

# Inductive link prediction facilitates the discovery of missing links and enables cross-community inference in ecological networks

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## Abstract

Predicting species interactions (links) within ecological networks is crucial for advancing our understanding of ecosystem functioning and responses of communities to environmental changes. Transductive link prediction models are often used but are constrained by sparse, incomplete data and are limited to single networks. We address these issues using an inductive link prediction (ILP) approach. We evaluated the model performance on 538 networks across four community types: plant-seed disperser, plant-pollinator, host-parasite, and plant-herbivore. By pooling data across communities and applying transfer learning, our model predicts interactions within and between ecological networks. The ILP model achieved higher precision and F1 scores than transductive models. However, cross-community prediction efficacy varies, with plant-seed disperser and host-parasite networks performing better than plant-pollinator and plant-herbivore networks as training and test sets. Moreover, the nature of ecological interactions (mutualistic versus antagonistic) did not strongly influence the predictability of missing links. Finally, leveraging ILP's generalizability, we developed a pre-trained model that ecologists can readily use to make instant predictions for their networks. This study highlights ILP's potential to improve the prediction of ecological interactions, enabling generalization across diverse ecological contexts and bridging critical data gaps.

## Introduction

Ecosystem services, such as pollination, are vital for human societies and arise from interactions among organisms in ecological communities [1]. For instance, crop pollination depends on interactions between wild bees and non-crop flowers [2]. These dependencies can be modeled as networks, where nodes represent species and links represent interactions. Fully sampling these networks would improve our understanding of indirect dependencies between species, but the resources needed to confirm each interaction are immense [3–5]. Consequently, ecological networks are often under-sampled, limiting structural analysis. Predicting probable but unconfirmed links offers a solution [6–12] and aids in anticipating network responses to anthropogenic changes, such as disease host prediction and local-invasive species interactions [11]. However, predicting links is challenging due to network sparsity and because current models typically predict interactions only within the network on which they were trained, limiting their applicability across different ecological contexts or communities.

Ecological link prediction models can use information on species traits [9,13] and phylogeny [12,14], but these are often difficult to obtain or biased because some taxonomic groups are far more studied than others. Instead, it is possible to rely on the topology of the known part of the network to predict the unknown [7,8]. For instance, a parasite with a broad host range (high node degree) will likely infect a new host. Such approaches that predict missing links based on known properties within the same network are called *transductive* link prediction [15]. While this is the primary approach used in ecology so far [7,8], its performance is hindered in networks where only a few links are known or where some parts are known much better than the others [16], two issues prevalent in ecological networks.

To overcome these issues, we take an approach called inductive link prediction (ILP), in which links in one network are predicted by learning the structure of others [17]. ILP harnesses the principle of universality [18–21], reflected in ecological networks by cross-system topological similarities [22,23]. Indeed, despite system differences (e.g., mutualistic vs. antagonistic networks), processes such as spatiotemporal distributions, evolutionary history, and neutral dynamics produce recurring non-random patterns across networks [24–29]. For example, heavy-tail degree distributions, with most nodes having few links and a few having many [30–32] and structures like nestedness and modularity are common in multiple types of ecological networks [22,33–36].

Cross-system similarities in macroscopic patterns provide an opportunity to increase training data by using multiple networks within an ILP framework [37,38]. In addition, it enables transfer learning, where a model trained on data from one domain is applied to predict outcomes in another. The idea of using transfer learning in ecology has been recently proposed [39], but its application has been very limited [13,14]. For example, Caron et al. [13] used transfer learning to predict links in food webs in one area (e.g., Europe) based on knowledge in another (e.g., Serengeti). Using trait-based predictions, they found that pairwise interactions are better predicted using a model trained on the same food web than with models trained on other food webs. However, they did not pool data from different food webs for training. Moreover, no ecological study so far has applied ILP within a machine learning framework, a powerful approach that offers remarkable flexibility and scalability for leveraging data across diverse systems.

Given the structural similarities between networks from different ecological communities, we hypothesize that ILP is more effective than transductive link prediction for predicting links in ecological

networks. Further evidence for the plausibility of this hypothesis comes from two recent studies that showed that it is challenging to identify the type of network (e.g., plant-pollinator, host-parasite) based solely on its structure [23,40]. This is because variation in network structure is similar within and between different types of ecological networks. On the one hand, this observation means that it would be challenging to predict the ecological community of a network based on its structure [23]. On the other hand, the structural similarity of different community types can be utilized to improve link prediction by training models on a large dataset of diverse communities.

We tested this hypothesis using an ILP model we developed, which further enables cross-community prediction. We find that our model outperforms transductive link prediction models in predicting links, but that prediction accuracy varies by community type. Specifically, plant-pollinator networks weaken cross-community predictions while host-parasite networks enhance them. The developed ILP model is publicly available as an online tool to facilitate broader use and further testing by ecologists (<https://ecomplab-eco-ilp.hf.space>).

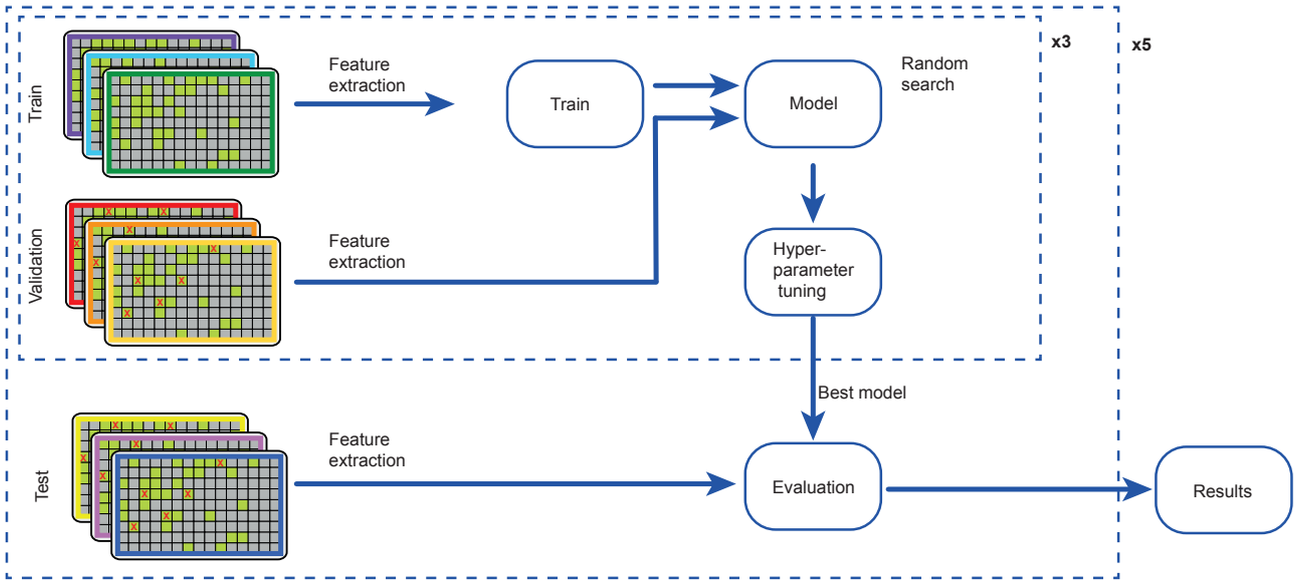
## Results

### Data

We used the data set compiled by Michalska-Smith and Allesina [23], later used by [40]. We used networks with  $\geq 25$  species and with connectance  $\geq 0.1$  (Table S1, Figs. S1, S2). This data set includes 205 plant-seed disperser networks (PSD), 217 plant-pollinator networks (PP), 84 host-parasite networks (HP), and 32 plant-herbivore networks (PH) (538 networks in total). A potential limitation of the data set could be the non-equal number of networks per community type. To test this, we included the type of community as a feature in the model, and it was not an important feature in the prediction. In addition, communities represented by more networks did not necessarily perform better as train or test sets (see results below), indicating that the number of networks is not a limitation. As previously shown for this data set [23,40], variation in network structure was not higher between than within networks (Fig. S3).

### Overview of the link prediction pipeline

We developed a pipeline to evaluate the performance of ILP models on multi-network data sets using nested grouped cross-validation (Fig. 1). We split the data into training, validation and test sets, ensuring that instances (links) from the same network are always together in the set. We predicted links based on network properties, which we used as features in the model. Before training, we used feature selection to mitigate correlations, reduce dimensionality, prevent over-fitting, and identify informative features for link prediction (see feature selection in Methods). In the inner loop, we used the training and validation sets to tune hyperparameters and select the best model to perform link prediction on the test set. We performed 5-fold cross-validation in the outer loop. We trained our model to optimize the F1 score, which balances model’s ability to predict links and non-links, using popular machine learning models: logistic regression, random forest, and XGboost. The results did not qualitatively change between models (Supplementary note S1.1), and we present results for Random Forest. We considered the imbalance between the number of observed and unobserved interactions in our evaluations [41] (Methods).



**Fig. 1: Pipeline overview.** We used nested cross-validation, with 3 and 5 folds in the inner and outer loops, respectively. We split the data into train, validation, and test sets and calculated network properties (feature extraction). The validation and test sets contained withheld links (red x's). We used the inner loop to tune the hyperparameters (using random search) and selected the best model to predict links in the test set in the outer loop. Model evaluation was based on a confusion matrix with TP, TN, FP, and FN values gathered from the outer loop's five folds. When splitting the data, we ensured that each network was included entirely in the set (i.e., we split the data by networks, not by links) and that each network appeared at least once in the test set. See details in Methods.

## Evaluation metrics

Our evaluation is based on metrics derived from a confusion matrix, which summarizes the counts of true positives (TP), true negatives (TN), false positives (FP), and false negatives (FN). From an ecological perspective, the primary interest lies in accurately identifying missing links—interactions that exist but are challenging to detect empirically. Consequently, we first focus on recall, defined as  $TP/(TP + FN)$ , which measures the proportion of actual interactions correctly identified by the model. A high recall value indicates a model's strong ability to uncover potential new interactions in the field. However, because it is impractical for field ecologists to validate every predicted interaction, recall alone may lead to inefficient fieldwork. Therefore, we complement recall with precision, defined as  $TP/(TP + FP)$ , representing the likelihood that a predicted link truly exists. A model achieving high precision ensures that ecologists focus their validation efforts on predictions with greater reliability. Given the typical imbalance in ecological interaction data, precision and recall can be effectively combined into the F1 score, calculated as the harmonic mean of these two metrics. This combined metric provides a balanced evaluation, simultaneously accounting for both false positives and false negatives, which is particularly suitable for ecological applications.

These metrics require setting a threshold for link probability to decide when a link is considered present. Following standard practice, we used a threshold of 0.5, representing a random guess. However, recognizing that alternative thresholds might offer insights into model performance, we also assessed the precision-recall tradeoff across various thresholds. Specifically, we computed the area under the precision-recall curve (PR AUC). The PR tradeoff effectively reflects a model's capability to maximize recall while minimizing false positives, making it particularly relevant for evaluating imbalanced ecological datasets. Although these metrics primarily evaluate our ability to identify links, it is

also valuable to quantify the model’s accuracy in detecting the absence of interactions. Therefore, we provide additional indices such as specificity and balanced accuracy in the supplementary information to comprehensively assess model performance. For more details, see Supplementary Note S1.2.

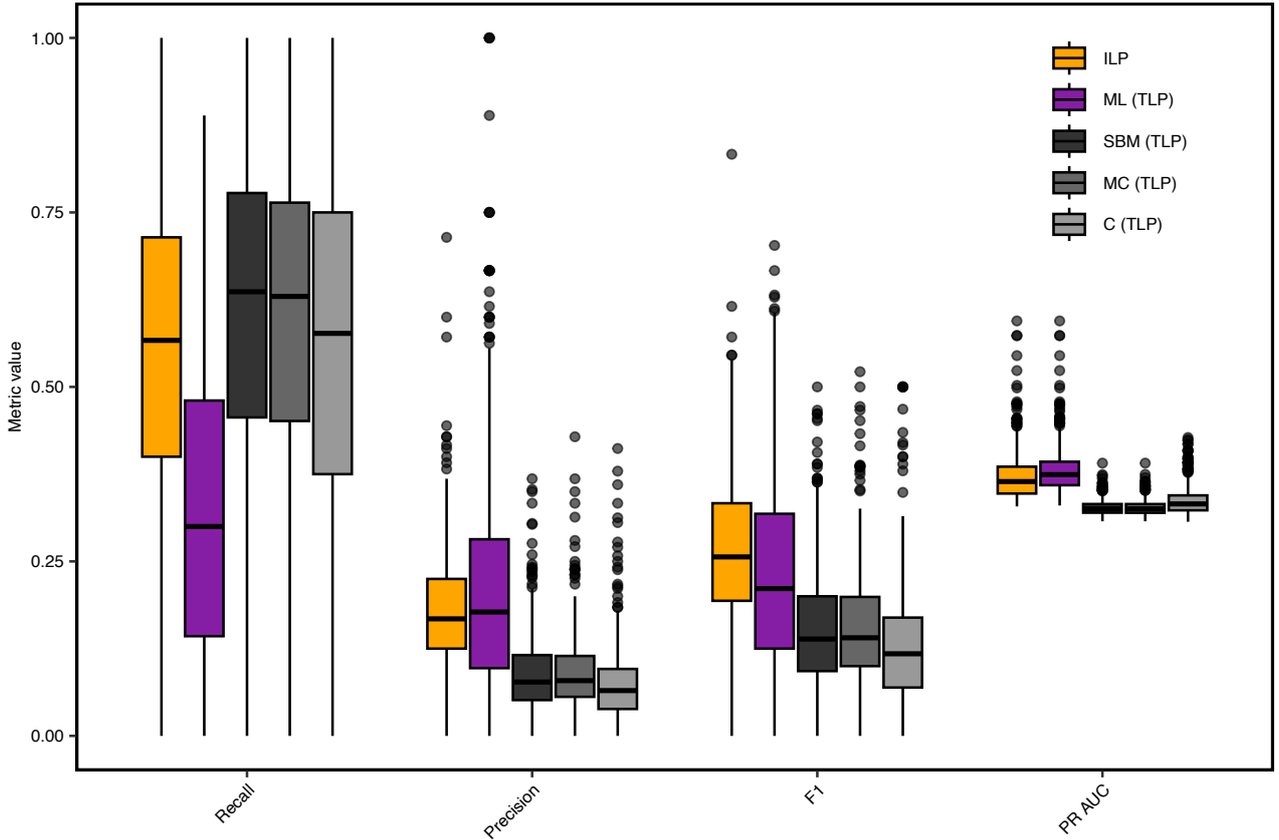
### **Inductive prediction better guides missing link detection**

To test the hypothesis that collating data across communities would improve link prediction, we compared our model performance to three transductive models that train and test on a single network at a time: the stochastic block model (SBM), connectance, and matching centrality models, which were previously published and are not based on machine learning [8]. While ILP predicts links with similar recall to that of transductive link prediction (TLP) models, it does so with higher precision. The area under the precision-recall (PR) curve was also significantly higher in the ILP model (Fig. 2). However, the TLP models were not based on machine learning techniques. To ensure a more fair comparison we also trained a transductive machine learning model. The ability of that model to recover links (recall) was much lower than ILP, though its precision was similar. Therefore, overall, the ILP model outperformed the transductive ones (Fig. 2, Fig. S4).

To better understand the differences in model performance, we investigated how model predictions varied with node degree (Fig. 3). Specifically, each predicted link connects two species from the low and high trophic levels, and their node degrees served as key model features. We focused on node degree because it emerged among the most influential predictive features (Fig. S5). Compared to the non-machine learning transductive models, which exhibited consistently high recall yet low precision across node degrees, the ILP model showed a clear improvement by maintaining high precision across the entire range of node degrees while also achieving comparable recall at intermediate and higher degrees. Consequently, ILP consistently obtained better F1 scores relative to these models. When compared to the machine learning-based transductive model—which performed poorly in recall but attained higher precision than the other transductive models—the ILP model was again superior, displaying similar precision but substantially higher recall. This improved recall of ILP implies greater utility for guiding targeted field sampling efforts. The advantage of ILP was particularly pronounced at node degree extremes (both low and high degrees), areas that typically pose significant prediction challenges. Low-degree nodes represent specialist species interactions, which are inherently rare and challenging to detect, while high-degree nodes involve fewer species and thus offer limited information for accurate predictions (Fig. S6). Therefore, the ILP model specifically enhanced performance precisely in those areas most critical for effective ecological link prediction (Fig. 3, Fig. S7).

