

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

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## Introduction

Understanding which and how species interact underlies many questions in evolutionary biology and ecology, but also is an increasing imperative to both mitigate the anthropogenic change on Earth's biodiversity (Jordano 2016a; Makiola *et al.* 2020) and to predict potential spillover of zoonotic disease (Becker *et al.* 2021). Over the past few decades biodiversity data has become increasingly available: remote-sensing has enabled collection of data on spatial scales and resolutions previously unimaginable, automation of in-situ sensing (Stephenson 2020), and adoption of open data practices (Kenall *et al.* 2014) have substantially amount of data available to ecologists. Still, widespread data about species *interactions* remains elusive (**TIM\_DATASET\_PAPER?**). Often, observing an interaction between two species requires human sampling, typically because remote sampling methods can only detect co-occurrence, and this itself is not necessarily indicative of interaction (Blanchet *et al.* 2020). This constraint induces biases on species interaction data subject to the spatial and temporal scales that humans can feasibly sample.

*Sampling effort* and its impact on the resulting data collected from ecosystems has encouraged a long history of discourse. The recorded number of species in a sample depends on the total number of observations (Walther *et al.* 1995; Willott 2001), as do estimates of population abundance (Griffiths 1998). This 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 to measure species richness across large scales (Carlson *et al.* 2020). In the context of interactions, the initial concern was the compounding effects of limited sampling effort combined with the amalgamation of data (across both study sites and across taxonomic scales) could lead any empirical set of observations to inadequately reflect the reality of how species interact (Paine 1988). Martinez *et al.* (1999) showed that in a plant-endophyte trophic network, network connectance is robust to sampling effort, but this done in the context of a system for which observation of 62,000 total interactions derived from 164,000 plant-stems 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.

Because we cannot feasibly observe all (or even many) of the interactions that occur in nature, our samples end up capturing only a small fraction of those interactions. This means we can be reasonably confident two species actually interact if we have a record of it, but not at all confident that two 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* interact) from *false-negatives* (two species interact in some capacity, but we have not observed it). This is then amplified as the interaction data we have is geographically toward the usual suspects (Poisot *et al.* 2021a), This noise in data has practical consequences for

answering questions about species interactions (de Aguiar *et al.* 2019)—these false-negatives could go on to effect the inferences we make about network properties and relations among species, and our predictions about how species will interact in the future.

This is compounded by semantic confusion about the definition of “interaction”. Here distinguish between: a species *occurring*, a species being *observed occurring*, two species being observed *co-occurring*, and two species being observed *interacting* (fig. ??). In this manuscript, we refer to species either as “interacting”—two species co-occur (and, at least sometimes, interact)—or “not-interacting” (two species that, regardless of whether they co-occur, neither exhibits any meaningful effect on the biomass of the other). In fig. ?? we see that, under our definition, observing two species co-occurring is a prerequisite for observing an interaction between two species.

But species are not observed with equal probability but instead in proportion to their relative biomass—you are much more likely to observe a species of high relative abundance than one of very low relative abundance (Poisot *et al.* 2015). 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 here we show increasing strength of associations leads to increasing probability of false-negatives in interaction data. Further observed co-occurrence is often equated with meaningful interaction strength, but this is not necessarily the case (Blanchet *et al.* 2020; Strydom *et al.* 2021). Bears and salmon *interact*—a bear and the microbes in the soil of a dens interact, but less so.

Here, we show that the probability of observing a actual “non-interaction” between species depends on sampling effort, and suggest that surveys of species interactions can benefit from simulation modeling of detection probability (Jordano 2016b). We demonstrate that the realized false-negative rate of interactions is directly related the relative abundance of a particular species, relationship between total sampling effort (the total count of all individuals of all species seen) and false-negative rate. questions we pose and attempt to answer are: 1) How many times do you have to observe a non-interaction between two species to be confident in saying that is a true negative? 2) How “wrong” are the measurements of network structure as a function of false-negative probability? and lastly 3) How do false-negatives impact our ability to make reliable predictions about interactions? We show that positive associations in co-occurrence data can increase realized probability of false negatives, and demonstrate these positive associations are present in two spatially-replicated systems. We conclude by suggesting that simulation of sampling effort and species occurrence can and should be used to help design surveys of species diversity (Moore & McCarthy 2016), and by advocating use of null models like those presented here as a tool for guiding design of surveys of species interactions, and for modeling detection error in predictive ecological models.

