# The missing link: discerning true from false negatives when sampling species interaction networks

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Last revision: January 19, 2023

**Abstract:** Ecosystems are composed of networks of interacting species. These interactions allow communities of species to persist through time through both neutral and adaptive processes. Despite their importance, a robust understanding of (and ability to predict and forecast) interactions among species remains elusive. This knowledge-gap is largely driven by a shortfall of data—although species occurrence data has rapidly increased in the last decade, species interaction data has not kept pace, largely due to the effort required to sample interactions. This means there are many interactions between species that occur in nature, but we do not know these interactions occur because we have never observed them. These so-called "false-negatives" bias data and hinder inference about the structure and dynamics of interaction networks. Here, we show the realized number of false-negatives in data can be quite high, even in thoroughly sampled systems, due to variation in abundances in a community. We provide a null model of occurrence detection to estimate the false-negative rate in a given dataset. We also show how to directly incorporate uncertainty due to observation error into model-based predictions of interactions between species. One hypothesis is interactions between "rare" species are themselves rare because these species are less likely to encounter one-another than species of higher relative abundance, and this can (in part) explain the common pattern of nestedness in bipartite interaction networks. However, we demonstrate that across several datasets of spatial/temporally replicated networks, there are positive associations between species co-occurrence and interactions, which suggests these interactions among "rare" species actually exist but simply are not observed. Finally, we assess how false negatives influence various models of network prediction, and recommend directly accounting for observation error in predictive models. We conclude by discussing how the understanding of false-negatives can inform how we design monitoring schemes for species interaction surveys.

**Keywords:** species interactions, network ecology, sampling effort, spatial ecology, null models

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## **Introduction**

Species interactions drive many processes in evolution and ecology. A better understanding of species interactions is an imperative to understand the evolution of life on Earth, to mitigate the impacts of anthropogenic change on biodiversity (Makiola et al. 2020), and for predicting zoonotic spillover of disease to prevent future pandemics (Becker et al. 2021). At the moment we lack sufficient data to meet these challenges (Poisot et al. 2021), largely because species interactions are hard to sample (Jordano 2016). Over the past few decades biodiversity data has become increasingly available through remotely collected data and adoption of open data practices (Kenall et al. 2014; Stephenson 2020). Still, interaction data remains relatively scarce because sampling typically requires human observation. This induces a constraint on the amount, spatial scale, and temporal frequency of resulting data that it is feasible to 10 collect by humans. Many crowdsourced methods for biodiversity data aggregation (e.g. GBIF, eBird) still 11 rely on automated identification of species, which does not easily generalize to interaction sampling. 12 There is interest in using remote methods for interaction sampling, which primarily detect co-occurrence 13 and derive properties like species avoidance from this data (Niedballa et al. 2019). However, co-occurrence itself is not necessarily indicative of an interaction (Blanchet et al. 2020). This is an example of semantic confusion around the word "interaction"—for example one might consider competition a type of species 16 interaction, even though it is marked by a lack of co-occurrence between species, unlike other types of 17 interactions, like predation or parasitism, which require both species to be together at the same place and time. Here we consider interaction in the latter sense, where two species have fitness consequences on 19 one-another if (and only if) they are in the sample place at the same time. In addition, here we only 20 consider direct (not higher-order) interactions. 21 We cannot feasibly observe all (or even most) of the interactions that occur in an ecosystem. This means 22 we can be confident two species actually interact if we have a record of it (assuming they are correctly identified), but not at all confident that a pair of species do not interact if we have no record of those species observed together. In other words, it is difficult to distinguish true-negatives (two species never 25 interact) from false-negatives (two species interact sometimes, but we do not have a record of this interaction). For a concrete example of a false-negative in a food web, see fig. 1. Because even the most highly sampled systems will still contain false-negatives, there is increasing interest in combining 28 species-level data (e.g. traits, abundance, range, phylogenetic relatedness, etc.) to build models to predict

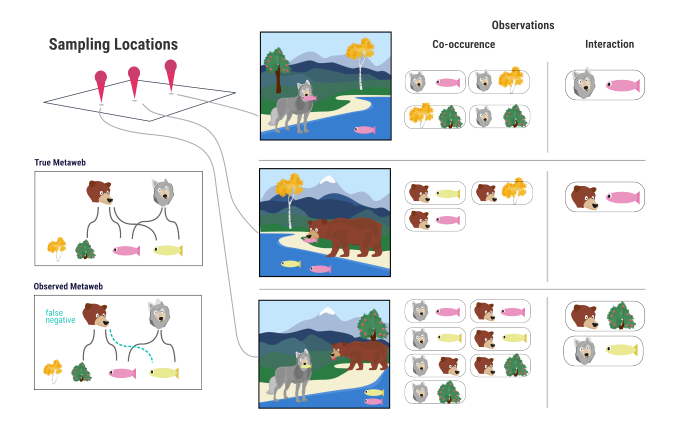


Figure 1: This conceptual example considers a sample of the trophic community of bears, wolves, salmon (pink fish), pike (yellow fish), berry trees, and aspen trees. The true metaweb (all realized interactions across the entire spatial extent) is shown on the left. In the center is what a hypothetical ecologist samples at each site. Notice that although bears are observed co-occurring with both salmon and pike, there was never a direct observation of bears eating pike, even though they actually do. Therefore, this interaction between bears and pike is a false-negative.

- interactions between species we haven't observed together before (Strydom et al. 2021). However, the
- 31 noise of false-negatives could impact the efficacy of our predictive models and have practical
- consequences for answering questions about interactions (de Aguiar et al. 2019). This data constraint is
- amplified as the interaction data we have is geographically biased toward the usual suspects (Poisot et al.
- <sup>34</sup> 2021). We therefore need a statistical approach to assessing these biases in the observation process and
- their consequences for our understanding of interaction networks.
- 36 The importance of sampling effort and its impact on resulting ecological data has produced a rich body of
- 37 literature. The recorded number of species in a dataset or sample depends on the total number of
- observations (Walther et al. 1995; Willott 2001), as do estimates of population abundance (Griffiths 1998).
- This relationship between sampling effort, spatial coverage, and species detectability has motivated more