### **Prediction ability varies by community type**

Interactions can vary in the ecological and evolutionary processes that determine them. Therefore, we expect variation in our ability to predict links in different communities. Therefore, we first assessed predictive performance within each community individually. Precision-recall (PR) curves indicated notable variation, with the highest predictive performance observed in host-parasite networks and the lowest in plant-pollinator networks (Fig. 4A). Because co-evolutionary constraints shape interaction networks, we further anticipated that our ability to predict interactions of the same type would be similar. Contrary to this expectation, the ILP model predicted plant-seed disperser interactions with similar effectiveness as host-parasite interactions, and plant-pollinator interactions comparably to plant-herbivore interactions (Fig. 4B). Moreover, the ILP model demonstrated significantly greater

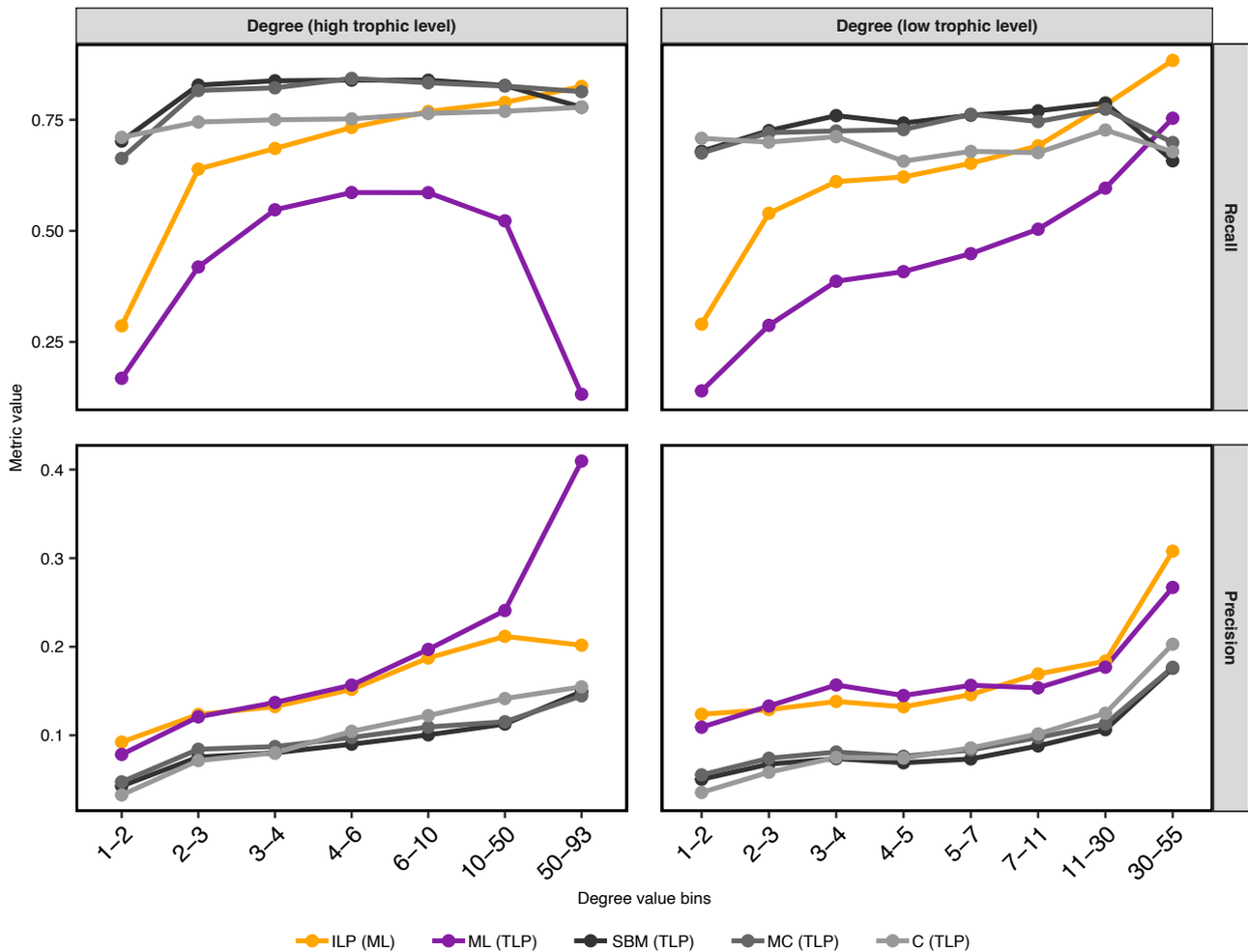


**Fig. 2: Comparison of model performance.** We compared ILP to three transductive (TLP) previously published models (stochastic block model, connectance (C) and matching centrality (MC) [8]), and to a machine learning (ML) model that we trained. The inductive link prediction model was trained and tested on networks from all communities. Each boxplot is a distribution of an evaluation metric using a 0.5 classification threshold, besides the PR AUC. The boxplots show the interquartile range (box), median (central line), data within 1.5 times the interquartile range (whiskers), and potential outliers (individual points beyond whiskers). Each data point is a network and boxplots contain networks from all the five outer folds ( $n = 538$  networks in all the box plots). See definitions of evaluation metrics in the Methods and additional metrics in (Fig. S4).

predictive capability for plant-seed disperser interactions than for plant-pollinator interactions (Fig. 4B, Fig. S8, Table S2). Although the recall and PR AUC were significant in 4 out of the 6 comparison, the precision was only different when comparing the prediction of pollination to that of seed dispersal. Therefore, overall, our findings suggest that the nature of ecological interactions (mutualistic versus antagonistic) does not strongly influence the predictability of missing links.

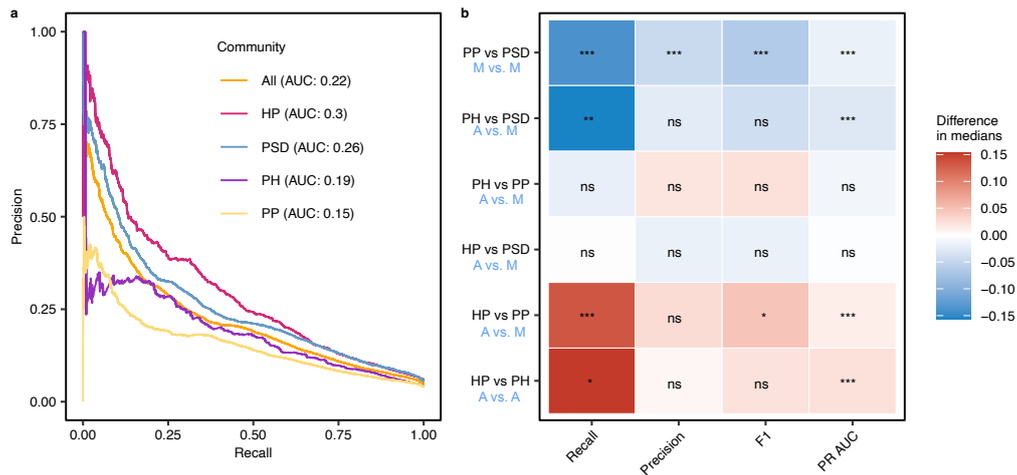
The main advantage of the ILP model is in transfer learning, allowing us to perform cross-community link prediction. We trained the model using networks from a given community type to predict links in the same community or in any other (Fig. 5). To create unbiased comparisons, we ensured that the number and identity of the networks in the training set were the same across experiments. We expected prediction to be more robust within community types. This prediction was confirmed only for host-parasite and plant-herbivore networks (the diagonal in Fig. 5). In other communities, interactions are predicted similarly well or better when trained on communities that are not the predicted ones.

We further expected that pooling data across all networks (All) for training will improve predictions (i.e., that the ‘All’ column will have higher values than within-community predictions). Contrary

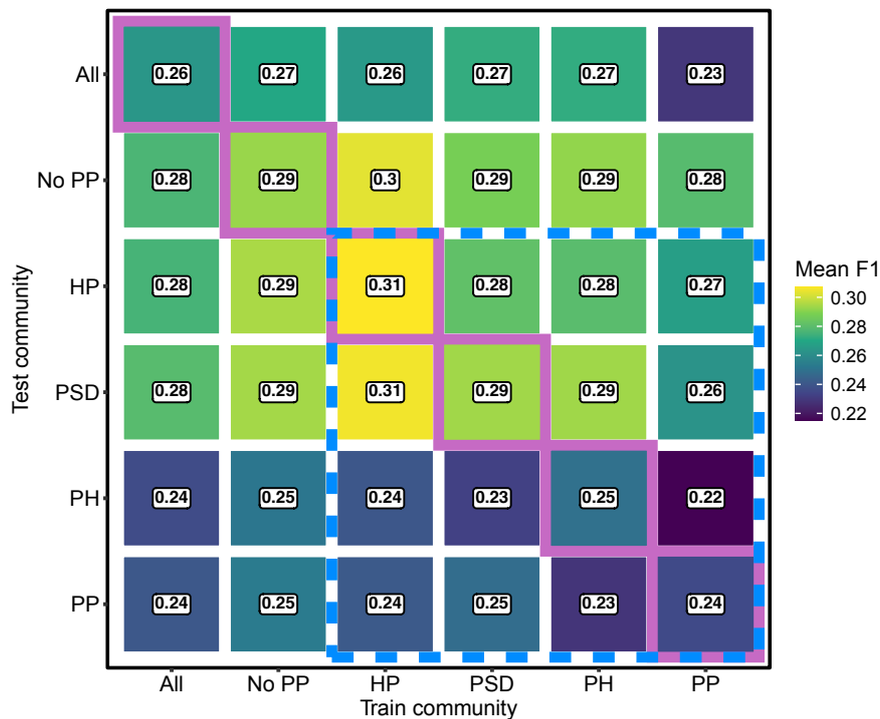


**Fig. 3: Comparison of model performance across node degrees.** The ILP model provides more effective guidance for discovering missing links compared with non-machine-learning transductive models by consistently maintaining higher precision, ensuring that identified potential interactions are more likely to be validated in the field. While achieving comparable recall at intermediate to high node degrees, the ILP model excels at extremes of node degree—specialist interactions at low degrees and interactions with limited information at high degrees. Compared with the machine-learning-based transductive (TLP) model, the ILP model offered similar precision but substantially higher recall, further enhancing its practical utility for targeted ecological sampling efforts.

to this expectation, pooling data worsened the predictions. This result can stem from the fact that plant-pollinator networks are highly sparse and consistently under-perform (Fig. 4A), affecting overall predictions. Indeed, removing plant-pollinator networks from the training set (No PP) improved predictions. This may happen because while numerous instances existed in the data set, plant-pollinator networks had the lowest connectance [17]. Low connectance leads to a significant imbalance between existing and non-existing links, which can decrease the precision and recall scores.



**Fig. 4: Prediction within community types.** The model was trained on networks from all communities and tested separately for each community type. **a** Precision-recall curves across thresholds. The AUC values are noted in the legend. **b** Pairwise comparison between communities (rows) for multiple evaluation metrics (columns). For each metric, we tested differences in medians across all communities using Kruskal-Wallis, and then performed a Dunn posthoc test with Bonferroni correction for pairwise comparisons. Positive values indicate that the median was higher in the second community (e.g., PSD had higher precision than PP networks). The type of interaction (A: antagonistic, M: mutualistic) is noted below each comparison. \*:  $p$ -value < 0.05, \*\*:  $p$ -value < 0.01, \*\*\*:  $p$ -value < 0.001, ns: not significant. PSD: plant-seed disperser, PP: plant-pollinator, HP: host-parasite, PH: plant-herbivore.

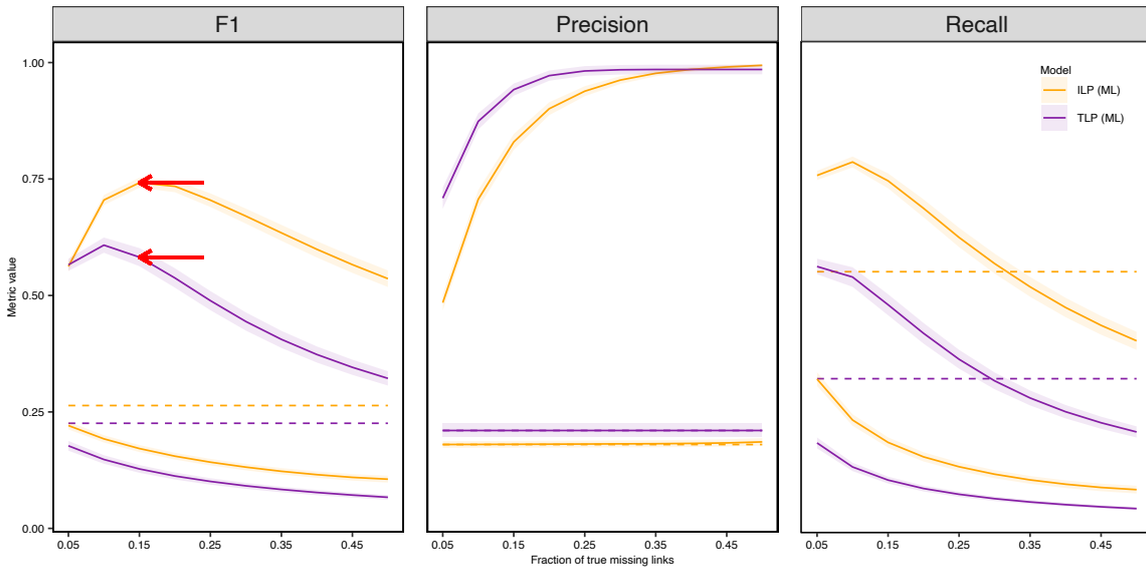


**Fig. 5: Link prediction within and between community types.** Each cell in the heatmap is the median of F1, calculated across all networks for a model that was trained on data from a certain community (columns) and tested on another (rows). The diagonal depicts results for models trained and tested on the same community type. The red dashed border encloses models trained and tested using single community types. PSD: plant-seed disperser, PP: plant-pollinator, HP: host-parasite, PH: plant-herbivore. 'All' is a model trained or tested using pooled data from all communities. 'No PP' is the same as 'All' but without plant-pollinator networks.

## Estimating the bounds of link prediction performance

One issue common to all link prediction studies is the lack of knowledge of the ground truth. That is, which of the non-existing links are truly missing (if we had this knowledge, we would not need link prediction). Moreover, it is likely that obtaining a fully complete network without setting a threshold may be extremely hard because there is typically a very long tail of very weak interactions. Ideally, link prediction models would guide the field sampling efforts of ecologists who want to complete their networks. In turn, field data can evaluate model predictions. Empirically estimating model performance in light of ground truth is necessary but would be extremely time-consuming because if links were missing in the first place, an intensive amount of sampling would likely be needed to sample even a few of them.

As a starting point, we take here an alternative computational approach. We calculate the bounds of the machine learning ILP and TLP model predictions across a theoretical range of proportion of true missing links. We devised two scenarios. In the best-case scenario, the false positive links of the models were indeed missing links. In the worst-case scenario, the predicted negative links are actually positive in nature (see Methods). The ILP model outperforms TLP in recovering missing links overall, as indicated by the consistently higher F1 in the best-case and worst-case scenarios (Fig. 6). For instance, if the true proportion of missing links in nature was 0.15, the ILP and TLP models' best-scenario F1 values would be  $\approx 0.74$  and  $\approx 0.58$ , respectively. This advantage stems from the large difference in recall compared to precision between these models (Fig. 6).

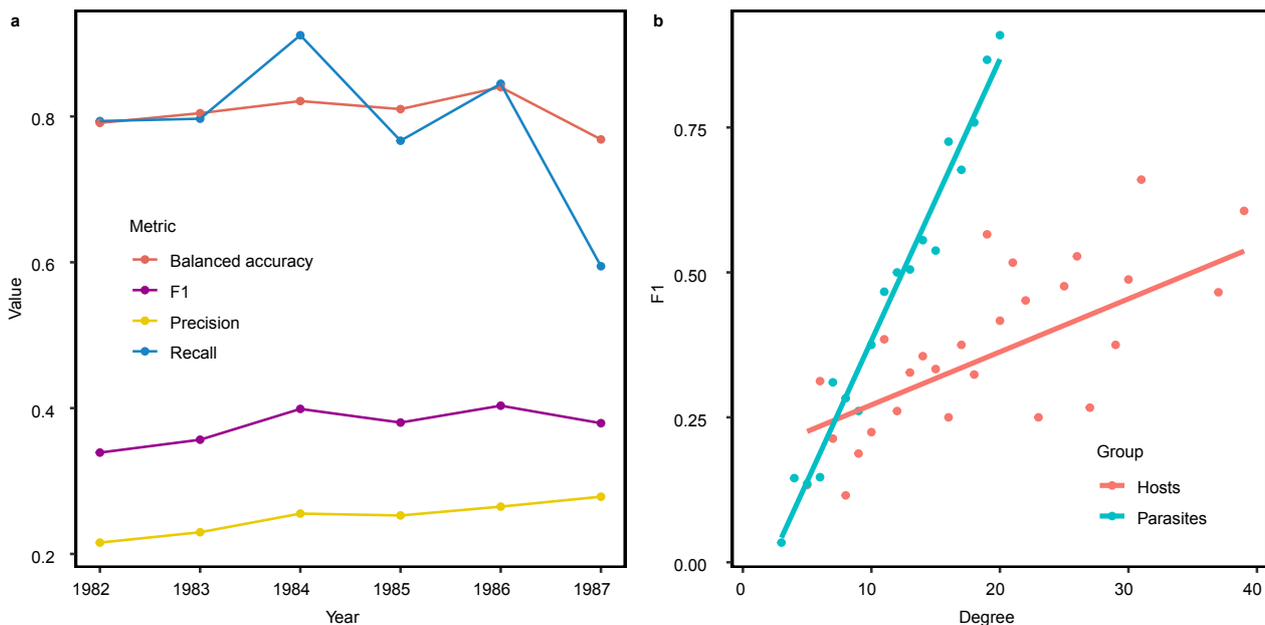


**Fig. 6: Bounds of model predictions across proportions of ground-truth missing links.** The horizontal dashed lines depict the current models' performance. The upper and lower curves (solid lines) represent the best-case (false positive links of the models were indeed missing links) and worst-case (predicted negative links are actually positive in nature) scenarios, respectively. The x-axis is the proportion of links that are missing (unobserved but exist in nature). An example for figure interpretation for the red arrows: If the true proportion of missing links in nature is 0.15, the ILP and TLP models' best-scenario F1 values would be  $\approx 0.74$  and  $\approx 0.58$ , respectively. Bounds were calculated as means, and their confidence intervals (light-colored ribbons) were calculated across all networks. The ILP model was one in which the train and test contained networks from all communities.