## 2

### How many observations of a non-interaction do we need to classify it as a true negative?

To answer the titular question of this section, we present a naive model of interaction detection: we assume that every interacting pair of species is incorrectly observed as a 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 (callend the geometric distribution, a special case of the negative-binomial distribution) is shown in fig. ?? for varying values of the false negative rate  $p_{fn}$ . This illustrates a fundamental link between our ability to reliably say an interaction doesn’t exist— $p_{tn}$ —and the number of times we have observed a given species. In addition, note that there also 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-interaction  $N$  we have,  $p_{tn} < 1$ .

From fig. 1 (A) (and general intuition) it is clear that the more times we see two species *occurring*, but *not* interacting, the more likely the interaction is a true negative. But how does one decide what this threshold of number of observations should be when planning to sample a given system? If false-negative rates presented in fig. 1 seem unrealistically high, consider that species are not observed independent of their relative abundance. In the next section we demonstrate that distribution of abundance in ecosystems can lead to realized values of  $p_{fn}$  similar to those in fig. ?? for species with low relative abundance, simply as a artifact of sampling effort.

**2.1. False-negatives as a product of relative abundance** Here we show the realized false-negative rate of species interactions changes drastically with sampling effort, largely due to the intrinsic variation of abundances 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 (Banville *et al.* 2021) as well as random food-webs generated using the niche model (Williams & Martinez 2000). Our neutral model of observation assumes each observed species is drawn from the distribution of those species' abundances at that place and time. Although there is no shortage of debate as to the processes that govern this distribution of abundances within a community, this abundance distribution can be reasonably well described by a log-normal distribution (Volkov *et al.* 2003) (Note that in addition to the log-normal distribution, we also tested the case where the abundance distribution is derived from power-law scaling  $Z^{(T_i-1)}$  where  $T_i$  is the trophic level of species  $i$  and  $Z$  is a scaling coefficient. (Savage *et al.* 2004), which yields the same qualitative behavior, *supplement figure 1*). The practical consequence of abundance distributions spanning many orders of magnitude is seeing two “rare” species interacting requires two low probability events: observing two rare species *at the same time*.

To simulate the process of observation, for an ecological network  $A$  with  $S$  species, we sample abundances for each species from a standard log-normal distribution. For each true interaction in  $A$  (i.e.  $A_{ij} = 1$ ) we estimate the probability of observing both species  $i$  and  $j$  at given place and time by simulating  $n$  observations of individuals, where the species of the individual observed at the 1, 2, ...,  $n$ -th observation is drawn from the generated log-normal distribution of abundances. For each pair of species  $(i, j)$ , if both  $i$  and  $j$  are observed within the  $n$  observations, the interaction is tallied as a true positive if  $A_{ij} = 1$  and a false positive otherwise. Similarly, if only one of  $i$  and  $j$  are observed—but *not both*—in these  $n$  observations, but  $A_{ij} = 1$ , this is counted as a false-negative, and a true-negative otherwise.