- quantitatively robust approaches to account for error in sampling data in many contexts: to determine if a given species is extinct (Boakes et al. 2015), to determine sampling design (Moore & McCarthy 2016), and 41 to measure species richness across large scales (Carlson et al. 2020). In the context of interactions, an 42 initial concern was the compounding effects of limited sampling effort combined with the amalgamation of data (across both study sites, time of year, and taxonomic scales) could lead any empirical set of observations to inadequately reflect the reality of how species interact (Paine 1988) or the structure of the 45 network as a whole (Martinez et al. 1999; McLeod et al. 2021). Martinez et al. (1999) showed that in a 46 plant-endophyte trophic network, network connectance is robust to sampling effort, but this was done in the context of a system for which observation of 62,000 total interactions derived from 164,000 plant-stems 48 was feasible. In some systems (e.g. megafauna food-webs) this many observations is either impractical or infeasible due to the absolute abundance of the species in question. The intrinsic properties of ecological communities create several challenges for sampling: first, species are 51 not observed with equal probability—we are much more likely to observe a species of high abundance than one of very low abundance (Poisot et al. 2015). Canard et al. (2012) presents a null model of food-web structure where species encounter one-another in proportion to each species' relative-abundance. This assumes that there are no associations in species co-occurrence due to an interaction (perhaps because this interaction is "important" for both species; Cazelles et al. (2016)), but in this paper we later show 56 increasing strength of these associations leads to increasing probability of false-negatives in interaction 57 data, and that these positive associations are common in existing network data. Second, observed co-occurrence is often equated with meaningful interaction strength, but this is not necessarily the case (Blanchet et al. 2020)—a true "non-interaction" would require that neither of two species, regardless of 60 whether they co-occur, ever exhibit any meaningful effect on the fitness of the other. So, although co-occurrence is not directly indicative of an interaction, it is a precondition for an interaction.
- Here, we illustrate how our confidence that a pair of species never interacts highly depends on sampling
  effort. We demonstrate how the realized false-negative-rate of interactions is related to the relative
  abundance of the species pool, and introduce a method to produce a null estimate of the false-negative-rate
  given total sampling effort (the total count of all interactions seen among all species-pairs) and a method
  for including uncertainty into model predictions of interaction probabilities to account for observation
  error. We then confront these models with data, by showing that positive associations in co-occurrence
  data can increase the realized number of false-negatives and by showing these positive associations are

- 70 rampant in network datasets. We conclude by recommending that the simulation of sampling effort and
- species occurrence can and should be used to help design surveys of species interaction diversity (Moore &
- mcCarthy 2016), and by advocating use of null models like those presented here as a tool for both guiding
- design of surveys of species interactions and for including detection error into predictive models.

## 74 Accounting for false-negatives in species interactions

- In this section, we demonstate how differences in species' relative-abundance can lead to many
- 76 false-negatives in interaction data. We also introduce a method for producing a null estimate of the
- 77 false-negative-rate in datasets via simulation. Because the true false-negative-rate is latent, we can never
- actually be sure how many false-negatives are in our data. However, here we outline an approach to deal
- vith this fact—first by using simulation to estimate the false-negative-rate for a dataset of a fixed size
- 80 using neutral models of observation. We then illustrate how to incorporate uncertainty directly into
- predictions of species interactions to account for observation error based on null estimates of both the
- false-positive rate (as an a priori estimate of species misidentification probability) and false-negative rate
- 83 (as generated via the method we introduce).

#### 84 How many observations of a non-interaction do we need to be confident it's a true

#### 85 negative?

- We start with a naive model of interaction detection: we assume that every interacting pair of species is
- incorrectly observed as not-interacting with an independent and fixed probability, which we denote  $p_{fn}$
- and subsequently refer to as the False-Negative-Rate (FNR). If we observe the same species not-interacting
- N times, then the probability of a true-negative (denoted  $p_{tn}$ ) is given by  $p_{tn} = 1 (p_{fn})^N$ . This relation
- 90 (the cumalitive-distribution-function of geometric distribution, a special case of the negative-binomial
- distribution) is shown in fig. 2(a) for varying values of  $p_{fn}$  and illustrates a fundamental link between our
- <sup>92</sup> ability to reliably say an interaction doesn't exist— $p_{tn}$ —and the number of times N we have observed a
- given species. In addition, note that there is no non-zero  $p_{fn}$  for which we can ever *prove* that an
- interaction does not exist—no matter how many observations of non-interactions N we have,  $p_{tn} < 1$ .
- From fig. 2(a) it is clear that the more often we see two species co-occurring, but not interacting, the more
- 96 likely the interaction is a true-negative. This has several practical consequences: first it means negatives

taken outside the overlap of the range of each species aren't informative because co-occurrence was not possible, and therefore neither was an interaction. In the next section we demonstrate that the distribution of abundance in ecosystems can lead to very high realized values of FNR  $(p_{fn})$  simply as an artifact of sampling effort. Second, we can use this relation to compute the expected number of total observations needed to obtain a "goal" number of observations of a particular pair of species (fig. 2(b)). As an example, if we hypothesize that A and B do not interact, and we want to see species A and B both co-occurring and *not interacting* 10 times to be confident this is a true negative, then we need an expected 1000 observations of all species if the relative abundances of A and B are both 0.1.

#### 55 False-negatives as a product of relative abundance

We now show that the realized FNR changes drastically with sampling effort due to the intrinsic variation 106 of the abundance of individuals of each species within a community. We do this by simulating the process of observation of species interactions, applied both to 243 empirical food webs from the Mangal database 108 (Banville et al. 2021) and random food-webs generated using the niche model, a simple generative model 109 of food-web structure that accounts for allometric scaling (Williams & Martinez 2000). Our neutral model 110 of observation assumes each observed species is drawn in proportion to each species' abundance at that 111 place and time. The abundance distribution of a community can be reasonably-well described by a 112 log-normal distribution (Volkov et al. 2003). In addition to the log-normal distribution, we also tested the 113 case where the abundance distribution is derived from power-law scaling  $Z^{(log(T_i)-1)}$  where  $T_i$  is the 114 trophic level of species i and Z is a scaling coefficient (Savage et al. 2004), which yields the same 115 qualitative behavior. The practical consequence of abundance distributions spanning many orders of 116 magnitude is that observing two "rare" species interacting requires two low probability events: observing two rare species at the same time. 118 To simulate the process of observation, for an ecological network M with S species, we sample relative abundances for each species from a standard-log-normal distribution. For each true interaction in the 120 adjacency matrix M (i.e.  $M_{ij} = 1$ ) we estimate the probability of observing both species i and j at a given 121 place and time by simulating n observations of all individuals of any species, where the species of the 122 individual observed at the  $\{1, 2, ..., n\}$ -th observation is drawn from the generated categorical distribution of abundances. For each pair of species (i, j), if both i and j are observed within the n-observations, the 124 interaction is tallied as a true positive if  $M_{ij} = 1$ . If only one of i or j are observed—but not both—in these