## Model implementation and a case study

Because ILP enables prediction in networks on which the model was not trained, we developed an online implementation that includes a pre-trained model. The application allows users to upload an existing network, and instantly obtain probabilities for the links (Fig. S9). Users can also choose to withhold links and obtain model evaluations (e.g., F1, balanced accuracy). The model is available online (<https://ecomplab-eco-ilp.hf.space>). In addition, we released the model as a Python package [https://github.com/Ecological-Complexity-Lab/eco\\_ILP/tree/main/package](https://github.com/Ecological-Complexity-Lab/eco_ILP/tree/main/package).

To demonstrate a use case, we analyze a host-parasite network (that was not part of the training set), representing the infection of 22 small mammalian host species by 56 ectoparasite species during six consecutive summers in Siberia (1982–1987)[42,43]. In each of the six networks, we predicted links (withholding 20% of interactions). The prediction ability did not vary greatly across years (except for recall, Fig. 7A). We therefore aggregated the network (unifying links across years) and predicted links in the aggregated network. We find that F1 increases with the degree of species but faster in parasites (Fig. 7B; see Fig. S10 for yearly correlations). Therefore, for the same degree, it is easier to predict links for parasites than for hosts. This pattern was not affected by host or parasite taxonomy (Fig. S11).



**Fig. 7: Case study.** We use our model to predict a host-parasite network (that was not part of the training set) representing the infection of 22 small mammalian host species by 56 ectoparasite species during 6 consecutive summers in Siberia (1982–1987). In each prediction we withheld 20% of the links. **(A)** Overall prediction metrics per year. **(B)** A correlation between parasite and host degree and F1.

## Discussion

Species interactions are the backbone of ecosystem functioning. Link prediction helps us improve our knowledge of species interaction structure and gain insights into how interactions would change in response to perturbations. However, network data are often incomplete and biased. In this paper, we take a step forward using ILP, which allowed us to pool data across networks. The ILP approach also enables predicting interactions in entirely new networks (based on their existing structure) via

transfer learning, an advantage we leveraged to develop implementations that allow researchers to obtain instant predictions for existing networks.

Beyond enhancing the detection of missing links, ILP represents a conceptual advancement by offering ecologists a new lens through which to approach the challenge of link prediction in ecological networks. For example, our pretrained model provides an accessible tool for generating immediate predictions, allowing researchers to explore potential interactions in their networks with minimal effort. This capability can help prioritize fieldwork by highlighting the most likely missing interactions, thereby improving the efficiency of data collection. While it is crucial that model predictions are validated through field sampling in well-studied systems where missing and forbidden links can be assessed with high confidence, such systems are rare. Instead, field ecologists can iteratively use model prediction and field sampling, which will enable prediction validation in real time.

By explicitly framing the link prediction task as an inductive learning problem, our approach also leads to new research directions that were previously limited by models lacking generalizability. For example, researchers can use ILP to predict how interactions might change in time [44] or under environmental shifts, such as habitat loss, climate change, or species introductions. This can be done by providing a baseline network and potential changes such as removal/addition of species or interactions under the new conditions (or time points), allowing the model to infer changes in interaction patterns from the baseline to the new scenarios. By forecasting dynamic responses, ILP offers valuable insights for anticipating ecosystem reorganization, guiding conservation efforts, and addressing both theoretical and pressing ecological challenges.

ILP models overcome the limitations of sparse and incomplete data by pooling information across different systems. Indeed, the model we developed generally outperformed transductive models. Moreover, our theoretical evaluation of model performance showed that ILP would retrieve missing links better than TLP regardless of the true proportion of missing links. Nevertheless, our model can be improved, and we hope researchers build upon our approach; for instance, by incorporating additional ecological information, such as trait-based data, phylogenetic signals, or environmental contexts [9,12,45], enhancing its predictive accuracy and ecological relevance.

Using networks from multiple communities allowed us to perform cross-community predictions, an approach previously tested only on four food webs using trait data [13]. Unlike that study, we found that predictions were not necessarily better within each community. For instance, plant-pollinator and plant-herbivore interactions were predicted equally well regardless of the training community, suggesting that different kinds of ecological networks can share similar assembly patterns or co-evolutionary dynamics [46], leading to comparable structures and constraints on interactions [23]. These findings reinforce that the structural similarities across ecological networks enable effective cross-community link prediction, even amidst taxonomic differences. They further align with our finding that network type was not an important predictor of links. Previous studies also showed that distinguishing between different types of ecological networks can be challenging because structural variation is similar within compared to between network types [23,40].

On the other hand, host-parasite and plant-seed disperser networks had better predictive performance than plant-pollinator and plant-herbivore networks overall, and were better predicted when trained on the same community. These differences between communities in train/test quality are an avenue for future research. An obvious question is whether those differences are solely due to data quality

or whether there are ecological mechanisms underlying interaction probabilities. For instance, host-parasite and plant-seed disperser networks exhibit higher connectance, further facilitating predictions (Table S1). Along with that, host-parasite and seed-disperser networks often involve more specialized animal groups, possibly rendering their topology more predictable. In contrast, the diverse and heterogeneous animal species in plant-pollinator networks, each with distinct behaviors and roles, can reduce predictability. Therefore, variation between communities in their quality as training data underscores the need to consider each community type’s unique characteristics. Incorporating features specific to community types, such as traits [9,12,13], structural properties [47] or environmental features [45], may aid in addressing such open questions and simultaneously increase model performance.

In conclusion, our study demonstrates the potential of ILP models to enhance ecological research. By shifting from TLP to ILP, our approach provides a more generalizable framework, broadening the applicability of link prediction across ecological contexts. We further provide an online application and a Python package for instant predictions via a pretrained model that can guide efficient field sampling, and we hope researchers will build on this open-source code to further explore and improve ILP. Therefore, in addition to filling gaps in data, generalizable ILP models such as ours provide a scalable solution for predicting links in rapidly changing ecosystems, potentially enabling identification of interactions involving invasive species and novel disease hosts to support proactive conservation and management strategies.

## Methods

### Nested cross validation and hyperparameters

Nested cross-validation allows optimization of hyperparameters along with an unbiased estimation of the model’s performance. This reduces the risk of over-fitting by ensuring that the model evaluation is conducted on data not seen during the hyperparameter tuning phase [48]. Hyperparameters control the learning process rather than being learned from the data, and are set before the training process begins [49]. The optimal values for the hyperparameters that maximize the machine learning model’s performance are selected through hyperparameter tuning during that inner loop of the nested cross-validation (Table S3). We optimized hyperparameters using a random search, which is a computationally efficient tuning technique. The process involved randomly sampling values from a predefined search space, followed by training and evaluating model performance on those values using the training and validation sets. We used the F1-score as the performance metric to evaluate the hyperparameter combinations. The average F1-score across all folds was calculated for each random set of hyperparameters. We repeated this procedure for multiple random sets of hyperparameters, and the set that yields the highest F1 score was chosen as the optimal solution for the current iteration of the outer loop.

Because the data includes multiple networks, we used grouped cross-validation to prevent information leakage between training, validation, and test sets [50]. This method ensures each fold contains entire networks, increasing model robustness by preventing instances from the same network from appearing in the training, validation and test sets simultaneously.

We built and evaluated the machine learning models using the `Scikit-learn` package [51], with the addition of the `xgboost` package [52]. We executed hyperparameter optimization through randomized search using the `scikit-learn`’s `RandomizedSearchCV` [51]. We used randomized optimization due to

its efficiency in exploring the hyperparameter space with fewer iterations compared to traditional grid search method. We optimized the model’s hyperparameters using the F1-score.

## Link withholding

Because the goal is to predict missing links (possible yet unobserved interactions [53]), link prediction requires generating a ground truth. That is, a lack of link (0 in the matrix), of which we are certain the link actually exists. A common way to do so is by withholding some existing links randomly [8,54]. Hence, to emulate real-world under-sampled networks, each network in the validation and test sets was sub-sampled by randomly withholding 20% of the existing links. We chose this proportion to balance sufficient data for training while ensuring robust evaluation of model performance. However, we ran a sensitivity analysis withholding 5-30% of the links. In addition, we tested three withholding strategies (Supplementary note S1.3). Withholding creates three types of links:

- *Existing links*: links that exist in the network and were not sub-sampled.
- *Non-existing links*: links that did not exist in the network. These may be missing or forbidden (interactions not possible due to some ecological, morphological or other constraints [53]).
- *Withheld links*: links that existed in the network and are now missing because they were removed. Because we want to predict instances in which non-existing links should have existed, withheld links were relabeled as existing after feature extraction.

## Feature extraction

Each instance in the data set, representing an interaction (or lack of an interaction), between each two species, constitutes a vector of topological features. Our features encompassed three levels as follows:

- Network-level features: Defined for a whole network (e.g., nestedness). Hence, all the instances that are related to a network will get the same values for those features. We also included the type of network (e.g., plant-pollinator).
- Link-level features: Defined for a pair of nodes (e.g., preferential attachment: the multiplication of both node degrees).
- Node-level features: Defined for each node (e.g., centrality, degree). Each instance will have two different versions of the feature, one for each node in the pair (higher and lower trophic levels).

Feature extraction is done once on all the data, and is not related to the fold. We rescaled numerical features to a range of [0, 1] to ensure that no particular feature dominates others during the learning process. We calculated feature importance based on the average decrease in Gini impurity across all trees. Features that are more important are used more frequently and result in improvements in node purity. For the analysis of ecological networks and extraction of topological features (network properties), we used the `networkx` package [55] in Python and the `igraph` [56] and `bipartite` [57] R packages. We handled data manipulation through `numpy` [58] and `Pandas` [59,60] in Python. A complete list of the features we used and their descriptions are provided on the GitHub repository accompanying this article ([https://github.com/Ecological-Complexity-Lab/eco\\_ILP/blob/main/results/final/features.csv](https://github.com/Ecological-Complexity-Lab/eco_ILP/blob/main/results/final/features.csv)).

While some of the features we used were inevitably correlated [27], the overall correlation was not high (generally below 0.5). We present a summary of the correlation values in Fig. S12, as well as the detailed correlation matrices between features at the network (Fig. S13), node (Figs. S14, S15) and link (Fig. S16) levels.

## Feature selection

Prior to training, we explored correlations using a correlation matrix. While some of the features we used were inevitably correlated [27], the overall correlation was not high (generally below 0.5, see feature extraction in Methods). Moreover, given that the study’s goal was to perform a large-scale pattern detection rather than to identify features that drive specific patterns, we did not a-priori remove features. In addition, we mitigated correlations by feature selection, which is a common process in machine learning. Feature selection also allowed us to reduce dimensionality, mitigate over-fitting, and identify the most informative features for link prediction.

We implemented feature selection as an integral part of the modeling process using a pipeline-based approach, employing mutual information as the selection criterion to capture both linear and non-linear relationships between features and the target variable (link presence/absence). We used scikit-learn’s SelectKBest with `mutual_info_classif` as the scoring function. This method evaluates the mutual dependence between each feature and the target variable without assuming a specific relationship type. We chose mutual information over alternative metrics (e.g., correlation coefficients or chi-squared tests) for its robustness in capturing non-linear dependencies. The number of selected features was treated as a hyperparameter, optimized through cross-validation alongside other model parameters using `RandomizedSearchCV`. We tested several feature set sizes ( $k \in \{10, 20, 30, 40, 50\}$ ), allowing the cross-validation process to determine the optimal number of features based on the specific dataset and model configuration.

In the inductive model, feature selection was performed once per each outer fold, and the selected features were consistently applied to all predictions of that fold. Cross-validation ensured the stability of feature selection across different data splits. In contrast, the transductive model applied feature selection independently for each network, enabling the model to adapt its feature set to the unique characteristics of each network. This approach integrated feature selection directly into the prediction task, ensuring flexibility and precision across different ecological contexts.

## Dealing with class imbalance

In machine learning, class imbalance is a situation in which the distribution of classes in the training data is highly skewed, where one or more classes have considerably fewer samples compared to others. This imbalance can significantly impact the training and evaluation of prediction models [41]. Classifiers developed with such skewed data tend to favor the majority class, which can lead to subpar performance when identifying instances of the minority class. This issue is particularly prevalent in ecological networks, characterized by their sparsity (i.e., low connectance). In binary classification tasks, this sparsity creates a disparity between the small number of existing links (positive class) and the much larger set of non-existing links (negative class) [11,41]. To overcome this problem we incorporated cost-sensitive learning (Supplementary note S1.4) [51]. Specifically, to make the importance of both classes equal, we computed their weights inversely proportionally to the frequency of the respective classes [61,62].

## Comparison to transductive models

We compared our model to four transductive link prediction models: The SBM, connectance, and matching-centrality, which were previously published [8] and are not based on machine learning, and to a transductive machine learning model we developed. All four models train and test on each network separately and do not separate links between the train and test set.

In *SBM*, nodes are partitioned into blocks or groups, and the probability of a link between any two nodes depends on the blocks to which they belong [63]. The probability of a link between two nodes is higher if they belong to the same block, reflecting community structure. The stochastic block model is a degree corrected bipartite SBM algorithm, which accounts for the degree heterogeneity within the blocks [64]. The *connectance model* is a model used to describe the pattern of interactions in ecological networks. It assigns a connectivity value to each species, reflecting its propensity to interact with others. The connectivity values are estimated by using maximum likelihood optimization, which adjusts the parameters to best fit the observed interaction data [8]. The *matching-centrality model* combines a trait-based framework with connectivity values to predict interactions in ecological networks. Each species is assigned a latent trait value, representing an unmeasured characteristic influencing its interaction propensity. The model calculates the probability of an interaction between two species as a function of their trait similarity, with higher probabilities assigned to species pairs with similar traits. Additionally, it incorporates connectivity values to account for species' interaction frequencies. Parameters are estimated through maximum likelihood optimization, balancing both trait similarity and species connectivity to fit observed interaction data [8]. We used the R package *cassandRa* [65] to predict links using these two models. The TLP model we developed uses random forest with a standard 3-fold cross validation with hyper-parameters tuning (maximizing f1 score). The model balances the weights of the classes by computing weights inversely proportionally to the frequency of the respective classes.

## Estimating the bounds of link prediction performance

We evaluate the models' ability to predict links under a range of true missing link values. For a theoretical proportion  $p$  (range 0.05-0.5) of ground-truth missing links, the **number** of missing links is calculated as  $L_m = p \times L_{ne}$ , where  $L_{ne}$  is number of zeros (non-existing links) in the original matrix. In the *best case scenario*, we choose  $L_m$  FP links and change them to TP. This simulates a scenario in which the model indeed predicted links that were not observed. If  $L_{FP} < L_m$  ( $L_{FP}$  is the number of FP links) we convert a remainder of  $L_m - L_{FP}$  TN links to FN to simulate the scenario in which the links actually existed in nature. In the *worst-case scenario*, we choose  $L_m$  TN links and convert their class to FN. This simulates a scenario in which the links actually existed in nature but the model failed to retrieve them. If  $L_{TN} < L_m$  ( $L_{TN}$  is the number of TN links) we convert a remainder of  $L_m - L_{TN}$  FP links to TP. After the conversion of links, we recalculate each evaluation metric (e.g., F1, recall) for each of the two scenarios, to form the upper and lower bounds of model performance per network.

## Code and data availability

The data are available in the repository set up in original publication <https://osf.io/my9tv/>. The full code and technical descriptions on how to run the pipeline are available on the GitHub repository [https://github.com/Ecological-Complexity-Lab/eco\\_ILP](https://github.com/Ecological-Complexity-Lab/eco_ILP).