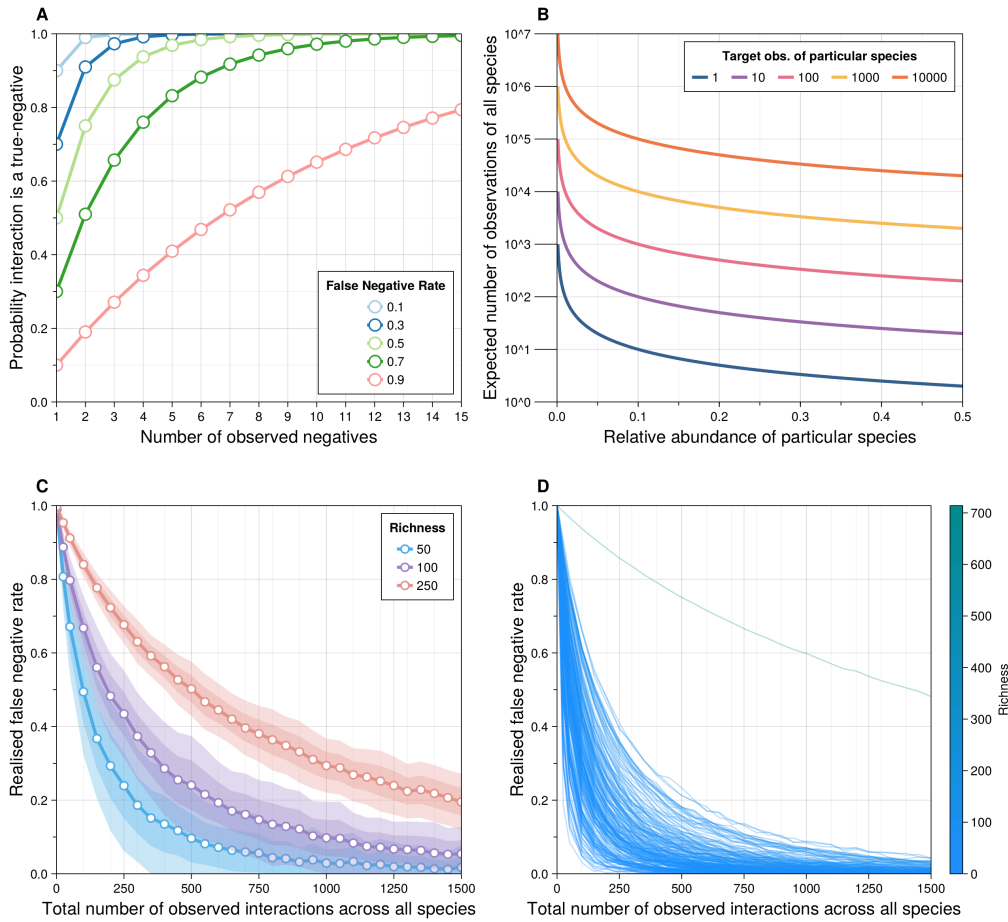
In fig. 1 (C) and (D) we see this model of observation applied to networks generated using the niche model (Williams & Martinez 2000) across varying levels of species richness, and in (b) applied to 243 food-webs from the Mangal database. For all niche model simulations in this manuscript, for a given number of species  $S$  the number of interactions is drawn from the flexible-links model fit to Mangal data (MacDonald *et al.* 2020), effectively drawing the number of 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 MAP estimate of  $(\mu, \phi)$  applied to Mangal data from MacDonald *et al.* (2020) is  $(\mu = 0.086, \phi = 24.3)$ . All simulations were done with 500 independent replicates per unique number of observations  $n$ . All analyses presented here are done in Julia v1.6 (Bezanson *et al.* 2015) using both EcologicalNetworks.jl v0.5 and Mangal.jl v0.4 [Banville *et al.* (2021); ZENODO link TODO]. Note that the empirical data also is, due to the phenomena described here, very likely to *already* have many false negatives, which is why we are interested in prediction of networks in the first place—we'll revisit this in the final section.

In panel (c) of fig. ??, we show the expected number of total observations needed to obtain a “goal” number of observations (colors) of a particular “focal” species. 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 negative (a la fig. ??), then we need an expected 10,000 observations of all species if the relative abundance of  $A$  is 0.00125.

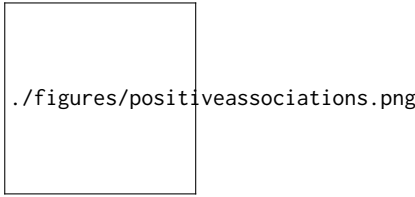
Empirical data on interactions are subject to the practical limitations of funding and human-work hours, and therefore existing data tend to fall on the order of 100s or 1000s observations of individuals per site (Nielsen & Bascompte 2007; Schwarz *et al.* 2020; Resasco *et al.* 2021). Clear aggregation of this data has proven difficult to find and a meta-analysis of network data and sampling effort seems both pertinent and necessary, in addition to the effects of aggregation of interactions across taxonomic scales (Gauzens *et al.* 2013; Giacomuzzo & Jordán 2021). Further, from fig. ?? it is evident that the number of species considered in a study is inseparable from the false-negative rate in that study, and this effect should be taken into account when designing samples of ecological networks in the future.

