- 1 Title: Estimating species interactions from observational data with Markov networks
- 2 Author: David J. Harris: Population Biology; 1 Shields Avenue, Davis CA, 95616
- 3 Abstract: Estimating species interactions from observational 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, two competing species would ordinarily correlate negatively in space, but this effect
- 7 can be reversed in the presence of a third species that is capable of outcompeting both of
- 8 them when it is present. Here, I apply models from statistical physics, called Markov
- 9 networks or Markov random fields, that can predict the direct and indirect consequences of
- any possible species interaction matrix. Interactions in these models can be estimated from
- observational data via maximum likelihood. Using simulated landscapes with known
- pairwise interaction strengths, I evaluated Markov networks and several existing approaches.
- 13 The Markov networks consistently outperformed other methods, correctly isolating direct
- 14 interactions between species pairs even when indirect interactions or abiotic environmental
- 15 effects largely overpowered them. Two computationally efficient approximations, based on
- linear and generalized linear models, also performed well. Indirect effects reliably caused a
- common null modeling approach to produce incorrect inferences, however.
- 18 Key words: Ecological interactions; Occurrence data; Species associations; Markov network;
- Markov random field; Ising model; Biogeography; Presence-absence matrix; Null model

20 Introduction

- 21 If nontrophic species interactions, such as competition, are important drivers of community
- assembly, then ecologists might expect to see their influence in species composition data
- ²³ (MacArthur 1958, Diamond 1975). Despite decades of work and several major controversies,

however (Lewin 1983, Strong et al. 1984, Gotelli and Entsminger 2003, Connor et al. 2013), existing methods for detecting competition's effects on community structure are unreliable 25 (Gotelli and Ulrich 2009). In particular, species' effects on one another can become lost in the complex web of direct and indirect interactions in real assemblages. For example, the competitive interaction between the two shrub species in Figure 1A can become obscured by their shared tendency to occur in unshaded areas (Figure 1B). While ecologists have long known that indirect effects can overwhelm direct ones at the landscape level (Dodson 1970, Levine 1976), the vast majority of our methods for drawing inference from observational data 31 do not control for these effects (e.g. Diamond 1975, Strong et al. 1984, Gotelli and Ulrich 2009, Veech 2013, Pollock et al. 2014). To the extent that indirect interactions like those in Figure 1 are generally important (Dodson 1970), existing methods will thus not generally provide much evidence regarding species' direct effects on one another. The goal of this 35 paper is to resolve this long-standing problem. While competition doesn't reliably leave the expected pattern in community structure (low 37 co-occurrence rates at the landscape level), it nevertheless does leave a signal in the data (Figure 1C). Specifically, among shaded sites, there will be a deficit of co-occurrences between the competing shrub species, and among unshaded sites, there will also be such a deficit. More generally, we can obtain much better estimates of the association between two species from their conditional relationships (i.e. by controlling for other species in the network) than we could get from their overall co-occurrence rates. This kind of precision is difficult to obtain from null models, which begin with the assumptions that all the pairwise interactions are zero and thus don't need to be controlled for. Nevertheless, null models have dominated this field for more than three decades (Strong et al. 1984, Gotelli and Ulrich 2009).

Following recent work by Azaele et al. (2010) and Fort (2013), this paper shows that Markov networks (undirected 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 detecting them from observed presence-absence matrices. Markov networks have been used in many scientific fields for decades in similar 51 contexts, 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 introduce Markov networks and show how they can be used to simulate landscape-level data or to make exact predictions about the direct and indirect consequences of possible interaction matrices. Then, 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

65 Methods

Markov networks. Markov networks provide a framework for translating back and forth
between the conditional 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 be used to determine the how groups of species can co-occur. Methods for

on species composition (Kissling et al. 2012, Pollock et al. 2014).

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} , as

 $p(\vec{y}; \alpha, \beta) \propto exp(\sum_i \alpha_i y_i + \sum_{i \neq j} \beta_{ij} y_i y_j).$

Here, α_i is an intercept term determining the amount that the presence of species i 74 contributes to the log-probability of \vec{y} ; it directly controls the prevalence of species i. 75 Similarly, β_{ij} is the amount that the co-occurrence of species i and species j contributes to the log-probability; it controls the conditional relationship between two species, i.e. the 77 probability that they will be found together, after controlling for the other species in the network (Figure 2A, Figure 2B). For example, β_{ij} might have a value of +2 for two mutualists, indicating that the odds of observing one species are e^2 times higher in sites 80 where its partner is present than in comparable sites where its partner is absent. Because the relative probability of a presence-absence vector increases when positively-associated species 82 co-occur and decreases when negatively-associated species co-occur, the model tends—all else equal—to produce assemblages that have many pairs of positively-associated species and relatively few pairs of negatively-associated species (exactly as an ecologist might expect). 85 Of course, if all else is *not* equal (e.g. Figure 1, where the presence of one competitor is 86 associated with release from another competitor), then species' marginal association rates can 87 differ from this expectation. Determining the marginal relationships between species from their conditional interactions entails summing over the different possible assemblages (Figure 2B). This becomes intractable when the number of possible assemblages is large, though several methods beyond the scope of this paper can be employed to keep the calculations feasible (Salakhutdinov 2008, Lee and Hastie 2012). Alternatively, as noted below, some