## Acknowledgments

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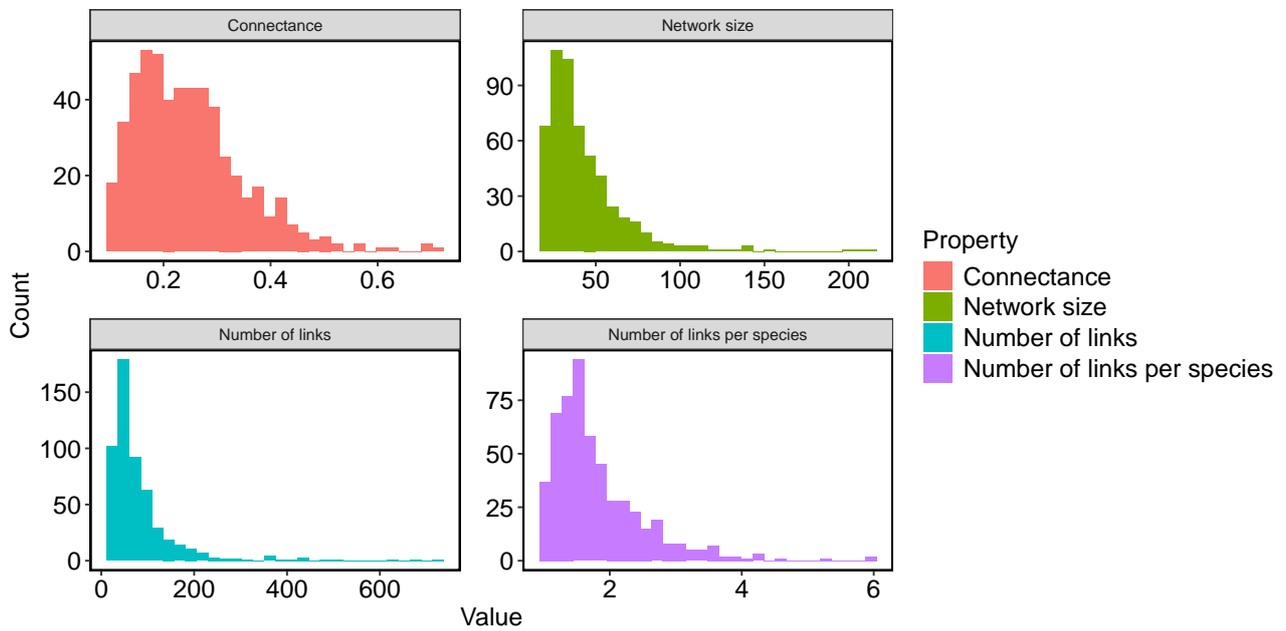
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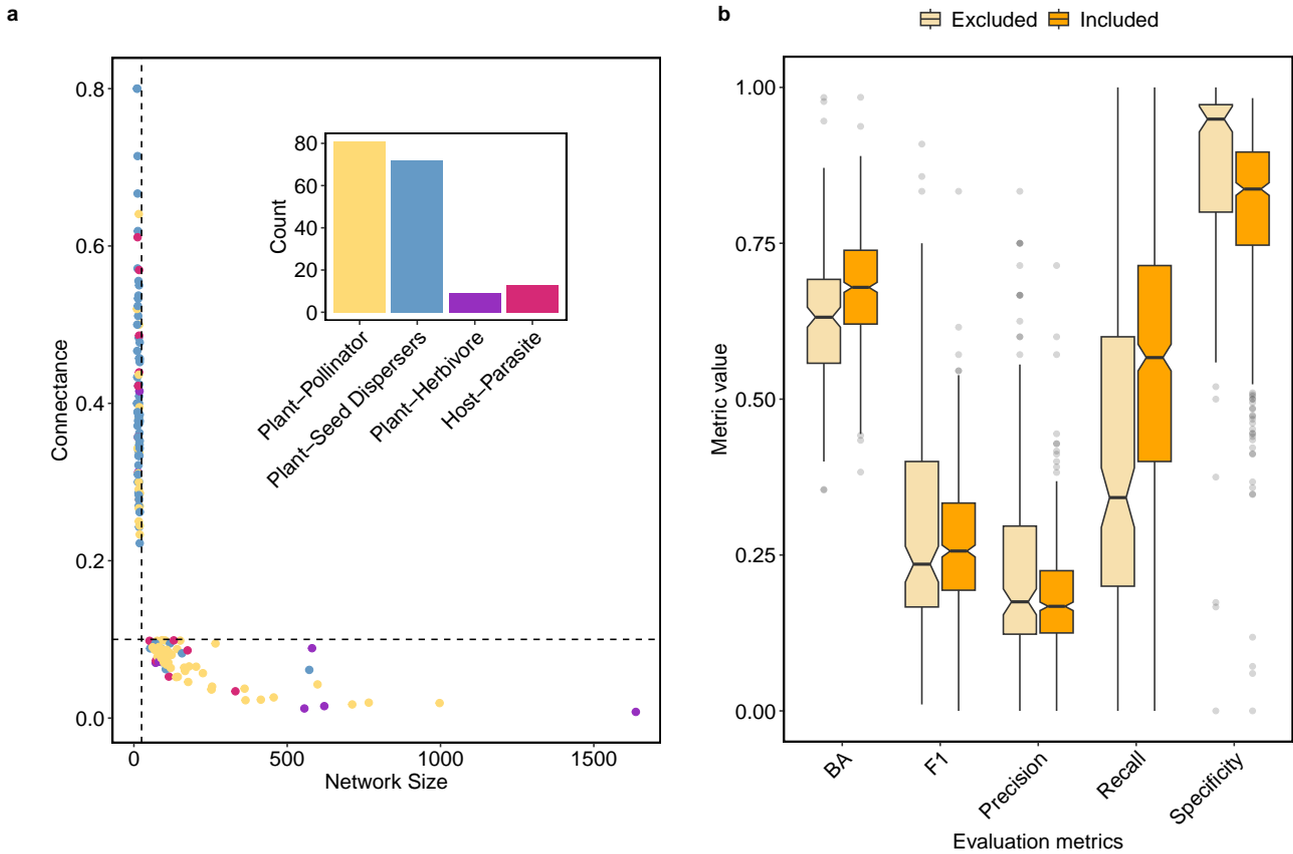
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# Supplementary Information

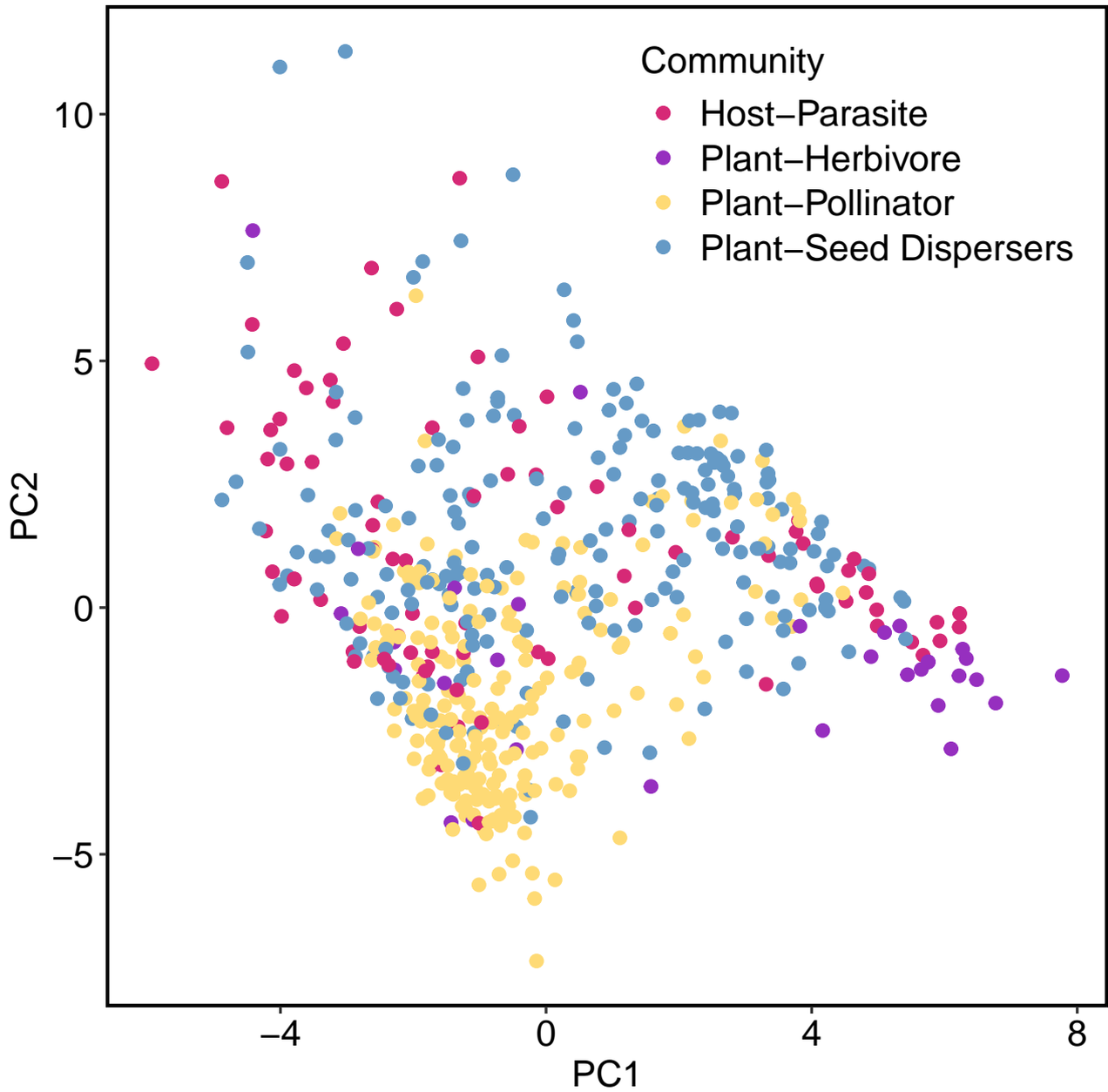
## Supplementary figures and tables



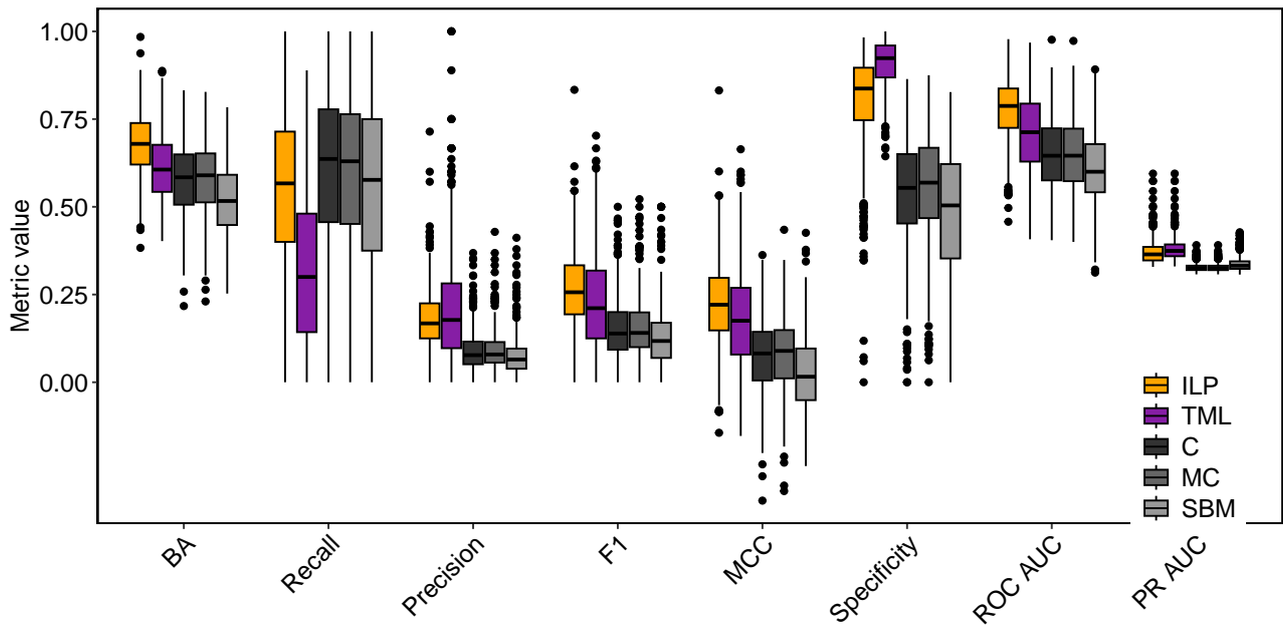
**Fig. S1: Distributions of basic network properties of the networks included in the analysis.** Number of links is the average number of links per species.



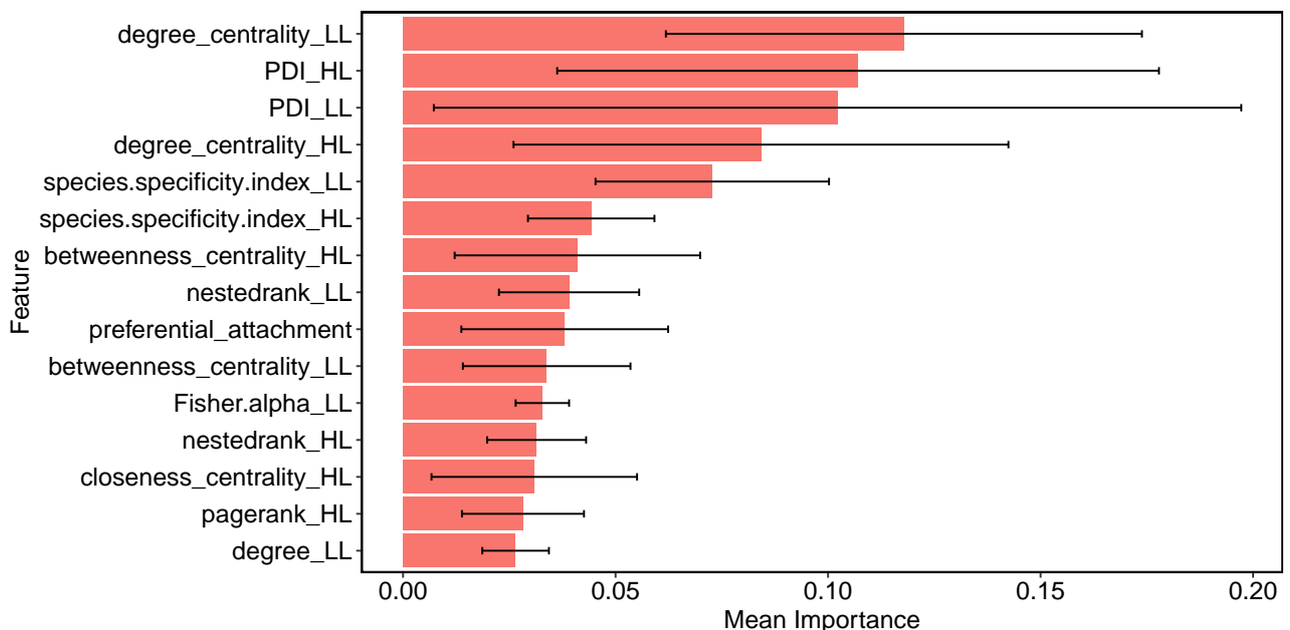
**Fig. S2: Networks excluded from analysis. (A)** The relationship between network size (x-axis) and connectance (y-axis) for networks (data points) excluded from the training set. Networks were excluded if their connectance was  $< 0.1$  (below the horizontal line) or their size was  $< 25$  species (to the left of the vertical line). The inset summarizes the number of excluded networks. Including the large amount of plant-pollinator networks would reduce the model's performance due to their sparseness, as demonstrated in the manuscript. **(B)** Comparison of test performance between included and excluded networks. Data for included networks are as presented in Fig. 2 of the main text. The boxplots show the interquartile range (box), median (central line), data within 1.5 times the interquartile range (whiskers), and potential outliers (individual points beyond whiskers). Each data point is a network and boxplots contain networks from all the five outer folds ( $n = 538$  networks in all the box plots). A significant difference in medians is indicated by non-overlapping notches. Overall, the excluded networks exhibit poorer test performance, with higher specificity attributed to their extreme sparsity, facilitating the detection of true negatives (0's).



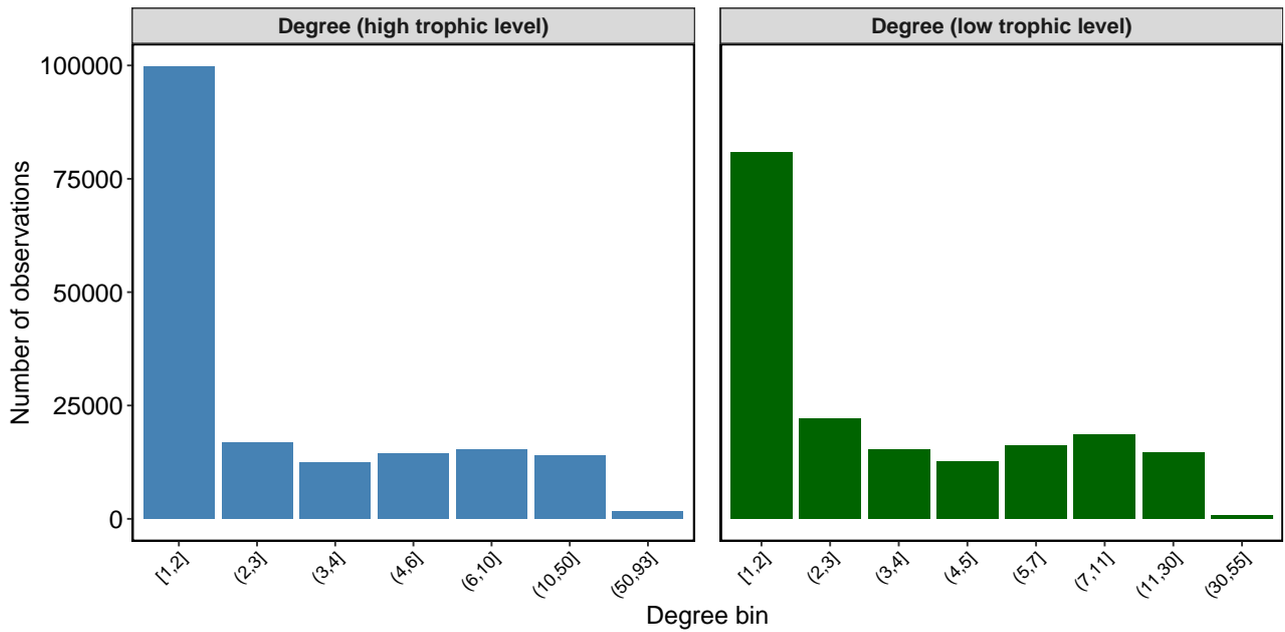
**Fig. S3: Variation in network structure.** The PCA shows no clear subgrouping of networks by their class, reflecting previous studies using the same data set [1,2]. The PCs were calculated using the topological features we used in the link prediction model.