We conclude this section by advocating for the use of neutral models similar to above to generate expectations about the number of false-negatives in a data set of a given size. This could prove fruitful both for designing surveys of interactions (Canard *et al.* 2012), but also because we may want to incorporate models of observation error into predictive models (Joseph 2020). Additionally, one must consider the context for sampling—is the goal to detect a particular species  $A$  (as in fig. ?? (c)), or to get a representative sample of interactions across the species pool? This argument is well-considered when sampling species (Willott 2001), but has not yet been internalized for designing samples of communities.

**2.2. Positive associations can increase the probability of false-negatives** This model above doesn't consider the possibility that there are positive or negative associations which shift the probability of



**Figure 1** A) The probability 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 cdf of the negative-binomial distribution with  $r = 1$ . (B) The expected needed observations of all individuals of all species (y-axis) required to obtain a goal number of observations (colors) of a particular species, and a function of the relative abundance of that focal species (x-axis). (C) and (D): 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 Williams2000SimRul at varying levels of species richness (colors) with connectance drawn according to the flexible-links model MacDonald2020RevLin as described in the main text. For each draw from the niche model, 200 sets of 1500 observations are simulated, for which each the mean false negative rate at each observation-step is computed. Means denoted with points, with  $1\sigma$  in the first shade and  $2\sigma$  in the second. B: empirical food webs from Mangal database in teal, applied to the same process as the A. The outlier on panel B is a 714 species food-web



**Figure 2** Top: Hadfield, Bottom: NZ Stream Foodwebs. Effectively a version of Cazelles *et al.* (2016) figure 1 panel A. Both distributions have  $\mu \neq 0$  with  $p < 10^{-50}$

observing two species together due to their interaction (Cazelles *et al.* 2016). However, here we demonstrate that the probability of observing a false negative can be *higher* if there is some positive association between occurrence of species  $A$  and  $B$ .

If we denote the probability that we observe an interaction we know exists between  $A$  and  $B$  as  $P(AB)$ , and if there is *no* association between the marginal probabilities of observing  $A$  and observing  $B$ , denoted  $P(A)$  and  $P(B)$  respectively, then the probability of observing the interaction  $P(AB) = P(A)P(B)$ . In the other case where there *is* some positive strength of association between observing both  $A$  and  $B$  because this interaction is “important” for each species, 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 not independent and instead are positively correlated, *i.e.*  $P(AB) > P(A)P(B)$ . In this case, the probability of observing a false negative in our naive model from fig. ?? is  $p_{fn} = 1 - P(AB)$  which due to the above inequality implies  $p_{fn} \geq 1 - P(A)P(B)$  which indicates increasingly greater probability of a false negative as  $P(AB) \rightarrow P(AB) \gg P(A)P(B)$ .

However this does not consider variation in species abundance in space and time, (Poisot *et al.* 2015). If positive or negative associations between species structure variation in the distribution of  $P(AB)$  across space/time, then the spatial/temporal biases induced by data collection would further impact the realized false negative rate, as the probability of false negative would not be constant for each pair of species across sites. To test for this association empirical data, we use two datasets: a set of host-parasite interactions sampled across 51 sites with 327 total taxa (Hadfield *et al.* 2014) and a set of 18 New Zealand freshwater stream food webs with 566 total taxa (Thompson & Townsend 2000). We simply compute the empirical marginal distribution of species occurrence, and compare the product of the marginals,  $P(A)P(B)$ , to the empirical joint distribution  $P(AB)$ .

