- ¹ Title: Inferring species interactions from co-occurrence data with Markov networks
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- 3 Abstract: Inferring species interactions from co-occurrence data is one of the most
- 4 controversial tasks in community ecology. One difficulty is that a single pairwise interaction
- 5 can ripple through an ecological network and produce surprising indirect consequences. For
- 6 example, the negative correlation between two competing species can be reversed in the
- 7 presence of a third species that outcompetes both of them. Here, I apply models from
- 8 statistical physics, called Markov networks or Markov random fields, that can predict the
- 9 direct and indirect consequences of any possible species interaction matrix. Interactions in
- these models can be estimated from observed co-occurrence rates via maximum likelihood,
- controlling for indirect effects. Using simulated landscapes with known interactions, I
- evaluated Markov networks and six existing approaches. Markov networks consistently
- outperformed the other methods, correctly isolating direct interactions between species pairs
- even when indirect interactions or abiotic factors largely overpowered them. Two
- 15 computationally efficient approximations, which controlled for indirect effects with partial
- correlations or generalized linear models, also performed well. Null models showed no
- evidence of being able to control for indirect effects, and reliably yielded incorrect inferences
- when such effects were present.
- 19 Key words: Ecological interactions; Occurrence data; Species associations; Markov network;
- Markov random field; Ising model; Biogeography; Presence-absence matrix; Null model

21 Introduction

- To the extent that nontrophic species interactions (such as competition) affect community
- 23 assembly, ecologists might expect to find signatures of these interactions in species

composition data (MacArthur 1958, Diamond 1975). Despite decades of work and several major controversies, however (Lewin 1983, Strong et al. 1984, Connor et al. 2013), existing 25 methods for detecting competition's effects on community structure are unreliable (Gotelli and Ulrich 2009). In particular, species' effects on one another can become lost in a web of indirect effects. For example, the competitive interaction between the two shrub species in Figure 1A is obscured by their shared tendency to occur in unshaded areas (Figure 1B). While ecologists have long known that indirect effects can overwhelm direct ones (Levine 1976), most methods for drawing inferences from co-occurrence data do not control for these 31 effects (e.g. Diamond 1975, Strong et al. 1984, Gotelli and Ulrich 2009, Veech 2013, Pollock et al. 2014). As a result, ecologists do not have tools that allow them to isolate direct interactions from indirect effects. While competition doesn't reliably reduce co-occurrence rates at the whole-landscape 35 level (as most methods assume), it does still leave a signal in the data (Figure 1C). After controlling for the presence of the tree species (e.g. by splitting the data set into shaded and unshaded sites or by using a model that estimates conditional relationships amongs species), the two shrubs do have a negative association, with fewer co-occurrences in unshaded areas than would be expected if they occurred independently of one another. Following Azaele et al. (2010), this paper shows that Markov networks (undirected 41 graphical models also known as Markov random fields; Murphy 2012) can provide a framework for understanding the landscape-level consequences of pairwise species interactions, and for estimating them from observed presence-absence matrices. Markov networks have been used in diverse scientific fields for decades, from physics (where nearby particles interact magnetically; Cipra 1987) to spatial statistics (where adjacent grid cells have correlated values; Harris 1974, Gelfand et al. 2005). While community ecologists

explored some related approaches in the 1980's (Whittam and Siegel-Causey 1981), they
used severe approximations that led to unintelligible results (e.g. "probabilities" greater than
one; Gilpin and Diamond 1982).

Below, I demonstrate Markov networks' ability to produce exact predictions about the
direct and indirect consequences of an interaction matrix, and also to make inferences about
the interaction matrix based on co-occurrence rates. Using simulated data sets where the
"true" interactions are known, I compare this approach with several existing methods.

Finally, I discuss opportunities for extending the approach presented here to other problems
in community ecology, e.g. quantifying the overall effect of species interactions on occurrence
rates (Roughgarden 1983) and disentangling the effects of biotic versus abiotic interactions
on species composition (Harris 2016).

59 Methods

Markov networks. Markov networks provide a framework for translating back and forth between the conditional (all-else-equal) relationships among species (Figure 1C) and the kinds of species assemblages that these relationships produce. Here, I show how a set of conditional relationships can determine species composition. Methods for estimating conditional relationships from data are discussed in the next section.