Figure 2: (a) The probability that an observed interaction is a true negative (y-axis) given how many times it has been sampled as a non-interaction (x-axis). Each color reflects a different value of  $p_{fn}$ , the false-negative-rate (FNR)—this is effectively the cumulative distribution function (cdf) of the geometric distribution. (b) The expected number of total observations needed (colors) to observe 10 co-occurrences between a species with relative abundance P(A) (x-axis) and a second species with relative abundance P(Y). (c): false-negative-rate (y-axis) as a function of total sampling effort (x-axis) and network size, computed using the method described above. For 500 independent draws from the niche model (Williams & Martinez (2000)) at varying levels of species richness (colors) with connectance drawn according to the flexible-links model (MacDonald *et al.* (2020)) as described in the main text. For each draw from the niche model, 200 sets of 1500 observations are simulated, for which the mean false-negative-rate at each observation-step is computed. Means denoted with points, with 1 in the first shade and 2 in the second. (d): Same as (c), except using empirical food webs from Mangal database, where richness. The outlier on (d) is a 714 species food-web.

*n* observations, but  $M_{ij} = 1$ , this is counted as a false-negative, and a true-negative otherwise. For each pair of species (i, j), if both i and j are observed within the n-observations, the interaction is tallied as a 127 true positive if  $M_{ij} = 1$ . If only one of i or j are observed—but not both—in these n observations, but 128  $M_{ij} = 1$ , this is counted as a false-negative, and a true-negative otherwise ( $M_{ij} = 0$ ). This process is illustrated conceptually in fig. 3(a). 130 In fig. 2(c) we see this model of observation applied to niche model networks across varying levels of 131 species richness, and in fig. 2(d) the observation model applied to Mangal food webs. For all niche model 132 simulations in this manuscript, for a given number of species S the number of interactions is drawn from 133 the flexible-links model fit to Mangal data (MacDonald et al. 2020), effectively drawing the number of 134 interactions L for a random niche model food-web as

$$L \sim \text{BetaBinomial}(S^2 - S + 1, \mu\phi, 1 - \mu\phi)$$

where the maximum a posteriori (MAP) estimate of  $(\mu, \phi)$  applied to Mangal data from (MacDonald et al.

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2020) is ( $\mu = 0.086$ ,  $\phi = 24.3$ ). All simulations were done with 500 independent replicates of unique niche 137 model networks per unique number of total interactions observed n. All analyses presented here are done 138 in Julia v1.8 (Bezanson et al. 2015) using both EcologicalNetworks.jl v0.5 and Mangal.jl v0.4 (Banville et 139 al. 2021) and are hosted on Github). Note that the empirical data, for the reasons described above, very 140 likely already contains many false-negatives, we'll revisit this issue in the final section. From fig. 2(c) it is evident that the number of species considered in a study is inseparable from the 142 false-negative-rate in that study, and this effect should be taken into account when designing samples of 143 ecological networks in the future. We see a similar qualitative pattern in empirical networks (fig. 2(d)) where the FNR drops off quickly as a function of observation effort, mediated by total richness. The 145 practical consequence of the bottom row of fig. 2 when conducting an analysis is whether there are 146 enough total number of observed interactions (the x-axis) for the threshold FNR we deem acceptable (the y-axis) is feasible. This raises two points: first, empirical data on interactions are subject to the practical 148 limitations of funding and human-work hours, and therefore existing data tend to fall on the order of 149 hundreds or thousands observations of individuals per site. Clear aggregation of data on sampling effort has proven difficult to find and a meta-analysis of network data and sampling effort seems both pertinent 151 and necessary, in addition to the effects of aggregation of interactions across taxonomic scales (Gauzens et

al. 2013; Giacomuzzo & Jordán 2021). This inherent limitation on in-situ sampling means we should optimize where we sample across space so that for a given number of samples, we obtain the maximum 154 information possible. Second, what is meant by "acceptable" FNR? This raises the question: does a shifting 155 FNR lead to rapid transitions in our ability inference and predictions about the structure and dynamics of networks, or does it produce a roughly linear decay in model efficacy? We explore this in the final section. 157 We conclude this section by advocating for the use of neutral models similar to above to generate 158 expectations about the number of false-negatives in a dataset of a given size. This could prove fruitful both 159 for designing surveys of interactions but also because we may want to incorporate models of imperfect 160 detection error into predictive interactions models, as Joseph (2020) does for species occurrence modeling. 161 Additionally, we emphasize that one must consider the context for sampling—is the goal to detect a 162 particular species (as in fig. 2(c)), or to get a representative sample of interactions across the species pool? 163 These arguments are well-considered when sampling individual species (Willott 2001), but have not yet 164 been adopted for designing samples of communities.

#### Including observation error in interaction predictions

Here we show how to incorporate uncertainty into model predictions of interaction probability to account 167 for imperfect observation (both false-negatives and false-positives). Models for interaction prediction typically yield a probability of interaction between each pair of species,  $p_{ij}$ . When these are considered 169 with uncertainty, it is usually model-uncertainty, e.g. the variance in the interaction probability prediction 170 across several cross-validation folds, where the data is split into training and test sets several times. The 171 method we introduce adjusts the value of a model's predictions to produce a distribution of interaction 172 probabilities corrected by a given false-negative-rate  $p_{fn}$  and false-positive-rate  $p_{fp}$  (outlined in figure 173 fig. 3). First we describe how to sample from this distribution of adjusted interaction probabilities via 174 simulation, and show that this distribution can be well-approximated analytically. To get an estimate of each interaction probability that accounts for observation error, we resample the 176 output prediction from an arbitrary model for each interaction  $p_{ij}$  by simulating a set of several 'particles.' 177 Each particle is a realization of an interaction actually occurring assuming  $p_{ij}$  is a correct estimate of the 178 probability of an interaction being observed. Each particle starts as being drawn as true or false according 179 to  $p_{ij}$ , and then adjusting this by the rate of observation error given by  $p_{fp}$  and  $p_{fn}$  to yield a single