common linear and generalized linear methods can also be used as computationally efficient approximations to the full network (Lee and Hastie 2012, Loh and Wainwright 2013). 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 possible species assemblages. In practice, however, ecologists will often need to estimate the 97 parameters from an observed co-occurrence matrix (i.e. from a matrix of ones and zeros indicating which species are present at which sites). When the number of species is reasonably small, one can compute exact maximum likelihood estimates for all of the α and β coefficients given a presence-absence matrix by optimizing $p(\vec{y}; \alpha, \beta)$. Fully-observed Markov 101 networks like the ones considered here have unimodal likelihood surfaces (Murphy 2012), 102 ensuring that this procedure will always converge on the global maximum. This maximum represents the unique combination of α and β coefficients that would be expected to produce 104 exactly the observed co-occurrence frequencies on average (i.e. it matches the sufficient 105 statistics of the model distribution to the sufficient statistics of the data; Murphy 2012). I 106 used the rosalia package (Harris 2015a) for the R programming language (R Core Team 107 2015) to optimize the Markov network parameters. The package was named after Santa Rosalia, the patron saint of biodiversity, whose supposedly miraculous healing powers played 100 an important rhetorical role in the null model debates of the 1970's and 1980's (Lewin 1983). 110 Simulated landscapes. In order to compare different methods, I simulated two sets of 111 landscapes using known parameters. The first set included the three competing species shown 112 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). 114 Each of the methods described below was then evaluated on its ability to correctly infer that 115

the two shrub species competed with one another, despite their frequent co-occurrence.

I also simulated a second set of landscapes using a stochastic community model based on generalized Lotka-Volterra dynamics, as described in Appendix 2. In these simulations, each species pair was randomly assigned to either compete for a portion of the available carrying capacity (negative interaction) or to act as mutualists (positive interaction). Here, mutualisms operate by mitigating the effects of intraspecific competition on each partner's death rate. For these analyses, I simulated landscapes with up to 20 species and 25, 200, or 1600 sites (50 replicates per landscape size; see Appendix 2).

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 newtork models, as described 127 above. For the analyses with 20 species, I added a very weak logistic prior distribution on the α and β terms with scale 2 to ensure that the model estimates were always finite. The 129 bias introduced by this prior should be small: the 95% credible interval on β only requires that one species' effect on the odds of observing a different species to be less than a factor of 131 1500 (which is not much of a constraint). The logistic distribution was chosen because it is 132 convex and has a similar shape to the Laplace distribution used in LASSO regularization (especially in the tails), but unlike the Laplace distribution it is differentiable everywhere and 134 does not force any estimates to be exactly zero. To confirm that this procedure produced 135 stable estimates, I compared its estimates on 50 bootstrap replicates (Appendix 4). 136

I also evaluated five alternative methods from the existing literature (plus a novel variant of the fifth method). The first two alternative interaction measures were the sample

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correlations and the partial correlations between each pair of species' data vectors on the
    landscape (Albrecht and Gotelli 2001, Faisal et al. 2010). In the context of non-Gaussian
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    data, the partial correlation can be thought of as a computationally efficient approximation
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   to the full Markov network model (Loh and Wainwright 2013). This sort of model is very
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    common for estimating relationships among genes and gene products (Friedman et al. 2008).
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   Because partial correlations are undefined for landscapes with perfectly-correlated species
    pairs, I used a regularized estimate based on James-Stein shrinkage, as implemented in the
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    corpcor package's pcor.shrink function with the default settings (Schäfer et al. 2014).
      The third alternative, generalized linear models (GLMs), can also be thought of as a
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   computationally efficient approximation to the Markov network (Lee and Hastie 2012).
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   Following Faisal et al. (2010), I fit regularized logistic regression models (Gelman et al. 2008)
    for each species, using the other species on the landscape as predictors. To avoid the
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   identifiability problems associated with directed cyclic graphs (Schmidt and Murphy 2012), I
    then symmetrized the relationships within species pairs via averaging.
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      The fourth alternative method, described in Gotelli and Ulrich (2009), involved simulating
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   new landscapes from a null model that retains the row and column sums of the original
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    matrix (Strong et al. 1984). I used the Z-scores computed by the Pairs software described in
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    Gotelli and Ulrich (2009) as my null model-based estimator of species interactions.
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      The last two estimators used the latent correlation matrix estimated by the BayesComm
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   package (Golding and Harris 2015) in order to evaluate the recent claim that the correlation
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   coefficients estimated by "joint species distribution models" provide an accurate assessment
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    of species' pairwise interactions (Pollock et al. 2014, see also Harris 2015b). In addition to
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    using the posterior mean correlation (Pollock et al. 2014), I also used the posterior mean
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Evaluating model performance. For the simulated landscapes based on Figure 1, I

partial correlation, which might be able to control for indirect effects.