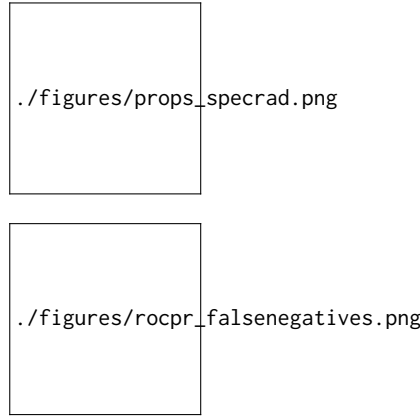
In fig. 2, both host-parasite system (top) and food-web (bottom) exhibit these positive associations. There is no reason to expect the strength of this association to be the same in different systems. At the moment, computing this metric for all of the networks in the Mangal database proves challenging as most data sets use different taxonomic identifiers, often at different resolutions. These particular datasets (Thompson & Townsend 2000; Hadfield *et al.* 2014) were usable because they already have been sorted to have a fixed taxonomic backbone (as part of EcologicalNetworks.jl (Banville *et al.* 2021)). Applying this in bulk to Mangal food-webs presents the difficulty of resolving different taxon identifiers across spatial samples of species with to different resolutions, which is why we can’t simply apply this to the whole Mangal database—this highlights a general problem of resolving taxonomic identifiers which use different names and different resolutions in different ecological datasets, which is a problem that needs to be addressed for computational approaches to scale up to the world of big-ecological-data we hope to build, although this is a task that may be aided via natural-language-processing methods.

### 3

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

We now transition toward assessing the effects of false negatives in data on the properties of the networks which we derive from this interaction data, and their effect on models for predicting interactions in the future.

**3.1. Effects of false-negatives on network properties** Here we simulate the process of observation with error to generate synthetic data with a known proportion of false negatives, and compare the computed network properties of the original “true” network to the computed properties of the “observed” network with added false-negatives. In fig. 3 we see the mean-squared error of connectance, mean degree-centrality, and spectral radius, computed across 2000, 2000, and 300 replicates respectively at each value of the false negative rate  $p_{fn}$ . All replicates use random food-webs simulated using the



**Figure 3** The mean-squared error (y-axis) of various network properties (different colors) across various simulated false-negative rates (x-axis). Means denoted with points, with  $1\sigma$  in the first shade and  $2\sigma$  in the second.

**Figure 4** Receiver-operating-characteristic (left) and precision-recall (right) curves for the model on varying levels of false-negatives in the data (colors). For each value of FNR, we run 30 random training/test splits on 80/20 percent of the data. Replica of figure 1 in Strydom *et al.* (2021)

niche model (Williams & Martinez 2000) with 100 species and connectance drawn from the flexible-links model (MacDonald *et al.* 2020) as before.

We consider three properties: connectance, mean-degree-centrality, and spectral radius, indicative of local, meso, and global structure. Connectance is effectively a node-level property, a proxy for the degree distribution. Degree-centrality captures a different aspect of network structure than connectance, more indicative of meso-level properties that describe local ‘regions’ of nodes interact. Spectral radius (equivalent to the magnitude of the largest eigenvalue of  $A$ ) is a measure of global structure, and demonstrates the most variability in response to false-negatives. For example, if a false-negative splits a metaweb into two components, spectral-radius becomes the largest eigenvalue of each of those two components. Also note that the form of this error function varies little as species richness changes (*supplemental figure 2*). Practically, fig. 3 shows us that different scales of measuring network structure vary in their response to false negatives—connectance responds roughly linearly to false negatives, whereas mean-degree-centrality decisively does not. This implies that false-negatives adversely could effect indirect interactions (Williams *et al.* 2002).

**3.2. Effects of false negatives on ability to make predictions** Here, we assess the effect of false negatives in data on our ability to make predictions about interactions. The prevalence of false-negatives in data is the catalyst for interaction prediction in the first place, and as a result methods have been proposed to counteract this bias (Stock *et al.* 2017; Poisot *et al.* 2021b). However, it is feasible this could induce too much noise for an interaction prediction model to detect the signal of interaction chance from to the latent properties of each species derived from the empirical network if the number of false-negatives in a dataset becomes too overwhelming.