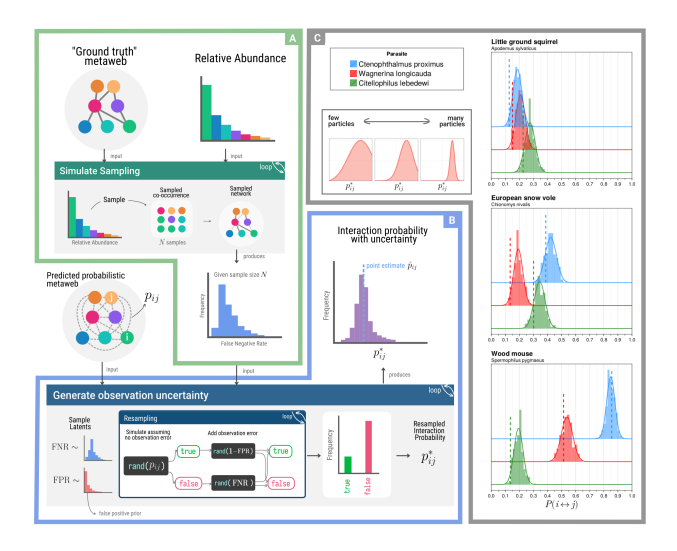


Figure 3: (a) The process for estimating the false-negative-rate (FNR) for an interaction dataset consisting of N total observed interactions. (b) The method for resampling interaction probability based on estimates of false-negative and false-positive rates. (c) The method for interaction probability resampling applied to three mammals and three parasites from the Hadfield  $et\ al.$  (2014) dataset. The original probability  $p_{ij}$  is indicated with a vertical dashed line. The histogram is simulated from the resampling process, and the line indicates the gaussian approximation to this distribution. Both resampling simulations and the gaussian approximation is applied with  $n_p=150$ 

boolean outcome for each particle ("Resampling" within fig. 3(b)). Across of many particles, the resulting frequency of 'true' outcomes is a single resample of the probability  $p_{ij}^*$  that the interaction actually 182 occurred, not just that it was observed. Across several samples each of several particles, this forms a 183 distribution of probabilities which are adjusted by the true and false-negative-rates. There is also an analytic way to approximate this distribution using the normal approximation to binomial. 185 As a reminder, as the total number of samples N from a binomial distribution for n trials with success probability p from approaches infinity, the sum of total successes across all samples approaches a normal 187 distribution with mean np and variance np(1-p). For notation, here we refer to a normal distribution 188 with mean  $\mu$  and standard-deviation  $\sigma$  as  $\mathcal{N}(\mu, \sigma)$ . We can use this to correct the estimate  $p_{ij}$  based on the 189 expected false-negative-rate  $p_{fn}$  and false-positive rate  $p_{fp}$  to obtain the limiting distribution as the 190 number of resamples approaches infinity for the resampled  $p_{ij}^*$  for a given number of particles  $n_p$ . We do 191 this by first adjusting for the rates of observation error to get the mean resampled probability,  $\mathbb{E}[p_{ij}^*]$ , as 192

$$\mathbb{E}[p_{ij}^*] = p_{ij}(1 - p_{fp}) + (1 - p_{ij})p_{fn}$$

193 This yields the normal approximation

$$\sum_{i=1}^{n_p} p_{ij}^* \sim \mathcal{N} \bigg( n_p \cdot \mathbb{E}[p_{ij}^*], \sqrt{n_p \mathbb{E}[p_{ij}^*] (1 - \mathbb{E}[p_{ij}^*])} \bigg)$$

which then can be converted back to a distribution of frequency of successes to yield the final
 approximation

$$p_{ij}^* \sim \mathcal{N}\left(\mathbb{E}[p_{ij}^*], \sqrt{\frac{\mathbb{E}[p_{ij}^*](1 - \mathbb{E}[p_{ij}^*])}{n_p}}\right) \tag{1}$$

We can then further truncate this distribution to remain on the interval (0, 1), as the output is a probability, although in practice often the probability mass outside (0, 1) is extremely low except for  $p_{ij}$  values very close to 0 or 1. As an example case study, we use a boosted-regression-tree to predict interactions in a host-parasite network (Hadfield *et al.* 2014) (with features derived in the same manner as Strydom *et al.* (2021) derives features on this data) to produce a set of interaction predictions. We then applied this method to a set of a few resampled interaction probabilities between mammals and parasite

species shown in figure fig. 3(c).

Why is this useful? For one, this analytic method avoids the extra computation required by simulating 203 samples from this distribution directly. Further, it enables the extension of the natural analogue between 204  $n_p$  (the number of particles) and the number of observations of co-occurrence for a given pair of 205 species—the fewer the particles, the higher the variance of the resulting approximation. The normal 206 approximation is undefined for 0 particles (i.e. 0 observations co-occurrence), although as  $n_p$  approaches 0 207 the approximated normal (once truncated) approaches the uniform distribution on the interval (0, 1), the 208 maximum entropy distribution where we have no information about the possibility of an interaction. 209 This also has implications for what we mean by 'uncertainty' in interaction predictions. A model's 210 prediction can be 'uncertain' in two different ways: (1) the model's predictions may have high variance, or 211 (2) the model's predictions may be centered around a probability of interaction of 0.5, where we are the 212 most unsure about whether this interaction exists. Improving the incorporation of different forms of 213 uncertainty in probabilistic interaction predictions seems a necessary next step toward understanding 214 what pairs of species we know the least about, in order to prioritize sampling to provide the most new 215 information possible.