A Markov network defines the relative probability of observing a given vector of species-level presences (1s) and absences (0s), \vec{y} at a site, as

$$p(\vec{y}; \alpha, \beta) \propto exp(\sum_{i} \alpha_{i} y_{i} + \sum_{\langle ij \rangle} \beta_{ij} y_{i} y_{j}),$$

where the second sum is over all $\frac{1}{2}n(n-1)$ pairs of n species. In this model, α_i is an intercept term determining the amount that the presence of species i contributes to the log-probability of \vec{y} ; it directly controls the prevalence of species i. Similarly, β_{ij} is the amount that the

co-occurrence of species i and species j contributes to the log-probability; it determines the conditional relationship between two species, i.e. the probability that they will be found 72 together, after controlling for the other species in the network (Figure 2A, Figure 2B). For 73 example, if $\beta_{ij} = +2$, then each species' odds of occurrence would be e^2 times higher when the other one is present (as compared with otherwise equivalent sites). The relative probability of a presence-absence vector increases when positively-associated species co-occur and decreases when negatively-associated species co-occur. As a result, the model tends—all 77 else equal—to produce assemblages where many positively-associated species pairs co-occur 78 and few negatively-associated pairs do (just as an ecologist might expect). Estimating α and β coefficients from presence-absence data. In the previous section, the values of α and β were known and the goal was to make predictions about 81 possible species assemblages. In practice, however, ecologists will often need to estimate the 82 parameters from an observed co-occurrence matrix (i.e. from a set of independent \vec{y} vectors indicating which species are present at each site on the landscape). When Equation 1 can be normalized (Figure 2B), one can find exact maximum likelihood estimates for α and β by numerically optimizing $p(\vec{y}|\alpha,\beta)$. Fully-observed networks like the ones considered here have unimodal likelihood surfaces (Murphy 2012), so optimizers will always find the global 87 optimum. When the number of species is larger than about 30, noralizing Equation 1 can become 89 intractable, and researchers will either need to approximate it (Lee and Hastie 2012) or 90 approximate its gradient (Harris 2016) if they want to fit a Markov network model. For the 91 analyses presented here, where the number of species did not exceed 20, approximations were not necessary. Instead, I used the rosalia package (Harris 2015a) for the R programming language (R Core Team 2015) to calculate $p(\vec{y}; \alpha, \beta)$ and its gradients exactly (Murphy

- ⁹⁵ 2012); the package passes these functions to the "BFGS" method in R's general-purpose optimizer, which finds values for α and β .
- Simulated landscapes. I simulated several sets of landscapes using known parameters so
 that model estimates could be compared with "true" values. The first set of landscapes
 included the three competing species shown in Figure 1. For each of 1000 replicates, I
 generated a landscape with 100 sites by sampling from a probability distribution defined by
 the figure's interaction coefficients (Appendix 1). Each of the methods described below was
 then evaluated on its ability to correctly infer that the two shrub species competed with one
 another, despite their frequent co-occurrence in unshaded areas.

I then simulated landscapes with up to 20 interacting species at 25, 200, or 1600 sites using three increasingly complex models (50 replicates for each combination of size and model; Appendix 2). The simplest set of simulated landscapes were generated with Gibbs samples from Equation 1. For each replicate, I randomly drew the "true" β coefficient magnitudes from an exponential distribution with rate 1 so that most species pairs interacted negligibly but a few interactions were strong enough to propagate through the newtwork. I randomly assigned 25% of the interactions to be positive; the remainder were negative.

The next set of landscapes provided a way to assess each method's ability to identify direct interactions in the presence of environmental heterogeneity. Here, the β coefficients were calculated as above, but each species' α value depended linearly on two environmental factors, which were drawn from independent Gaussians for each site. Once the local α values were calculated, independent Gibbs samplers determined the species composition for each site based on Equation 1.

In the final set of landscapes, I simulated each species' abundance (instead of just presence/absence); furthermore, interactions between species occurred on a per-capita basis

in these simulations (i.e. each species' effect on the others is proportional to its abundance).

To prevent runaway mutualisms leading to infinite abundance, all interaction coefficients

were negative in these simulations. Good performance on these landscapes would indicate

some robustness to the mechanistic details of species interactions.