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 165 significance using its Z-score. For the Markov network, I used the Hessian matrix to 166 generate approximate confidence intervals and noted whether these intervals included zero. 167 I then evaluated the relationship between each method's estimates and the "true" 168 interaction strengths among all of the species pairs from the larger simulated landscapes. 169 This determined which the methods provide a consistent way to know how strong species 170 interactions are—regardless of which species were present in a particular data set or how 171 many observations were taken. Because the different methods mostly describe species 172 interactions on different scales (e.g. correlations versus Z scores versus regression 173 coefficients), I used linear regression through the origin to rescale the different estimates produced by each method so that they had a consistent interpretation. After rescaling each 175 method's estimates, I calculated squared errors between the scaled interaction estimates and 176 "true" interaction values across all the simulated data sets. These squared errors determined the proportion of variance explained for different combinations of method and landscape size 178 (compared with a null model that assumed all interaction strengths to be zero).

180 Results

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Three species. As shown in Figure 1, the marginal relationship between the two shrub 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 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 conclusions were statistically significant. The partial correlation and Markov network 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 intervals overlapped zero in most replicates).

Twenty species. Despite some variability across contexts (Figure 3A), the four methods
that controlled for indirect effects clearly performed the best: the Markov network explained
the largest portion of the variance in the "true" interaction coefficients (35% overall),
followed by the generalized linear models (30%), partial correlations from the raw
presence-absence data (28%), and partial correlations from BayesComm, the joint species
distribution model (26%). The benefit of choosing the full Markov network over the other
three methods was largest on the smaller landscapes, which are also the ones that are most
representative of typical analyses in this field (Gotelli and Ulrich 2009).

The three methods that did not attempt to control for indirect interactions all explained 198 less than 20% of the variance. Of these, the sample correlation matrix based on the raw data 199 performed the best (19%), followed by the null model (15%) and BayesComm's correlation 200 matrix (11%). Although these last three methods had different R^2 values, there was a close 201 mapping among their estimates (especially after controlling for the size of the simulated 202 landscapes; Figure 3B). This suggests that the effect sizes from the null model (and, to a 203 lesser extent, the correlation matrices from joint species distribution models) only contain 204 noisy versions of the same information that could be obtained more easily and interpretably 205 by calculating correlation coefficients between species' presence-absence vectors. 206

Bootstrap resampling indicated that the above ranking of the different methods was

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robust (Appendix 3). In particular, the 95% confidence interval of the bootstrap distribution indicated that the Markov network's overall R^2 value was between 14 and 18 percent higher than the second-most effective method (generalized linear models) and between 2.12 and 2.38 times higher than could be achieved by the null model (Pairs). Bootstrap resampling of a 200-site landscape also confirmed that the rosalia package's estimates of species' conditional relationships were reasonably robust to sampling variation (Appendix 4).

214 Discussion

The results presented above show that Markov networks can reliably recover species' 215 pairwise interactions from observational data, even for cases where a common null modeling 216 technique reliably fails. Specifically, Markov networks were successful even when direct 217 interactions were largely overwhelmed by indirect effects (Figure 1). For cases where fitting a 218 Markov network is computationally infeasible, these results also indicate that partial 210 covariances and generalized linear models (the two methods that estimated conditional 220 relationships rather than marginal ones) can both provide useful approximations. The 221 partial correlations' success on simulated data may not carry over to real data sets, however; 222 Loh and Wainwright (2013) show that the linear approximations can be less reliable in cases where the true interaction matrix contains more structure (e.g. guilds or trophic levels). 224 Similarly, the approximation involved in using separate generalized linear models for each 225 species can occasionally lead to catastrophic overfitting with small-to-moderate sample sizes 226 (Lee and Hastie 2012). For these reasons, it will usually be best to fit a Markov network 227 rather than one of the alternative methods when one's computational resources allow it. 228 It's important to note that none of these methods can identify the exact nature of the 229 pairwise interactions (e.g. which species in a positively-associated pair is facilitating the