To test this, we use the same predictive model and dataset as in Strydom *et al.* (2021) to predict a metaweb from various empirical slices of the species pool observed across space. This dataset from Hadfield *et al.* (2014) describes host-parasite interaction networks sampled across 51 sites. We partition the data into 80-20 training-test split, and then seed the training data with false negatives varying rates, but crucially do nothing to the test data. We use the same model, a neural-network with 3 feed-forward layers to predict outputs based on features extracted from co-occurrence (see Strydom *et al.* (2021) for more details). The single modification we make to the model is not enforcing a number of positives in the training data as this constraint is eventually impossible for increasing FNR. In fig. 4, we show receiving-operating-characteristic (ROC) and precision-recall (PR) curves for the model with varying levels of synthetic false-negatives added to the data.

Interestingly, the performance of the model from Strydom *et al.* (2021) changes little with many added false-negatives, which is good evidence in favor neural-networks as a class of model for interaction detection. Again, similar to our caveat in the previous section, this data is *already* likely to have many false-negatives, so the effects of adding more as we do in this illustration might be mitigated because there are already non-simulated false-negatives in the original data which impact the models performance, even in the  $p_{fn} = 0$  case.

We conclude by proposing that simulating the effects of false negatives in this way can serve as an additional validation tool when aiming to detect structural properties of networks using generative null models (Connor *et al.* 2017), or when evaluating the robustness of a predictive model.



## Discussion

Here, we have demonstrated that we expect false-negatives in species interaction datasets purely due to the distribution of abundances within a community. Positive associations between species occurrence (Cazelles *et al.* 2016) can increase the realized false-negative rate if the sampling effort is limited, and we have presented evidence of this non-random structure of co-occurrence in two sets of spatially-replicated ecological network samples. We have also shown that false-negatives can cause varying responses in our measurements of network properties and further could impact our ability to reliably predict interactions, which highlights the need for further research into methods for correcting this bias in existing data (Stock *et al.* 2017). A brief caveat here is that we do not consider the rate of false-positives—in large part false-positives can be explained by misidentification of species, although this could be a relevant consideration in some cases.

What does the future hold for this research? A better understanding of how false-negatives impact our analyses and prediction of ecological networks is a practical necessity. False-negatives could pose a problem for many forms of inference in network ecology. For example, if we aim to measure structural or dynamic stability of a network, or to infer indirect interactions (Williams *et al.* 2002), these estimates could be prone to error if the observed network is not sampled “enough”. What exactly “enough” means is then specific to the application, and should be assessed via methods like those here when designing samples. Further, predictions about network rewiring (Thompson & Gonzalez 2017) due to a changing climate could be error-prone without accounting for interactions that have not been observed but that still may become climatically infeasible.

This highlights the need for a quantitatively robust approach samples design: for interactions (Jordano 2016b) and otherwise (Carlson *et al.* 2020). The primary takeaway is that when planning the sampling effort across sites, it is necessary to take both the size of the species pool into account. Further, simulating the process of observation could be a powerful tool for planing study design which takes relative abundance into account, and provide a null baseline for detection of interaction strength. A model similar to that here can and should be used to provide a neutral expectation of true-negative probability given a number of observations of individuals at a given place and time.

As we derive from fig. ??, we can never guarantee there are no false-negatives in data. In recent years, there has been interest toward explicitly accounting for false-negatives in models (Stock *et al.* 2017; Young *et al.* 2021), and toward a predictive approach toward interactions —rather than expect that our samples can fully capture all interactions, we know that some interactions between species will not be observed due to finite sampling capacity, and instead we must impute the true metaweb of interactions given a set of samples (Strydom *et al.* 2021). As a result, better predictive approaches are needed for interaction networks (Strydom *et al.* 2021), and building models that explicitly account for observation error is a necessary step forward for predictive ecological models (Johnson & Larremore 2021; Young *et al.* 2021). Neural networks, like the one used to predict interactions in the above section, have been used to reflect hidden states which account for detection error in occupancy modeling (Joseph 2020), and could be integrated in the predictive models of the future.

A better conceptual framework for designing surveys and monitoring networks, and incorporating sequential observations over time is clearly needed (Carlson *et al.* 2020), combined with a meta-analysis of sampling effort and taxonomic resolution in existing data. Incorporating a better understanding of sampling effects and bias on both the future design of biodiversity monitoring systems, and the predictive models we wish to apply to this data, is imperative in making actionable predictions about the future of ecological interactions on our planet.

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