brachyptera* and *Carcharhinus melanopterus*, (D) epiphytic bromeliad (fam. Bromeliaceae), and (E) *Amphiprion percula* and *Entacmea quadricolor*; examples of mutualism for (F) shelter-defense interaction between *Pseudomyrmex ferruginea* and *Cecropia peltata*, of (G) pollination of *Heliconia caribaea* by *Eulampis jugularis* and of (H) pollination of *Stenocereus thurberi* by *Leptonycteris curasoae*; competition between (I) *Panthera leo* and *Crocuta crocuta* and between (J) *Swietenia mahagoni* individuals; amensalism produced by (K) *Penicillium expansum*, and parasitism of (L) virus of genus *Ebolavirus* and (M) *Anopheles gambiae* mosquito, which is itself host for *Plasmodium falciparum*.

detectability of interactions partly explains why different interactions have received different attention. For example, in the past 20 years competition attracted ~50% of all citations in the biotic interactions literature, with commensalism and amensalism being almost residual (Box 1). Species can also interact in more than one way, making the classification of interaction particularly complex. For example, barnacles facilitate the establishment of mussels in the rocky intertidal, while simultaneously competing with them for space. Whether +,– versus –,– interactions are prevalent will depend on the net effect of the interactions, which will in turn determine the resulting pattern of attraction or repulsion [20]. Focusing on easily inferable direct interactions as a starting point (e.g., predator–prey interactions) can help because it will resolve the major interacting groups of the network.

(iii) *The strength of the interaction*

The direct effect that species have on each other's demography is termed interaction strength. Determining the strength of interactions between species is essential to understanding network dynamics, stability, and robustness [21]. Approaches for measuring strength of interactions will vary with the specific types of interactions involved. Whereas interaction strength in a plant–pollinator network can be measured based on the frequency of visits between species pairs [22], in a food-web it can be measured by the amount of energy ingested by the predator [23]. More generally, interaction strength can be measured as the total effect of a population on another (*per population*) or the effect of each individual of a population on another (*per capita*). Most attempts to estimate strength of interactions in the field will mix both. We focus here on the strength of interactions at the population level.

(iv) *Asymmetry of interactions*

The degree to which the strength of the interaction between two species differs between the two is termed asymmetry. Interactions are generally asymmetric [13,22,24] such that if one species A depends strongly on species B, then species B depends weakly on species A. For instance, plants and pollinators vary in their degrees of mutual dependence, with pollinators being typically more specialized than plants [22]. Another example is provided by predator–prey interactions, wherein only a fraction of the prey biomass consumed by the predator is converted into predator biomass. Failure to acknowledge these asymmetries will necessarily lead to overestimating the importance of particular links.

## How to infer interactions?

### *The proxies used in inference*

When direct information about biotic interactions is unavailable, we must resort to indirect information or proxies to obtain insight about them. Three classes of proxies can help with inferring interactions between species: traits, phylogenies, and geographical data (for a review of examples see Table 1). Traits are usually defined as morphological,

physiological, phenological, or behavioral characteristics of species that directly impact on their fitness [25]. However, they are also expected to influence how species interact with one another. When information on traits is unavailable, or is incomplete, one option is to use phylogenies measuring evolutionary relationships among species as a surrogate for trait similarity. This approach is useful provided that there are phylogenetic signals in the traits of interest [26]. Strong phylogenetic discrimination is more likely when lineages are examined across broader geographical scales of extent or large environmental gradients, but strong discrimination is insufficient to guarantee the existence of strong phylogenetic signals between lineages and the traits of interest. Finally, in addition to information on traits, the geographical co-distribution of species, complemented with analysis of species–environment/species–habitat relationships, can also be considered.

The three classes of proxies are not expected to provide equally valuable information (Table 1). Traits ultimately determine if a given interaction is probable, or even possible, because they mediate biotic interactions among species and the interactions between species and the environment. For example, different species might occupy a particular microhabitat differently, and habitat occupancy can determine whether they are likely to compete for a particular resource. When trait information is lacking, phylogenies and geography can help to fill the gaps; for example, by determining which species are likely to co-occur in the same parts of the world. We illustrate below how these proxies (e.g., body size, abiotic requirements, phylogenetic distance, spatial co-occurrence) can be utilized to infer species interactions.