# Positive associations in co-occurrence increase the false-negative-rate

The model above doesn't consider the possibility that there are positive or negative associations which shift the probability of species cooccurrence away from what is expected based on their relative abundances due 219 to their interaction (Cazelles et al. 2016). However, here we demonstrate that the probability of having a 220 false-negative can be higher if there is some positive association in the occurrence of species A and B. If 221 we denote the probability that we observe the co-occurrence of two species A and B as P(AB) and if there 222 is no association between the marginal probabilities of observing A and observing B, denoted P(A) and 223 P(B) respectively, then the probability of observing their co-occurrence is the product of the marginal 224 probabilities for each species, P(AB) = P(A)P(B). In the other case where there is some positive strength 225 of association between observing both A and B because this interaction is "important" for each species, 226 then the probability of observation both A and B, P(AB), is greater than P(A)P(B) as P(A) and P(B) are 227 not independent and instead are positively correlated, i.e. P(AB) > P(A)P(B). In this case, the probability of observing a single false-negative in our naive model from fig. 2(a) is  $p_{fn} = 1 - P(AB)$ , which due to the

above inequality implies  $p_{fn} > 1 - P(A)P(B)$ . This indicates an increasingly greater probability of a false negative as the strength of association gets stronger,  $P(AB) \rightarrow P(AB) \gg P(A)P(B)$ . However, this still does 231 not consider variation in species abundance in space and time (Poisot et al. 2015). If positive or negative 232 associations between species structure variation in the distribution of P(AB) across space/time, then the 233 spatial/temporal biases induced by data collection would further impact the realized false-negative-rate, as 234 the probability of false negative would not be constant for each pair of species across sites. 235 To test for these positive associations in data we scoured Mangal for datasets with many spatial or temporal replicates of the same system, which led the tresulting seven datasets set in figure fig. 4. For each 237 dataset, we compute the marginal probability P(A) of occurrence of each species A across all networks in 238 the dataset. For each pair of interacting species A and B, we then compute and compare the probability of co-occurrence if each species occurs independently, P(A)P(B), to the empirical joint probability of 240 co-occurrence, P(AB). Following our analysis above, if P(AB) is greater than P(A)P(B), then we expect 241 our neutral estimates of the FNR above to underestimate the realized FNR. In fig. 4, we see the difference between P(AB) and P(A)P(B) for the seven suitable datasets with enough spatio-temporal replicates and a 243 shared taxonomic backbone (meaning all individual networks use common species identifiers) found on 244 Mangal to perform this analysis. Further details about each dataset are reported in tbl. 1. In each of these datasets, the joint probability of co-occurrence P(AB) is decisively greater than our 246 expectation if species co-occur in proportion to their relative abundance P(A)P(B). This suggests that 247 there may not be as many "neutrally forbidden links" (Canard et al. 2012) as we might think, and that the reason we do not have records of interactions between rare species is probably due to observation error. 249 This has serious ramifications for the widely observed property of nestedness seen in bipartite networks 250 (Bascompte & Jordano 2007)—perhaps the reason we have lots of observations between generalists is 251 because they are more abundant, and this is particularly relevant as we have strong evidence that 252 generalism drives abundance (Song et al. 2022a), not vice-versa. 253

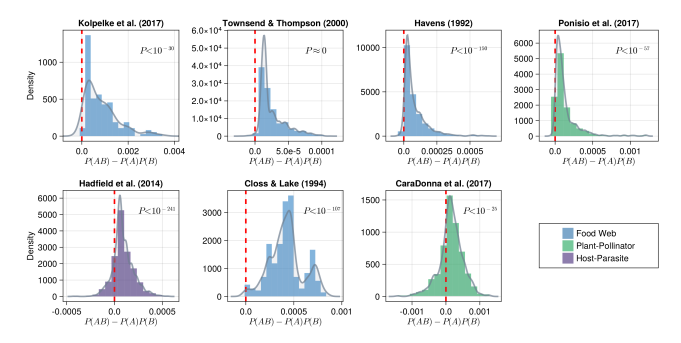


Figure 4: The difference between joint-probability of co-occurrence (P(AB)) and expected probability of co-occurrence under independence (P(A)P(B)) for interacting species for each dataset. The red-dashed line indicates 0 (no association). Each histogram represents a density, meaning the area of the entire curve sums to 1. The continuous density estimate (computed using local smoothing) is shown in grey. The p-value on each plot is the result of a one-sided t-test comparing the mean of each distribution to 0.

Table 1: The datasets used in the above analysis (Fig 2). The table reports the type of each dataset, the total number of networks in each dataset (N), the total species richness in each dataset (S), the connectance of each metaweb (all interactions across the entire spatial-temporal extent) (C), the mean species richness across each local network  $\bar{S}$ , the mean connectance of each local network  $\bar{C}$ , the mean  $\beta$ -diversity among overlapping species across all pairs of network species ( $\bar{\beta}_{OS}$ ), and the mean  $\beta$ -diversity among all species in the metaweb ( $\bar{\beta}_{WN}$ ). Both metrics are computed using KGL  $\beta$ -diversity (Koleff *et al.* 2003)

Network	Туре	N	S	C	$ar{S}$	$ar{C}$	$ar{eta}_{OS}$	$ar{eta}_{WN}$
Kopelke <i>et al.</i> (2017)	Food Web	100	98	0.037	7.87	0.142	1.383	1.972
Thompson & Townsend (2000)	Food Web	18	566	0.014	80.67	0.049	1.617	1.594
Havens (1992)	Food Web	50	188	0.065	33.58	0.099	1.468	1.881
Ponisio et al. (2017)	Pollinator	100	226	0.079	23.0	0.056	1.436	1.870
Hadfield et al. (2014)	Host-Parasite	51	327	0.085	32.71	0.337	1.477	1.952
Closs & Lake (1994)	Food Web	12	61	0.14	29.09	0.080	1.736	1.864
CaraDonna et al. (2017)	Pollinator	86	122	0.18	21.42	0.312	1.527	1.907