Recovering species interactions from simulated data. I compared seven techniques for determining the sign and strength of the associations between pairs of species from simulated data (Appendix 3). First, I used the rosalia package (Harris 2015a) to fit Markov network models, as described above. For the analyses with 20 species, a weakly-informative regularizer (equivalent to a logistic prior with location 0 and scale 2) ensured that the estimates were always finite (Appendix 3).

I also evaluated six alternative methods: five from the existing literature, plus a novel 129 combination of two of these methods. The first alternative interaction metric was the sample 130 correlation between species' presence-absence vectors, which summarizes their marginal 131 association. Next, I used partial correlations, which summarize species' conditional 132 relationships. This approach, which is closely related to linear regression, is common in 133 molecular biology (Friedman et al. 2008), but is rare in ecology (see Albrecht and Gotelli 134 (2001) and Faisal et al. (2010) for two exceptions). In the context of non-Gaussian data, the 135 partial correlation (or partial covariance) can be thought of as a computationally efficient 136 approximation to the full Markov network model (Loh and Wainwright 2013). Partial 137 correlations are undefined for landscapes with perfectly-correlated species pairs, so I used the 138 regularized estimate provided by the corpcor package's pcor.shrink function with the 139 default settings (Schäfer et al. 2014). 140

The third alternative, generalized linear models (GLMs), also provide a computationally efficient approximation to the Markov network (Lee and Hastie 2012). Following Faisal et al.

(2010), I fit regularized logistic regression models (Gelman et al. 2008) for each species, using the other species as predictors. This produced two interaction estimates for each species pair (one for the effect of species i on species j and one for the reverse). These two estimates were very tightly correlated (mean Pearson correlation of 0.95, Appendix 3); their arithmetic mean provided a consensus estimate of the overall interaction.

The next method used the Pairs software described in Gotelli and Ulrich (2009). This
program simulates new landscapes from a null model that retains the row and column sums
of the original matrix (Strong et al. 1984) and calculates Z-scores to summarize a species
pair's deviation from this null.

The last two estimators used the latent correlation matrix estimated by the BayesComm package (Golding and Harris 2015) in order to evaluate the recent claim that the correlation coefficients estimated by "joint species distribution models" provide an accurate assessment of species' pairwise interactions (Pollock et al. 2014, see also Harris 2015b). In addition to using the posterior mean correlation (Pollock et al. 2014), I also used the posterior mean partial correlation, which should control better for indirect effects.

Evaluating model performance. For the simulated landscapes based on Figure 1, I
assessed whether each method's test statistic indicated a positive or negative relationship
between the two shrubs (Appendix 1). For the null model (Pairs), I calculated statistical
significance using its Z-score. For the Markov network, I used the Hessian matrix to
generate approximate confidence intervals.

For the larger landscapes, I evaluated the relationship between each method's estimates and the "true" interaction strengths. To ensure that the different test statistics (e.g. correlations versus Z scores) were on a common scale, I rescaled them using linear regression through the origin. I then calculated the proportion of variance explained for

different combinations of model type and landscape size (compared with a baseline model that assumed all interaction strengths to be zero).

For the null model and the Markov network, the probability of rejecting the null hypothesis of zero interaction was estimated across a range of "true" interaction strengths using a kernel smoother (Appendix 4). The probability of rejection when the "true" value of β was zero was defined as the Type I error rate. Because the coefficients' interpretation is different for the abundance simulations than for the other two types, its error rates were not analyzed in this way.

175 Results

Three species. As shown in Figure 1, the marginal relationship between the two shrub 176 species was positive—despite their competition for space at a mechanistic level—due to indirect effects of the dominant tree species. As a result, the correlation between these 178 species was positive in 94% of replicates, and the randomization-based null model falsely reported positive associations 100% of the time. Worse, more than 98% of these false 180 conclusions were statistically significant. The partial correlation and Markov network 181 estimates, on the other hand, each correctly isolated the direct negative interaction between the shrubs from their positive indirect interaction 94% of the time (although the confidence 183 intervals overlapped zero in most replicates). Twenty species. In general, each model's performance was highest for large landscapes 185 with simple assembly rules and no environmental heterogeneity (Figure 3). Despite some 186 variability across contexts, the rank ordering across methods was very consistent. In 187 particular, the four methods that controlled for indirect effects (the Markov network, the 188 generalized linear models, and the two partial correlation-based methods) always matched or 189 outperformed those that did not. The Markov network consistently performed best of all. As