### *Building the backbone of interaction networks*

We propose a hierarchical approach to infer biotic interactions from proxies that is analogous to reconstructing large phylogenies based on collating smaller phylogenies on the top of a structuring backbone (i.e., ‘super-trees’ or ‘meta-trees’). Broad groups of interacting species would, ultimately, form a ‘super-network’ or ‘meta-web’ of interactions. The analogy with a super-tree or meta-tree is not new because it is implicit in the compilation of several networks of interactions [27], but it formalizes what is typically done for network construction.

The procedure for building the backbone of an interaction network starts with the identification of species more likely to share similar interactions. The concept is similar to that of modules [28], but we avoid this terminology because modules are usually determined *a posteriori* and can also refer to simple assemblages of species such as linear food chains or apparent competition [29]. Instead, we define interacting groups based on *a priori* expectations of interactions. The concept is also analogous to that of guilds [30]. Guilds, however, are restricted to species sharing similar resources, and thus do not encompass non-consumptive interactions such as competition or niche construction [27]. A flexible definition of interacting groups based on traits, phylogenies, and geographical distributions would enable combination of heterogeneous information. Such flexibility can be crucial when inferring interactions with information that is not required across



**Table 1. Examples of proxies (traits, phylogenies, and geography) used in inferences of biotic interactions<sup>a</sup>**