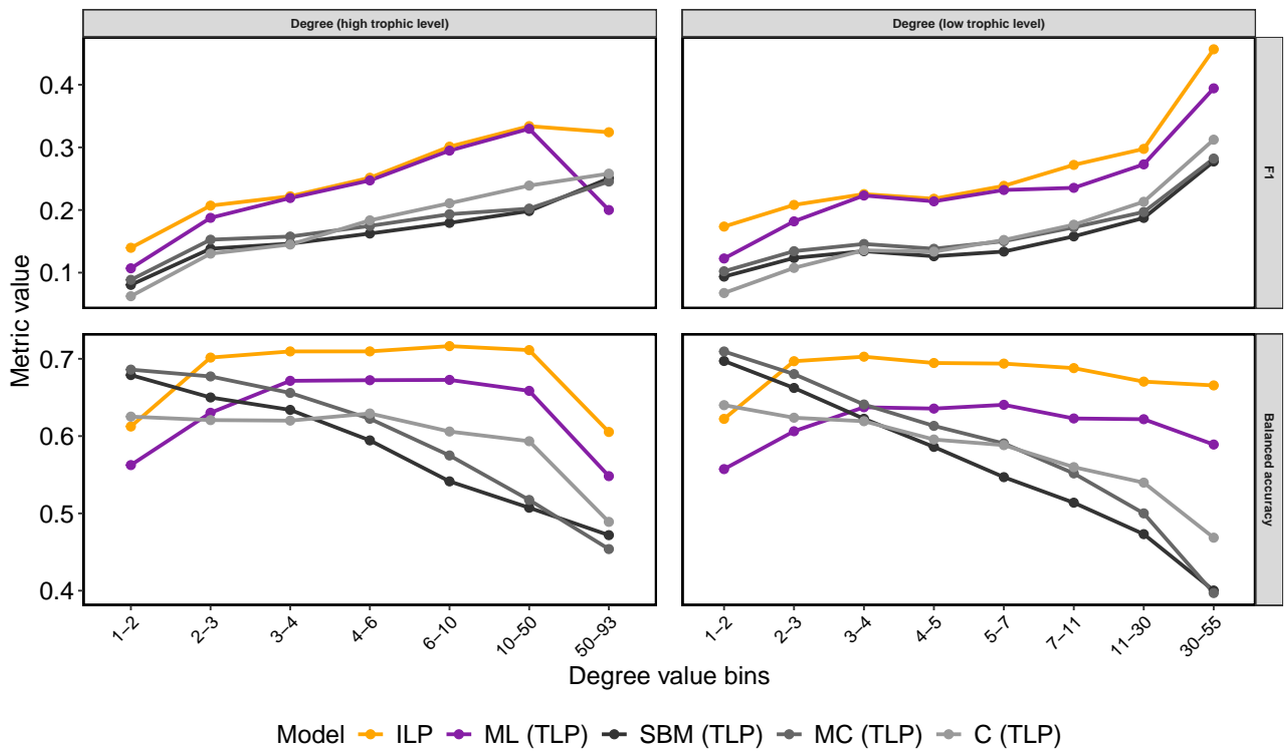
**Fig. S4: Comparison of model performance.** The inductive link prediction model was trained and tested on networks from all communities. We compared ILP to three transductive previously published models (stochastic block model, connectance (C) and matching centrality (MC)<sup>3</sup>), and to a transductive machine learning model that we trained (TLP). Each boxplot is a distribution of an evaluation metric using a 0.5 classification threshold, besides the ROC and PR AUC. The boxplots show the interquartile range (box), median (central line), data within 1.5 times the interquartile range (whiskers), and potential outliers (individual points beyond whiskers). Each data point is a network and boxplots contain networks from all the five outer folds. BA is balanced accuracy. See definitions of evaluation metrics in S1.2 Exploration of model evaluation.



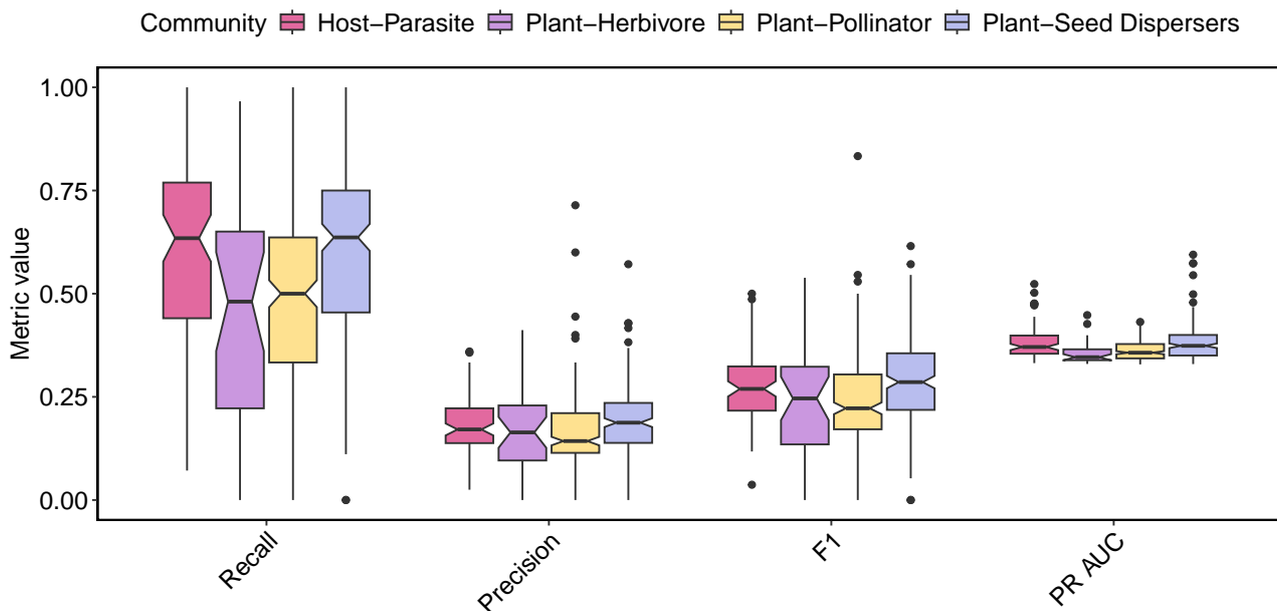
**Fig. S5: Feature importance.** Each bar presents the average value of feature importance across the five folds for the random forest model. Error bars depict minimum and maximum. HL are higher-level trophic species that consume resources (pollinators, parasites, seed dispersers, herbivores). LL are lower-level trophic species (resources) (plants, hosts, seeds). Features are invariably related to degree. The link-level feature preferential attachment (the multiplication of the two interacting species' degrees). Feature importance was computed based on the average decrease in Gini impurity across all trees.



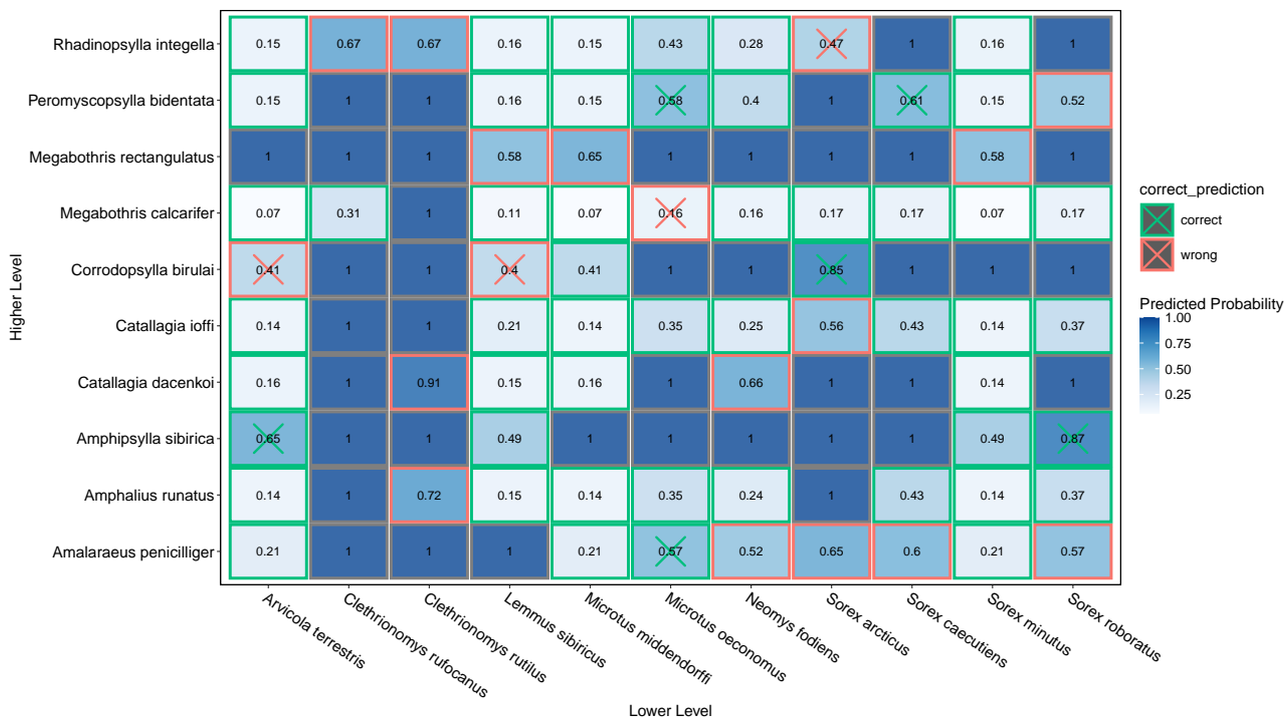
**Fig. S6: Counts of observations in species degrees.** This figure presents the underlying counts for the bins presented in Fig. 3.



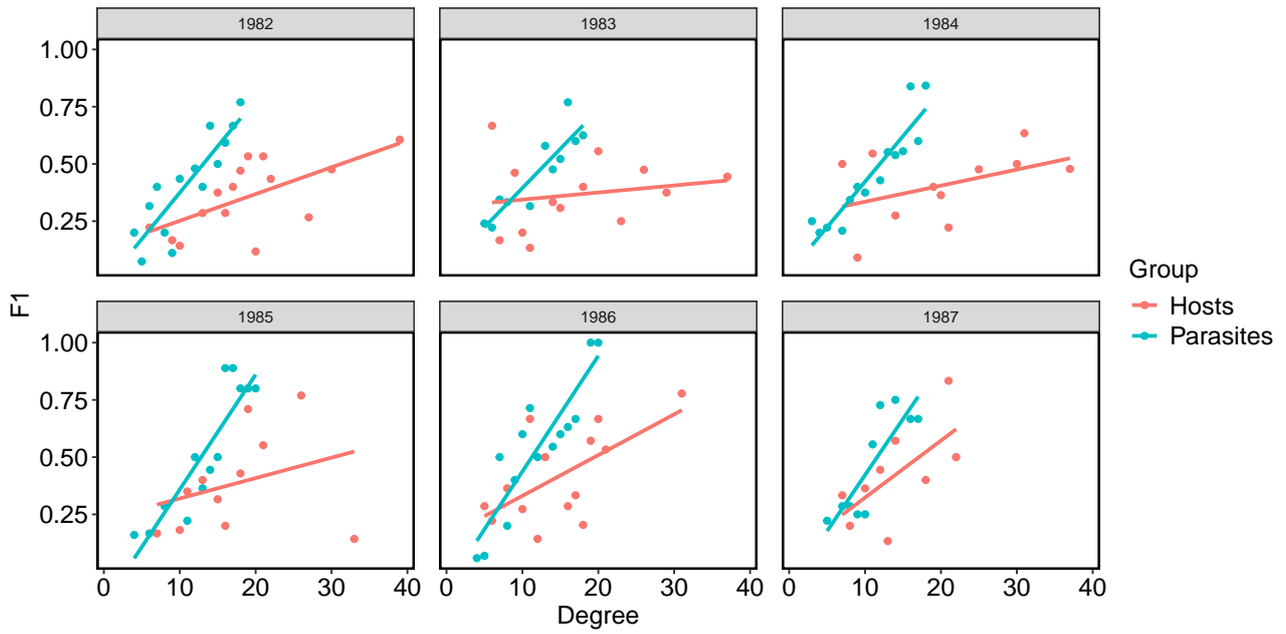
**Fig. S7: Comparison of model performance across node degrees.** We present two evaluation metrics that complement those in Fig. 3.



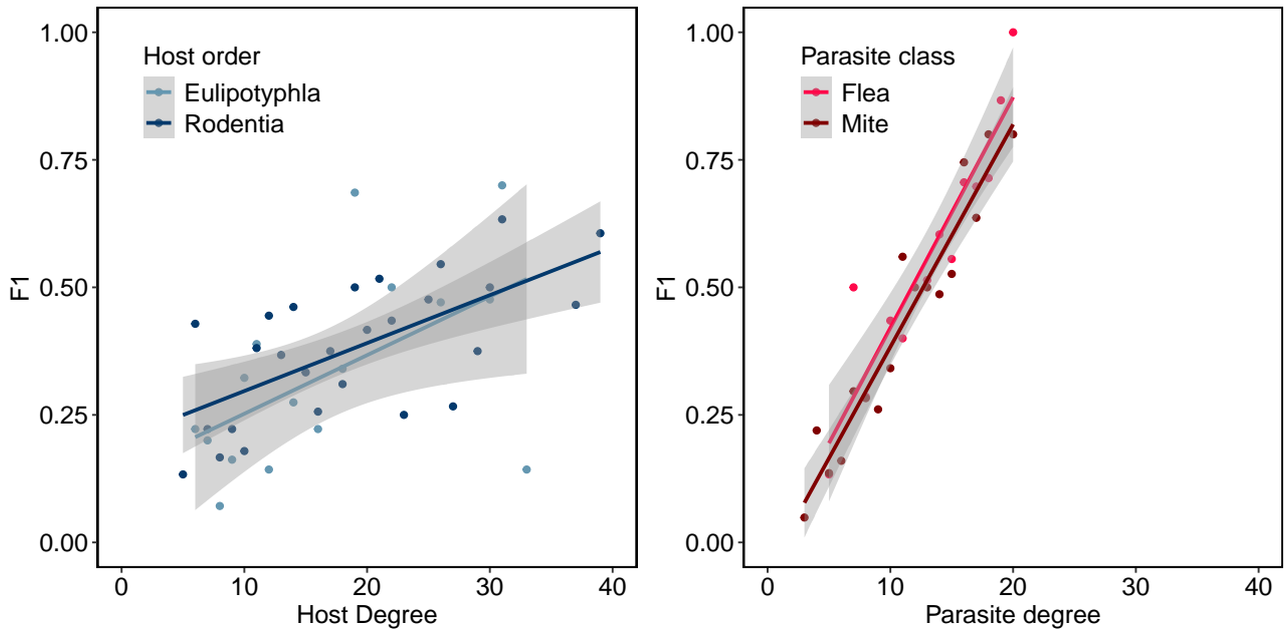
**Fig. S8: Distributions of evaluation metrics per test community type.** The model was trained on networks from all communities, and tested separately on each community type. Each data point is a network and box plots contain networks from all the five outer folds. The boxplots show the interquartile range (box), median (central line), data within 1.5 times the interquartile range (whiskers), and potential outliers (individual points beyond whiskers). Sample sizes (across 5 folds): Host-Parasite: 336, Plant-Herbivore: 128, Plant-Pollinator: 868, Plant-Seed Dispersers: 820.



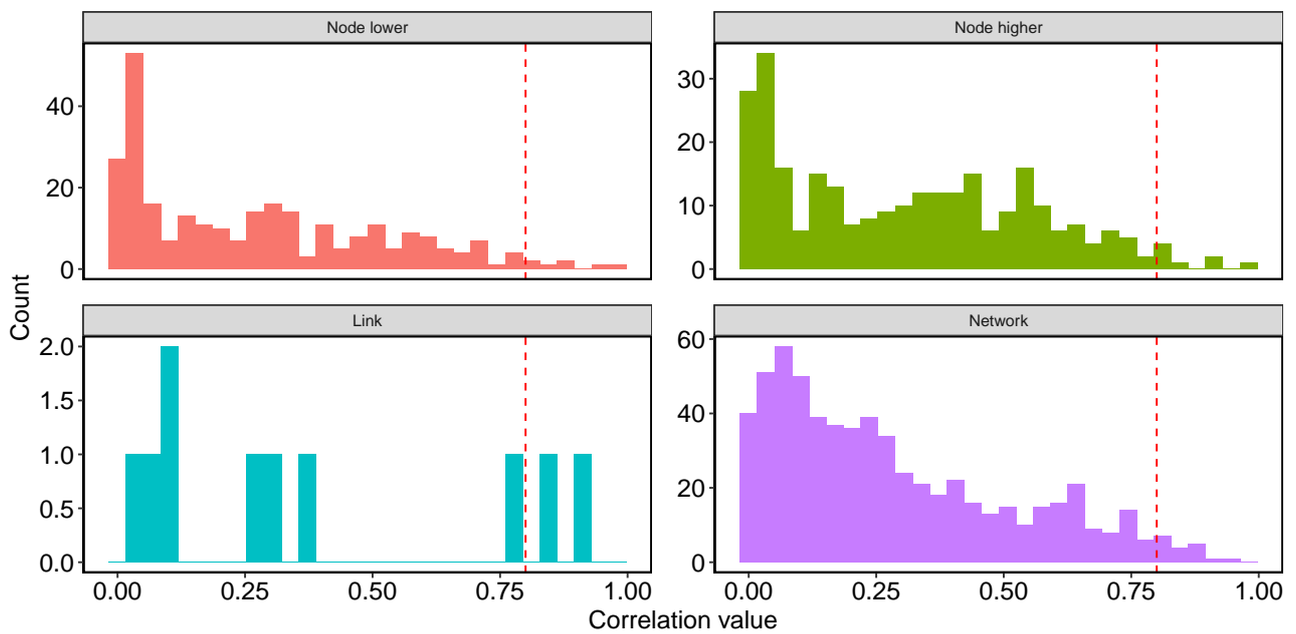
**Fig. S9: Link prediction example for a host-parasite network.** Values inside matrix cells are the predicted link probabilities. The sub-sampled links are marked with X. Cells above the 0.5 threshold are classified as links. Correct and wrong classifications are colored green and red, respectively. Therefore, true positives are green with X, true negatives are green, false positives are red (no X) and false negatives are red with X.



**Fig. S10: Correlation between degree and F1 per year.** The plots qualitatively reflect the results in Fig 7.



**Fig. S11: Correlation between degree and F1.** The plots reflect the results in Fig 7, separately for host and parasite groups.