anticipated by Lee and Hastie (2012), generalized linear models closely approximated the 191 Markov network estimates (Figure 4A), especially when the data sets were very large (Figure 192 3). As reviewed in Gotelli and Ulrich (2009), however, most analyses in this field of ecology 193 involve fewer than 50 sites; in this context, the gap between the methods was larger. As 194 shown in Appendix 4, the standard errors associated with the estimates in Figure 3 are small 195 (less than 0.01), so the differences among methods should not be attributed to sampling error. Of the methods that did not control for indirect effects, Figure 3 shows that simple 197 correlation coefficients provided a more reliable indicator of species' true interaction strengths 198 than either the joint species distribution model (BayesComm) or the null model (Pairs). 95% 199 of the variance the Pairs test statistic was explained by correlation coefficients (controlling for landscape size; Figure 4B); much of the remaining variance is due to sampling error. 201 Finally, we can evaluate the models' inferential statistics (focusing on the first two 202 simulation types, where the interaction coefficients are easiest to interpret). The Markov 203 network's Type I error rate was 0.02 for simulations that matched the model's assumptions, and 0.14 for simulations that included environmental heterogeneity (see Appendix 4 for 205 confidence interval coverage across a range of β_{ij} values). In contrast, the null model's Type 206 I error rates were 0.30 and 0.51, respectively—far higher than the nominal 0.05 rate. Figure 207 4C shows, across a range of true interaction strengths, the probability that the null model or 208 the Markov network will predict the wrong sign of the interaction with 95% confidence. The 209 null model makes such errors more than 8 times as often as the Markov network, even 210 though it only rejects the null hypothesis twice as often overall (Appendix 4). The Markov 211 network's errors were also more concentrated around 0, as it never misclassified strong 212 interactions like the null model did (Figure 4C).

Discussion

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The results presented above show that Markov networks can reliably recover species' pairwise interactions from species composition data, even for cases where environmental 216 heterogeneity and indirect interactions cause ecologists' typical null modeling approaches to 217 reliably fail. Partial correlations and generalized linear models can both provide 218 computationally efficient approximations, but with somewhat lower accuracy (especially for 219 typically-sized data sets with small numbers of sites; Gotelli and Ulrich 2009). The difference in accuracy may be even larger for real data sets than for the simulated landscapes in Figure 221 3; linear approximations to the Markov network make larger errors when the interaction 222 matrix is structured (e.g. due to guilds or trophic levels; Loh and Wainwright 2013). Similarly, the separate generalized linear models for each species can severely overfit in some 224 cases (Lee and Hastie 2012). The full Markov network should thus be preferred to the approximations when it is computationally tractable. 226

Compositional data only contains enough degrees of freedom to estimate one interaction
per species pair (Schmidt and Murphy 2012), so none of these methods can identify the
exact nature of the pairwise interactions (e.g. which species in a positively-associated pair is
facilitating the other). To estimate asymmetric interactions, such as commensalism or
predation, ecologists could use time series, behavioral observations, manipulative
experiments, or natural history. These other sources of information could also be used to
augment the likelihood function with a more informative prior distribution, reducing
ecologists' error and uncertainty relative to Figure 3's results.

Markov networks have enormous potential to improve our understanding of species interactions. In particular, they make many fewer errors than existing approaches, and can make precise statements about the conditions where indirect interactions will overwhelm direct ones. They also provide a simple answer to the question of how competition should