## The impact of false-negatives on network properties and prediction

Here, we assess the effect of false-negatives on our ability to make predictions about interactions, as well as their effect on network structure. The prevalence of false-negatives in data is the catalyst for interaction 256 prediction in the first place, and as a result methods have been proposed to counteract this bias (Stock et 257 al. 2017; Poisot et al. 2022). However, it is feasible that the FNR in a given dataset is so high that it could induce too much noise for an interaction prediction model to detect the signal of possible interaction 259 between species. 260 To test this we use the dataset from Hadfield et al. (2014) that describes host-parasite interaction networks 261 sampled across 51 sites, and the same method as Strydom et al. (2021) to extract latent features for each 262 species in this dataset based on applying PCA to the co-occurrence matrix. We then predict a metaweb 263 (equivalent to predicting true or false for an interaction between each species pair, effectively a binary 264 classification problem) from these species-level features using four candidate models for binary 265 classification—three often used machine-learning (ML) methods (Boosted Regression Tree (BRT), 266 Random Forest (RF), Decision Tree (DT)), and one naive model from classic statistics (Logistic Regression 267 (LR)). Each of the ML models are bootstrap aggregated (or bagged) with 100 replicates each. We partition 268 the data into 80-20 training-test splits, and then seed the training data with false negatives at varying rates, 269 but crucially do nothing to the test data. We fit all of these models using MLJ.jl, a high-level Julia 270 framework for a wide-variety of ML models (Blaom et al. 2020). We evaluate the efficacy of these models 271 using two common measures of binary classifier performance: the area under the receiver-operator curve 272 (ROC-AUC) and the area under the precision-recall curve (PR-AUC), for more details see Poisot (2022). 273 Here, PR-AUC is slightly more relevant as it is a better indicator of prediction of false-negatives. The results of these simulations are shown in fig. 5(a & b). 275 One interesting result seen in fig. 5(a & b) is that the ROC-AUC value does not approach random in the same way the PR-AUC curve does as we increase the added FNR. The reason for this is that ROC-AUC is 277 fundamentally not as useful a metric in assessing predictive capacity as PR-AUC. As we keep adding more 278 false-negatives, the network eventually becomes a zeros matrix, and these models can still learn to predict 279 "no-interaction" for all possible species pairs, which does far better than random guessing (ROC-AUC = 280 0.5) in terms of the false positive rate (one of the components of ROC-AUC). This highlights a more broad 281 issue of label class imbalance, meaning there are far more non-interactions than interactions in data. A



Figure 5: **(a)** The area-under the receiver-operator curve (ROC-AUC) and **(b)** The area-under the precision-recall curve (PR-AUC; right) for each different predictive model (colors/shapes) across a spectrum of the proportion of added false-negatives (x-axis). **(c)** The mean trophic-level of all species in a network generated with the niche model across different species richnesses (colors). For each value of the FNR, the mean trophic level was computed across 50 replicates. The shaded region for each line is one standard-deviation across those replicates.

full treatment of the importance of class-balance is outside the scope of this paper, but is explored in-depth in Poisot (2022). Further we see, if anything, gradual decline in the performance of the model until we 284 reach very high FNR levels (i.e.  $p_{fn} > 0.7$ ). This is consistent with other recent work (Gupta *et al.* 2023), 285 although it must be considered that the empircal data on which these models are trained already are almost certain to already contain false-negatives. 287 Although these ML models are surprisingly performant at link prediction given their simplicity, there 288 have been several major developments in applying deep-learning methods to many tasks in network inference and prediction—namely graph-representation learning (GRL, Khoshraftar & An (2022)) and 290 graph convolutional networks (Zhang et al. 2019). At this time, these advances can not yet be applied to 291 ecological networks because they require far more data than we currently have. We already have lots of features that could be used as inputs into these models (i.e. species level data about occurrence, genomes, 293 abundance, etc.), but our network datasets barely get into the hundreds of local networks sampled across 294 space and time (tbl. 1). Once we start to get into the thousands, these models will become more useful, but 295 this can only be done with systematic monitoring of interactions. This again highlights the need to 296 optimize our sampling effort to maximize the amount of information contained in our data given the 297 expense of sampling interactions. 298 We also consider how the FNR affects network properties. In fig. 5(c) we see the mean trophic level across 299 networks simulated using the niche model (as above), across a spectrum of FNR values. In addition to the 300 clear dependence on richness, we see that mean trophic level, despite varying widely between niche model 301 simulations, tends to be relatively robust to false-negatives and does not deviate widely from the true value 302 until very large FNRs. This is not entirely unsurprising. Removing links randomly from a food-web is 303 effectively the inverse problem of the emergence of a giant component (more than half of the nodes are in 304 a connected network) in random graphs (see Li et al. (2021) for a thorough review). The primary 305 difference being that we are removing edges, not adding them, and thus we are witnessing the dissolution 306 of a giant component, rather than the emergence of one. Further applications of percolation theory (Li et 307 al. 2021) to the topology of sampled ecological networks could improve our understanding of how 308 false-negatives impact the inferences about the structure and dynamics on these networks.

#### Discussion

Species interactions enable the persistence and functioning of ecosystems, but our understanding of 31 interactions is limited due to the intrinsic difficulty of sampling them. Here we have provided a null 312 model for the expected number of false-negatives in an interaction dataset. We demonstrated that we 313 expect many false-negatives in species interaction datasets purely due to the intrinsic variation of 314 abundances within a community. We also, for the first time to our knowledge, measured the strength of 315 association between co-occurrence and interactions (Cazelles et al. 2016) across many empirical systems, 316 and found that these positive associations are both very common, and showed algebraically that they increase the realized FNR. We have also shown that false-negatives could further impact our ability to 318 both predict interactions and infer properties of the networks, which highlights the need for further 319 research into methods for correcting this bias in existing data. 320 A better understanding of how false-negatives impact species interaction data is a practical 321 necessity—both for inference of network structure and dynamics, but also for prediction of interactions by 322 using species level information. False-negatives could pose a problem for many forms of inference in 323 network ecology. For example, inferring the dynamic stability of a network could be prone to error if the 324 observed network is not sampled "enough." What exactly "enough" means is then specific to the 325 application, and should be assessed via methods like those here when designing samples. Further, 326 predictions about network rewiring (Thompson & Gonzalez 2017) due to range shifts in response to 327 climate change could be error-prone without accounting for interactions that have not been observed but 328 that still may become climatically infeasible. As is evident from fig. 2(a), we can never guarantee there are 329 no false-negatives in data. In recent years, there has been interest toward explicitly accounting for 330 false-negatives in models (Stock et al. 2017; Young et al. 2021), and a predictive approach to 331 networks—rather than expecting our samples to fully capture all interactions (Strydom et al. 2021). As a 332 result, better models for predicting interactions are needed for interaction networks. This includes 333 explicitly accounting for observation error (Johnson & Larremore 2021)—certain classes of models have 334 been used to reflect hidden states which account for detection error in occupancy modeling (Joseph 2020), 335 and could be integrated in the predictive models of interactions in the future. 336 This work has several practical consequences for the design of surveys for species' interactions. 337 Simulating the process of observation could be a powerful tool for estimating the sampling effort required