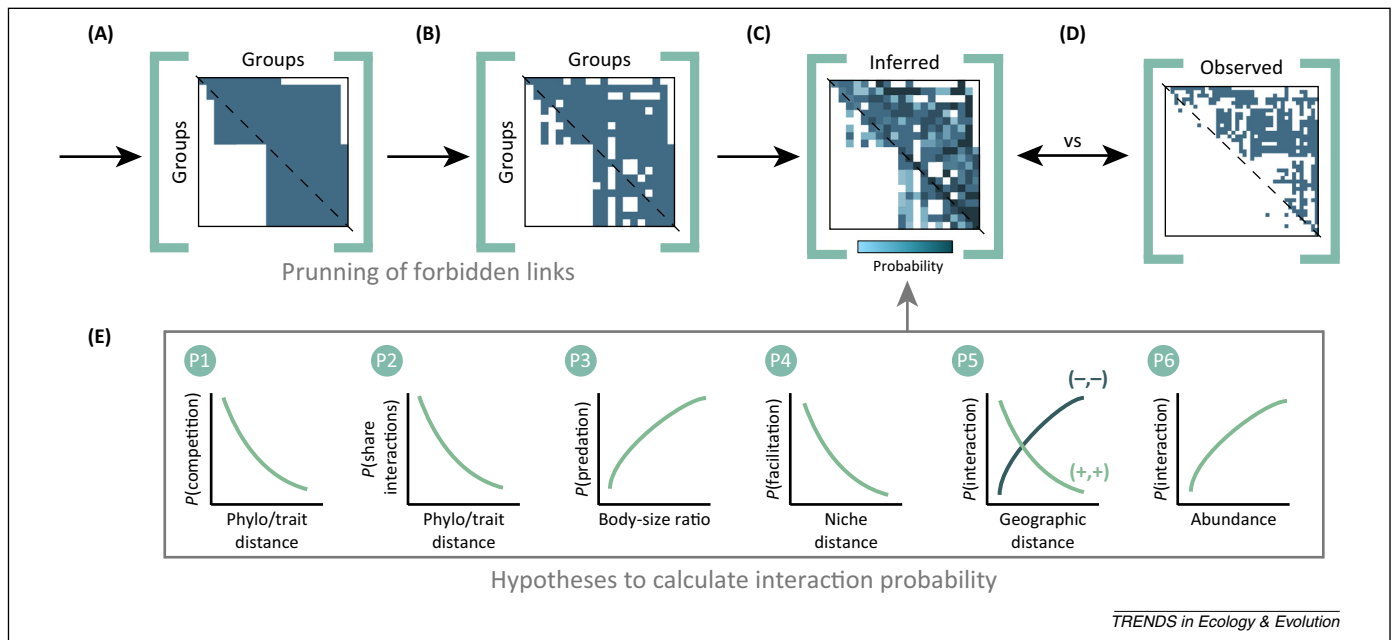
Interaction type	Traits	Phylogeny	Geography
Mutualism (+,+)	Functional composition of subalpine grasslands is related to facilitation when resources are scarce [59]*  Behavioral trait composition can turn the interactions of spider <i>Anelosimus studiosus</i> into mutualism or commensalism [60]*  Flower size can determine the number of pollinator species [61]**	Phylogenetic history plays a relevant role in structuring plant–animal mutualistic networks [62] and could be used to predict co-extinctions [54]***	Distribution of woodpeckers improves species distribution models of boreal owls [10]***  Positive co-occurrence can be used to infer facilitative interactions [63]*  The probability of geographic co-occurrence is higher for (+,+) interactions [20]*
Predation /parasitism (+,-)	Body size is a determinant of consumer–resource interactions in aquatic food-webs [64–66]***, and can also determine interaction strength [67]***  Shifts in body size affect type and strength of predator–prey interactions [68]*  Body temperature can have major implications of our understanding of how thermal stress modulates predator–prey relationships under field conditions [69]*	Phylogenetic data has commonly been used to infer predation in paleontological data [70]***  The phylogenetic signal in host range can be used to predict which plant species are likely to be susceptible to a particular pest or pathogen [71]***  Phylogeny as a proxy for unmeasured trait information explains much of food-web structure [67,72]***	Spatial distribution can affect disease interactions for the foot-and-mouth livestock disease [73]**  In very specialized predator–prey interactions, prey geographic distribution might enhance predictions for the distribution of the predator [48]*
Competition (-,-)	Functional composition of subalpine grasslands is related to facilitation when there is no resource limitation [59]*  The prevalence of competition versus environmental filtering can be inferred based on functional traits [74]**	Plant interactions turn from facilitation into competition with increasing phylogenetic relatedness [75]*  Competition is assumed in community phylogenetics when communities are overdispersed [76,77]**  The prevalence of competition versus environmental filtering is not affected by phylogenetic relationships for tree species [74]**	Pairwise negative residual spatial co-variation indicates potential competitive interactions [78]**  Negative co-occurrence patterns can be used to infer competitive interactions [63,79]*
Commensalism (+,0)	Behavioral trait composition can turn the interactions of spider <i>Anelosimus studiosus</i> into mutualism or commensalism [60]*	Phylogenetic relationships do not seem to affect network structure in an epiphyte–phorophyte network [80]*	Host plant distributions can improve the distribution model for the butterfly <i>Parnassius mnemosyne</i> [9]***  Species abundance, species spatial overlap, and host size largely predicted pairwise interactions and several network metrics [80]*
Amensalism (-,0)	Behavioral trait composition can be related to amensalism of spider <i>Anelosimus studiosus</i> with interacting species [60]*		

<sup>a</sup>The scales at which examples were conducted are annotated as follows: local scale including lab experiments and simulations (\*); regional scale (\*\*); and biogeographic or global scale (\*\*\*).

all species (e.g., foliage chemical defense compounds are not relevant to infer predator–prey interactions).

To illustrate the implementation of these ideas, we infer the backbone of food-webs for two ecosystems: the Serengeti [31]; and the Cuban coral reef [32] (Figure 1). Interacting groups of species are defined *a priori* to simplify the removal of forbidden links. The groups were defined based on the trophic hierarchy of the different species within each ecosystem (e.g., primary producers, grazers, small and large carnivores). This process of trophic classification of species led to identification of forbidden links and removal of ~30% of all potential direct links in the coral reef, and ~22% in the Serengeti (e.g., herbivores eating predators; Figure 1).