**Fig. S12: A histogram of feature correlation values.** We only kept one of the features when the correlation was above the 0.8 threshold (indicated with a dashed red line. Overall, correlations were below 0.5.

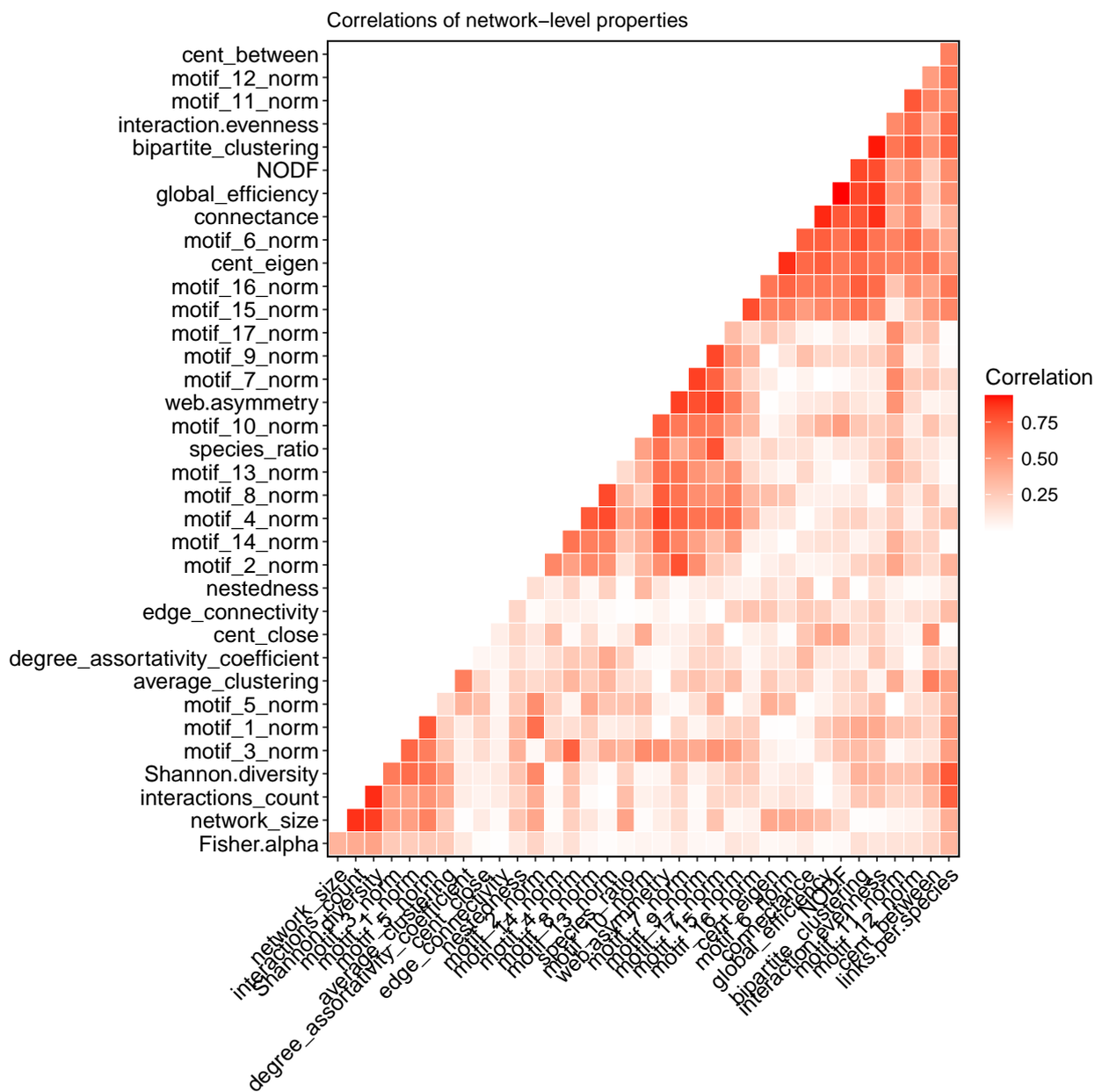
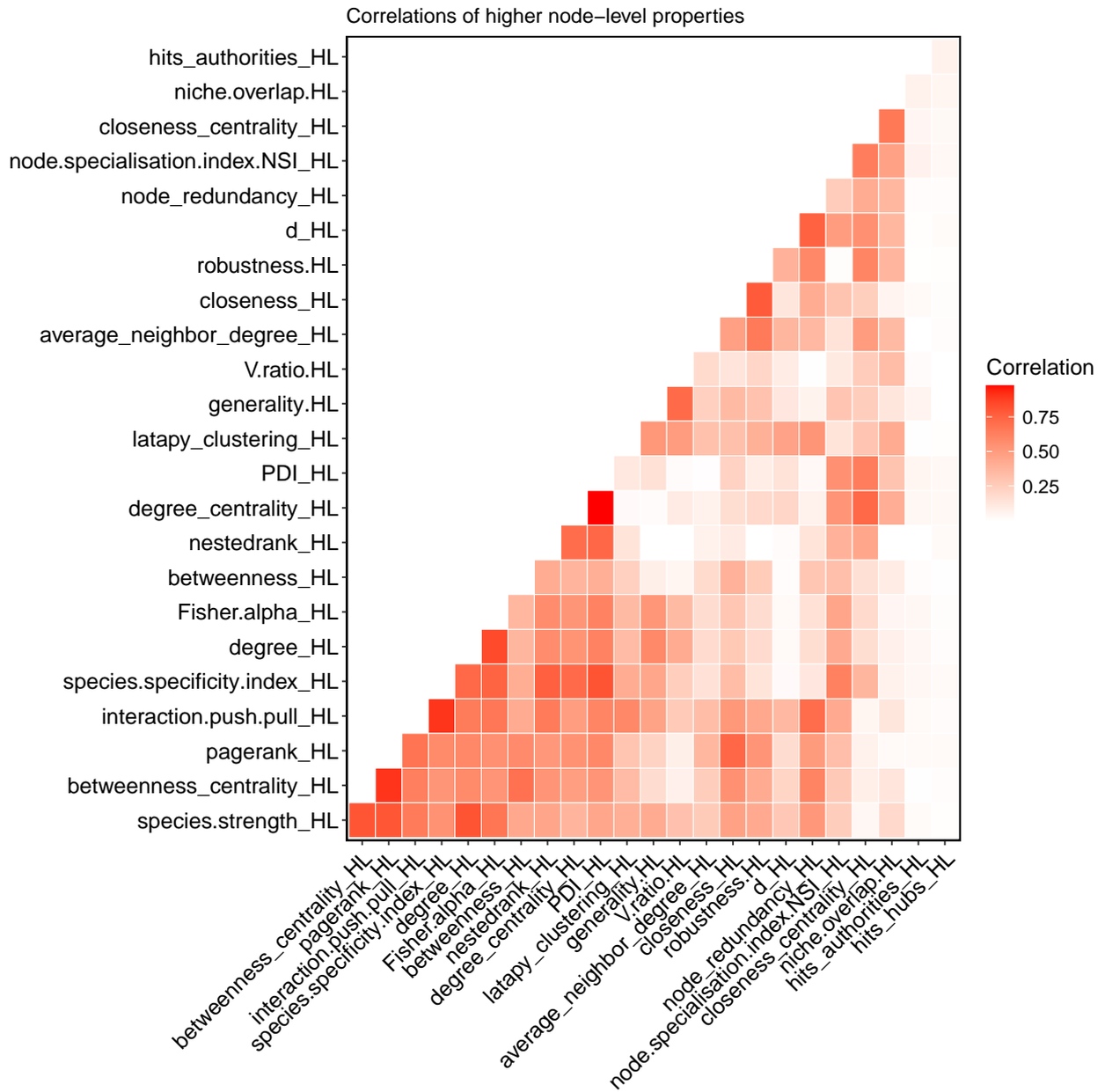
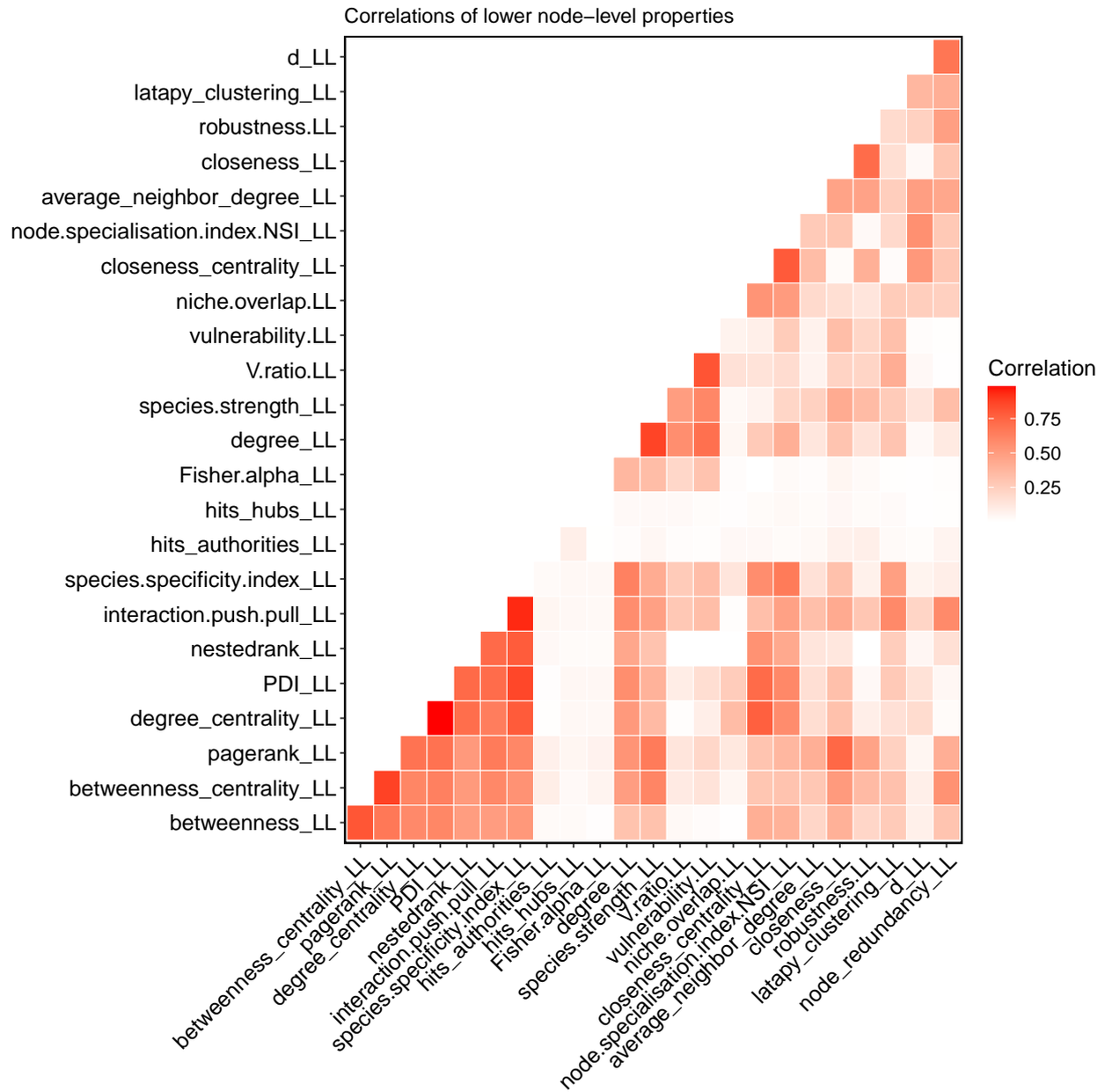


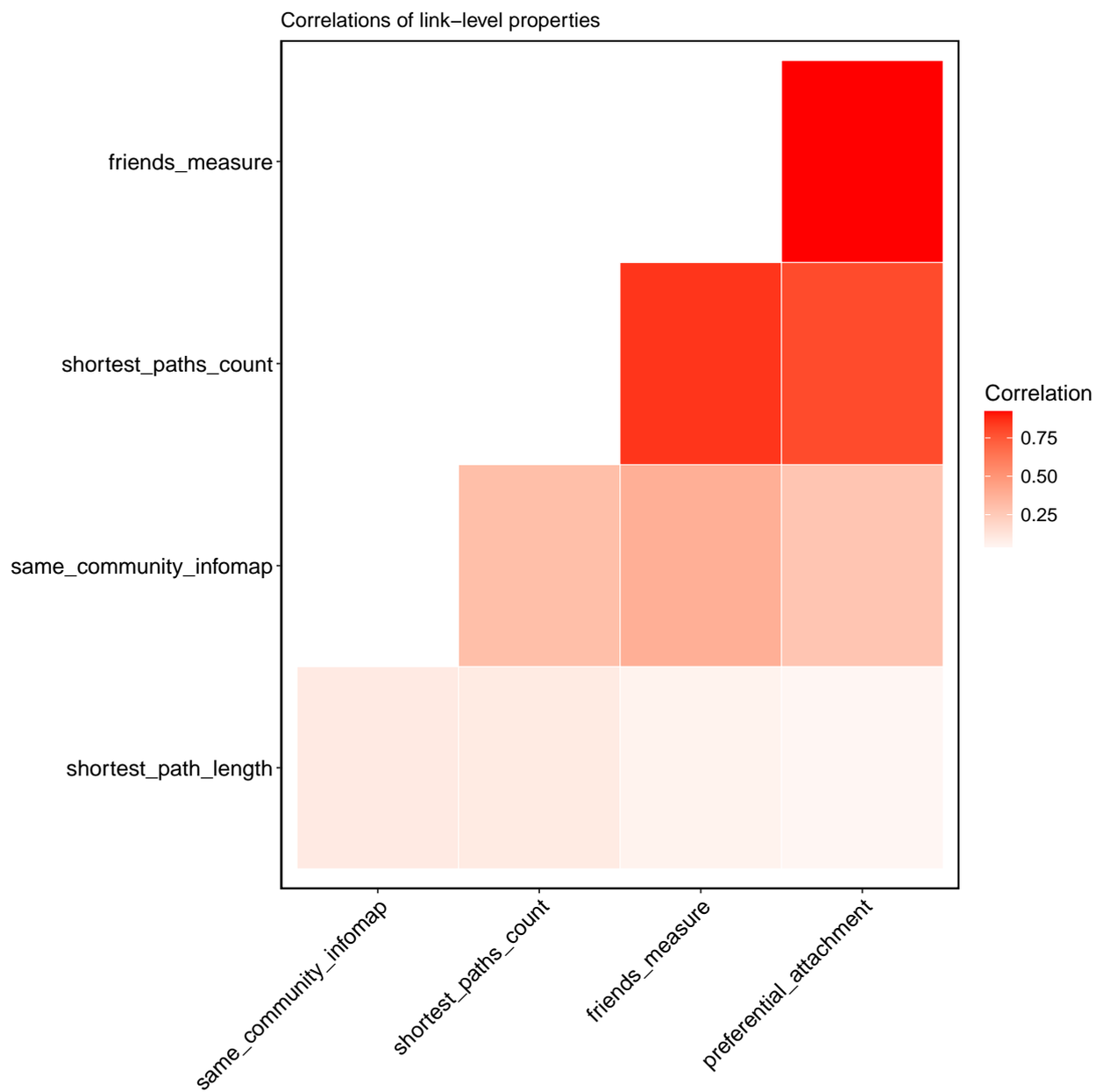
Fig. S13: Correlation between features at the network level.



**Fig. S14: Correlation between features at the node level for the higher-trophic level (e.g., parasites, pollinators).**



**Fig. S15: Correlation between features at the node level for the lower-trophic level (e.g., hosts, plants).**



**Fig. S16: Correlation between features at the link level.**

**Table S1: Summary of network properties.** The table provides an overview of the network properties for different ecological communities, detailing each variable's mean, median, and range values.  $N$ : number of nodes;  $L_e$ : number of existing links;  $L_{ne}$ : number of nonexisting links;  $C$ : connectance,  $C = L_e/(L_e+L_{ne})$

Variable	Community	Mean	Median	Range
$N$	Host-Parasite	37.58	33	20-137
	Plant-Herbivore	42.53	37	21-78
	Plant-Pollinator	43.36	36	20-205
	Plant-Seed Dispersers	43.96	36	20-213
$L_e$	Host-Parasite	82.86	63	19-490
	Plant-Herbivore	67.69	42.5	22-294
	Plant-Pollinator	72.94	52	21-631
	Plant-Seed Dispersers	98.59	72	21-720
$L_{ne}$	Host-Parasite	1622.08	991.5	349-18279
	Plant-Herbivore	1997.22	1332.5	409-5996
	Plant-Pollinator	2394.52	1260	363-41600
	Plant-Seed Dispersers	2585.2	1238	362-44687
$C$	Host-Parasite	0.28	0.25	0.11-0.61
	Plant-Herbivore	0.2	0.165	0.1-0.47
	Plant-Pollinator	0.22	0.2	0.1-0.42
	Plant-Seed Dispersers	0.28	0.27	0.1-0.71

**Table S2:** Kruskal-Wallis and Dunn’s test results for pairwise comparisons of metrics between communities. The KW test was performed for each metric separately, with communities as factor levels. All Comparisons were significant, and we therefore performed a Dunn post-hoc test (p value adjusted using Bonferroni correction) for all of them. Note that the median values are close for some comparisons despite being statistically significant. A visualization of the values and their distributions is in the main text, in Fig. 4B.