by a study that takes relative abundance into account, and provides a null baseline for expected FNR. It is necessary to take the size of the species pool into account when deciding how many total samples is 340 sufficient for an "acceptable" FNR (fig. 2(c & d)). Further the spatial and temporal turnover of interactions 341 means any approach to sampling prioritization must be spatiotemporal. We demonstrated earlier that observed negatives outside of the range of both species aren't informative, and therefore using species 343 distribution models could aid in this spatial prioritization of sampling sites. 344 We also should address the impact of false-negatives on the inference of process and causality in community ecology. We demonstrated that in model food webs, false-negatives do not impact the measure 346 of total trophic levels until very high FNR (figure fig. 5(c)), although we cannot generalize this further to other properties. This has immediate practical concern for how we design what taxa to sample—does it matter if the sampled network is fully connected? It has been shown that the stability of subnetworks can 349 be used to infer the stability of the metaweb paper beyond a threshold of samples (Song et al. 2022b). But 350 does this extend to other network properties? And how can we be sure we are at the threshold at which we 351 can be confident our sample characterizes the whole system? We suggest that modeling observation error 352 as we have done here can address these questions and aid in the design of samples of species interactions. 353 To try to survey to avoid all false-negatives is a fool's errand. Species ranges overlap to form mosaics, 354 which themselves are often changing in time. Communities and networks don't end in space, and the 355 interactions that connect species on the 'periphery' of a given network to species outside the spatial extent 356 of a given sample will inevitably appear as false-negatives in practical samples. The goal should instead be 357 to sample a system enough to have a statistically robust estimate of the current state and empirical change over time of an ecological community at a given spatial extent and temporal resolution, and to determine 359 what the sampling effort required should be prior to sampling. 360

# 361 Acknowledgements

AG & MDC acknowledge the support of the Liber Ero Chair for Biodiversity conservation and NSERC.

## References

- Banville, F., Vissault, S. & Poisot, T. (2021). Mangal.jl and EcologicalNetworks.jl: Two complementary 364 packages for analyzing ecological networks in Julia. Journal of Open Source Software, 6, 2721. 365
- Bascompte, J. & Jordano, P. (2007). Plant-Animal Mutualistic Networks: The Architecture of Biodiversity. 366 Annual Review of Ecology, Evolution, and Systematics, 38, 567–593. 367
- Becker, D.J., Albery, G.F., Sjodin, A.R., Poisot, T., Bergner, L.M., Dallas, T.A., et al. (2021). Optimizing 368 predictive models to prioritize viral discovery in zoonotic reservoirs. 369
- Bezanson, J., Edelman, A., Karpinski, S. & Shah, V.B. (2015). Julia: A Fresh Approach to Numerical 370 Computing. 371
- Blanchet, F.G., Cazelles, K. & Gravel, D. (2020). Co-occurrence is not evidence of ecological interactions. 372 Ecology Letters, 23, 1050–1063. 373
- Blaom, A.D., Kiraly, F., Lienart, T., Simillides, Y., Arenas, D. & Vollmer, S.J. (2020). MLJ: A Julia package 374 for composable machine learning. Journal of Open Source Software, 5, 2704. 375
- Boakes, E.H., Rout, T.M. & Collen, B. (2015). Inferring species extinction: The use of sighting records. 376 *Methods in Ecology and Evolution*, 6, 678–687. 377
- Canard, E., Mouquet, N., Marescot, L., Gaston, K.J., Gravel, D. & Mouillot, D. (2012). Emergence of 378 Structural Patterns in Neutral Trophic Networks. PLOS ONE, 7, e38295. 379
- CaraDonna, P.J., Petry, W.K., Brennan, R.M., Cunningham, J.L., Bronstein, J.L., Waser, N.M., et al. (2017).
- Interaction rewiring and the rapid turnover of plantpollinator networks. *Ecology Letters*, 20, 385–394. 381
- Carlson, C.J., Dallas, T.A., Alexander, L.W., Phelan, A.L. & Phillips, A.J. (2020). What would it take to 382 describe the global diversity of parasites? Proceedings of the Royal Society B: Biological Sciences, 287,
- 20201841. 384

383

- Cazelles, K., Araújo, M.B., Mouquet, N. & Gravel, D. (2016). A theory for species co-occurrence in 385 interaction networks. Theoretical Ecology, 9, 39–48. 386
- Closs, G.P. & Lake, P.S. (1994). Spatial and Temporal Variation in the Structure of an Intermittent-Stream 387 Food Web. *Ecological Monographs*, 64, 1–21. 388

- de Aguiar, M.A.M., Newman, E.A., Pires, M.M., Yeakel, J.D., Boettiger, C., Burkle, L.A., et al. (2019).
- Revealing biases in the sampling of ecological interaction networks. *PeerJ*, 7, e7566.
- Gauzens, B., Legendre, S., Lazzaro, X. & Lacroix, G. (2013). Food-web aggregation, methodological and
- <sup>392</sup> functional issues. *Oikos*, 122, 1606–1615.
- Giacomuzzo, E. & Jordán, F. (2021). Food web aggregation: Effects on key positions. Oikos, 130,
- 394 2170–2181.
- Griffiths, D. (1998). Sampling effort, regression method, and the shape and slope of sizeabundance
- relations. *Journal of Animal Ecology*, 67, 795–804.
- Gupta, A., Figueroa H., D., O'Gorman, E., Jones, I., Woodward, G. & Petchey, O.L. (2023). How many
- predator guts are required to predict trophic interactions? *Food Webs*, 34, e00269.
- Hadfield, J.D., Krasnov, B.R., Poulin, R. & Nakagawa, S. (2014). A Tale of Two Phylogenies: Comparative
- Analyses of Ecological Interactions. *The American Naturalist*, 183, 174–187.
- 401 Havens, K. (1992). Scale and Structure in Natural Food Webs. Science, 257, 1107–1109.
- Johnson, E.K. & Larremore, D.B. (2021). Bayesian estimation of population size and overlap from random
- subsamples.
- Jordano, P. (2016). Sampling networks of ecological interactions. Functional Ecology, 30, 1883–1893.
- Joseph, M.B. (2020). Neural hierarchical models of ecological populations. *Ecology Letters*, 23, 734–747.
- <sup>406</sup> Kenall, A., Harold, S. & Foote, C. (2014). An open future for ecological and evolutionary data? *BMC*
- Evolutionary Biology, 14, 66.
- Khoshraftar, S. & An, A. (2022). A Survey on Graph Representation Learning Methods.
- Koleff, P., Gaston, K.J. & Lennon, J.J. (2003). Measuring beta diversity for presenceabsence data. Journal
- of Animal Ecology, 72, 367–382.
- 411 Kopelke, J.-P., Nyman, T., Cazelles, K., Gravel, D., Vissault, S. & Roslin, T. (2017). Food-web structure of
- willow-galling sawflies and their natural enemies across Europe. *Ecology*, 98, 1730–1730.
- 413 Li, M., Liu, R.-R., Lü, L., Hu, M.-B., Xu, S. & Zhang, Y.-C. (2021). Percolation on complex networks:
- Theory and application. *Physics Reports*, Percolation on complex networks: Theory and application,
- 415 907, 1–68.