Refinement of the species groupings was achieved by considering the characteristics of the consumer species (e.g., distinguishing small versus large carnivores in the Serengeti example, or separating invertebrate feeders, omnivorous, and carnivorous fish in the coral reef example; Figure 1). Geography was then used to subdivide the producer groups, defining environments where only particular consumer species feed (e.g., grassland, woodland, and shrubland are differentiated in the Serengeti, and coral reefs and seagrasses are distinguished in the Cuban network). This step led to a reduction of up to 50% for the remaining links (Figure 1). Notice that the proportion of links removed by the implementation of the framework was similar despite



**Figure 2.** Overview of the hierarchical framework to infer biotic interactions from proxies. (A) Constrained matrix after removal of forbidden links based on trait information. (B) Constrained matrix after removal of forbidden links based on geographical considerations. (C) Filtering of the matrix based on estimated probabilities of interaction (see below) derived from traits, phylogenies, and/or species geographical distributions. (D) Comparison of the inferred interaction matrix against the actual (observed) interaction matrix to evaluate the accuracy of the inference. (E) Probabilities of interaction between groups are derived from the following predictions. P1: The strength of competitive interactions between two species is inversely related to their proximity in trait or phylogenetic space. Functionally similar, or closely related, species are more likely to compete with each other. P2: Closely related species are more likely to share interactions with other species than distantly related species. P3: The probability that a species predares on another in particular food-webs (i.e., marine food-webs) is likely to increase with an increase of their body-size ratio. P4: The more similar the ecological niches of two species, the greater the probability that they will co-occur and hence the higher their interaction probability. P5: Related to P4, interactions will also affect co-distribution. The current understanding of co-occurrence is that negative interactions will generate repulsion; positive interactions aggregation and exploitative interactions (+, -) will generate asymmetric co-occurrence (where one species is most often associated to the other, and the other one less often). P6: Interaction strength will be proportional to the product of relative abundances of the two species (neutral expectation).

the distinct nature of the ecosystems and the reversed structure of the trophic networks (Figure 1).

There are few attempts in the literature at using trait-matching constraints to infer links. For example, Gravel *et al.* [25] used predator–prey body-size relationships to calibrate the niche model [26] and infer potential predator–prey interactions among pelagic Mediterranean fishes. Eklof *et al.* [33] showed that usually fewer than five dimensions were required to represent the structure of 200 networks of different types. When direct measurements of traits are unavailable, latent traits or phylogenetic information can also be used [27]. The ability to establish empirical relationships between traits (measured trait, latent traits, or phylogenetic relationships) and interactions will be key to predicting whether different species coming into contact will interact or not. This would be the case when spread of exotic species leads them to colonize new environments or when species shift their ranges in response to climate change.

Even though we illustrated our framework with examples of food-webs (Figure 1), the procedure is general and can be applied to other types of interactions. For example, after direct trophic interactions are mapped into the backbone of the network, potential indirect interactions, such as exploitative and apparent competition [34,35] or trophic cascades [36] can be inferred [27].

We note that theoretical [37] and experimental [38] studies have shown that pairwise interactions and network structure can be constrained by the environment, and this leads to significant spatial variability in network

structure [39]. In addition, phylogenetic signals in networks can increase with increasing environmental stress – as shown by parasitoids narrowing their host-genotype niche breadth when temperature increases [40]. Further studies will need to address the relationship between the environment and network structure because it is essential to predict feedbacks among species co-distributions, biotic interactions, and environmental change.

### Assigning interaction probabilities

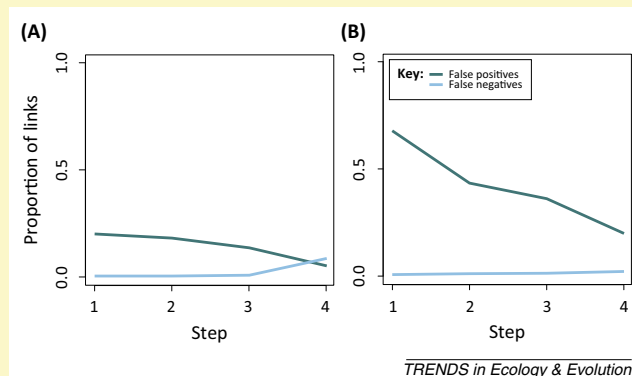
Identifying forbidden links in a network is relatively straightforward compared to the more subtle exercise of assigning probabilities of interactions between species. The null expectation is that, all other things being equal, the probability of interaction between two species is given by their prevalence (i.e., the proportion of an area occupied by the species in a given geographical region) [20,41]. Departures from this null expectation should arise whenever interaction strength between species is different from zero. In such cases, traits or the interaction between traits and the environment should modulate interaction probabilities between species, thus helping to refine the prediction of links between species within the backbone of the interactions network (Figure 2).