Metric	KW statistic	Community 1	Community 2	Median 1	Median 2	P value (adjusted)
Recall	39.03	HP	PH	0.63	0.48	0.019
Recall	39.03	HP	PP	0.63	0.50	0.001
Recall	39.03	HP	PSD	0.63	0.64	1.000
Recall	39.03	PH	PP	0.48	0.50	1.000
Recall	39.03	PH	PSD	0.48	0.64	0.004
Recall	39.03	PP	PSD	0.50	0.64	0.000
Precision	20.54	HP	PH	0.17	0.16	1.000
Precision	20.54	HP	PP	0.17	0.14	0.060
Precision	20.54	HP	PSD	0.17	0.19	1.000
Precision	20.54	PH	PP	0.16	0.14	1.000
Precision	20.54	PH	PSD	0.16	0.19	0.415
Precision	20.54	PP	PSD	0.14	0.19	0.000
Specificity	53.43	HP	PH	0.79	0.91	0.000
Specificity	53.43	HP	PP	0.79	0.87	0.000
Specificity	53.43	HP	PSD	0.79	0.82	1.000
Specificity	53.43	PH	PP	0.91	0.87	0.088
Specificity	53.43	PH	PSD	0.91	0.82	0.000
Specificity	53.43	PP	PSD	0.87	0.82	0.000
F1	27.82	HP	PH	0.27	0.25	0.611
F1	27.82	HP	PP	0.27	0.22	0.023
F1	27.82	HP	PSD	0.27	0.29	1.000
F1	27.82	PH	PP	0.25	0.22	1.000
F1	27.82	PH	PSD	0.25	0.29	0.099
F1	27.82	PP	PSD	0.22	0.29	0.000
BA	9.12	HP	PH	0.68	0.65	1.000
BA	9.12	HP	PP	0.68	0.66	1.000
BA	9.12	HP	PSD	0.68	0.69	1.000
BA	9.12	PH	PP	0.65	0.66	1.000
BA	9.12	PH	PSD	0.65	0.69	0.275
BA	9.12	PP	PSD	0.66	0.69	0.040
PR AUC	47.86	HP	PH	0.37	0.35	0.000
PR AUC	47.86	HP	PP	0.37	0.36	0.000
PR AUC	47.86	HP	PSD	0.37	0.37	1.000
PR AUC	47.86	PH	PP	0.35	0.36	0.820
PR AUC	47.86	PH	PSD	0.35	0.37	0.000
PR AUC	47.86	PP	PSD	0.36	0.37	0.000

## S1 Supplementary notes on methods

### S1.1 Models applied

Selecting a machine learning model a priori can be challenging<sup>4</sup>. There is no one-size-fits-all solution; different models have different strengths and weaknesses, and knowing which model will perform better is often impossible. Some models may perform well on particular data types, while others may perform better on different ones. Hence, model selection often requires some experimentation. In practice, trying multiple models and comparing their performance is often a good idea before deciding on a final model. Furthermore, using a model ensemble—a technique in which multiple models are combined—can often lead to better results than a single model<sup>5</sup>. In this study, we tried multiple models and their ensemble. For the ensemble, we averaged the probabilities for each prediction. The models performed similarly overall, and their ensemble did not improve the results (Fig. S17). Therefore, in the main text, we present results for random forest. In this section, we describe the models we used. We present model hyperparameters in Table S3.

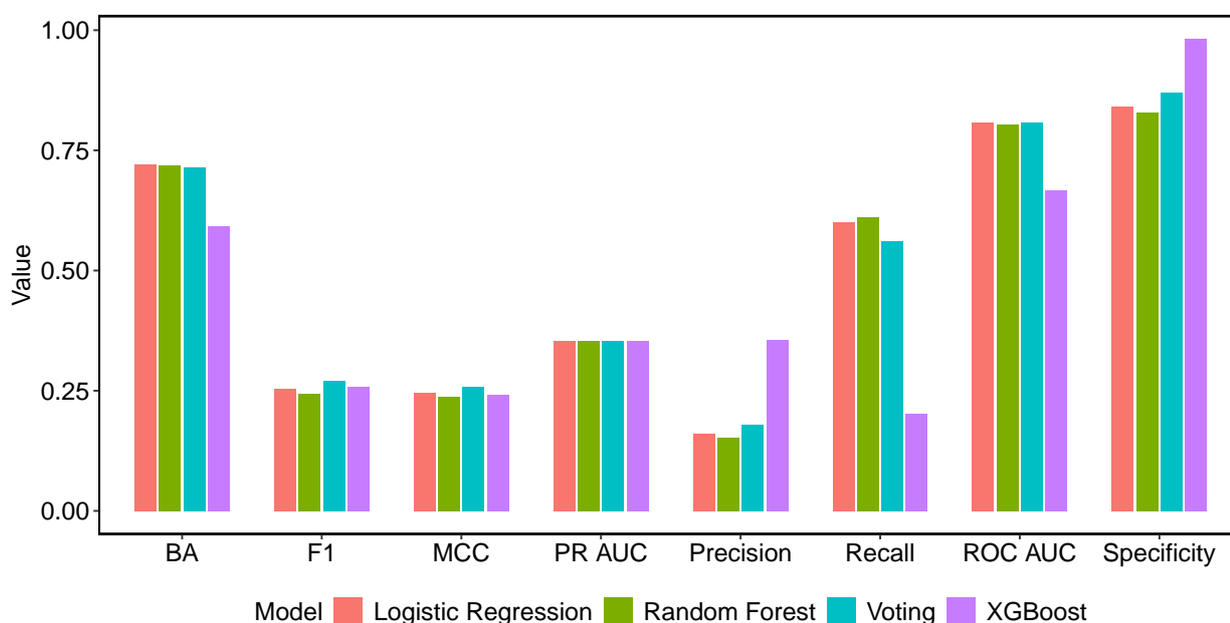
To understand how these models work, it is necessary to first explain the terms bagging and boosting<sup>6</sup>, two popular ensemble learning techniques. The key difference between bagging and boosting lies in how they combine multiple models to make predictions. Bagging, which stands for bootstrap aggregating, is a parallel ensemble technique. In bagging, multiple base models (multiple instances of the same algorithm) are trained independently on different random subsets of the training data using bootstrap sampling (sampling with replacement). The outputs of the individual models are then combined, often through a voting mechanism or by taking the average of their predictions, to produce a single output. Bagging is often used with decision trees as it has been repetitively shown to outperform other models. Boosting, on the other hand, is a sequential ensemble technique. Boosting works by iteratively training multiple weak models on modified versions of the training data, with each subsequent model trying to correct the errors of the previous models, focusing on the examples that the previous models misclassified. The training data is re-weighted at each iteration so that the misclassified examples receive higher weights and are given more importance in subsequent iterations. The final prediction of the boosted model is a weighted combination of the predictions of all the individual models, with the weights determined by the performance of each model on the training data. More weight is given to models that achieved higher performance.

**Logistic regression.** This generalized linear model uses a logistic function to model a binary dependent variable. The logistic function maps the linear combination of input variables to a discrete binary value between 0 and 1, which is interpreted as the probability of the input belonging to a particular class. During training, the logistic regression algorithm finds the input variables' best parameters (coefficients) by minimizing the error between the predicted probabilities and the actual class labels in the training data. Logistic regression is easy to interpret and implement without necessarily compromising performance<sup>7</sup>.

**Random Forest.**<sup>8,9</sup> This is a specific version of a bagging method with decision trees. A random forest model creates a collection of decision trees and combines their predictions to produce a final output. Decision tree models create a tree-like model of decisions and their possible consequences, with each internal node representing a feature, each branch representing a decision rule, and each leaf node representing the outcome class label. The algorithm starts at the root of the tree. It recursively

splits the data into subsets using a feature that provides the most information gain at that stage until it reaches a leaf node, representing the predicted class label for the input data. Each tree in the random forest ensemble is trained on a different random subset of the data and features, and the final output is the combination of the predictions of all the decision trees, usually made by either averaging the results for regression tasks or by taking a majority vote for classification tasks.

**XGBoost.** This gradient-boosting tree-based algorithm is designed for speed and performance<sup>10</sup>. Gradient boosting<sup>11</sup> differs from other boosting algorithms in the way it calculates the weights of the models. Specifically, gradient boosting uses the gradient descent optimization algorithm to minimize the model’s loss function. At each iteration, the algorithm calculates the (negative) gradient of the loss function with respect to the predictions of the previous ensemble. It uses this gradient to adjust the weights of the new model. The negative gradient represents the direction of the steepest descent for the loss function, which is the direction that will reduce the loss the most. By fitting a new tree to the negative gradient, gradient boosting focuses on the examples misclassified in the previous ensemble and attempts to correct those errors.



**Fig. S17: Model comparison.** We evaluated link prediction using various models. Average is a model ensemble. Overall, model choice did not qualitatively affect the results. Therefore, we present results of random forest throughout the main text.

**Table S3: Summary of model hyperparameters.** The table provides an overview of the best values for different parameters across five folds for each model. We optimized hyperparameters using a random search. We used the F1-score as the performance metric to evaluate the hyperparameter combinations. RFC: RandomForestClassifier; LR: LogisticRegression; XGB: XGBClassifier.

Model	Parameter	Fold 1	Fold 2	Fold 3	Fold 4	Fold 5	Range
RFC	n_estimators	15	10	15	50	15	[10, 15, 20, 50, 100, 300]
RFC	min_samples_split	5	1	10	1	5	[1, 2, 3, 4, 5, 10]
RFC	min_samples_leaf	12	4	3	10	7	[3, 4, 5, 6, 7, 8, 9, 10, 12, 15, 20]
RFC	max_samples	0.6	0.1	0.4	0.7	0.4	[0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9]
RFC	max_leaf_nodes	128	16	16	16	32	[2, 4, 8, 16, 32, 64, 128]
RFC	max_features	log2	sqrt	log2	log2	log2	['sqrt', 'log2']
RFC	max_depth	30	60	20	None	3	[3, 5, 7, 10, 20, 30, 40, 50, 60, None]
RFC	criterion	gini	entropy	entropy	gini	gini	['gini', 'entropy']
RFC	bootstrap	True	True	True	True	True	[True]
RFC	feature_selector_k	70	40	70	50	70	[10, 20, 30, 40, 50, 70]
LR	solver	saga	saga	saga	saga	saga	['newton-cg', 'lbfgs', 'sag', 'saga', 'liblinear']
LR	penalty	l2	l2	l2	l2	l1	['l1', 'l2', 'elasticnet', 'none']
LR	max_iter	300	300	300	300	300	[300]
LR	C	0.001	0.001	0.001	0.001	0.001	[100, 10, 1.0, 0.7, 0.5, 0.3, 0.1, 0.01, 0.001]
LR	feature_selector_k	50	20	40	40	50	[10, 20, 30, 40, 50, 70]
XGB	tree_method	hist	hist	hist	hist	hist	['hist']
XGB	subsample	0.5	0.2	0.5	0.9	0.8	[0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9, 1.0]
XGB	reg_lambda	100	0.1	100	0	1e-05	[0, 1e-05, 0.01, 0.1, 1, 100]
XGB	reg_alpha	100	0	1	100	1	[0, 1e-05, 0.01, 0.1, 1, 100]
XGB	objective	binary:logistic	binary:logistic	binary:logistic	binary:logistic	binary:logistic	['binary:logistic']
XGB	n_estimators	30	50	60	60	10	[10, 20, 30, 40, 50, 60, 70, 80, 90, 100]
XGB	max_depth	7	11	15	None	5	[1, 3, 5, 7, 9, 11, 13, 15, 17, 19, None]
XGB	learning_rate	0.001	0.001	0.1	0.01	0.01	[0.001, 0.01, 0.05, 0.1, 0.2, 0.3]
XGB	gamma	0	0.1	0.3	1	0.5	[0, 0.1, 0.3, 0.5, 1]
XGB	colsample_bytree	0.2	0.4	0.2	0.1	0.3	[0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9, 1.0]
XGB	booster	gbtree	gbtree	gbtree	gbtree	gbtree	['gbtree']
XGB	feature_selector_k	30	30	70	40	50	[10, 20, 30, 40, 50, 70]

## S1.2 Exploration of model evaluation

In binary classification, the terms “positive” and “negative” refer to the two possible outcomes, where the positive class is typically used to denote the class of interest. Here, the positive class is the presence of a link. There are four possible prediction outcomes defined for binary classification models, computed by comparing the predicted and actual values of the model’s outputs (Fig. S9):

- **True Positives (TP)**: instances correctly predicted as the positive class. That is, sub-sampled links that were correctly predicted as existing links.
- **True Negatives (TN)**: instances correctly predicted as the negative class. That is, non-existing links that were correctly predicted as non-existing links.
- **False Positives (FP)**: instances falsely predicted as the positive class. That is, non-existing links that were incorrectly predicted as existing links. This is also known as a Type I error.
- **False Negatives (FN)**: instances falsely predicted as the negative class. That is, sub-sampled links that were incorrectly predicted as non-existing links. This is also known as a Type II error.

The output of machine learning models is a probability of a link (Fig. S9). This continuous output is transformed into a categorical (positive / negative) prediction via a threshold. The threshold is typically 0.5, with values  $> 0.5$  considered a link (and values below as a no-link). Therefore, classification into the four categories above depends on the threshold applied. The relationship between true positives, false positives, false negatives, and true negatives are summarized in a *confusion matrix* from which the following metrics are calculated.

**Recall or sensitivity**: The proportion of correctly predicted positive instances out of all actual positive instances. Recall is important when false negatives are costly. For example, not predicting missing links can have consequences for conservation.

$$\frac{TP}{TP + FN} \tag{S1}$$

**Precision**: Recall is typically complemented by precision. Precision is the proportion of correctly predicted positive instances out of all positive predictions:

$$\frac{TP}{TP + FP} \tag{S2}$$

Precision is important when false positives are costly. For example, falsely predicting an interaction can lead ecologists to spend research efforts trying to validate a forbidden link in nature.

**PR AUC**: There is a tradeoff between precision and recall, which depends on the prediction threshold. To evaluate this tradeoff, the area under the PR curve provides a single number that summarizes the overall performance of a model across all possible classification thresholds. Like the ROC-AUC curve, the PR curve is calculated across all thresholds.

**F1-score**: Another way to evaluate the precision-recall tradeoff is via a balanced measure of performance that reflects the classifier’s ability to identify true positive instances while avoiding false positives. The F1-score is defined as the harmonic mean between precision and recall:

$$F1 = \frac{2 * precision * recall}{precision + recall} \quad (S3)$$

**Specificity:** The proportion of correctly predicted negative instances out of all actual negative instances:

$$\frac{TN}{TN + FP} \quad (S4)$$

While recall and precision focus on true positives, specificity focuses on the retrieval of true negatives.

**Balanced accuracy:** Balanced accuracy aims to balance the prediction ability for links and no-links, particularly in situations where the classes are imbalanced. It is the average of the true positive rate (sensitivity or recall) and the true negative rate (specificity):

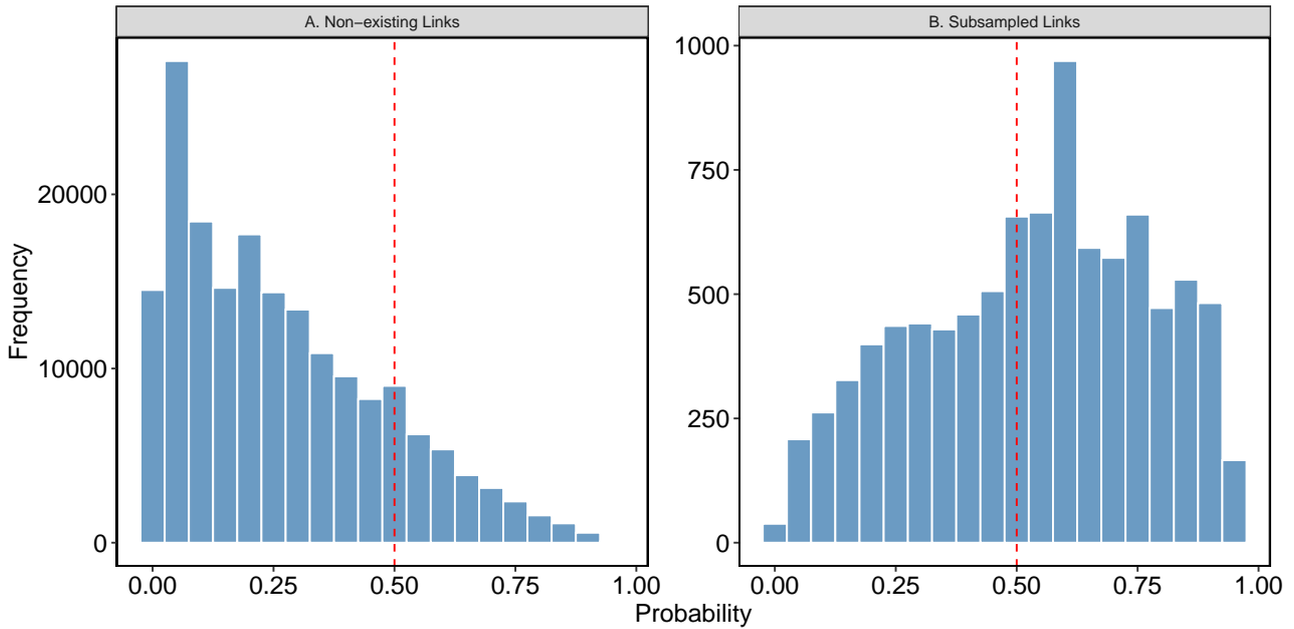
$$\frac{recall + specificity}{2} \quad (S5)$$

**MCC:** The Matthews Correlation Coefficient (MCC) takes into account the balance ratios of the four confusion matrix categories (TP, TN, FP, FN), and is, therefore, a balanced measure that can be used even if the classes are of very different sizes<sup>12</sup>.