other; Schmidt and Murphy 2012), particularly when real pairs of species can reciprocally influence one another in multiple ways simultaneously (Bruno et al. 2003); with 232 compositional data, there is only enough information to provide a single number describing 233 each species pair. To estimate asymmetric interactions, such as commensalism or predation, 234 ecologists would need other kinds of data, as from time series, behavioral observations, 235 manipulative experiments, or natural history. These other sources of information could also be used to augment the likelihood function with an informative prior distribution, which 237 could lead to better results on some real data sets than was shown in Figure 3A. 238 Despite their limitations, Markov networks have enormous potential to improve ecological 239 understanding. In particular, they are less vulnerable than some of the most commonly-used 240 methods to mistakenly identifying positive species interactions between competing species, and can make precise statements about the conditions whre indirect interactions will 242 overwhelm direct ones. They also provide a simple answer to the question of how competition should affect a species' overall prevalence, which was a major flashpoint for the 244 null model debates in the 1980's (Roughgarden 1983, Strong et al. 1984). Equation 1 can be 245 used to calculate the expected prevalence of a species in the absence of biotic influences $(\frac{1}{1+e^{-\alpha}};$ Lee and Hastie 2012). Competition's effect on prevalence in a Markov network can then be calculated by subtracting this value from the observed prevalence (cf Figure 2D). This kind of insight would have been difficult to obtain without a generative model that 240 makes predictions about the consequences of species interactions; null models (which 250 presume a priori that interactions do not exist) have no way to make such predictions. Markov networks—particularly the Ising model for binary networks—have been studied in 252 statistical physics for nearly a century (Cipra 1987), and the models' properties, capabilities,

and limits are well-understood in a huge range of applications. Using the same framework for species interactions would thus allow ecologists to tap into an enormous set of existing 255 discoveries and techniques for dealing with indirect effects, stability, and alternative stable 256 states. Numerous other extensions are possible: for example, the states of the interaction 257 network can be modeled as a function of the local abiotic environment (Lee and Hastie 258 2012), which would provide a rigorous and straightforward approach to the difficult and important task of incorporating whole networks of biotic interactions into species 260 distribution models (Kissling et al. 2012, Pollock et al. 2014), leading to a better 261 understanding of the interplay between biotic and abiotic effects on community structure. 262 There are even methods (Whittam and Siegel-Causey 1981, Tjelmeland and Besag 1998) 263 that would allow one species to affect the sign or strength of the relationship between two other species, tipping the balance between facilitation and exploitation (Bruno et al. 2003). 265 Finally, the results presented here have important implications for ecologists' continued 266 use of null models for studying species interactions. Null and neutral models can be useful 267 for clarifying our thinking about the numerical consequences of species' richness and 268 abundance patterns (Harris et al. 2011, Xiao et al. 2015), but deviations from a particular null model must be interpreted with care (Roughgarden 1983). Even in small networks with 270 three species, it may simply not be possible to implicate individual species pairs or specific ecological processes like competition by rejecting a general-purpose null (Gotelli and Ulrich 272 2009), especially when the test statistic is effectively just a correlation coefficient (Figure 3B). 273 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 275 the range of results shown in Figure 3A, scientists in this field could often double their explanatory power by switching from null models to Markov networks (or increase it

- substantially by 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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286 References:

287 Figure captions

- 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
 occurrence vectors positively correlated (circled). C. Controlling for the tree species'
 presence with a conditional method such as a partial covariance or a Markov network leads
 to correct identification of the negative shrub-shrub interaction (circled).
- Figure 2. A. A small Markov network with two species. The abiotic environment favors
 the occurrence of both species ($\alpha > 0$), particularly species 2 ($\alpha_2 > \alpha_1$). The negative β coefficient linking these two species implies that they co-occur less than expected under
 independence. B. Relative probabilities of all four possible presence-absence combinations
 for Species 1 and Species 2. The exponent includes α_1 whenever Species 1 is present ($y_1 = 1$),

- but not when it is absent $(y_1 = 0)$. Similarly, the exponent includes α_2 only when species 2 is
- present $(y_2 = 1)$, and β only when both are present $(y_1y_2 = 1)$. The normalizing constant Z,
- ensures that the four relative probabilities sum to 1. In this case, Z is about 18.5. C. We
- can find the expected frequencies of all possible co-occurrence patterns between the two
- species of interest. D. If β_{12} equaled zero (e.g. if the species no longer competed for the same
- resources), then the reduction in competition would allow each species to increase its
- occurrence rate and the co-occurrence deficit would be eliminated.
- Figure 3. A. Proportion of variance in interaction coefficients explained by each versus
- number of sampled locations. **B.** The Z-scores produced by the null model ("Pairs") for each
- pair of species are very predictable from the correlation between the presence-absence vectors
- of those same species and from the number of sites on the landscape.
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