Given that probabilities of interactions depend on the strength of the interactions between species, and their asymmetry, calculating these metrics is important. The strength of interactions between pairs of species can be inferred, indirectly, from theoretical models or, directly, by field or laboratory experiments. Examples of the latter

## Box 2. Refining the backbone of an inferred interaction network by assigning probabilities and testing inferred interactions

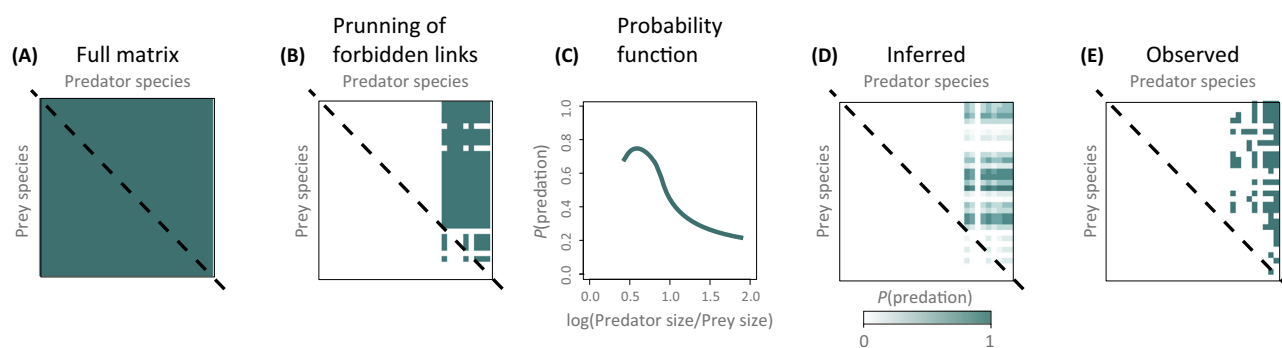
Estimating probabilities of interactions between pairs of species is challenging, but using proxies coupled with simple sets of hypotheses, or rules, can help (see Figure 2 in main text). As an example, we use a simple allometric hypothesis to infer the probabilities of interaction for a subset of mammalian species from the Serengeti food-web and fish species from the Cuban coral reefs food-web. Following the procedure described in Figure 1 in main text, we removed forbidden links from the full matrix of potential interactions (Figure 1A,B,F,G). Next, we removed unlikely interactions based on inferences of probabilities of interaction between pairs of species in the interaction matrix (Figure 1C,D,H,I). To compute probabilities we used a modification of the niche model of food-web structure [81] and considered body size as a main niche axis. We fitted a probabilistic function describing the relationship between predator–prey body sizes (describing the optimum and the range of predator niches) and used it to infer interaction probabilities across all pairs of predator and prey species. We parameterized the model with the observed log-sizes of predator and prey species based on the realized interactions. We show in Figure 1C,H the probability density function across different values of predator–prey body-size ratio for an average predator. We fitted the model to a subset of the data and assigned a probability to each remaining link to obtain the inferred matrix of interactions (Figure 1D,I). Although we restrict this example to allometric relationships, additional hypotheses (e.g., phylogenetic distance, niche distance; see Figure 2 in main text) could be incorporated as subsequent steps. Once interactions are inferred, testing of inferences is carried out against observed interactions (E,J). In this example we show how pruning of forbidden links coupled with

a simple model to infer probabilities of interaction contributed to decrease the false-positive error rate, while leaving false negatives relatively unchanged (Figure 1I). Although the fourth step involving inferences of probabilities of interaction based predator–prey body-size relationships did not enhance the inferences for the Serengeti mammals (Figure 1I), it did slightly improve accuracy for Cuban coral reefs (Figure 1J). Failure to reduce false positives in the Serengeti implies that different hypotheses might be necessary to guide the assignment of probabilities.