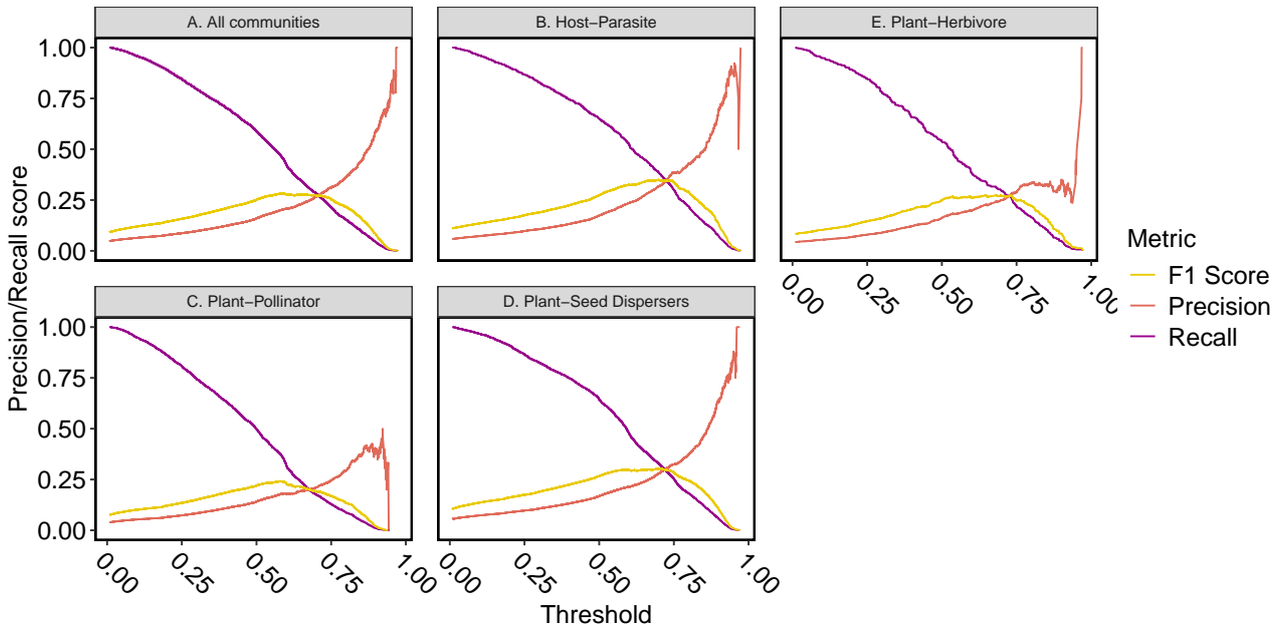
$$MCC = \frac{TP \times TN - FP \times FN}{\sqrt{(TP + FP)(TP + FN)(TN + FP)(TN + FN)}} \quad (S6)$$

MCC values range from  $-1$  to  $+1$ . A coefficient of  $+1$  indicates a perfect prediction,  $0$  indicates no better than a random prediction, and  $-1$  indicates total disagreement between prediction and observation.

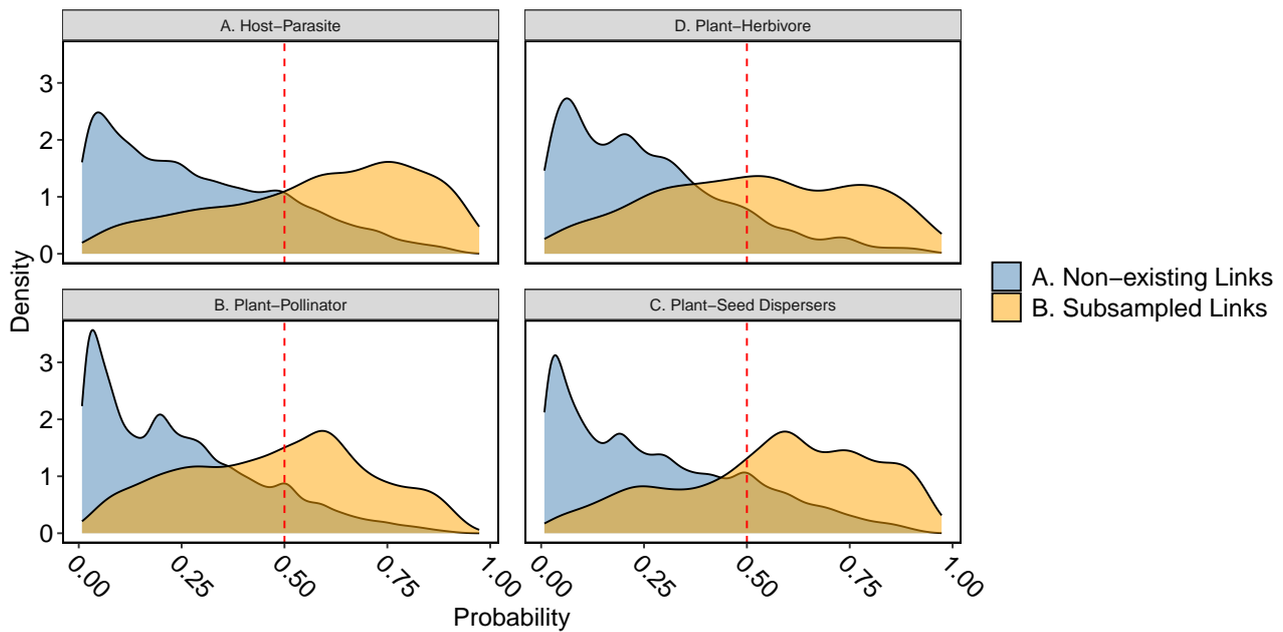
Given ecological networks' imbalanced and noisy nature, we explored how the classification threshold affects the PR tradeoff. As a first step, we plotted the link probabilities generated by the model. The non-existing links were generally correctly classified ( $\approx 0.83\%$  out of 183K non-links; Fig. S18A). This is expected in imbalanced data sets. However, for the withheld links, only  $\approx 0.63\%$  out of 9K were classified correctly (Fig. S18B). This mismatch underlies the tradeoff between precision and recall. Further exploration of the precision-recall tradeoff (Fig. S19) indicated that there is no apparent threshold to choose from, and so we decided to use the common value of  $0.5$ . In a more detailed examination of each community separately, as shown in Fig. S20, we observed a pronounced right-skewed distribution of the sub-sampled links. Specifically, there was a more apparent separation between the two link types in both host-parasite and plant-seed disperser networks.



**Fig. S18: Distribution of link probabilities obtained from the model.** The histograms depict the distribution of the predicted probabilities for our binary classification task, representing the two classes in the test set: non-existing links (A) and sub-sampled links (B). The x-axis represents the predicted probabilities ranging from 0 to 1, while the y-axis represents the frequency of the observations. The dashed red line represents the decision threshold of 0.5.



**Fig. S19: The precision-recall tradeoff as a function of classification threshold.** The tradeoff is presented when testing on all communities (A) or per community type (B-D). Each data point on the curve corresponds to a distinct cutoff threshold value (x-axis).



**Fig. S20: Distribution of link probabilities across different ecological communities.** The kernel density estimate (KDE) curves of the three communities: (A) Host-parasite networks, (B) Plant-pollinator networks, and (C) plant-speed disperser networks. The distributions for non-existing links and sub-sampled links are depicted in blue and orange, respectively. The x-axis represents the predicted probabilities ranging from 0 to 1, while the y-axis denotes the density estimation of the observations. A dashed red line marks the decision threshold of 0.5. The overlap between the blue and orange distributions represents areas of prediction ambiguity. In regions where non-existing links (blue) surpass the decision threshold, false positives emerge, negatively influencing precision. Conversely, in the region where sub-sampled links (orange) fall below the threshold, it results in false negatives, negatively influencing recall.

### S1.3 Exploration of link withholding

Link removal strategies could affect evaluation outcomes. Due to the binary nature of our data we tested three link removal scenarios: (1) *Uniform removal*: This scenario assumes equal probability of removal for all interactions, as implemented in the initial submission. (2) *High-degree biased sampling*: This scenario preferentially retains interactions involving species with many partners. It reflects ecological survey biases where interactions of abundant or generalist species are more likely to be observed due to greater detectability or sampling effort. (3) *Low-degree biased sampling*: This scenario preferentially retains interactions involving species with fewer partners, simulating sampling efforts that might focus on rare or specialized interactions. The high-degree biased sampling has lower performance than the other two scenarios because it has a lower recall (Fig. S21). Therefore, when missing links are those of generalist species, the model will have lower prediction abilities. This result is logical because those interactions contain most of the information in the network. However, such scenario is unlikely because typically generalist species are those that are more easily observed. Therefore, our model does perform well under the typical low-degree biased sampling. These additional analyses provide a more comprehensive assessment of our model’s performance.

A detailed description of these scenarios is as follows:

**Uniform sampling:** Our baseline approach implements random uniform sampling, where each link has an equal probability of being included in the sampled network. This strategy assumes that missing interactions are randomly distributed throughout the network, which may occur when sampling effort is uniformly distributed across species pairs but insufficient to detect all interactions.

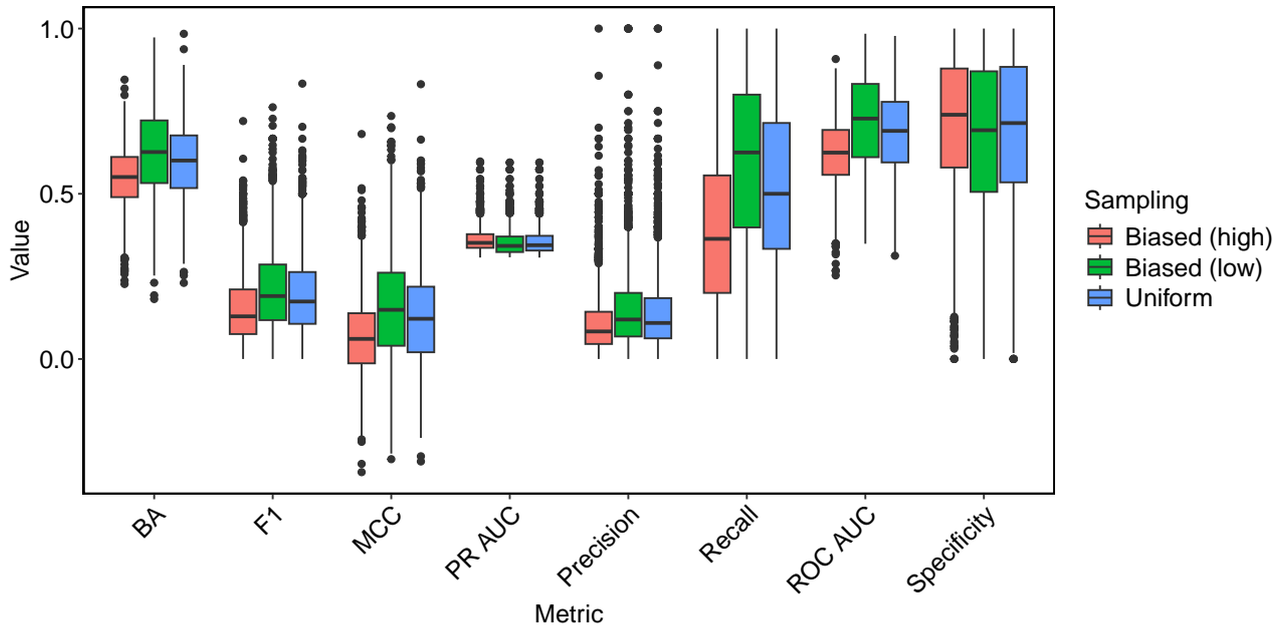
**Low-degree bias:** The low-degree biased sampling preferentially retains interactions involving species with fewer partners. For a bipartite network with species sets  $A$  and  $B$ , the probability of selecting species  $i \in A$  and  $j \in B$  is:  $P(i) \propto \frac{1}{k_i+1}$  for  $i \in A$ , and  $P(j) \propto \frac{1}{k_j+1}$  for  $j \in B$ , where  $k_i$  and  $k_j$  are the degrees of species  $i$  and  $j$  respectively. The probability of sampling an interaction between species  $i$  and  $j$  is then:  $P(i, j) \propto P(i) \times P(j)$ . This strategy simulates scenarios where rare or specialized interactions are more likely to be observed, which might occur when sampling efforts focus on less common species or when common interactions are overlooked.

**High-degree bias:** Conversely, the high-degree biased sampling preferentially retains interactions involving species with many partners. The sampling probabilities are proportional to the degrees:  $P(i) \propto k_i$  for  $i \in A$ , and  $P(j) \propto k_j$  for  $j \in B$ , with the interaction probability again being:  $P(i, j) \propto P(i) \times P(j)$ .

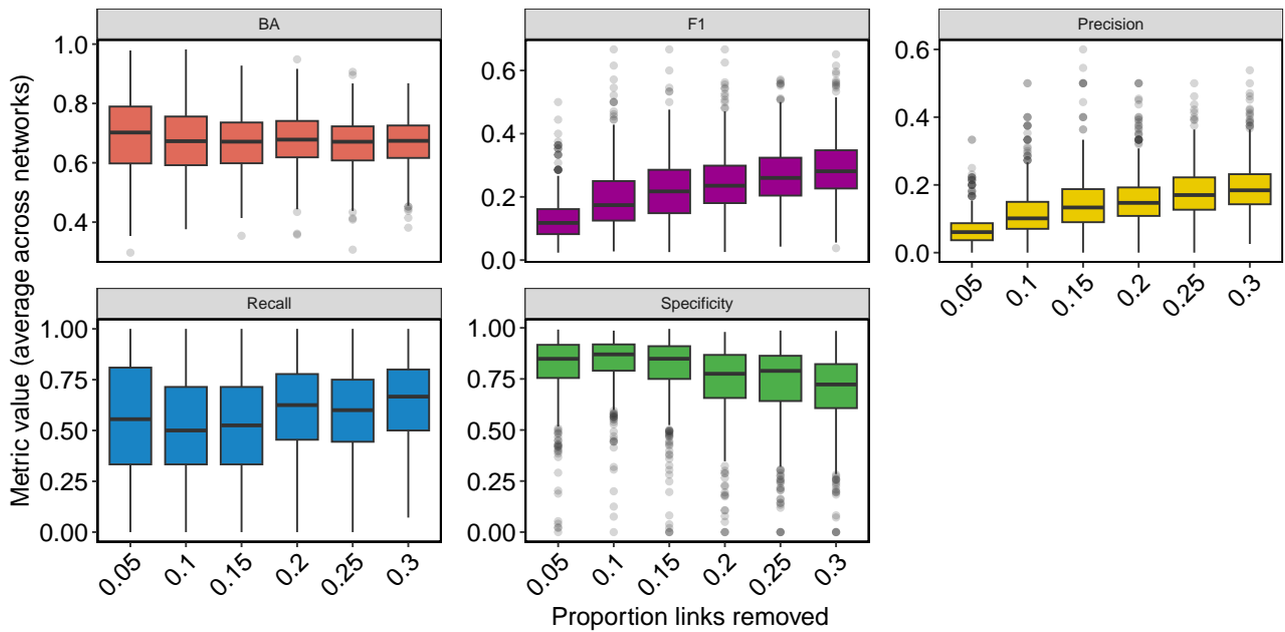
This approach simulates scenarios where common or generalist species and their interactions are more likely to be observed, which often occurs in ecological surveys due to the greater detectability of abundant species or sampling bias toward easily observable interactions.

**Implementation** For each sampling strategy, we: (1) Calculate the target number of interactions to retain based on the desired sampling fraction  $f$ . (2) For degree-biased sampling: Calculate degrees for species in each level of the network, compute normalized sampling probabilities for each species based on their degrees and then iteratively sample species pairs using their respective probabilities until reaching the target number of interactions. (3) For uniform sampling we directly sample the target number of interactions with uniform probability.

**Withholding proportion** For the uniform strategy, which we present in the main text, we also conducted sensitivity analysis, removing 5-30% of the links (Fig. S22). In the main text we present results for 20%, which balances the imbalance in non-links and links with choosing a higher proportion that would artificially improve the results.



**Fig. S21: Comparison of sampling strategies.** *Uniform removal:* This scenario assumes equal probability of removal for all interactions. (2) *High-degree biased sampling:* This scenario preferentially retains interactions involving species with many partners. It reflects ecological survey biases where interactions of abundant or generalist species are more likely to be observed due to greater detectability or sampling effort. (3) *Low-degree biased sampling:* This scenario preferentially retains interactions involving species with fewer partners, simulating sampling efforts that might focus on rare or specialized interactions.



**Fig. S22: Link removal sensitivity analysis.** Balanced accuracy does not change with withheld proportion (only its variation). F1 generally increases with withheld proportion because increasing the proportion of sub-sampled links reduces the imbalance in the train and test sets, improving precision. Therefore, the choice of 20% balances imbalance with choosing a higher proportion that would artificially improve the results.

#### S1.4 Cost-sensitive learning

Cost-sensitive learning uses a parameter that the algorithm considers during the learning process. The penalty is the cost, which is minimized during the training process. This approach allows dealing with imbalance while ignoring less samples<sup>13</sup>. The scikit-learn package<sup>14</sup> has a built-in support for cost-sensitive learning. This implementation provides custom weights to a model's classes or samples during training. Depending on the learning algorithm, the loss function is modified to penalize mistakes according to the provided weights, such that higher weights lead to higher penalizing. Because in our study, the minority class is the existing links, these are assigned higher importance by using higher weights. Specifically, each class automatically gets a weight of 1 but can be set with a higher weight. For instance, a class with a weight 2 will have double importance. We aimed for 'balanced' weights, which are proportional to the classes' frequency. For instance, if the training set contains 20 more times non-existing than existing links, then existing links will get 20 times more importance.

#### Appendix References

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