- MacDonald, A.A.M., Banville, F. & Poisot, T. (2020). Revisiting the Links-Species Scaling Relationship in
  Food Webs. *Patterns*, 1.
- Makiola, A., Compson, Z.G., Baird, D.J., Barnes, M.A., Boerlijst, S.P., Bouchez, A., *et al.* (2020). Key

  Questions for Next-Generation Biomonitoring. *Frontiers in Environmental Science*, 7.
- Martinez, N.D., Hawkins, B.A., Dawah, H.A. & Feifarek, B.P. (1999). Effects of Sampling Effort on

  Characterization of Food-Web Structure. *Ecology*, 80, 1044–1055.
- McLeod, A., Leroux, S.J., Gravel, D., Chu, C., Cirtwill, A.R., Fortin, M.-J., *et al.* (2021). Sampling and asymptotic network properties of spatial multi-trophic networks. *Oikos*, 130, 2250–2259.
- Moore, A.L. & McCarthy, M.A. (2016). Optimizing ecological survey effort over space and time. *Methods*in Ecology and Evolution, 7, 891–899.
- Niedballa, J., Wilting, A., Sollmann, R., Hofer, H. & Courtiol, A. (2019). Assessing analytical methods for detecting spatiotemporal interactions between species from camera trapping data. *Remote Sensing in Ecology and Conservation*, 5, 272–285.
- Paine, R.T. (1988). Road Maps of Interactions or Grist for Theoretical Development? *Ecology*, 69, 1648–1654.
- Poisot, T. (2022). Guidelines for the prediction of species interactions through binary classification.
- Poisot, T., Bergeron, G., Cazelles, K., Dallas, T., Gravel, D., MacDonald, A., *et al.* (2021). Global knowledge gaps in species interaction networks data. *Journal of Biogeography*, 48, 1552–1563.
- Poisot, T., Ouellet, M.-A., Mollentze, N., Farrell, M.J., Becker, D.J., Brierly, L., *et al.* (2022). Network embedding unveils the hidden interactions in the mammalian virome.
- Poisot, T., Stouffer, D.B. & Gravel, D. (2015). Beyond species: Why ecological interaction networks vary through space and time. *Oikos*, 124, 243–251.
- Ponisio, L.C., Gaiarsa, M.P. & Kremen, C. (2017). Opportunistic attachment assembles plantpollinator networks. *Ecology Letters*, 20, 1261–1272.
- Savage, V.M., Gillooly, J.F., Brown, J.H., West, G.B. & Charnov, E.L. (2004). Effects of Body Size and

  Temperature on Population Growth. *The American Naturalist*, 163, 429–441.

- Song, C., Simmons, B.I., Fortin, M.-J. & Gonzalez, A. (2022a). Generalism drives abundance: A computational causal discovery approach. *PLOS Computational Biology*, 18, e1010302.
- Song, C., Simmons, B.I., Fortin, M.-J., Gonzalez, A., Kaiser-Bunbury, C.N. & Saavedra, S. (2022b). Rapid monitoring for ecological persistence.
- Stephenson, P. (2020). Technological advances in biodiversity monitoring: Applicability, opportunities
- and challenges. Current Opinion in Environmental Sustainability, Open issue 2020 part A: Technology
- Innovations and Environmental Sustainability in the Anthropocene, 45, 36–41.
- Stock, M., Poisot, T., Waegeman, W. & De Baets, B. (2017). Linear filtering reveals false negatives in species interaction data. *Scientific Reports*, 7, 45908.
- Strydom, T., Catchen, M.D., Banville, F., Caron, D., Dansereau, G., Desjardins-Proulx, P., *et al.* (2021). A roadmap towards predicting species interaction networks (across space and time). *Philosophical*
- Transactions of the Royal Society B: Biological Sciences, 376, 20210063.
- Thompson, P.L. & Gonzalez, A. (2017). Dispersal governs the reorganization of ecological networks under environmental change. *Nature Ecology & Evolution*, 1, 1–8.
- Thompson, R.M. & Townsend, C.R. (2000). Is resolution the solution?: The effect of taxonomic resolution on the calculated properties of three stream food webs. *Freshwater Biology*, 44, 413–422.
- Volkov, I., Banavar, J.R., Hubbell, S.P. & Maritan, A. (2003). Neutral theory and relative species abundance in ecology. *Nature*, 424, 1035–1037.
- Walther, B.A., Cotgreave, P., Price, R.D., Gregory, R.D. & Clayton, D.H. (1995). Sampling Effort and
  Parasite Species Richness. *Parasitology Today*, 11, 306–310.
- 462 Williams, R.J. & Martinez, N.D. (2000). Simple rules yield complex food webs. *Nature*, 404, 180–183.
- Willott, S.j. (2001). Species accumulation curves and the measure of sampling effort. *Journal of Applied Ecology*, 38, 484–486.
- Young, J.-G., Valdovinos, F.S. & Newman, M.E.J. (2021). Reconstruction of plantpollinator networks from observational data. *Nature Communications*, 12, 3911.
- Zhang, S., Tong, H., Xu, J. & Maciejewski, R. (2019). Graph convolutional networks: A comprehensive review. *Computational Social Networks*, 6, 11.