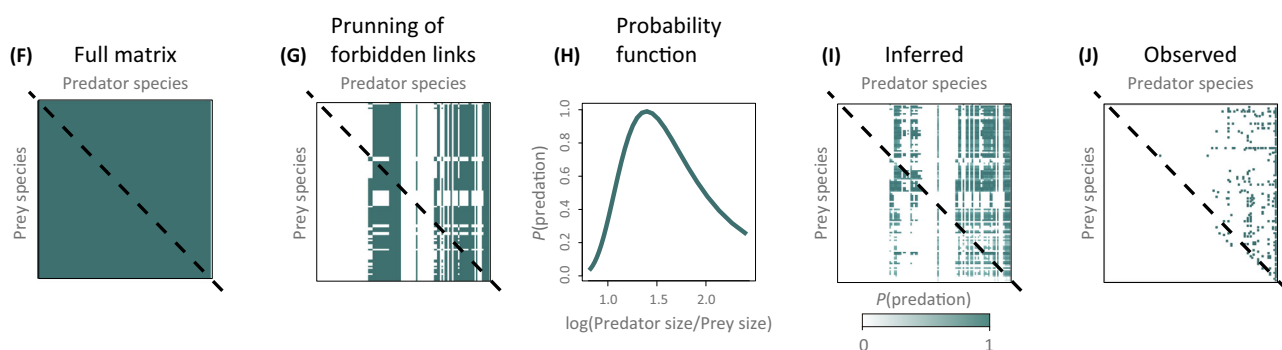


**Figure 1I.** False-positive and false-negative error rates achieved in different steps of the implementation of our framework. (A) Serengeti; (B) Cuban reef.

### Serengeti mammals ( $n = 32$ )



### Cuban coral reef fish ( $n = 116$ )



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**Figure 1.** Predicting probabilities interactions for a subset of 32 mammal species of the Serengeti (A–E) and for 116 fish species present in the Cuban coral reefs dataset (F–J). Forbidden links are initially removed by grouping species based on their trophic level, their traits (size or diet) and their geography (B,G) (as for Figure 1 in main text). Probability density functions for predator–prey body-size ratios (C,H) are then applied to estimate the probability of interaction of each link in the inferred matrices (D,I). Finally, inferred matrices of interaction are compared against the observed interactions (E,J). (see Figure 2 in main text).

have involved examining the role of species in mutualistic networks [42], or changes in the structural complexity of the habitat [43,44] in parasitoid–host networks. Predator or competitor removal experiments [45] have also provided estimates of *per* population interaction strength; *per capita* interactions would then be estimated providing that the predator density is known. Mass-balance models can also be used to estimate interaction strength. For example, de Ruiter *et al.* [46] parameterized Lotka–Volterra equations based on prior identification of trophic links in soil food-webs, measurements of population size, and estimates of natural mortality rates. At equilibrium, the total energy consumed should balance mortality and predation, and therefore it should be possible to derive interaction strength by deduction.

There are also several examples of indirect approaches for inference of interaction strength. For example, predator–prey body-size ratio is often used to estimate *per capita* effect of predators on their prey [47], or to derive this effect from metabolic scaling relationships [48]. In addition to population size, interaction strength has also been associated with the frequency at which actual interactions occur. Neutral models of interactions were shown to predict relatively well the interaction strength between pairs of species within trophic guilds based on their local population sizes [49,50]. In neutral models, the probability of species interacting together is proportional to the product of their relative abundances. Neutral forbidden links arise because two rare species are very unlikely to interact [41]. The network properties are consequently the direct result of frequency distributions of abundance of the different guilds, which in turn respond to changing environmental conditions [43,44]. Increasing understanding of environmental-driven non-random changes in network structure will allow the probabilities of interaction to be adjusted and more accurate predictions to be generated.