affect a species' overall prevalence, which has important implications for community-level modeling (Strong et al. 1984). Specifically, Equation 1 can be used to calculate the expected 240 prevalence of a species in the absence of biotic influences as $e^{\alpha_i}/(e^0 + e^{\alpha_i})$. Competition's effect on prevalence can then be estimated by comparing this value with the observed 242 prevalence (e.g. comparing Figure 2D with Figure 2C). This novel quantitative result 243 undermines most of our null models, which unreasonably assume that prevalence would be the exactly same in the absence of competition as it is in the observed data (Roughgarden 245 1983). 246 Markov networks—particularly the Ising model for binary networks—are very well 247 understood, having been studied for nearly a century (Cipra 1987). Tapping into this framework would thus allow ecologists to take advantage of into a vast set of existing 249 discoveries and techniques for dealing with indirect effects, stability, and alternative stable 250 states. Numerous extensions to the basic network are possible as well. For example, the 251 states of the interaction network can be modeled as a function of the local abiotic 252 environment (Lee and Hastie 2012, Harris 2016), which would lead to a better understanding of the interplay between biotic and abiotic effects on community structure. Alternatively, 254 models could allow one species to alter the relationship between two other species (Whittam 255 and Siegel-Causey 1981, Tjelmeland and Besag 1998, cf Bruno et al. 2003). Finally, the results presented here have important implications for ecologists' continued 257 use of null models for studying species interactions. When the non-null backdrop is not 258 controlled for, Type I error rates can skyrocket, the apparent sign of the interaction can 259 change, and null models can routinely produce misleading inferences (Figure 1, Figure 4C, 260 Gotelli and Ulrich (2009)). Null and neutral models can be useful for clarifying our thinking 261 (Harris et al. 2011, Xiao et al. 2015), but deviations from a given null model must be

interpreted with care (Roughgarden 1983). Even in small networks with three species, it may simply not be possible to implicate specific ecological processes like competition by rejecting a general-purpose null (Gotelli and Ulrich 2009), especially when the test statistic is effectively just a correlation coefficient (Figure 4B).

Controlling for indirect effects via simultaneous estimation of multiple ecological
parameters seems like a much more promising approach: to the extent that the models'
relative performance on real data sets is similar to the range of results shown in Figure 3,
scientists in this field could often triple their explanatory power by switching from null
models to Markov networks (or increase it nearly as much with linear or generalized linear
approximations). Regardless of the methods ecologists ultimately choose, controlling for
indirect effects could clearly improve our understanding of species' direct effects on one
another and on community structure.

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282 References:

Figure 1: A. A small network of three competing species. The tree (top) tends not to

co-occur with either of the two shrub species, as indicated by the strongly negative

coefficient linking them. The two shrub species also compete with one another, but more

weakly (circled coefficient). B. In spite of the competitive interactions between the two

shrub species, their shared tendency to occur in locations without trees makes their 287 occurrence vectors positively correlated (circled). C. Controlling for trees with a 288 conditional (all-else-equal) approach such as a partial correlation or a Markov network 289 leads to correct identification of the negative shrub-shrub interaction (circled). See 290 Appendix 1 and the results for "three species" for more details. 291 **Figure 2:** A. A small Markov network, defined by its α and β values. The abiotic environment favors the occurrence of each species ($\alpha > 0$), particularly species 2 ($\alpha_2 > \alpha_1$). 293 The negative β_{12} coefficient is consistent with competition between the two species. **B.** 294 The coefficients determine the probabilities of all four possible presence-absence 295 combinations for Species 1 and Species 2. α_1 is added to the exponent whenever Species 1 296 is present $(y_1 = 1)$, but not when it is absent $(y_1 = 0)$. Similarly, the exponent includes α_2 297 only when species 2 is present $(y_2 = 1)$, and includes β_{12} only when both are present 298 $(y_1y_2=1)$. The normalizing constant Z, ensures that the four probabilities sum to 1. In 299 this case, Z is about 18.5. C. The expected frequencies of all possible co-occurrence 300 patterns between the two species of interest, as calculated in the previous panel. **D.** 301 Without competition (i.e. with $\beta_{12} = 0$, each species would occur more often. 302 Figure 3: Proportion of variance in interaction coefficients explained by each method versus 303 number of sampled locations across the three simulation types. For the null model (Pairs), 304 two outliers with |Z| > 1000 were manually adjusted to |Z| = 50 to mitigate their 305 detrimental influence on \mathbb{R}^2 (Appendix 5). 306 **Figure 4: A.** The Markov network's estimated interaction coefficients were generally very 307 similar to the GLM estimates. B. The null model's estimates typically matched the 308 (negative) correlation coefficient, after controlling for landscape size. C. For any given 309 interaction strength, the null model was much more likely to misclassify its sign with 95% 310

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