The asymmetry of interactions is the most understudied of the factors affecting interaction probabilities. Most modeling studies use a constant across all species to represent consumption inefficiency [51], but theoretical or empirical support for such an assumption is limited. Experimental studies are also usually conducted in a single direction; measuring the effect of predator removal on the prey biomass in the field [52] is often achieved, but it is more challenging to evaluate the effect of prey removal on the predator (but see [53]). The asymmetry of interactions has also been related to co-evolutionary dynamics [54], and it is likely that further indirect inferences of asymmetry will rely on phylogenetic relationships.

Based on the above, we list six simple hypotheses to guide inference about probabilities of interaction between pairs of species (Figure 2). To illustrate the implementation of these hypotheses, we apply one of them (i.e., increasing probability of predation with increasing predator–prey body-size ratio) to infer probability of interaction for the subsets of consumer species within each of our two examples (Serengeti mammals and Cuban fish) (see Figure I in Box 2).

### Testing inferences about interactions

The usefulness of a theoretical model is partly dependent on it being successfully tested. However, inferences of

biotic interactions by models are not easily tested because reliable data on absence of interactions are generally unavailable. Similar problems exist in the literature on modeling species distributions [4,55], with the consequence that inferences of interactions must necessarily be interpreted as potential rather than realized. Indeed, observed interactions will typically constitute a small subset of all realized interactions, and these are themselves a subset of all potential interactions within a given species pool. The consequence is that false positives (interactions predicted but not recorded) might not be an error at all; they often characterize existing but undetected interactions, or potential interactions that have yet not been realized. Although the meaning of false positives is often difficult to ascertain, false negatives (observed interactions not predicted by the model) provide a clear indication that the inferences by the models are inaccurate. Our implementations of the proposed framework show how reductions of false positives are achieved by stepwise implementation of the proposed framework up to estimating the probabilities of interaction based on one rule (i.e., predator–prey body-size ratios), while false negatives remain stable and at low levels (see Figure II in Box 2).

In most cases, inferred interactions will never be observed for several reasons, including low detectability, low abundances, dispersal barriers, and lack of overlap between ecological niches of the two species [19]. Detection of interactions will increase if sufficiently coarse spatial and temporal scales of observation are allowed. Likewise, non-interacting species might start interacting if novel circumstances prompt them to do so (e.g., stochastic long distance dispersal enabling contact of otherwise disjoint species; reshuffling of species geographic distributions due to shifts in the distributions of ecological niches).

In other cases, species might co-occur but interact only in some parts of their ranges. This is the case of Australian tiger sharks, whose diet shifts geographically, selecting different (but equally available) prey species in different locations [56]. The Cuban Coral reef network example shows a substantial number of predicted links below the diagonal for which there is no empirical evidence (see Figure II,J in Box 2). The question arises as to whether these false positives could be realized in other locations; an issue gaining particular prominence for the study of the study of novel ecological communities under climate change [57].

The sheer number of potential interactions that exist in nature poses additional difficulties to testing inferences of interactions with experimental or observational approaches. Nevertheless, not all links affect properties of interactions networks equally and there is scope for simplification if only the most relevant species in the network, for example those with more or stronger connections, are examined [22].

### Concluding remarks

We have proposed a framework for inferring biotic interactions based on stepwise removal of forbidden links and calculation of the probabilities of interaction for the remaining links. With such a process one is able to establish the backbone of an interactions network occurring in a



given regional species pool. The pruning of the network is made using rules derived from the analysis of functional traits, phylogenies, and geographical proxies. To provide an illustration we implemented the framework with food-webs in two ecosystems: terrestrial (the Serengeti) and marine (the Cuban coral reef). We have demonstrated how basic understanding of traits enables accurate predictions of the overall structure of the food-webs with startlingly regional species pools (Figure 1 and Box 2). While further testing of these ideas is warranted, several conceptual challenges remain. Among them, it will be crucial to look more closely at how emerging properties such as mean trophic level, modularity, and nestedness change along the pruning sequence. This issue is reminiscent of the intense debates in the food-web literature about the impact of sampling intensity on network properties [18]. Full understanding of the web of life seems distant, but sequentially building super-networks or meta-webs of biotic interactions networks will help in unraveling key interactions and their potential effects on the distribution of life on earth.

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